Auditory hallucinations and brain structure in schizophrenia: voxel-based morphometric study

Igor Nenadic, Stefan Smesny, Ralf G. M. Schlösser, Heinrich Sauer and Christian Gaser

Summary

We applied voxel-based morphometry to high-resolution magnetic resonance images of 99 participants with schizophrenia. Voxel-wise correlations with a score of auditory hallucination severity identified areas in the left and right superior temporal cortices and subcortical or cerebellar clusters correlated with severity of auditory hallucinations related to modality-specific superior temporal areas including primary and secondary auditory cortices.

Declaration of interest

None.

Method

We analysed high-resolution images acquired on a 1.5 T Philips ASCII MRI scanner (Philips Medical Systems, Best, The Netherlands): repetition time 13 ms, echo time 5 ms, flip angle 250, 256 sagittal slices, voxel dimensions 1 mm × 1 mm × 1 mm. Participants were 99 patients with schizophrenia (57 men and 42 women, aged 18–65 years, mean 36.2 years, s.d. = 11.2) treated at our department and diagnosed according to both ICD–10 and DSM–IV using semi-structured interviews. None of the patients had a second psychiatric, neurological or major medical condition, or history of traumatic brain injury. All participants were stable in terms of both medication and clinical symptoms before undergoing scanning and rating with the Scales for Assessment of Positive/Negative Symptoms (SAPS/SANS).8 The sample was partly based on a previously studied cohort of 85 patients.7,9 All patients gave written informed consent to a study protocol approved by the ethics committee of the University of Jena medical school. For MRI analysis we applied VBM2 (http://dbm.neuro.uni-jena.de/vbm), implemented as a toolbox in SPM (Institute of Neurology, London, UK), using hidden Markov random fields to increase the signal-to-noise ratio. Images were segmented and normalised onto a previously created study-specific custom template. We computed an auditory hallucinations score as a sum of SAPS single items for auditory hallucinations, voices conversing and voices commenting. For statistics, we used a general linear model entering the auditory hallucinations score as a variable of interest (regressor) and removing effects of age, gender and SAPS total score (without auditory hallucinations) to obtain voxel-wise correlations of each cerebral grey matter voxel with auditory hallucination severity. Based on the previous findings and the resulting anatomical hypotheses, we applied an uncorrected height threshold of P<0.01 as well as an extent threshold of k = 723 voxels (expected number of voxels according to random field theory).

Results

The distribution of hallucination scores is given in online Fig. DS1. Voxel-based analyses (Fig. 1) revealed six cortical (but no subcortical or cerebellar) clusters correlated with severity of auditory hallucination (maximum voxel coordinates and t-values in parentheses): left primary/secondary auditory cortex (−58, −11, 9; t = 3.30); left anterior superior temporal cortex (−57, 3, −10; t = 3.82); right superior temporal gyrus including the primary/secondary auditory cortex (64, −19, 14; t = 3.75) and posterior cingulate cortex. This study extends previous region-of-interest studies demonstrating main effects of auditory hallucinations related to modality-specific superior temporal areas including primary and secondary auditory cortices.

Fig. 1 Voxel-based morphometry analysis of voxel-wise correlations of grey matter with auditory hallucination severity score in the patient cohort. Significant results with P<0.01 (uncorrected) and extent threshold k = 723 voxels are projected onto the cortical surface.
mid-superior temporal gyrus area (55, −14, 0; \( t = 3.53 \)); left angular gyrus (51, −51, 18; \( t = 4.18 \)); left postcentral gyrus (−12, −38, 75; \( t = 3.27 \)); and left posterior cingulate cortex extending towards the precuneus (−14, −53, 12; \( t = 3.16 \)).

Discussion

We found several temporal and parietal brain areas to be correlated with severity of auditory hallucinations in schizophrenia. This finding offers three important new insights. First, the superior temporal gyrus is affected bilaterally to different extents in specific subregions. Importantly, this includes the primary and secondary auditory areas, long suspected to be a modality-specific correlate of this symptom. These areas coincide with findings from functional MRI studies, as well as potential target areas of repetitive transcranial magnetic stimulation therapy for treatment-resistant auditory hallucinations. To our best knowledge our study is the largest VBM study of its kind, and is thus likely to provide higher statistical power than previous studies of hallucinations in schizophrenia with smaller sample sizes. However, its cross-sectional nature precludes identification of the timing of these changes. A study of auditory hallucinations in bipolar disorder, in which this symptom is much less frequent, found a similar correlation for a slightly more inferior cluster in the middle temporal cortex. Second, our findings suggest that this specific association of superior temporal/Heschl’s gyrus alterations and auditory hallucinations is not categorical, but rather reflects an increasingly higher load of structural changes to be associated with symptom expression. This implies a dimensional rather than categorical association. Some previous MRI studies have been restricted to people with persistent auditory hallucinations, which gives some advantage for phenotypic stability. In contrast, our correlational approach takes into account patients with a low probability of having (or having had) auditory hallucinations and also those without current hallucinations. Considering other disease-related sources of variation, the findings are thus less likely to reflect changes due to selection of a subgroup of patients with schizophrenia in whom auditory hallucinations persist over longer periods or are treatment-refractory. Third, we found several areas outside the superior temporal gyrus to be correlated with auditory hallucinations as well. Although the inferior lateral parietal cluster (located somewhat posterior to a supramarginal cluster of our previous study) may be related to verbal aspects of hallucinations, we did not replicate our previous finding of right prefrontal changes. Also, the postcentral changes, although described in another VBM study of auditory hallucinations, have not been included in any of the current hypotheses (either misattribution of inner speech or modality-specific sensory cortical dysfunction); hence, their significance remains unclear.

A few limitations should be considered. First, VBM does not disclose the nature of underlying neuropathological changes. Although reduction of neuropil would be a potential substrate, we cannot exclude effects of local cerebral blood volume or flow. As with most studies correlating a state-related variable (such as psychopathological symptoms) with a rather trait-related variable (such as brain structure), findings might be prone to type 2 errors, as symptoms are more dynamic than structural changes. However, it is important to note the stable psychopathological state of our patients, and that our superior temporal gyrus findings in particular are consistent with previous findings using region-of-interest or VBM methods. Also, we statistically removed variance related to SAPS total score (without auditory hallucination items), which makes it less likely that results reflect significant correlations with positive symptoms overall. Finally, we need to consider the effect of antipsychotic medication, since we were not able to correct for lifetime antipsychotic exposure, which might vary in relation to symptom profiles.

Taken together, our findings provide refined anatomical mapping of auditory hallucinations in schizophrenia using the largest sample studied so far, and have identified a set of superior temporal cortical areas, including primary and secondary auditory cortex, which provide a modality-specific structural correlate of this frequent symptom of schizophrenia.

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References

Fig. DS1  Distribution of auditory hallucination scores in participants with a score of at least 1 (n = 38). Data for participants without auditory hallucinations (n = 61) are not shown but are used in the morphometric analysis.
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