Electrophysiological correlates of individual differences in perception of audiovisual temporal asynchrony

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Abstract

Sensitivity to the temporal relationship between auditory and visual stimuli is key to efficient audiovisual integration. However, even adults vary greatly in their ability to detect audiovisual temporal asynchrony. What underlies this variability is currently unknown. We recorded event-related potentials (ERPs) while participants performed a simultaneity judgment task on a range of audiovisual (AV) and visual-auditory (VA) stimulus onset asynchronies (SOAs) and compared ERP responses in good and poor performers to the 200 ms SOA, which showed the largest individual variability in the number of synchronous perceptions. Analysis of ERPs to the VA200 stimulus yielded no significant results. However, those individuals who were more sensitive to the AV200 SOA had significantly more positive voltage between 210 and 270 ms following the sound onset. In a follow-up analysis, we showed that the mean voltage within this window predicted approximately 36% of variability in sensitivity to AV temporal asynchrony in a larger group of participants. The relationship between the ERP measure in the 210–270 ms window and accuracy on the simultaneity judgment task also held for two other AV SOAs with significant individual variability − 100 and 300 ms. Because the identified window was time-locked to the onset of sound in the AV stimulus, we conclude that sensitivity to AV temporal asynchrony is shaped to a large extent by the efficiency in the neural encoding of sound onsets.

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1. Introduction

Temporal proximity is one of the determining factors for integrating multisensory, and more specifically audiovisual, stimuli into a coherent percept (Stein and Meredith, 1993). Importantly, a consistent finding in research on audiovisual integration is that the perception of multisensory temporal synchrony does not require that auditory and visual stimuli occur at exactly the same time. Instead, we perceive audiovisual information as synchronous as long as the onsets of the two modalities fall within a certain temporal distance from each other, termed the temporal binding window (TBW) (for reviews, see Keetels and Vroomen, 2012; Vatakis and Spence, 2010; Vroomen and Keetels, 2010).

Arguably, one of the key features of the TBW is that its size is not constant and is influenced by many factors. It is typically larger for visual-auditory (VA) sequences of stimuli compared to auditory-visual (AV) ones (Bushara et al., 2001; Dixon and Spitz, 1980; Grant et al., 2004; Lewkowicz, 1996; van Wassenhove et al., 2007).

It is larger for speech and other complex stimuli compared to simpler non-speech stimuli (Vatakis and Spence, 2010; Vroomen and Stekelenburg, 2011); it can be reduced through perceptual training (Powers III et al., 2009; Stevenson et al., 2013); it depends on one’s expertise with specific audiovisual stimuli (e.g., Petrini et al., 2009); and it is affected by the task used to measure it (Stevenson and Wallace, 2013; van Eijk et al., 2008).

In most studies, the size of the TBW is evaluated at a group level. Significantly less research has been conducted on individual variability in sensitivity to audiovisual temporal asynchrony and its causes. The significance of individual variability in TBW is underlined by a number of findings. First, impairment in the ability to detect audiovisual temporal correspondences (and, as a result, a much broader than typical TBW) has been reported for multiple neurodevelopmental disorders (for a comprehensive review, see Wallace and Stevenson, 2014), such as dyslexia (Hairston et al., 2005), specific language impairment (SLI) (Grondin et al., 2007; Kaganovich et al., 2014), and autism (Foss-Feig et al., 2010; Kwakye et al., 2011; Stevenson et al., 2014). Importantly, at least in some of these studies, precision with which participants perceive audiovisual asynchrony predicted the degree of language and other cognitive impairments. For example, Donohue and colleagues (Donohue et al., 2012) examined a correlation between the degree...
of self-reported symptoms of autism in general population and the temporal relationship between auditory and visual stimuli that leads to the most salient perception of synchrony. They found that individuals with the greater number of autism traits consistently reported as simultaneous those stimuli in which the auditory modality slightly preceded the visual one – the pattern that is opposite to the one seen in individuals with fewer autism traits. In an earlier study from our laboratory (Kaganovich et al., 2014), those children with a history of SLI who were worse at detecting asynchrony at long stimulus-onset asynchronies (SOAs) (400–500 ms) also had lower core language scores as determined by the Clinical Evaluation of Language Fundamentals (CELF-4; Semel et al., 2003). Second, in healthy adults, the smaller size of the TBW was linked to greater susceptibility to the McGurk illusion (Stevenson et al., 2012) and to better comprehension of degraded audiovisual speech (Conrey and Pisoni, 2006). The relationship between the TBW and susceptibility to the McGurk illusion has also been replicated in children with autism (Stevenson et al., 2014). Together, these findings suggest that sensitivity to audiovisual temporal relationships may contribute to successful development of at least some cognitive and linguistic skills.

The TBW is typically measured in a simultaneity judgment task (SJT), in which audiovisual stimuli are presented in a range of SOAs, and participants have to identify each stimulus presentation as audiovisually synchronous or asynchronous. The number of synchronous perceptions is then plotted as a function of SOA, and the results are fitted to a sigmoid function, separately for AV and VA SOAs. The TBW is determined as an estimated SOA at which participants detect asynchrony with a specific degree of certainty (typically on 50–75% of trials, depending on the study). The TBW thus provides a single measure of sensitivity to audiovisual synchrony and is a succinct description of individuals’ performance on the SJT. However, the synergistic nature of this measure may also be its weakness under some circumstances. More specifically, the shape of the sigmoid function (and, as a consequence, the size of the TBW) may be determined to a greater degree by some SOAs than by others. Yet, this information is, for the most part, lost in a single TBW measure. Careful examination of published work and research in our own laboratory show that while at short (e.g., 100 ms or less) and long1 (e.g., 400 ms or more) SOAs adults largely agree in their simultaneity judgment, medium-sized SOAs lead to significant individual variability.

Several neuroimaging studies have examined the neural correlates of sensitivity to audiovisual temporal synchrony/asynchrony at a group level and reported activations in a network of regions that include both well-established multisensory areas, such as parts of the superior temporal cortex, as well as auditory and visual sensory cortices (Macaluso et al., 2004; Powers III et al., 2012; Stevenson et al., 2010) and the right insula (Bushara et al., 2001). In an insightful addition to this literature, Powers and colleagues (Powers III et al., 2012, 2009) have demonstrated that the reduction in the size of TBW following perceptual training leads to decreased activation in the key elements of the network, such as posterior superior temporal sulcus and auditory and visual cortices, as well as enhanced connectivity among them. This line of research reveals the complexity of the neural mechanisms engaged during audiovisual temporal processing and suggests that individual variability in any number of neural functions – from early sensory encoding to actual multisensory integrative mechanisms – may potentially contribute to observed individual differences in sensitivity to audiovisual temporal asynchrony.

One other aspect of audiovisual temporal processing deserves a special mention because of its relevance to the findings of the current study – namely, neural activity underlying temporal audiovisual processing can be modified not only by the physical properties of stimuli (e.g., whether the auditory and visual components of a stimulus in fact occurred at the same time) but also by the subjective perception of such properties (e.g., whether audiovisual stimuli were perceived as synchronous or asynchronus). This distinction was clearly demonstrated by the study of Stevenson and colleagues (Stevenson et al., 2011), who presented their participants with ambiguous audiovisual stimuli, which were perceived as synchronous in approximately half of all trials and asynchronous in another half. They identified two distinct areas of the multisensory superior temporal cortex (mSTC) that responded differently to physical synchronicity and to perceptual fusion – the synchrony-defined mSTC was activated by true audiovisual synchrony regardless of how it was perceived, while the bimodal mSTC responded significantly only to subjective perception of synchrony, regardless of whether the stimulus that elicited the perceptual fusion was synchronous or asynchronous.

In the current study, we combined the SJT with event-related potential recordings (ERPs) in order to focus on the timing of the neural processes engaged during the detection of audiovisual temporal asynchrony. More specifically, we asked at which point in time brain responses of individuals who are better detectors of asynchrony (i.e., good performers) differ from brain responses of those individuals who are worse detectors of asynchrony (i.e., poor performers), with the expectation that the outcome of this comparison would be informative as to the perceptual and cognitive processes that underlie individual variability in sensitivity to temporal asynchrony. Earlier ERP studies of audiovisual integration reported the attenuation of the auditory N1 and/or P2 component to audiovisual as compared to the sum of auditory only and visual only stimuli (Baart et al., 2014; Besle et al., 2004; Kaganovich and Schumaker, 2014; Knowland et al., 2014; Stekelenburg and Vroomen, 2007; van Wassenhove et al., 2005). However, given significant design differences between the SJT and the above studies, focusing on just N1 and P2 in our analyses was not justifiable. Instead, in an approach similar to the region of interest analyses used in fMRI research, we first defined our temporal windows of interest based on ERPs elicited by auditory only (a pure tone) and visual only (a flash of light) stimuli. These windows included all visible ERP components elicited by the onset of the stimuli. We then used these windows to analyze ERPs to an audiovisually asynchronous presentation of the same stimuli at the SOA that led to the largest variability in synchronous perceptions (200 ms). We compared ERPs elicited in good and poor performers in a series of t-tests conducted on each consecutive measurement point within the window of interest and used the false discovery rate (FDR) correction to control for type I error due to multiple comparisons. Following this initial step, we extended our analyses in two ways. First, in order to determine that the identified ERP differences between good and poor performers did in fact relate to their ability to detect asynchrony on a SJT, we conducted a linear regression analysis on a larger group of participants with the ERP measure as a predictor and the number of synchronous perceptions at a 200 ms SOA as an outcome. Second, to ascertain that our finding can be generalized to other SOAs with substantial individual variability, we conducted similar regression analyses between ERP measures and the number of synchronous perceptions for the 100 and 300 ms SOAs.

1 The length of what may be considered “short” and “long” SOAs will of course depend to some degree on the stimuli used. The numbers given are not meant to be absolute values but serve as an example based on our own work with non-speech stimuli.
The stimuli and design of this study were identical to those described in two previous publications from our laboratory (Kaganovich, 2016; Kaganovich et al., 2014). A 2 kHz pure tone and a flash of light (shaped as a cartoon explosion, see Fig. 1) served as stimuli. Both were 200 ms in duration. The tone was presented at 60 dB SPL via a sound bar located at the bottom of the computer monitor. In order to avoid visual after-effects, the explosion-shaped figure was shown at a slightly different location on the monitor on consecutive trials. On each trial, the auditory and visual stimuli appeared either synchronously (the SYNC condition) or at one of the following stimulus-onset asynchronies – 100, 200, 300, 400, or 500 ms. In half of all audiovisual trials, the sound preceded the flash of light (AV trials), while in another half, the flash of light preceded the sound (VA trials). Additionally, on some trials only the auditory stimulus (A) or only the visual stimulus (V) was presented. Each participant completed 10 blocks of trials, with each block containing 5 repetitions of 13 different trial types (synch, A, V, 5 AV SOAs, and 5 VA SOAs) presented in a random order. This yielded 50 responses for each trial type.

The task was presented as a game because we were collecting data from both children and adults. In this study, we focus on adult data only. Children’s performance on the task was described in earlier studies from our laboratory (Kaganovich, 2016; Kaganovich et al., 2014). At the start of the session, participants watched a video with instructions and practiced the task until it was clear. They saw a dragon in the middle of the screen and a boy and a girl with futuristic-looking weapons at the top left and right corners of the screen. Participants were told that the dragon lives on the planet Cabula, where dragons are friendly but like to raid apple orchards. The only way to keep dragons away is to use special weapons—one of them shoots lights while the other shoots sounds. If the light and the sound “hit” the dragon at the same time, the dragon runs away. However, if either the sound or the light arrives first, even by the tiniest amount, or if one of the children forgets to shoot his/her weapon (and we only see the light or only hear the sound), the dragon does not get scared away. The images of children were present at the start of each block prior to the onset of trial presentation. To avoid excessive eye movement and the shift of attention away from the dragon and the visual stimulus, there were no images of children during trial presentation. All participants sat approximately 4 feet from the monitor inside a dimly-lit sound-attenuating booth. Presentation of trials was controlled by the Presentation software (www.neurobs.com). Participants pressed one button if the dragon was scared away (i.e., the sound and the visual stimulus were perceived as occurring synchronously, or the button if the dragon was not scared away (i.e., either the sound and the explosion figure were perceived as asynchronous or only one of the two stimuli was detected (A and V trials)). Responses were recorded within a 2200 ms response window.

2 Originally, our participants were split into two groups – avid video gamers and non-gamers. Because video gamers were predominantly male, we recruited mostly male participants into the non-gamer group. However, the two groups did not differ either in behavioral or ERP measures. We therefore pooled them together in order to increase the overall number of participants and evaluate individual variability in sensitivity to audiovisual asynchrony regardless of the video gaming status.

2.3. EEG recordings

The EEG data were recorded from the scalp at a sampling rate of 512 Hz using 32 active Ag-AgCl electrodes secured in an elastic cap (Electro-Cap International Inc., USA). Electrodes were positioned over homologous locations across the two hemispheres according to the criteria of the International 10–20 system (American Electroencephalographic Society, 1994). The specific locations were as follows: midline sites: Fz, Cz, Pz, and Oz; mid-lateral sites: Fp1/Fp2, AF3/AF4, F3/F4, FC1/FC2, C3/C4, CP1/CP2, P3/P4, PO3/PO4, and O1/O2; and lateral sites: T7/T8, FC5/FC6, T7/T8, CP5/CP6, and P7/P8; and left and right mastoids. EEG recordings were made with the Active-Two System (BioSemi Instrumentation, Netherlands), in which the Common Mode Sense (CMS) active electrode and the Driven Right Leg (DRL) passive electrode replace the traditional “ground” electrode (Metting van Rijn et al., 1990). During recording, data were displayed in relationship to the CMS electrode and then referenced offline to the average of the left and right mastoids (Luck, 2014). The Active-Two System allows EEG recording with high impedances by amplifying the signal directly at the electrode (Van Bergen et al., 2006). In order to monitor eye movement, additional electrodes were placed over the right and left outer canthi (horizontal eye movement) and below the left eye (vertical eye movement). Horizontal eye sensors were referenced to each other, while the sensor below the left eye was referenced to Fpz in order to create electrooculograms. Prior to data analysis, EEG recordings were filtered between 0.1 and 30 Hz. Individual EEG recordings were visually inspected to exclude trials containing excessive muscular and other non-ocular artifacts. Additionally, the trials on which blinks occurred during the visual stimulus presentation were excluded. Ocular artifacts were corrected by applying a spatial filter (EMSE Data Editor, Source Signal Imaging, USA). Similar to the Independent Component Analysis (ICA), this method is able to separate ocular artifacts from brain activity by constructing a head model and without subtracting ocular activity from EEG channels. However, unlike the ICA, it bypasses the need to subjectively determine which components need to be excluded by relying on a representative set of artifacts. The mean data segments, ERPs were epoching starting at 200 ms pre-stimulus and ending at 1000 ms post-stimulus onset. On trials in which the auditory and visual stimuli were not completely synchronized, the stimulus onset was the onset of the first stimulus in a pair (for example, when the sound preceded the appearance of the explosion-shaped figure, ERPs were averaged to the onset of the sound). The 200 ms prior to the recording onset served as a baseline. 4. Data analysis

2.4.1. Primary analyses

Our data analysis included several distinct steps. First, in agreement with earlier reports, different SOAs led to different degrees of variability in participants’ performance (see Table 1). We selected the 200 ms SOA for both AV and VA trials as the primary focus of our analyses because it elicited the largest range in the number of synchronous perceptions and the largest standard error of the mean. We ranked all participants according to the number of synchronous perceptions, separately for the AV200 and the VA200 SOAs, with the lower number indicating greater sensitivity to audiovisual temporal asynchrony. Ten participants with the lowest number of synchronous perceptions formed the good performers group (for AV200, mean 15.4%, range 2–24%; for VA200, mean 26%, range 12–36%) while ten participants with the highest number of synchronous perceptions formed the poor performers group (for AV200, mean 76.2%, range 66–86%; for VA200, mean 70.2%,
VA indicates a stimulus in which the explosion figure preceded the tone, and AV indicates a stimulus in which the tone preceded the explosion figure. The number following the VA and AV abbreviations reflects the SOA. For example, in the VA200 stimulus, the onset of the explosion figure occurred 200 ms prior to the onset of tone. Range indicates the difference between the smallest and the largest number of reported synchronous perceptions in the group of 31 participants. Stimuli with the largest range and standard error of the mean are shown in bold.

Table 1

<table>
<thead>
<tr>
<th>Stimulus</th>
<th>Mean percent of synchronous perceptions</th>
<th>Range</th>
<th>Standard error of the mean</th>
</tr>
</thead>
<tbody>
<tr>
<td>VA500</td>
<td>1.03</td>
<td>4</td>
<td>0.24</td>
</tr>
<tr>
<td>VA400</td>
<td>3.87</td>
<td>14</td>
<td>0.79</td>
</tr>
<tr>
<td>VA300</td>
<td>19.55</td>
<td>50</td>
<td>2.58</td>
</tr>
<tr>
<td>VA200</td>
<td>48.45</td>
<td>72</td>
<td>3.49</td>
</tr>
<tr>
<td>VA100</td>
<td>88.26</td>
<td>38</td>
<td>1.71</td>
</tr>
<tr>
<td>Sync</td>
<td>91.36</td>
<td>28</td>
<td>1.53</td>
</tr>
<tr>
<td>AV100</td>
<td>81.74</td>
<td>60</td>
<td>2.84</td>
</tr>
<tr>
<td>AV200</td>
<td>45.48</td>
<td>84</td>
<td>4.73</td>
</tr>
<tr>
<td>AV300</td>
<td>13.81</td>
<td>48</td>
<td>2.14</td>
</tr>
<tr>
<td>AV400</td>
<td>3.61</td>
<td>16</td>
<td>0.83</td>
</tr>
<tr>
<td>AV500</td>
<td>1.42</td>
<td>8</td>
<td>0.38</td>
</tr>
</tbody>
</table>

Range 60–84%. Participants included in the AV200 good or poor performers group only partially overlapped with the participants included in the VA200 good or poor performers group. 

Second, we selected the temporal windows over which to compare good and poor performers’ ERPs to the AV200 and the VA200 stimuli. Because we did not have an a priori assumption about the timing of possible group differences and in order to avoid the problem of multiple implicit comparisons (Luck, 2014, p. 328), we chose measurement windows based on ERPs to A and V stimuli, in an approach similar to the region of interest analysis (ROI) used in fMRI research. More specifically, we created grand average waveforms elicited by A and V stimuli in the entire group of participants (n=31) and selected the boundaries of time windows for analyses so that they included all clearly observable auditory or visual components following the stimulus onset in this group average, as shown in Figs. 2 and 3. Auditory stimuli elicited the typical auditory N1-P2-N2 sequence. These components were broadly distributed with the maximum amplitude over the fronto-central, central, and centro-parietal scalp (see Fig. 2 Panel B), in agreement with a multitude of earlier reports (for a review, see Pratt, 2011). The entire sequence of peaks occurred between 100 and 380 ms post-stimulus onset (see Fig. 2 Panel C). Visual stimuli elicited the P1-N1-P2 sequence of ERP components (see Fig. 3), with a clear maximum, as expected, over occipital and parieto-occipital sites (see Fig. 3 Panel B). This sequence spanned the time window of 80–400 ms post-stimulus onset (see Fig. 3 Panel C).

Third, the ERPs elicited by the AV200 and the VA200 stimuli in good and poor performers were down-sampled to 100 Hz (which resulted in one measurement point for each 10 ms of recording). The groups were then compared through a series of t-tests conducted on each measurement point between 100 and 380 following the onset of the auditory stimulus and between 80 and 400 ms following the onset of the visual stimulus. To minimize the number of overall comparisons, only the sites with the largest amplitudes of auditory and visual components were included in analyses. These were FC1/2, C3/4, CP1/2 and CZ for auditory analyses and P3/4, P03/4, O1/2, and OZ for visual analyses. In order to keep the number of sites equal for auditory and visual analyses, PZ was not included in the visual analyses. The false discovery rate (FDR) method was used to control for type I error due to multiple comparisons (Benjamini and Hochberg, 1995; Groppe et al., 2011), and all reported p-values are FDR corrected values with α=0.05. The outcome of this analysis provided us with a temporal window over which good and poor performers had significantly different brain responses to the 200 ms SOA.

Fourth, the outcome of the analyses described in step three above yielded a window of significant differences between good and poor performers only for the auditory responses elicited by the AV200 stimulus. In order to determine whether the ERP mean amplitude within this window is systematically related to individual sensitivity to audiovisual temporal asynchrony, we conducted a linear regression.

Fig. 2. Auditory responses. (A) Grand average ERPs elicited by the 2 kHz pure tone on A trials are shown for the entire group of participants (n=31). Electrodes FP1 and FP2 as well as the left and right mastoids are not shown. Negative is plotted up. (B) The distribution of voltage over the scalp at the peak amplitude of the auditory N1 component based on the grand average waveform (122 ms post-stimulus onset). Note that the auditory N1 was broadly distributed across the scalp, with the largest voltages over fronto-central, central, and centro-parietal sites (shown in red in panel A). (C) The auditory ERPs elicited by the onset of the pure tone and the window used for the auditory analysis (100–380 ms post-stimulus onset) are shown on the CZ site. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
analysis with the mean amplitude within the identified window as a predictor and the number of synchronous perceptions at the AV200 SOA as the outcome. All 31 study participants were included in this analysis.

Fifth, to make sure that the relationship between the ERP measure and sensitivity to audiovisual temporal asynchrony described in step four above is not specific to the AV200 SOA, we also measured the ERP mean amplitude elicited by the AV100 and AV300 SOAs (both of which still had significant individual variability in the number of synchronous perceptions — see Table 1) within the same window identified in step 4 above (see Results) and conducted a linear regression between this mean amplitude and the number of synchronous perceptions, separately for the AV100 and AV300 SOAs. Lastly, because all available trials to the AV200 stimulus were included into individual averages for good and poor performers, it is possible that the difference in ERPs to the AV200 stimulus reflected perception of synchrony on a trial by trial basis, rather than a more general individual difference in sensitivity to audiovisual temporal information. To tease these two variables apart (perception vs. individual sensitivity), we compared the mean amplitude between 210 and 270 ms in the ERPs elicited by the AV200 stimuli when such stimuli were perceived as synchronous and when they were perceived as asynchronous. To make sure that this comparison is conducted on a similar number of trials, we used ERP data from 11 individuals who were “average” performers — they were not included in either good or poor performer groups and perceived the AV200 stimulus both as synchronous and as asynchronous on a similar number of trials.

2.4.2. Follow-up analyses

After the primary analyses of our data were completed, we conducted several follow-up analyses in order to better position our findings within the existing literature on audiovisual temporal processing.

First, in our primary analyses, we examined a correlation between voltage within the 210–270 ms window following the auditory stimulus at the AV200 SOA and the number of synchronous perceptions at the same SOA to determine whether our electrophysiological measure is related to performance on the SJT. This left open the question of whether our ERP measure would relate to a more synergistic way of evaluating sensitivity to audiovisual temporal asynchrony — namely, the TBW. Given that intermediate SOAs have the largest inter-individual variability, they should influence the size of the TBW significantly more than shorter or longer SOAs. To confirm this, we conducted two additional analyses. (1) We used the glmmP function in MATLAB (MATHWORKS, Inc., Natick, MA) in order to fit a sigmoid function to the number of synchronous perceptions separately for the VA and AV SOAs, for each individual. The width of the right and left side of each function was then measured at 50% of its maximum. We performed a one-way ANOVA to see if good and poor performers (as defined by sensitivity to the AV200 SOA) differed in the size of the AV and VA TBW. (2) We conducted a linear regression analysis between mean voltage in the 210–270 ms time window at the AV200 SOA and the size of the AV TBW in the entire group of participants (n = 31).

Second, visual examination of the ERPs elicited by the VA200 stimulus suggested that good performers may have had significantly larger visual N1 compared to poor performers (see Fig. 6). Although the temporal interval included in the t-test analyses spanned the visual N1 window, we wanted to rule out with greater certainty that good and poor performers did not differ in the amplitude of visual N1. To this end, we measured its peak amplitude between 140 and 205 ms post-stimulus onset over PO3/4, O1/2, and OZ sites and conducted a repeated-measure ANOVA with group as a between-subject variable. Further, to ascertain that the amplitude of visual N1 did not play a role in behavioral performance on the SJT, we conducted a regression analysis with the visual N1 peak amplitude as a predictor and the percent of synchronous perceptions at the VA200 SOA as the outcome on the data from the entire group of participants.

Finally, the temporal window within which the ERP voltage was significantly correlated with sensitivity to the medium AV SOAs was time-locked to the onset of the auditory stimulus (see Results). However this window (210–270 ms post-auditory onset) did not include the auditory N1, which is known to be sensitive to sound onsets. To confirm this finding, we compared the peak amplitude and latency of the auditory N1 elicited by A only stimuli in good and poor performers (as defined by performance on the AV200 SOA) measured between 100 and 160 ms post-stimulus onset. Further, in order to investigate whether the peak amplitude of the auditory N1 is related to behavioral performance on the SJT, we conducted a linear regression analysis between its peak amplitude and the number of synchronous perceptions at two AV SOAs with the largest inter-individual variability — AV200 and AV300.
3. Results

3.1. Primary analyses

Fig. 4 shows the distribution of synchronous perceptions across the range of SOAs for the entire group of participants. As expected, the number of synchronous perceptions decreased with increasing SOAs. Error bars in Fig. 4 and ranges and standard errors included in Table 1 show individual variability for each SOA and reveal that the largest range of synchronous responses was to the 200 ms SOA.

For both the AV200 and VA200 SOAs, the good and poor performers’ responses had equivalent variances based on the Levene’s test (AV200, \( F(1,19) = 363.207, p < 0.001, r = 0.98 \); VA200: \( F(1,19) = 155.38, p < 0.001, r = 0.95 \)), with predictably fewer synchronous perceptions in the good performer group.

Fig. 5 shows ERPs to the AV200 stimulus and Fig. 6 ERPs to the VA200 stimulus overlaid for good and poor performers. Panel B in each figure specifies the time window over which consecutive \( t \)-tests were performed. None of the \( p \)-values resulting from the analysis of ERPs elicited by the VA200 stimulus survived the FDR correction. Only the comparison of the auditory components elicited by the AV200 stimulus in good and poor performers yielded results that remained significant after the FDR correction (see Fig. 5, panel C). The temporal points and sites that had significant group difference are shown in Table 2. All consecutive temporal points at which more than half of electrode sites showed FDR corrected \( p \)-values of less than 0.1 were included in the final temporal window for a regression analysis. This approach yielded a time window of 210–270 ms following the onset of the auditory stimulus.

In order to determine if the ERPs to the AV200 stimulus in the window separating good and poor performers were in fact related to individuals’ ability to detect audiovisual asynchrony, we conducted a linear regression analysis with the ERP mean amplitude (averaged across the 7 electrodes used for \( t \)-test analyses) as a predictor and the percent of synchronous perceptions at the AV200 SOA as an outcome. All 31 subjects were included in this analysis. The result is shown at the top of Fig. 7. The regression produced significant results (\( \beta = -0.6, R^2 = 0.6, F(1,30) = 16.272, p < 0.001 \)), with more positive voltage between 210 and 270 ms.
post-auditory onset being associated with better sensitivity to audiovisual temporal asynchrony. The mean voltage between 210 and 270 ms post-stimulus onset accounted for approximately 36% of variability in temporal synchrony judgment ($R^2 = 0.359$).

We also evaluated whether the above finding transfers to other SOAs with significant individual variability – namely, AV100 and AV300 (see Table 1 for ranges and standard errors). We conducted two separate linear regressions with the ERP mean amplitude between 210 and 270 ms post-stimulus onset to AV100 and AV300 stimuli as a predictor and the number of synchronous perceptions at each SOA as the outcome. The results are shown at the bottom of Fig. 7. Both regressions were significant and displayed the same relationship between ERP and behavioral data as in the case of the AV200 stimulus (AV100: $\text{Beta} = -0.37, R = 0.37, F(1,29) = 4.451, p = 0.044$; AV300: $\text{Beta} = -0.564, R = 0.56, F(1,30) = 13.544, p = 0.001$).

For all of the above regression analyses, we examined the presence of outliers by evaluating the standardized DFBeta function in the SPSS Statistics program, which tests the influence of individual cases on a regression model. When the regression model is stable, excluding any one case should not have a significant influence on the outcome. Predictor cases (ERP measures) with standardized DFBeta values over 1 were considered to have a significant influence over the model and were excluded from analyses (Field, 2009). Based on this criterion, only 1 case was excluded from the AV100 regression analysis.

Because the temporal window differentiating good and poor performers was time-locked to the auditory stimulus, one interpretation of the regression analyses might be that it was the

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### Table 2
Outcome of $t$-tests comparing good and poor performers’ auditory ERP responses within the context of the AV200 stimulus.

<table>
<thead>
<tr>
<th>Site/site</th>
<th>190-200</th>
<th>201-210</th>
<th>211-220</th>
<th>221-230</th>
<th>231-240</th>
<th>241-250</th>
<th>251-260</th>
<th>261-270</th>
<th>271-280</th>
<th>281-290</th>
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<td>FC1</td>
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The top row indicates time following the onset of the pure tone within the AV200 stimulus. The ERP data was down sampled to 100 Hz; therefore, each column represents one measurement point spanning 10 ms.

* indicates FDR corrected $p$ values of 0.1 or less.

** indicates FDR corrected $p$ values of 0.05 or less. Measurement points at which more than half electrode sites showed significance at either the 0.05 or 0.1 alpha levels were included in the final window for regression analysis (i.e., all values within a rectangle).
processing of the auditory stimulus by itself rather than in the context of the audiovisual task that influenced participants’ performance. To evaluate this possibility, we measured the mean amplitude of ERPs between 210 and 270 ms post-stimulus onset elicited by A trials (i.e., trials on which only the auditory stimulus was presented) in good and poor performers and compared the two groups in a repeated-measures ANOVA with site (FC1/2, C3/4, CP1/2, CZ) as a within-subject variable. The group effect fell somewhat short of significance (group, \(F_{(1,18)}=3.804, p=0.067, \eta^2_p=0.174\), and there was no group by site interaction (\(F_{(1,18)}=0.135\)).

Finally, “average” participants perceived the AV200 stimulus as synchronous on 44.9% of trials (range 26–60%, \(SD=11.3\)). Fig. 8 overlays the ERP average of the AV200 trials when the stimulus was perceived (incorrectly) as synchronous (mean=21.7, range=13–29, \(SD=5.7\)) with the average of trials on which it was perceived (correctly) as asynchronous (mean=26.6, range=21–34, \(SD=4.9\)) in the same group of 11 subjects. In absolute terms, in the 210–270 ms window following the auditory onset, the ERP response to trials perceived as synchronous was more negative than to trials perceived as synchronous. This pattern was opposite to that obtained from the comparison of good and poor performers, with good performers showing more positive ERPs during the same time window (compare Figs. 5 and 8). A statistical comparison of ERPs elicited by trials perceived as synchronous and as asynchronous by “average” performers over the FC1/2, C3/4, CP1/2, and CZ sites showed a small trend toward significance (\(F_{(1,10)}=3.275, p=0.1, \eta^2_p=0.247\)).

3.2. Follow-up analyses

The VA and AV TBW for good and poor performers at 50% of the function’s maximum are shown in Fig. 9 panel A. Good performers had a significantly smaller AV TBW window than poor performers (134 ms (\(SD=29\)) vs. 270 ms (\(SD=22\)), respectively); however, the groups did not differ in the size of the VA TBW (192 ms (\(SD=34\)) vs. 206 ms (\(SD=40\))) (modality order (VA vs. AV) by group (good vs. poor performers), \(F_{(1,18)}=34.204, p<0.001, \eta^2_p=0.655\); VA TBW, \(F_{(1,19)}<1, r=0.2\); AV TBW, \(F_{(1,19)}=135.068, p<0.001, r=0.94\)).

The size of the AV TBW in the entire group of participants (\(n=31\)) was significantly correlated with mean voltage between 210 and 270 ms post-sound onset at the AV200 SOA (see Fig. 9, panel B) (\(B = -12.668, R=0.488, F_{(1,30)}=9.061, p=0.005\), with more positive voltage being associated with better sensitivity to audiovisual temporal asynchrony.

Analysis of visual N1 showed that its peak amplitude tended to be larger in good compared to poor performers (as defined by accuracy at the VA200 SOA), although the group effect did not quite reach significance (\(F_{(1,18)}=3.615, p=0.073, \eta^2_p=0.167\)). Importantly, there was no relationship between the visual N1 peak amplitude and sensitivity to audiovisual asynchrony at the VA200 SOA (\(B = 1.868, R=0.298, F_{(1,30)}=2.82, p=0.104\)).

Finally, the N1 peak amplitude and peak latency elicited by the A only stimulus did not differ between good and poor performers (as defined by performance on the AV200 SOA) (N1 peak amplitude: group, \(F_{(1,18)}<1, \eta^2_p=0.029\); group by site, \(F(6,46.342)<1, \eta^2_p=0.016\); N1 peak latency: group, \(F_{(1,18)}=2.446, p=0.135, \eta^2_p=0.12\)). Furthermore, there was no correlation between either

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**Fig. 7.** Regressions. An outcome of the regression analyses with the mean voltage between 210 and 270 ms post sound onset as a predictor and the number of synchronous perception on the SJT as an outcome (\(n=31\)).

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<td>Beta</td>
<td>0.564, (R^2=0.318, F(1,30)=13.544, p=0.001)</td>
<td>0.37, (R^2=0.137, F(1,29)=4.451, p=0.044)</td>
<td>0.564, (R^2=0.318, F(1,30)=13.544, p=0.001)</td>
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the peak amplitude or the peak latency of N1 to A only stimulus and performance on either the AV200 or AV300 SOA (N1 peak amplitude: AV200, B = −0.965, R = 0.118, \( F(1,30) < 1 \); AV300, \( B = −0.238, R = 0.064, F(1,30) < 1 \); N1 peak latency: AV200, \( B = 733.9, R = 0.278, F(1,30) = 2.432, p = 0.13 \); AV300, \( B = 117.43, R = 0.098, F(1,30) < 1 \).

4. Discussion

We have determined that the ERP response between 210 and 270 ms following the auditory stimulus onset can predict up to 36% of individuals’ ability to detect temporal AV asynchrony at the intermediate (100–300 ms) SOAs. More specifically, those individuals who had more positive mean voltage within this window detected the audiovisual asynchrony with higher accuracy. In contrast, none of the analyses of ERPs to VA SOAs survived the FDR correction.

Because the temporal window predictive of performance at the AV SOAs was time-locked to the onset of the sound, the most logical explanation for our finding is that the accuracy with which individuals detect audiovisual asynchrony is tied to their processing of the auditory stimulus. A more nuanced understanding of the nature of the sensory and/or cognitive mechanisms unfolding during the 210–270 ms time window will require further study. One possibility is that the enhanced positivity in good performers reflects a more detailed neural representation of the sound onset, which can significantly facilitate judgment about its temporal overlap (or the lack thereof) with the visual stimulus. In agreement with this proposition, a comparison of ERPs to the A stimulus in good and poor performers showed a strong trend (\( p = 0.067 \)) for greater positivity during the 210–270 window in good performers, suggesting that our finding is not specific to audiovisual stimuli.

Alternatively, electrophysiological processes taking place during the identified window may reflect the encoding of the offset of the sound stimulus. However, because our stimulus was 200 ms in duration, the offset-elicited ERPs would have had the latency of just 1–70 ms, which is unlikely given earlier research on the latency of sound offset-elicited ERPs (Pratt et al., 2005). Nevertheless, to rule out this possibility more definitively, studies that use varying sound durations and/or examine a relationship between ERPs elicited by sound gaps and performance on the SJT are needed. It is noteworthy that in both visual and auditory modalities, the N1 component is thought to reflect sensory processing of stimulus onsets and is influenced by such basic properties as brightness and saturation in visual modality and intensity and frequency in auditory modality. Somewhat surprisingly, the peak amplitude of these components did not differentiate good and poor performers in our study and was not related to performance on the SJT, suggesting that early sensory encoding does not play a major role in the ability to detect audiovisual temporal asynchrony.

The strength of a relationship between ERP measures and sensitivity to audiovisual asynchrony was comparable for the AV200 and AV300 SOAs (\( R^2 = 0.359 \) and 0.318, respectively). This relationship was still significant but markedly weaker (\( R^2 = 0.137 \)) for the AV100 SOA. One reason for this finding likely lies in the less diffuse distribution of participants’ responses to the AV100 SOA. A close examination of Fig. 7 reveals that a large proportion of participants perceived this stimulus as synchronous on 80–100% of trials, with relatively few perceiving synchrony on fewer than 70% of trials. Therefore, responses to the AV100 SOA had overall
events) is shaped primarily by the strength of multisensory integrative mechanisms triggered by the VA stimuli. While, admittedly, this proposal requires direct testing, it fits well with multiple well-established findings in research on audiovisual temporal processing, which consistently show significant differences in behavioral and neurophysiological responses to AV and VA temporal sequences.

To start with, an asymmetry in sensitivity to AV and VA asynchrony at moderate SOAs is well-established and has been replicated numerous times (Bushara et al., 2001; Dixon and Spitz, 1980; Grant et al., 2004; Lewkowicz, 1996; van Wassenhove et al., 2007). Across different ranges of SOAs and types of stimuli, adults tend to be more accurate at detecting AV compared to VA asynchronies. The ability to detect multisensory temporal asynchrony has a prolonged developmental course, with even adolescents still performing worse than adults (Hillock-Dunn and Wallace, 2012). More importantly, perception of AV asynchrony matures before that of VA asynchrony, with 10–11 children significantly outperforming their 7–8 year old peers on a SJT, but only at AV SOAs (Kaganovich, 2016). Further, several studies have demonstrated that the size of the TBW is malleable and can be reduced through perceptual training (Powers III et al., 2012, 2009; Stevenson et al., 2013). Of note, however, is the fact that in all such studies, the reduction of the TBW was due to increased accuracy on the VA sequences of the SJT only. In other words, the perception of AV and VA asynchronies responds differently to training. In a similar vein, in the study by Stevenson and colleagues (Stevenson et al., 2012), only the size of the VA portion of the TBW significantly predicted individuals’ susceptibility to the McGurk illusion. Accounting for all of the above differences is beyond the scope of the current report. However, our finding contributes to the cumulative message of this literature by showing that the neural encoding of auditory stimuli also differs during the processing of AV and VA sequences.

While the number of studies on the neural correlates of audiovisual integration is quite large, only a few focused specifically on sensitivity to audiovisual temporal synchrony/asynchrony (Bushara et al., 2001; Powers III et al., 2012; Stevenson et al., 2010). These studies have not reported any major differences in the neural networks for detecting asynchrony in AV and VA stimuli – the result that, seemingly, does not agree with our finding. In one of the earliest studies, Bushara and colleagues (Bushara et al., 2001) reported a nearly identical set of activated brain regions when their participants made simultaneity judgments about AV and VA stimuli, with the only difference being the bilateral, rather than the right unilateral, activation of the inferior frontal gyrus. However, in this analysis, neural responses to both short and long SOAs were averaged together, potentially obscuring modality order differences that are most noticeable at intermediate SOAs. In an additional analysis, these authors also identified the brain region - namely, the right insula - whose activation correlated with the difficulty of the SJT. Unfortunately, in this case, the neural activity during AV and VA trials was averaged together and no interaction between the order of modalities and the degree of the right insula activation was reported. The studies by Stevenson et al. (2010) and by Powers III et al. (2012) both focused primarily on identifying brain regions that respond preferentially to either synchronous or clearly asynchronous audiovisual stimuli, with little detail provided about potential modality order differences. The report by Stevenson and colleagues (Stevenson et al., 2011) on different neural correlates of physical and perceived audiovisual temporal synchrony used only auditory-visual sequences as stimuli. In sum, recent findings have identified important components of the neural network underlying audiovisual temporal function. However, whether AV and VA sequences lead to similar patterns in the activation of this network remains undetermined.
When evaluating ERPs elicited by AV200 and VA200 stimuli in good and poor performers, we included all trials – the ones that were perceived as synchronous and the ones that were perceived as asynchronous – in our averages. Since good performers had more trials on which they subjectively perceived asynchrony, could the ERP differences we found for the AV200 SOA reflect the subjective perception of asynchrony rather than better encoding of the auditory stimulus as suggested above? A comparison of ERPs to the AV200 stimulus when it was perceived (incorrectly) as synchronous and when it was perceived (correctly) as asynchronous in the group of “average” performers helped us address this issue. Although this analysis did not reach significance at the alpha level of 0.05, its effect size was quite large ($\eta^2_p=0.247$). Because the perception-driven analysis was based on a relatively small number of trials and included only 11 participants, it is likely that it lacked sufficient power. Yet, its outcome is suggestive, especially because it yielded a pattern of results that was opposite in polarity to that seen in the analyses based on overall performance. More specifically, while good performers at the AV200 SOA had more positive voltage 210–270 ms post-auditory onset compared to poor performers, the correct perception of asynchrony at the AV200 SOA was associated with more negative voltage in the same window (compare Figs. 5 and 8). This finding suggests that the ERP indices of efficient encoding of stimulus onsets within multisensory contexts may dissociate from the ERP indices of subjective perception of audiovisual asynchrony, perhaps in a manner similar to that reported by Stevenson and colleagues for fMRI measures (Stevenson et al., 2011). However, more studies that carefully manipulate perception of audiovisual asynchrony across participants are needed to make more definitive conclusions about differences or similarities in the ERP correlates of objective vs. subjective audiovisual temporal synchrony.

Lastly, as all studies, our report has its limitations. The auditory stimuli used in the current paradigm were pure tones with a short rise and fall time. We know that sound complexity and its “speechness” can significantly affect both the amplitude and latency of auditory ERP components (e.g., Čepioniene et al., 2005). An important empirical question to address in future work is whether the time window identified in our study remains constant across different auditory stimulus types. Additionally, because our stimuli were 200 ms in duration, the amount of overlap between auditory and visual modalities varied for different SOAs. At longer SOAs (300–500 ms), the processing of both the onset and the offset of the first stimulus in a pair might have been completed prior to the onset of the second stimulus, potentially facilitating decision making at longer SOAs. Most of the earlier studies used significantly shorter stimuli in similar paradigms (~5–10 ms) (e.g., Grondin et al., 2007; Hillock-Dunn and Wallace, 2012). However, since many audiovisual events in real life last more than 10 ms, we believe our results are informative. Nevertheless, replicating our findings with shorter stimuli would help rule out modalities’ overlap as a contributing factor to audiovisual temporal sensitivity.

To conclude, we examined electrophysiological correlates of individual variability in sensitivity to audiovisual temporal asynchrony in a SJT. We reported that good and poor performers had significantly different ERPs to the AV200 stimulus 210–270 ms following the sound onset, with greater positivity in good performers. Furthermore, the mean amplitude of voltage in this window accounted for a significant amount of variability in performance not only at the AV200 SOA but also at the AV100 and AV300 SOAs in a larger group of participants. We suggested that the ability to detect temporal asynchrony in AV sequences depends, to a large extent, on the efficient encoding of auditory onsets.

Acknowledgements

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