Analysis of local gyrification index using a novel shape-adaptive kernel and the standard FreeSurfer spherical kernel – evidence from chronic schizophrenia outpatients

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ABSTRACT

Schizophrenia can be considered a brain disconnectivity condition related to aberrant neurodevelopment that causes alterations in the brain structure, including gyrification of the cortex. Literature findings on cortical folding are incoherent: they report hypogyria in the frontal, superior-parietal and temporal cortices, but also frontal hypergyria. This discrepancy in local gyrification index (LGI) results could be due to the commonly used spherical kernel (Freesurfer), which is a method of analysis that is still not spatially precise enough. In this study we would like to test the spatial accuracy of a novel method based on a shape-adaptive kernel (Cmorph). The analysis of differences in gyrification between chronic schizophrenia outpatients (n = 30) and healthy controls (n = 30) was conducted with two methods: Freesurfer LGI and Cmorph LGI. Widespread differences in the LGI between schizophrenia outpatients and healthy controls were found using both methods. Freesurfer showed hypogyria in the superior temporal gyrus and the right temporal pole; it also showed hypergyria in the rostral-middle-frontal cortex in schizophrenia outpatients. In comparison, Cmorph revealed that hypergyria is equally represented as hypogyria in orbitofrontal and central brain regions. The clusters from Cmorph were smaller and distributed more broadly, covering all lobes of the brain. The presented evidence from disrupted cortical folding in schizophrenia indicates that the shape-adaptive kernel approach has a potential to improve the knowledge on the disrupted cortical folding in schizophrenia; therefore, it could be a valuable tool for further investigation on big sample size.

1. Introduction

The core of psychopathological symptoms in schizophrenia is thought to be an effect of disrupted functional and structural connectivity in the brain that is related to aberrant neurodevelopment, which is believed to start at the same time as the formation of the brain tissues (Friston and Frith, 1995; Friston et al., 2016; Pettersson-Yeo et al., 2011). It is postulated that disconnectivity can have an impact on other neurodevelopmental processes, such as synaptic pruning and myelination (Cassoli et al., 2015; Rapoport et al., 2012). Moreover, tension-induced growth of white matter is considered able to modulate the folding pattern of the cortical surface (Garcia et al., 2018). In an attempt to better characterize neurodevelopmental underpinnings and identify connectivity alternations in schizophrenia, some recent studies have focused on cortical gyrification (Matsuda and Ohb, 2018), which can be defined as a process of cortical folding in which the initially smooth surface begins to buckle in order to fit into a relatively small space in the skull (Garcia et al., 2018). The most common measure of cortical folding is the gyrification index (GI), which is the ratio of the length of the outer folded surface of the brain to the length of the outer surface excluding sulci (Zilles et al., 1988). Briefly, brains with a higher degree of cortical folding have larger GI values. Initially, this method involved manual GI tracing, which relied on measurements collected by hand from coronal plane slices, often on post-mortem or 2D MRI brain slices (Zilles et al., 1988). Besides the fact that this was expensive and time-consuming, the main criticisms were related to the precision and accuracy of estimation of the architectural complexity of cortical folding because this was captured in only two dimensions (i.e. 2D-related underestimation tendency). Subsequently, the automated GI method was introduced (Moorhead et al., 2006), followed by the local gyrification index (LGI, Schaer et al., 2008).
The latter, in contrast to a single-point 2D measure, introduces the 3D spherical kernel concept and defines LGI as an area ratio between the outer hull and the pial surface. This method quickly gained popularity as it is more accurate, is accessible from open-source FreeSurfer software (http://surfer.nmr.mgh.harvard.edu/), and is widely used in clinical studies (Matsuda and Ohi, 2018). Importantly, some authors suggest that alterations in cortical folding measured with LGI may be regarded as a biomarker of schizophrenia (Matsuda and Ohi, 2018; White and Hilgetag, 2011). However, consistent conclusions are precluded by the discrepancies between the LGI results in the literature, indicating that abnormalities are more related to increased (hypergyria) or decreased (hypogyria) LGI (Nenadic et al., 2015; Palaniyappan et al., 2011; Sasabayashi et al., 2016, 2019). Previous studies on LGI in schizophrenia have mostly reported global suppression of cortical folding in comparison to healthy controls (White and Hilgetag, 2011). So far, hypogyria has been found mainly in the frontal, superior-parietal and temporal cortices (Cao et al., 2017; Kubera et al., 2018; Yan et al., 2019), as well as in the temporal lobes (Spalthoff et al., 2018), the anterior (Cao et al., 2017) and posterior cingulate cortex (Wheeler and Harper, 2007), the supramarginal gyrus (Cao et al., 2017; Kubera et al., 2018), the insula (Palaniyappan et al., 2013), the visual cortex (V1, V2, V5) (Schultz et al., 2012), and the multimodal association cortex (Palaniyappan and Liddle, 2012) importantly, gyri

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...lent factor (Matsuda and Ohi, 2018). Lastly, comparative research including other psychiatric diagnoses indicates that LGI changes are more severe than for, e.g., bipolar disorder (Cao et al., 2017; Nenadic et al., 2015) and exist in the broad spectrum of schizophrenic psychoses (Nanda et al., 2014; Sasabayashi et al., 2019).

On the other hand, along with the aforementioned hypogyria findings, some post-mortem (Vogeley et al., 2000), manual tracing or automated GI studies have revealed hygryrrification of the prefrontal cortex in chronic schizophrenia patients (Janssen et al., 2014; Schultz et al., 2010; Vogeley et al., 2000) and in non-affective first-episode psychosis subjects (Zuliani et al., 2018). Furthermore, using surface-based LGI methods, Palaniyappan and Liddle (2012) showed both hypogyryrrification and hygryrrification in the prefrontal cortex. Moreover, hygryrrgyria in the precuneus and the superior parietal cortex (Kubera et al., 2018) and in the visual cortex (Schultz et al., 2013) have been reported in schizophrenia with the same LGI method.

When summarizing the evidence and the inconsistency of the existing results in the literature, the following questions still seem to be insufficiently resolved: a) is schizophrenia particularly characterized by hypogyria, hypergyria, or both; b) are the methods used sufficiently spatially sensitive and morphologically exact to detect these cortical folding abnormalities accurately. To tackle the problem of the large amount of spatial (non-linear) data, researchers have incorporated kernel estimation into analyses in a broad range of scientific disciplines, e.g. robotics (Liu et al., 2016b; Sugiyama et al., 2008) and geodesy (Grazzini and Soille, 2009; Yuan et al., 2019). From this perspective, analysis of brain gyri

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The pial and white surfaces were checked (using Freeview v2.0.0) for the occurrence of any obvious errors (e.g. tissue fragments deleted) during the reconstruction process. No manual edits were made. One clinical subject was removed from the analysis due to insufficient reconstruction of the brain.

Secondly, the standard LGI (Schaer et al., 2008, 2012) implemented in FreeSurfer (i.e. the recon-all command with -localGI -qcache -measure pial_lgi options) was computed for all subjects. Oval-shaped regions of interest on the outer surface were matched with the corresponding regions of interest on the inner surface (as a result of using a 25mm spherical kernel at each vertex). The ratio of the cortical surface area to outer surface area was calculated and the values were then placed on the corresponding cortical surface vertices, thus creating a global heat map of the LGI.

Thirdly, a similar vertex-wise LGI map was calculated with the Cmorph v1.2 package (Lyu et al., 2018; https://github.com/ilwoolyu/LocalGyri fi cationIndex), MATLAB R2019a (MathWorks, Inc., Natick, Massachusetts, United States) and GNU parallel software (Tange, 2011; http://www.gnu.org/software/parallel). As the input we used FreeSurfer pial and white surfaces from each subject and hemisphere, converted to triangular 3D vtk format. The Cmorph method adapts the shape of the LGI kernel to elongate it along the sulcus/gyrus and keep it spherical near any plateaus. To obtain this effect, a tensor field from a geodesic distance (travel-time) map is created. The wavefront propagation along the tensor field enables the program to construct the desired shape-adaptive kernels for the LGI computations at each vertex of the input surfaces.

Then, before group comparison statistics, the average brain surface was constructed for all subjects' vertices by mapping them to common surface space corresponding to the surface locations of the individuals (fsaverage command).

Finally, two separate between-group Generalized Linear Model (GLM) analyses of both LGI measures (Cmorph and FreeSurfer) were performed by whole-brain vertex-wise analysis with permutation-based cluster correction for multiple comparisons (mri_glm fit -sim): 1. group with gender as a silent regressor in the group intercept GLM model; 2. group intercept contrast with gender and age as regressor (DODS option in FreeSurfer). Bidirectional contrasts were applied to both LGI methods (i.e. healthy controls > schizophrenia outpatients, healthy controls < schizophrenia outpatients). All contrasts for between group comparison were visualised by FreeView v.2.0 with smoothing 0 mm FWHM, vertex-wise threshold p = 0.01, and cluster-wise (CWP) threshold at p = 0.001. No additional formal statistical comparison between Cmorph and FreeSurfer methods was done, except this described by Lyu et al. (2018).

3. Results

Both LGI measures revealed various bilateral differences between schizophrenia and healthy controls (GLM contrast: group x gender x age).
However, essential differences in cluster sizes, localization, and the output of directional contrasts (i.e. Cluster Growing Summary by mri_surfcluster command) should be clearly pointed out. The spherical kernel LGI analysis (FreeSurfer) revealed 3 bilateral temporal region clusters that indicate hypogyria in schizophrenia and 5 clusters in frontal regions related to hypogyria; total size of hypogyria 1259 mm²; hypergyria 219 mm². At the same time, the cortex shape-adaptive kernel method (Cmorph) revealed widespread bidirectional differences, i.e. 23 hypogyria clusters (13 left hemisphere/10 right hemisphere) in 18 fronto-temporo-parietal brain regions in both hemispheres, and 25 hypergyria clusters (14 left hemisphere/11 right hemisphere) in 20 fronto-temporo-parieto-occipital brain regions in schizophrenia; total size of hypogyria 994 mm²; hypergyria 1184 mm². It is noteworthy that the adaptive kernel approach revealed smaller and more precise localized clusters within bidirectional contrasts as compared to the standard FreeSurfer output. Detailed statistics and labelling of cluster annotations are presented in Table 2. Visualization of schizophrenia-related hypogyria (red) and hypergyria (blue) clusters from both assessed LGI methods for group x gender x age GLM model are presented in Figure 1.

4. Discussion

To the best of our knowledge, this is the first study that has attempted to use the cortex shape-adaptive kernel approach in estimating cortical folding differences in schizophrenia. The results obtained from both methods (Cmorph and FreeSurfer) indicate widespread differences in LGI between schizophrenia outpatients and healthy controls. Both methods revealed consistent hypogyria findings in bilateral temporal and frontal hypergyria regions. Additionally, the clusters from the shape-adaptive kernel method are smaller and more spread out across the brain and create more diffuse and complex patterns of bidirectional changes on cortical folding in schizophrenia, i.e. fronto-temporo-parieto-occipital hypogyria and hypergyria. Overall, the presented results indicate that Cmorph LGI may be successfully used for LGI measurement in schizophrenia and other clinical populations.

4.1. Spherical kernel LGI

Firstly, the presented results from the spherical kernel LGI indicated hypogyriﬁcation only in the temporal regions (i.e. superior temporal gyrus) of both hemispheres and in the right temporal pole in schizophrenia outpatients. Similar structures were reported previously in people with a high clinical risk of schizophrenia (Damme et al., 2019), and in people with resistant auditory hallucinations (Cachia et al., 2008). Moreover, a big cluster of hypogyria but only in the right superior temporal gyrus in adolescents with schizophrenia was also identiﬁed (Palaniyappan and Liddle, 2012). Finally, the reported hypergyria in the precentral regions which would blend into one cluster in standard FreeSurfer LGI. The results obtained from both methods (Cmorph and FreeSurfer) indicate widespread differences in LGI between schizophrenia outpatients and healthy controls. Both methods revealed consistent hypogyria findings in bilateral temporal and frontal hypergyria regions. Additionally, the clusters from the shape-adaptive kernel method are smaller and more spread out across the brain and create more diffuse and complex patterns of bidirectional changes on cortical folding in schizophrenia, i.e. fronto-temporo-parieto-occipital hypogyria and hypergyria. Overall, the presented results indicate that Cmorph LGI may be successfully used for LGI measurement in schizophrenia and other clinical populations.

4.2. Hypogyria vs. hypergyria in schizophrenia

Hypogyria may be related to altered brain neurodevelopment and maturation. It is believed that axonal tension is not sufﬁcient to induce cortical folding, but tension-induced growth of white matter could modulate the folding of the cortex (Garcia et al., 2018). This is in line with studies on white matter in schizophrenia that show widespread disconnectivity in patients (Wheeler and Voinoskos, 2014). However, it only explains the source of the differences in cortical gyriﬁcation, but not its diversity. Other neurodevelopmental processes could have an impact on folding, for example, cell migration. Altered neurodevelopment has an inﬂuence on cortical thickness and some ﬁndings indicate thinning of the prefrontal cortex in schizophrenia (Nesvåg et al., 2008), which also shows hypergyriﬁcation in high genetic risk subjects (Dauvermann et al., 2012). Additionally, as suggested by some authors (Palaniyappan et al., 2013), regions which are involved in deﬁcits in schizophrenia patients, such as the multimodal association cortex, could have altered gyriﬁcation (e.g. the superior temporal gyrus, as shown in our study).

Yet, our results may shift the balance in the discussion towards the existence of both hypogyria and hypergyria in schizophrenia patients, both of which can be detected by the spherical-kernel method but are much more clearly seen with the cortex shape-adaptive kernel approach.

4.3. Cmorph – shape-adaptive kernel LGI

Cmorph methods revealed that hypergyria is equally represented in the schizophrenic brain as hypogyria, which is easily visible in a wide range of fronto-temporo-parieto-occipital brain regions. This may be related to two main factors which stand out when these two methods are compared in terms of their spatial accuracy in identiﬁcation of regions with altered brain gyriﬁcation: cluster size and biological mapping accuracy. In fact, standard LGI shows big clusters covering a few gyri, for example, in the right superior-temporal cortex (ca. 1200mm2) (see Figure 1), while clusters from Cmorph LGI are smaller, e.g. three clusters identiﬁed in the right superior-temporal region (ca. 105mm2). Moreover, the clusters from Cmorph seem to be more morphologically accurate and can ﬁt into a single gyrus/sulci and are thus distributed more broadly across the brain. Results from the shape-adaptive kernel LGI coincide with the spherical kernel LGI, but due to its more relevant measurements (smaller kernel size) Cmorph LGI is able to reveal small differences in regions which would blend into one cluster in standard FreeSurfer LGI.

As a result, differences can be detected that are coherent with other studies (Nesvåg et al., 2014; Schmitt et al., 2015; Schultz et al., 2010; Yan et al., 2019) despite the relatively small sample-size examined in our study. For example, the presented ﬁndings on hypogyria in the precentral gyrus and the middle frontal regions are complementary with previous studies (Schmitt et al., 2015; Yan et al., 2019) and ﬁndings on altered LGI in the right precuneus (Nesvåg et al., 2014) or supramarginal gyrus (Palaniyappan and Liddle, 2012). Finally, the reported hypergyria in the lateral orbitofrontal cortices, rostral middle frontal gyr, and occipital area regions are similar to those pointed out in a study on ﬁrst-episode schizophrenia patients (Sasabayashi et al., 2016).

4.4. Limitations and future directions

In summary, this is the ﬁrst Cmorph LGI study on schizophrenia, but some limitations preclude explicit conclusions concerning the unique patterns of cortex folding in schizophrenia. First of all, replication on a bigger sample size is needed and this work should be considered as an
Table 2. Comparison of the between group differences in the LGI obtained with FreeSurfer spherical kernel method and Cmorph shape-adaptive kernel method. Statistical analysis from mri_glmfit–sim utilized a whole-brain vertex-wise analysis with permutation-based cluster correction for multiple comparisons, controlled for age and gender, with smoothing 0 mm FWHM, vertex wise threshold \( p = 0.01 \) and cluster wise (CWP) threshold at \( p = 0.001 \). Differences are presented as: maximum value of the cluster (Max), maximum value of the vertex (Vtx max), size in mm\(^2\) with the MNI coordinates per cluster. Results are reported from both, unidirectional (positive) and bidirectional (absolute) contrasts (healthy controls > schizophrenia outpatients, healthy controls < schizophrenia outpatients).

| Spherical kernel LGI | Cortical shape-adaptive kernel LGI |
|----------------------|------------------------------------|
| Cluster Annotation   | Max (mm\(^2\)) | VtxMax | Size (mm\(^2\)) | MNI X | Y | Z |
|----------------------|----------------|--------|-----------------|------|---|---|
| Healthy controls > schizophrenia outpatients |
| Left hemisphere      | lateralorbitofrontal | 1.964 | 20593 | 17.11 | - | 27.9 | 29.4 | 1 |
|                      | medialorbitofrontal | 3.092 | 11982 | 9.34 | 17.68 | -8.8 | 10.6 | -15.5 |
|                      | rostralmiddlefrontal | 2.506 | 145943 | 18.17 | 37.24 | -21.9 | 53.1 | 7.4 |
|                      | superiorfrontal    | 2.133 | 886 | 16.58 | - | -28.9 | 48.1 | 2.4 |
|                      | precentral          | 3.975 | 131494 | 54.47 | 68.16 | -15 | 61.3 | 5.7 |
|                      | superiortemporal    | 2.822 | 34484 | 101.07 | 142.02 | -42 | -6.7 | 44.1 |
|                      | inferiortemporal    | 4.374 | 62831 | 92.28 | 134.2 | 38.2 | 22.2 | 39.1 |
|                      | fusiform            | 2.138 | 57856 | 9.22 | 29.93 | -55.9 | -79.6 | -8.9 |
|                      | precuneus           | 2.46 | 162404 | 42.6 | 138.78 | -26.5 | -64.5 | 6 |
|                      | superioparietal     | 2.32 | 135824 | 15.79 | - | -10.6 | -91.4 | 27.1 |
| Right Hemisphere     | caudalmiddlefrontal | 3.246 | 140460 | 92.28 | 134.2 | 38.2 | 22.2 | 39.1 |
|                      | superiorfrontal    | 2.448 | 123167 | 23.76 | - | 19.2 | 47.6 | 32.1 |
|                      | precentral          | 2.993 | 145515 | 96.84 | 144.37 | 39.9 | -4.9 | 44 |
|                      | paracentral         | 2.225 | 66777 | 17.91 | - | 14.9 | -26 | 45.2 |
|                      | postcentral         | 2.058 | 59788 | 10.01 | - | 23.3 | -31 | 70.4 |
|                      | superiortemporal    | 3.155 | 70390 | 66.24 | 97.71 | 60.2 | -1.4 | 5.8 |
|                      | inferiortemporal    | 2.735 | 94495 | 24.81 | 58.7 | 44.9 | 15.3 | -25.1 |
|                      | lingual             | 2.331 | 35907 | 22.33 | - | 22.1 | -54.2 | -7.3 |
| Healthy controls < schizophrenia outpatients |
| Left hemisphere      | medioorbitofrontal  | 2.319 | 10739 | 11.03 | - | 5.2 | 47.8 | -22.3 |
|                      | laterorbitofrontal  | 2.57 | 17304 | 32.35 | 81.4 | -25.7 | 23.2 | -17.6 |
|                      | parsorbitalis       | 2.076 | 51061 | 40.7 | - | -34.6 | 40.7 | -10.2 |
|                      | caudalmiddlefrontal | 2.57 | 47079 | 66.28 | 11.04 | -27.3 | 21.3 | 36 |
|                      | rostralmiddlefrontal | 2.425 | 106449 | 15.34 | 27.31 | -37.5 | 50.9 | 1 |
|                      | superiortemporal    | 3.301 | 73167 | 11.16 | - | 34.6 | 40.7 | 1 |
|                      | rostralantertiorcingulate | 2.728 | 83315 | 56.04 | 125.4 | -10.5 | 37.6 | 13.1 |
|                      | bank of superior temporal sulcus | 2.097 | 86620 | 50.02 | - | -50.1 | 40.2 | -5 |
|                      | fusiform            | 2.355 | 11016 | 11.32 | - | 34 | 44 | -20.6 |
|                      | precentral          | 5.326 | 63825 | 126.53 | 160.78 | -54.5 | -1.8 | 37 |
|                      | inferiorparietal    | 2.416 | 8339 | 15.74 | 34.45 | -51.2 | -58.4 | 27.8 |
|                      | lingual             | 3.312 | 11179 | 201.41 | 291 | -42.2 | 52.3 | 19.2 |
|                      | precuneus           | 2.501 | 53460 | 43.82 | 92.27 | -15.6 | -79.6 | -11.8 |
|                      | superioparietal     | 2.185 | 101451 | 23.4 | - | -14.7 | -51.3 | 33.2 |
| Right Hemisphere     | laterorbitofrontal  | 2.575 | 134238 | 27.99 | 105.38 | 31.6 | 30.8 | -11.3 |
|                      | rostralmiddlefrontal | 2.47 | 71009 | 63.04 | 16.92 | 37.9 | 42.6 | 10.3 |
|                      | rostralantertiorcingulate | 2.261 | 65649 | 35.28 | - | 23.5 | 51.6 | 20.6 |

(continued on next page)
inspiration for succeeding studies on the LGI abnormalities in schizophrenia. Nevertheless, besides the greater number of altered hypo- and hypergyria brain regions revealed by new LGI method (Cmorph) than standard FreeSurfer code, due to the limited sample size from our project and no formal statistical comparison of pipelines, we were unable to clearly state which method is more accurate at illustrating the actual cortical folding abnormalities in schizophrenia. In particular, the problem of normalization of individual brain size and chosen kernel size with the average of total cortical surface area over population is widely discussed and the code is still under development. Although, presented preliminary results may serve as valuable pioneering outcome as they point out the problems arising in the field of brain morphometry analyses and indicate the future direction of their development. More specifically, considering the debate over whether changes in LGI might become a biomarker of schizophrenia (Matsuda and Ohi, 2018; Sasabayashi et al., 2019; Veronese et al., 2013), precision in the mapping of brain regions should be a special concern (Bartholomeusz et al., 2013; Ronan et al., 2012). Aberrant LGI is a widely suggested potential biomarker for people with a genetic risk (Schmitt et al., 2015) or clinical high risk of developing schizophrenia (Dammé et al., 2019; Dauvermann et al., 2012; Stanfield et al., 2008), or in those who already have schizophrenia (Veronese et al., 2013) and may differentiate subjects’ gyriﬁcation changes from other psychiatric conditions (Cao et al., 2017; McIntosh et al., 2009). Thus, the application of shape-adaptive kernel Cmorph LGI in big data studies, like the ENIGMA consortium (https://enigma.ini.usc.edu), the Human Connectome Project (http://www.humanconnectomoproject.org/), or the SchizConnect project database (http://schizconnect.org/), could bring consistent results and allow the complex patterns of cortical folding in schizophrenia to be revealed. Furthermore, the shape-adaptive kernel approach could be useful in examining the relation between gyriﬁcation and psychopathology in schizophrenia, which has only been examined by a few studies so far that focused on the deficit and negative symptoms (Mihailev et al., 2017; Storvestre et al., 2019; Takayanagi et al., 2019). Finally, Cmorph seems to be a method which can be widely applied in morphometric studies concerning not only schizophrenia but also other clinical conditions related to altered brain structure.

Table 2 (continued)

| Cluster Annotation | Max | VtxMax | Size (mm²) positive contrast | Size (mm²) absolute contrast | MNI | X | Y | Z |
|--------------------|-----|--------|-----------------------------|-----------------------------|-----|---|---|---|
| Caudalmiddlefrontal | 2.099 | 124279 | 21.4 | - | 37.8 | 1.9 | 31.9 |
| Middletemporal     | 2.025 | 14059  | 18.18 | - | 24.8 | 5.3 | 46.8 |
| Precentral          | 2.074 | 1799   | 21.76 | - | 59.6 | -21.9 | -19.9 |
| Fusiform            | 2.464 | 82953  | 12.34 | - | 57.9 | 0.6 | 32.6 |
| Supramarginal       | 2.416 | 145233 | 23.38 | 56.45 | 36.1 | -20.1 | 52.8 |
| Lateraloccipital    | 3.712 | 101436 | 128.01 | 164.52 | 48.2 | -39.4 | 40.5 |

**Figure 1.** Comparison of between-group differences of the FreeSurfer gyriﬁcation index (top panels) and cortex-shape-adaptive kernel Cmorph method (bottom panels) depicted as non-inﬂated (A panel, on left) or inﬂated (B panel, on right) pial surfaces. Statistical analysis from mri_glmfit utilized a whole-brain vertex-wise shape analysis with smoothing 0 mm FWHM, vertex-wise threshold p = 0.01, and cluster-wise threshold p = 0.001. Images of cluster localizations (anterior, posterior, inferior, superior, sagittal lateral/middle plane) were obtained with FreeSurfer (http://surfer.nmr.mgh.harvard.edu/) and visualized with FreeView v.2.0. The colour scale (red: healthy controls > schizophrenia outpatients; blue: healthy controls < schizophrenia outpatients) is set the same for both methods. Coloured areas show the vertex-wise p-values of the difference between patients and controls as log10(p), so that dark colour (value = 1.3) means p = 0.05. The vertex-wise results are masked by the positions of signiﬁcant clusters.
5. Conclusions
In conclusion, the presented evidence from disrupted cortical folding in schizophrenia outpatients indicates that the cortex shape-adaptive kernel (Cmorph LGI) approach may be a valuable tool for further investigation, especially if a big sample size is involved, as well as in other clinical populations. Finally, presented preliminary results from novel LGI method (Cmorph) suggest that potential mechanism of altered cortical folding in schizophrenia is not related only to the hypogorgia, but rather with such changes in brain neurodevelopment, maturation and ageing with final settlement of alternate brain regions morphology (hypo- and hypergorgia) and its abnormal connectivity. However, the future investigation should be provided to resolve this question and to better determine biological origins of schizophrenic psychoses.

Declarations

Author contribution statement
Olga Plonka: Performed the experiments; Wrote the paper. 

Alicja Krzesińska: Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data.

Przemysław Adamczyk: Conceived and designed the experiments; Performed the experiments; Analyzed and interpreted the data.

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Competing interest statement
The authors declare no conflict of interest.

Additional information
No additional information is available for this paper.

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