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Mariam Al Harrach, François Rousseau, Samuel Groeschel, Xiaoyu Wang, Lucie Hertz-pannier, et al.. Alterations in Cortical Morphology after Neonatal Stroke: Compensation in the Contralesional Hemisphere?. *Developmental Neurobiology*, Wiley, 2019, 79 (4), pp.303-316. 10.1002/dneu.22679 . hal-02285917

HAL Id: hal-02285917

<https://hal-imt-atlantique.archives-ouvertes.fr/hal-02285917>

Submitted on 16 Sep 2019

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Alterations in Cortical Morphology after Neonatal Stroke:

Contralesional hemisphere Compensation?

(Neonatal stroke outcome on cortical morphology)

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Abstract

Although neonatal arterial ischemic Stroke is now well-studied, its complex consequences on long-term cortical brain development has not yet been solved. In order to understand the brain development after focal early brain lesion, brain morphometry needs to be evaluated using structural parameters. In this work, our aim was to study and analyse the changes in the morphometry of ipsilesional and contralesional hemispheres of seven-year-old children following neonatal stroke. Therefore, we used surface-based morphometry in order to examine the cortical thickness, surface area, cortical volume and local gyrification index in two groups of children that suffered from neonatal stroke in the left (n=19) and right hemispheres (n=15) and a group of healthy controls (n=30). Reduced cortical thickness, surface area and cortical volumes were observed in the ipsilesional hemispheres for both groups in comparison with controls. For the left lesioned group, an increase in the gyrification of the contralesional hemisphere underlined by an increase in surface area and cortical volume was observed primarily in the occipital region. As for the right lesioned group, higher gyrification was detected in two different clusters also in the contralesional hemisphere. However, these alterations in the gyrifications were not accompanied by a significant change in the cortical thickness, surface area or cortical volume in the occipital lobe. This is the first time that increase of the structural parameters is detected in the “healthy” hemisphere after unilateral neonatal stroke suggesting a compensatory phenomenon. Moreover, findings presented in this work suggest that lesion lateralization might have an influence on brain development and maturation.

Keywords: neonatal arterial ischemic stroke, cortical thickness, local gyrification index, surface area, FreeSurfer.

Abbreviations: NAIS = Neonatal Arterial Ischemic Stroke, LGI=Local Gyrification Index, HC= Healthy Controls, VBM = Voxel-Based Morphometry, VLSM = Voxel-Based Lesion-Symptom Mapping, DTI = Diffusion Tensor Imaging, MCA= Middle Cerebral Artery, LLP= Left lesioned Patients, LH= Left Hemisphere, RLP= Right Lesioned Children, RH= Right Hemisphere, TIV= Total Intracranial Volume

Introduction

Perinatal ischemic stroke, as defined as a “heterogeneous conditions with a focal cerebral arterial or venous occlusion, occurring between 20 weeks of foetal life through 28th postnatal day” (Raju *et al.*, 2007), appears more and more as an umbrella term which encompasses a variety of clinical entities. Neonatal Arterial Ischemic Stroke (NAIS) represents a unique model to study the impact of an early stroke on brain development, as the lesion and its timing are well circumscribed (Kirton, 2013; Grunt *et al.* 2015). Affected children will sustain cerebrovascular arterial insults occurring between birth and the 28th day of life (Stephan-Otto *et al.* 2017). According to Diffusion-weighted imaging, critical focal brain infarction is mostly located in a large artery; more precisely the left middle cerebral artery (Raju *et al.* 2007; Kirton & deVeber, 2013).

NAIS birth-prevalence in term or near-term new-borns varies from 6 to 17/100,000 live births (Darmency-Stamboul, Cordier, & Chabrier, 2017). Studies looking at NAIS show that approximately 60% of children exhibit some mild neurodevelopment disabilities at school-age (Kirton & deVeber, 2013). However, it has risen as a principal cause of perinatal brain injury, hemiparetic cerebral palsy, and lasting disability (Kirton *et al.* 2011). Furthermore, NAIS may trigger structural changes in both grey and white matter regions internal and remote to the infarct site (Dinomais *et al.*, 2015).

Neuroimaging studies on brain morphometry after early brain lesion provide interesting insights into the capacity, but also the limits, of brain (re)organization in the development course (Staudt, 2010a, 2010b). Research shows the potential of human brain to compensate for pre and perinatal focal brain lesions (Staudt, 2010a). This compensation is established through the contralesional hemisphere, where it will conduct motor functions in the paretic extremities (Staudt, 2010a, 2010b; Dinomais *et al.* 2015; Zewdie *et al.*, 2017). Moreover, in addition to motor system, different types of functional (re)organization were described in the literature including somatosensory and language systems (Staudt, 2010a, 2010b). On these accounts, studying the brain morphology after several years could help to better apprehend the long-term structural effect of NAIS.

Different approaches are adopted in the literature in order to study and explore the brain structural development. Most of these studies used classical approaches (Voxel-Based Morphometry (VBM), Voxel-Based Lesion-Symptom Mapping (VLSM) or Diffusion Tensor Imaging (DTI)) which do not allow an in-depth examination of the cortex (Van Der Aa *et al.*, 2013; Dinomais *et al.*, 2015). This research field has undergone recent methodological

advances where different approaches have been extensively used as tools to measure specific morphometric variables of the cortex including cortical thickness, surface area, cortical volume, complexity and gyrification (Kelly *et al.*, 2013; Gerrits *et al.*, 2016). These metrics have proven to be independently powerful as tools to evaluate differences or abnormalities in the brain structure in certain brain disorder cases like Autism (Libero *et al.*, 2014), Parkinson's disease (Gerrits *et al.*, 2016), schizophrenia (Schultz *et al.*, 2013) and bipolar disorder (Rimol *et al.*, 2012) or even in psychologically affected children (maltreatment, abuse and neglect) (Kelly *et al.*, 2013). However, to date, no information on the cortical shaping are available at long term after NAIS.

The aim of this study was to explore the impact of unilateral NAIS on the development of the cortex at age seven, using new approaches. This was done using cortical thickness, surface area, cortical volume and local gyrification for patients with lesions in the left and right hemisphere separately. The lesion lateralization was considered by separating the patients into two groups based on the side of the unilateral lesion. This was motivated by the brain asymmetry studies as well as the importance of laterality in different brain functions such as language (Staudt, 2010a). Accordingly, we recruited three groups of children: a control group of healthy children, a Left Lesioned Patients (LLP) with unilateral NAIS in the Left Hemisphere (LH) and a Right lesioned Patients (RLP) with unilateral NAIS at the Right Hemisphere (RH). Finally, to investigate the effects of NAIS on the previously mentioned parameters, we were also keen to explore the impact of the lesion lateralization on these brain biomarkers especially for the contralesional hemisphere.

Materials and methods

Participants

Patients belonged to the previously described AVCnn cohort (Accident Vasculaire Cerebral du nouveau-ne, that is, neonatal stroke—PHRC regional n80308052 and PHRC interregional n81008026 -eudract number 2010-A00329-30) (Dinomais *et al.*, 2015). In short, 100 term newborns with an arterial infarct, confirmed by brain imaging (CT and/or MRI), who were symptomatic in the neonatal period (thus matching the 2007 definition of NAIS (Raju *et al.*, 2007)) were consecutively enrolled between November 2003 and October 2006 from 39 neonatology and neuropsychiatric units distributed throughout mainland France. At the age of 7 years, a clinical follow-up visit was organized, and an MRI investigation was proposed to the 80 children who took part in the clinical evaluation. Of the 52 children who participated in the

MRI study, 38 had unilateral MRI lesions in the MCA territory. They constituted the patient population of this study (eudract number 2010-A00976-33). Patients with lesion in the left (LLP) and right hemisphere (RLP) were processed separately without masking out the brain lesions or projecting the lesioned hemisphere onto the left one as previously done (Dinomais *et al.*, 2015). By keeping the original orientation of the brain without flipping the image in the left-right direction we eliminate the ambiguity and inaccuracy that arises from the asymmetry of the brain (Kong *et al.*, 2018). However, after further examination we only kept 34 patients divided between LLP and RLP. Four patients were eliminated due to poor segmentation by reason of the position of the corresponding brain lesions (for more details please refer to section **MRI Acquisition and processing**). Along with the LLP and RLP patients we recruited 30 Healthy Controls (HC). These controls were chosen to be of similar age and gender distribution as the patients. General profile and characteristics of the participants are presented in Table 1.

Informed written consent respecting the declaration of Helsinki were obtained from participants/parents as well as approval from the ethical committee of the university hospital of Angers, France. Handedness was assessed according to the Edinburgh inventory (Oldfield, 1971).

MRI Acquisition and Processing

Images were acquired on a 3.0 Tesla scanner (MAGNETOM Trio Tim system, Siemens, Erlangen, Germany, 12 channel head coil) at Neurospin, CEA-Saclay, France. Imaging sequences included a high-resolution 3D T1- weighted volume using a magnetization-prepared rapid acquisition gradient-echo sequence [176 slices, repetition time (TR) 2300 msec, echo time (TE) 4.18 msec, field of view (FOV) 256 mm, flip angle= 9°, voxel size $1 \times 1 \times 1 \text{ mm}^3$], a 3D T2 FLAIR sequence [160 slices, TR 5,000 msec, TE 395 msec, FOV 230 mm, voxel size $0.9 \times 0.9 \times 1 \text{ mm}^3$]. MRI images were all inspected for abnormalities and inconsistencies by senior paediatric neuroradiologists and physical and rehabilitation medicine practitioners. For the NAIS patients, lesions were manually identified and outlined by two paediatric physical and rehabilitation medicine practitioners (MD and SG, co-authors). This was done on a slice by slice basis on the individual 3D T1 images in order to create a binary lesion mask using the MRIcron software (<http://www.mccauslandcenter.sc.edu/mricron>) (Rorden, Karnath, & Bonilha, 2007). For an optimum result, the co-registered 3D FLAIR images were used as a visual aid for accurate lesion delineation where the hyper-intense areas on FLAIR were defined as part of the lesion (Dinomais *et al.*, 2015). Afterwards, obtained lesion masks were double

checked by each practitioner and were spatially normalized. For in depth details about lesion delineation and localization please refer to (Dinomais *et al.*, 2015).

All T1 images were processed using the FreeSurfer suite, version 6.0.0 (<https://surfer.nmr.mgh.harvard.edu/>), on a single DELL workstation running ubuntu 16.04 LTS (Intel R Core TM i7-7820HQ CPU @ 2.9GHz × 8). The FreeSurfer processing pipeline carries out an automated cortical surface reconstruction process that includes various steps (for detailed description of the different surface-based pipeline please refer to (Dale, Fischl, & Sereno, 1999; Fischl *et al.*, 1999, 2000, 2001, 2002, 2004). Concisely, these steps contain motion correction, non-uniform intensity normalization, automated Talairach transformation, skull stripping, subcortical and cortical matter segmentation, tessellation and depiction of grey/white/pial boundaries (Dale, Fischl, & Sereno, 1999; Fischl *et al.*, 1999, 2001). In order to improve the segmentation and therefore the computing of the different parameters, we used the T2-weighted images as auxiliary to the T1 images. This additional contrast in the T2 images allows more accurate pial surfaces definitions especially in the presence of lesions. After the creation of the final surfaces, deformation operations are applied in order to inflate and register the surface to a spherical atlas (Fischl *et al.*, 2002). Then, parcellation of the cerebral cortex is performed and different statistics are computed (Fischl *et al.*, 2004). In this work, we used the Destrieux atlas (Fischl *et al.*, 2004) for the parcellation of the cortex into gyral regions.

Images of each participant (patients and controls) were inspected and visually checked after the segmentation process by a rehabilitation medicine practitioner (MD, last author). Four patients of the 38 were excluded from the study due to irrevocable errors in the segmentation results around and near the lesion despite attempts to manually edit and improve the results. Total Intracranial Volume (TIV) for all participants was estimated by FreeSurfer as part of the statistics.

Cortical thickness, surface area and cortical volume

Cortical thickness is computed at each vertex as the shortest distance between the pial surface and the white/grey boundary (Fischl & Dale, 2000). The obtained cortical thickness values are not restricted to the voxel resolution of the initial data. Accordingly, these maps have a sub-millimetre accuracy of around 150000 vertices for each surface mesh of the cerebral hemispheres which means a comparison at a sub-millimetre scale between the groups. The algorithm used by FreeSurfer for the computing of cortical thickness was validated and proven

to be reliable by various studies (Kuperberg *et al.*, 2003; Kelly *et al.*, 2013; Besteher, Gaser, Spalthoff, & Nenadic, 2017).

Surface area is estimated at the pial level. It is computed for each vertex of the grey matter by averaging the areas of the tessellation triangles in contact with that vertex. The cortical volume calculated by FreeSurfer is obtained at each vertex, by surface-based calculations in order to obtain better accuracy. All the measures were estimated in the native space.

Local Gyrification Index

In this work, cortical gyrification was computed using a surface-based approach that is available as auxiliary measure in FreeSurfer (Schaer *et al.*, 2008). This quantification of cortical folding is called Local Gyrification Index (LGI). It is a 3-D adaptation of the 2-D Gyrification index (GI) proposed by Zilles, Armstrong, Schleicher, & Kretschmann (1988). They computed the GI, at the coronal sections level, as the complex cortical surface proportion of the total brain perimeter (Schaer *et al.*, 2008). The LGI developed by Schaer *et al.* (2008) overcomes the limitation of the 2-D GI by taking into account the 3-D character of the cortical exterior compared to the GI that can suffer from inaccuracies due to buried sulci and slice orientation, not to mention the problems of manual tracing used in the 2-D approach. The LGI has been recently used in different studies in order to evaluate the gyrification abnormalities (Kelly *et al.*, 2013), folding deficit (Hyatt, Haney-Caron, & Stevens, 2012) or to quantify regional differences in gyrification (Palaniyappan, Mallikarjun, Joseph, White, & Liddle, 2011). This LGI is a new 3-D parameter that aims to compute and compare local cortical gyrification at thousands of points throughout the entire hemisphere using 3-D cortical reconstruction (Schaer *et al.*, 2008). It is the vertex wise ratio of the folded pial surface area, computed over a circular region of interest with 25 mm radius, to an outer surface. This outer surface area is defined as an outer hull layer that closely wraps around the pial surface and thus envelops the outer cortex perimeter. The obtained LGI maps describe the portion of cortex concealed in the sulcal folds of the brain at each pial surface vetted and therefore it is a direct measure of the cortical folding. For in depth description of the LGI computation algorithm please refer to (Schaer *et al.*, 2008).

Lesion Masks

Total lesion maps for LLP and RLP were computed as the sum of the normalized binary individual masks for each group separately (for detail of lesion masking, see (Dinomais *et al.*, 2015)). Thus, we obtained a LH and RH mask map for the LLP and RLP groups respectively

(see Figure 1). These maps were then transformed into surface maps using a FreeSurfer function that assigns values from the volume to each surface vertex. Afterwards, we applied a threshold of $n > 4$ and $n > 3$ for LLP and RLP groups respectively (Lo, Gitelman, Levy, Hulvershorn, & Parrish, 2010; Geva, S., Baron, Jones, Price, & Warburton, 2012) which is equivalent to $n > 20\%$ of the total participants. Finally, these maps were binarized in order to obtain the final masks for LLP and RLP. The block diagram of the lesion masks computing is detailed in Figure 1. The lesion masks represent the portion of the brain where lesions occurred for the previously described patients. Appropriately, it was important to compute these masks in order to study the effect of NAIS outside these regions by excluding them from the statistical tests.

Statistical Analysis

For all four parameters (cortical thickness, surface area, cortical volume and LGI), statistical analysis was carried using the general linear modelling built-in tool in FreeSurfer. Between group surface differences were run for each hemisphere separately using two-sample t-test models. First, all measurement values were resampled and mapped to the *fsaverage* subject provided by FreeSurfer and smoothed by an 8mm full-width-at-half-maximal Gaussian kernel. Then, vertex-wise comparisons between HC and each of the patient groups (LLP and RLP) are performed for each parameter and hemisphere at a time. These analyses were controlled for sex and handedness considering the account these parameters can have on the obtained clusters (Sowell *et al.*, 2006; Kong *et al.*, 2018). No significant group differences were detected between controls (HC) and patients (LLP and RLP) in age ($p=0.543$) or TIV ($p=0.127$) (Table 1). As a result, they were not included as covariates. We also have to mention that the final lesion masks for LLP and RLP groups (please refer to section **MRI Acquisition and Processing**) were added as parameter to the statistical analysis in order to exclude the lesioned area from the test. This ensures that the obtained clusters are external to the corresponding lesion mask in each case. Finally, between-group differences were corrected for multiple comparisons using a cluster wise correction using two-tailed Monte Carlo z-field simulation with $p < 0.05$. This approach of correction offered by FreeSurfer is similar to Gaussian Random Fields (GRF) though it uses a simulation instead of analytic equations for the corrected voxel-wise and cluster wise p-values estimation (Hagler Jr, Saygin, & Sereno, 2006).

Results

Figures 2 and 3 show sets of significant clusters for cortical thickness, surface area, cortical volume and LGI representing the following comparisons: HC versus LLP (Fig. 2) and HC versus RLP (Fig. 3). These results of vertex-by-vertex comparisons were all corrected for sex and handedness. Multiple comparisons were Monte Carlo corrected. The lesion regions delineated by the total LLP and RLP masks, previously described in detail (section **Lesion Masks**), were excluded from the statistical analysis in order to study the lesion impact on the brain accurately. These masks are displayed in Figures 2 and 3 (green colour). The details of the significant clusters are presented in Tables 2 and 3 for the two comparisons previously mentioned.

HC Versus LLP

The cortical thickness analysis for the HC versus LLP groups exhibits a significantly thinner cortex for the LLP subjects (Fig. 2A). These significantly inferior values of cortical thickness were observed only for the lesioned hemisphere (LH). The contralesional hemisphere (RH) in this case did not show any significant thinning compared to the controls. By observing the obtained clusters in the LH in Fig. 2A, multiple small clusters were localized around the lesion area (close to the lesion mask). Main clusters that survived whole brain correction were located, based on the Destrieux parcellation atlas (Fischl *et al.*, 2004), in the postcentral sulcus, superior segment of the circular sulcus of the insula and the inferior part of the precentral sulcus (Table 2).

For the surface area analysis, we obtained four major clusters (see Fig. 2B, Table 2) in the LH part of the brain. These clusters indicate a significantly reduced average surface area at the transverse temporal sulcus, the inferior segment of the circular sulcus of the insula, the precentral gyrus and the intraparietal sulcus and transverse parietal sulci for the LLP subjects in comparison with the healthy subjects (HC). These brain regions with reduced surface area values are encircling the total lesion mask. For the RH part of the brain, the surface area investigation determined one cluster where the surface area values of the LLP subjects were significantly higher than the HC group ones. These higher values of surface area of the contralesional hemisphere of LLP subjects were located, according to the Destrieux atlas, at the occipital pole. No other significant clusters survived whole brain cluster correction in the RH. Similar to the surface area analysis, cortical volume evaluation between LLP and HC groups revealed inferior cortical volume of the LLP subjects as to the HC group in the lesioned hemisphere (LH). This smaller cortical volume is located in nine different clusters (Fig. 2C,

Table 2). Main clusters that survived Monte Carlo null-z simulation correction are located at the inferior segment of the circular sulcus of the insula, the transverse temporal sulcus, the angular gyrus and the inferior part of the precentral sulcus. Smaller but significant clusters were found at the lingual and temporal gyrus areas of the brain. For the RH section of the brain, similarly to the surface area analysis, we identified one cluster that indicated a significantly higher cortical volume for the LLP subjects compared to the healthy subjects (Fig. 2C, Table 2). Based on the Destrieux atlas annotation, this cluster peak coordinate fell within the superior occipital gyrus.

Between group comparison for LLP and HC groups of LGI revealed two significant clusters in the LH with reduced gyrification in the LLP group compared to the HC groups and three significant clusters in the RH with more important gyrification in the LH for the same patients versus healthy subjects (Fig. 2D, Table 2). For the lesioned hemisphere (LH), automated annotation of the group structural data identified the first and biggest cluster as part of the subcentral gyrus and sulci regions of the brain. The second cluster that survived the Monte Carlo correction ($P < 0.05$) was located in the Anterior occipital sulcus and preoccipital notch. Regarding the RH region of the brain, the clusters with more important gyrification lied in the superior occipital gyrus, superior frontal gyrus and superior part of the precentral sulcus (Fig. 2D, Table 2). Table 2 summarizes the obtained cluster statistics for both LH and RH in the case of LLP versus HC.

HC Versus RLP

For the cortical thickness analysis, there were only small insubstantial clusters remaining in the RH after Monte Carlo null-z simulation correction and lesion masking by employing the RLP lesion mask (Table3). One of these small clusters is located in the postcentral gyrus and was reduced in size after adding the RLP lesion mask. In the contralesional hemisphere (LH), none of the clusters survived the whole brain cluster correction.

Surface area examination revealed inferior average surface area values for the RLP subjects with respect to the HC group in the RH of the brain (Fig. 3B and Table 3). Significantly lower surface area values were mainly observed in five clusters positioned at the posterior ramus of the lateral sulcus, angular gyrus, middle frontal sulcus, middle occipital gyrus and middle temporal gyrus (Table 3). Smaller but significant clusters were found in the suborbital sulcus, supramarginal gyrus and postcentral gyrus. For the healthy hemisphere (LH), no clusters were found after multiple comparison correction.

After analysis of the cortical volume, we observed complementary results to the surface area study. We found no significant difference after Monte Carlo correction between RLP and HC groups for the LH. However, for the lesioned hemisphere (RH) we identified seven more or less small clusters indicating inferior cortical volume for the RLP participants compared to the HC subjects (Fig. 3C, Table 3). The biggest identified cluster was located, according to the Destrieux atlas annotation, in the posterior ramus of the lateral sulcus. The other clusters were smaller in size and they were sitting in the angular gyrus, postcentral gyrus, anterior and middle occipital sulcus, inferior temporal gyrus and subcentral gyrus and sulci.

In the case of the LGI for the RLP versus HC groups, no significant differences in gyrification were detected for the RH. Nonetheless, for the healthy hemisphere (LH), group comparisons revealed higher gyrification for the RLP participants in comparison with healthy subjects. Two clusters survived the Monte Carlo null-z simulation correction ($p < 0.05$). These clusters were located in the Middle occipital gyrus and Precuneus (see Table 3).

Discussion

In this work, a surface-based analysis approach was used in order to study the structural brain abnormalities in seven-year-old children that sustained a NAIS. Taking into account the asymmetry of the brain (Kong *et al.*, 2018), structural changes were compared for the LLP and RLP groups separately. This allowed us to study the influence of the lesion lateralization on the obtained structural changes of the brain.

For the LLP group, we obtained lower cortical thickness, surface area, volume and LGI in the LH near the infarct site as presented (Fig. 2). Affected areas of the LH were found in the frontal, temporal and parietal lobes. For the surface area and cortical volume, principal clusters were found in the same regions of the brain at the inferior segment of the circular sulcus of the insula and transverse temporal sulcus (Table 2). Since cortical volume is considered as the product of cortical thickness and surface area and is affected by both measures in many studies (Winkler *et al.*, 2010; Gerrits *et al.*, 2016; Kong *et al.* 2018), it was expected that cortical volume and surface area results will be related. However, cortical thickness was proven to be independent from the surface area and volume. Therefore, the obtained clusters seem coherent especially since the clusters representing lower cortical thickness seem to be placed near and even at the edge of those of the surface area and cortical volume.

The degeneration in the cortical matter was anticipated since they fall into the MCA vascular territory (Kandel *et al.*, 2000). This indicates that damage do not only affect the lesion territory

but also sites distant from the injury based on the axotomy phenomenon that causes chain reactions including Wallerian degeneration and destruction of glial cells (Kandel *et al.*, 2000). Moreover, the clusters obtained in the LH of the LLP group seem to be associated with the language area. This can point out a correlation with language and speech outcomes. Especially since the language area is located in the LH.

One of the main findings in this study was obtained when comparing the contralesional hemisphere (RH) of LLP subjects to the RH of the HC group. We observed greater surface area and as follows cortical volume of the LLP group in the occipital pole and superior occipital gyrus respectively (Fig. 2). Although cortical thickness results did not reveal any significant clusters after Monte Carlo correction, surface area and cortical volume variations are normally driven by cortical thickness alteration (Rimol *et al.*, 2012). On that account, we found that we did have a significantly superior cortical thickness in the middle occipital gyrus (uncorrected $p < 0.05$) without surviving correction for multiple comparison (supplementary Fig.1). This provides evidence that in the case of NAIS with lesion in the LH, the RH part of the brain does not stay intact. This alteration of the contralesional hemisphere (RH) in the case of the LLP group is unexpected and it can be attributed to a number of reasons. The higher cortical surface, volume and LGI can be explained as a form of adjustment or compensation to the lower grey matter in the lesioned hemisphere. It can also be interpreted as a mechanical effect of the lesion on the growth of the opposite hemisphere. More precisely, it can mean that as a response to the lesion, the contralesional hemisphere can reflect an improved maturation in certain clusters (Remer *et al.*, 2017).

Gyrification analysis for the LLP group was consistent with reduced surface area and cortical volume where reduced cortical folding. The overlapping of the obtained LGI clusters with those of cortical volume and surface area is also found in the RH where we discovered more gyrification for LLP subjects relative to HC in the superior occipital gyrus. Two other clusters fell in the superior frontal gyrus and superior part of the precentral sulcus respectively. In the case of RLP group in comparison with the healthy subjects, we observed reduced surface area and cortical volume in the posterior ramus of the lateral sulcus, angular gyrus, postcentral gyrus and middle occipital gyrus of the lesioned hemisphere as well as other regions (Table 3). For the cortical thickness we found two very small clusters, the first representing less cortical thickness in the postcentral gyrus and the second representing extra cortical thickness at the pericallosal sulcus. The second cluster was assumed to be irrelevant due to its size. These clusters differ from the ones obtained in the LLP versus HC case for the lesioned hemisphere. In this case, clusters were found scattered all over the brain lobes and were not concentrated

near the lesion site like in the case of LLP and HC comparison clusters. Moreover, we found reduced surface area and cortical volume in the middle occipital gyrus for the RLP subjects in contrast to the increased grey matter observed for the LLP group in the RH. The thinning in the cortical matter at the presented regions of the brain can equally be explained by the Wallerian degeneration introduced in the LLP cases. In the case of the gyrification assessment in the RH of RLP, no significant differences were observed after multiple comparison correction. This may be explained by the fact that the individual normalized lesion maps in the RLP case were less overlapping than the LLP ones (Fig. 1).

Considering the group comparison between contralesional hemisphere of RLP group (RH) and the corresponding hemisphere of the HC group, no clusters have survived the Monte Carlo correction for the cortical thickness, surface area and cortical volume. Nonetheless, for the LGI two clusters, indicating higher gyrification for the RLP group, were obtained in the middle occipital gyrus and precuneus. Accordingly, even though we did not obtain the same transformation pattern as the one observed in the LLP case for the cortical thickness, surface area and cortical volume, local gyrification has detected a compensation type abnormality in different regions as the ones found for the LLP groups.

The interesting aspect found in the present findings is that the higher grey matter observed by surface area and cortical volume growth of occipital pole and gyrus regions in the healthy hemisphere is only observed for the LLP groups. This can be explained by different theories. One suggestion is that lesion lateralization have a decisive impact on the brain adaptation. This structural alteration is only observed in the contralesional hemisphere of the brain. Other theories may link the results variety between RLP group in comparison with the LLP group to the different lesion positions in the brain for the two groups with respect to the MCA territory (please refer to Fig. 1). Nevertheless, this last explanation does not justify the absence of variation for the surface area, cortical thickness and volume in the healthy hemisphere for the RLP case. Another analysis can consider the difference to be related to the moderate number of subjects per patient group ($n=19$ for LLP and $n=15$ for RLP). However, $n=15$ is considered to be the minimal number of sample subjects in order to conduct reliable statistical analysis of this kind (Kong *et al.*, 2018). Furthermore, in the case of uncorrected comparisons ($p < 0.001$) for both cases (LLP versus HC and RLP versus HC) we found similar results as the ones presented in this manuscript. This means that even in the case of uncorrected comparison, we did not observe higher cortical thickness, volume or surface area in the healthy hemisphere (LH) for the RLP group. Based on all of the above, the first theory suggesting that lesion lateralization plays a major role in the morphological abnormalities observed in children brains after NAIS,

seems to be the most reasonable. This theory can be further explained by considering the brain asymmetry and particularly the right frontal and left occipital petalia impressions. These impressions cause swelling of these regions in comparison with the opposite hemisphere (Toga and Thompson, 2003).

Another angle to consider in this study is the LGI metric. It seems that this particular parameter conveys more prominent results than the other markers especially for the contralesional hemisphere variation study in both LLP and RLP cases. The gyrification index is linked to the cerebral function and form and can allow to better understand the structural configuration, complexity and variability of the brain (Schaer *et al.*, 2008; White *et al.*, 2010). Thus, the question to ask is whether the LGI results presented in this study for the contralesional hemisphere indicate that the gyrification index is more sensitive to the cortical morphology impacted by a mechanical growth effect to counterbalance the deterioration of the opposite hemisphere.

Further work is required in order to find the link between the obtained findings and the clinical and neurological characteristics of patients from both groups (LLP and RLP). This can be done by studying the correlation between clinical scores and the four parameters presented in this study (cortical thickness, surface area, cortical volume and LGI). In addition, neurological examination of patients related to visual processing can help support and approve the results presented in this paper especially for the suggested adaptation theory in the case of LLP subjects. This can also help us explain and link the more important cortical matter in the occipital regions of the RH for the LLP to advanced visual tasks and compare these findings with the corresponding ones for the RLP group.

In this study we presented an original morphometric surface-based analysis of the human brain for seven-year-old children that sustained a NAIS. There is, however, some limitations in our approach that we need to point out. One of the main difficulties we had is related to the segmentation process in the case of the lesioned brains (LLP and RLP) which was more challenging for the subjects with larger lesions. In these cases, the segmentation was difficult around and near the lesion site even by manual correction. However, this did not affect the segmentation for the healthy hemispheres. Therefore, the findings obtained in the case of the healthy hemisphere studies (RH for LLP and LH for RLP) were not affected by this. We also have to mention that the lesion delineation was performed using an expert consensus approach (Dinomais *et al.*, 2015), which can introduce a certain subjectivity to the results.

Conclusion

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Funding

The research was supported by the University hospital of Angers (eudract number 2010-A00976-33), the University hospital of Saint-étienne (eudract number 2010-A00329-30), and the Fondation de l'Avenir (ET0-571). M Al Harrach, A Bohi, and X Wang were supported by grants obtained from the Fondation pour la Recherche Médicale (FRM-DIC20161236453). Sponsors of the study had no role in the study design data collection, data analysis, data interpretation, writing of the report, or decision to submit for publication.

Conflict of Interest

None declared.

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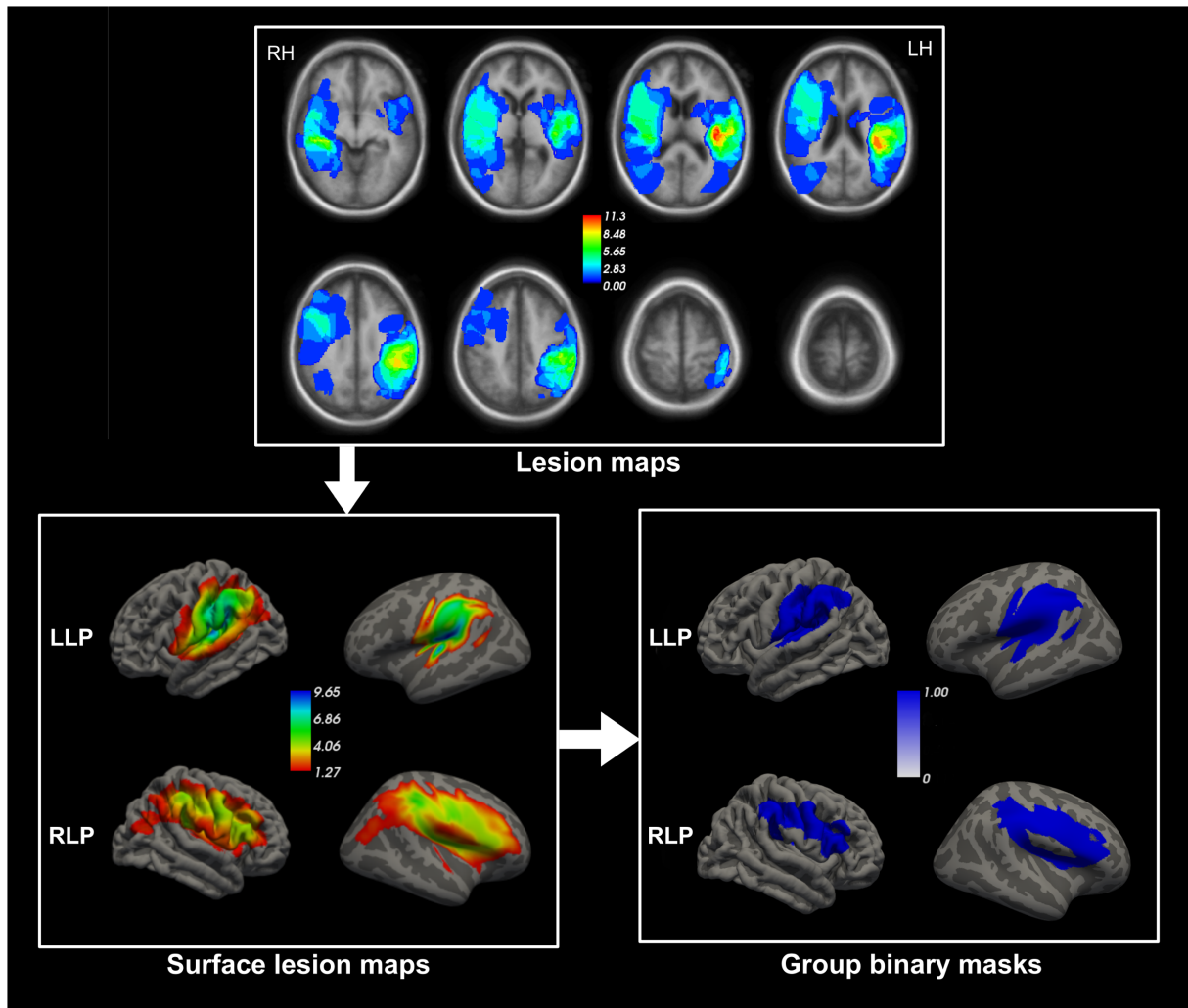


Figure 1: The pipeline used to obtain the surface lesion masks for LLP and RLP groups. The group lesion masks for LLP (LH) and RLP (RH) were obtained by summing the normalized individual lesion masks. The second step consisted on transforming these maps from volumetric to surface space. Final step was the thresholding and binarization of the surface maps in order to obtain the final binary masks for the LLP and RLP groups.

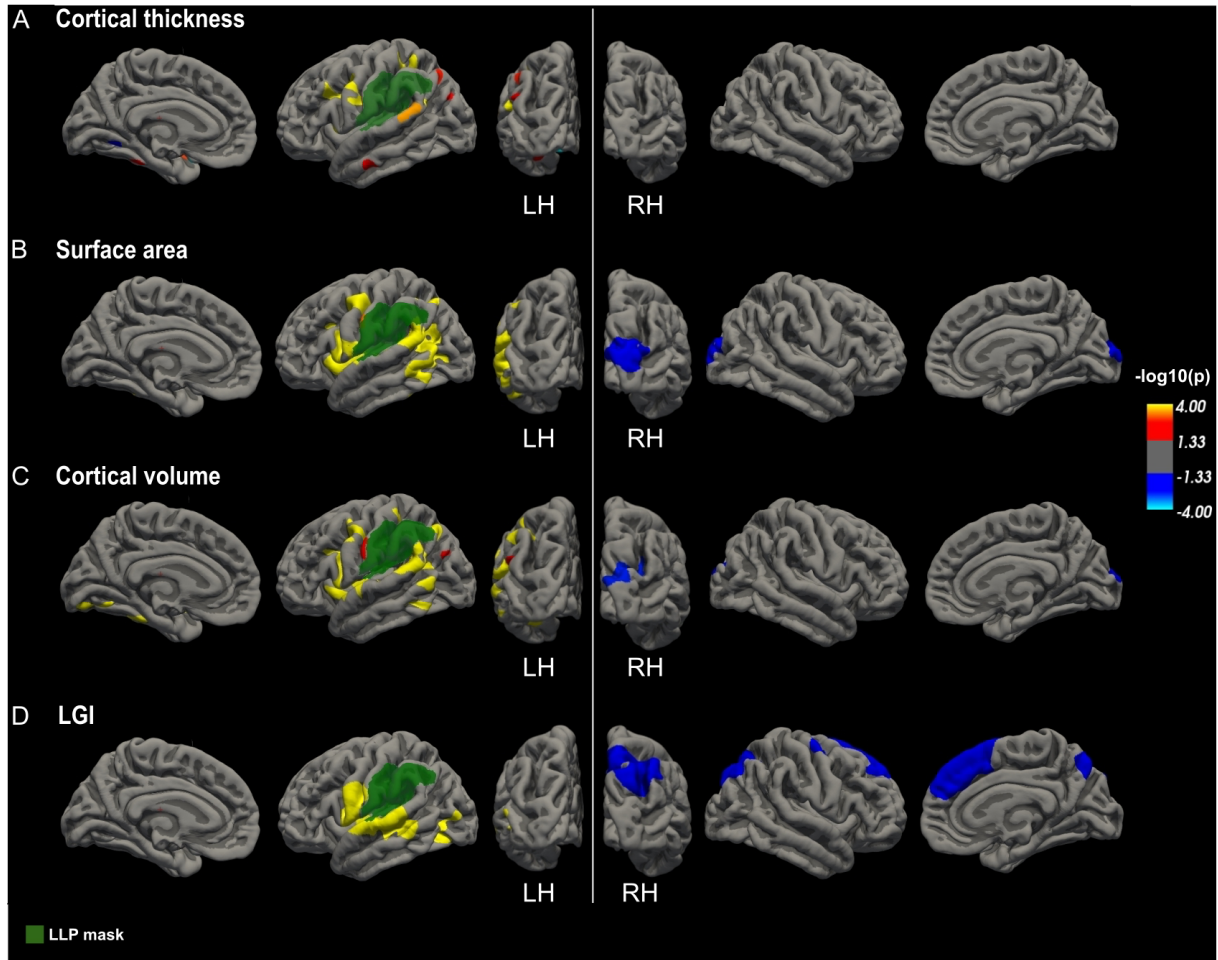


Figure 2: Between-group differences between HC and LLP ($p < 0.05$ cluster level corrected) in cortical thickness (A), surface area (B), cortical **volume (C) and LGI (D) projected onto the pial surface of the fsaverage in the medial, lateral and posterior view for the LH and the RH. Differences were computed separately for LH and RH. Clusters with red and yellow colours indicate brain regions where LLP have significantly lower cortical indexes values than the **HC** and clusters with blue colours indicate regions with increased values for LLP in comparison with HC. These clusters are defined in detail in Table 2. The final LLP mask is presented in green.**

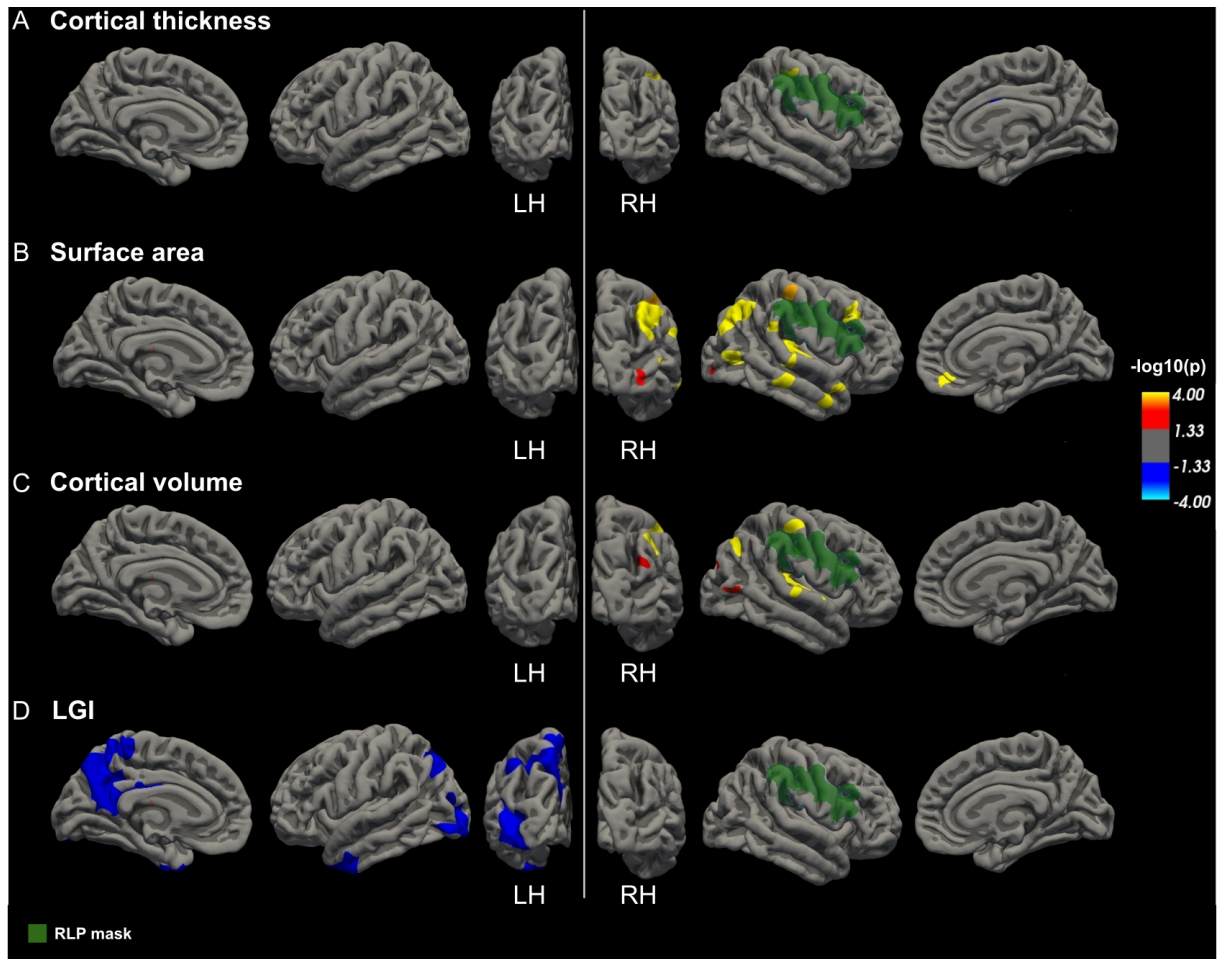
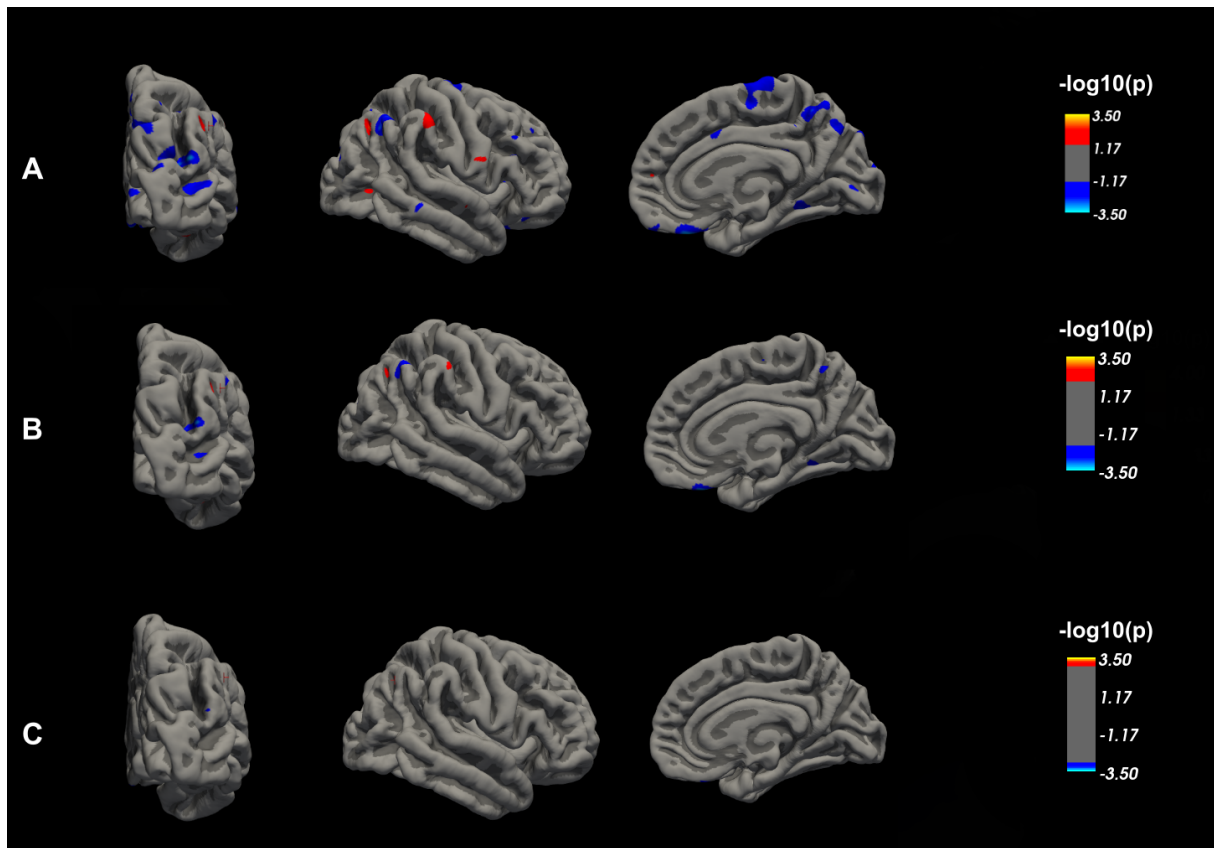


Figure 3: Between-group differences between HC and RLP ($p < 0.05$ cluster level corrected) in cortical thickness (A), surface area (B), cortical **volume (C) and LGI (D) projected onto the pial surface of the fsaverage in the medial, lateral and posterior view for the LH and the RH. Differences were computed separately for LH and RH. Clusters with red and yellow colours indicate brain regions where RLP have significantly lower cortical indexes values than the **HC** and clusters with blue colours indicate regions with increased values for RLP in comparison with HC. These clusters are defined in detail in Table 3. The final RLP mask is presented in green.**



Supplementary Figure 1: Between-group differences between HC and LLP in cortical thickness, (A) $p < 0.05$, (B) $p < 0.01$ and (C) $p < 0.001$ (Uncorrected). Clusters with red and yellow colours indicate brain regions where LLP have significantly lower cortical indexes values than the **HC** and clusters with blue colours indicate regions with increased values for LLP in comparison with HC. Clusters are projected onto the pial surface of the fsaverage in the medial, lateral and posterior view for the RH.

Table 1: General profile of the participants.

	HC	LLP	RLP	p-
	Mean (\pmstd) or n	Mean (\pmstd) or n	Mean (\pmstd) or n	value*
	(%)	(%)	(%)	
	n=30	n=19	n=15	
Age	7.71 (\pm 0.54)	7.23 (\pm 0.13)	7.28 \pm 0.20)	0.543
(years)				
Gender	Males: 14 (46.67%) Females: 16 (53.33%)	Males: 10 (52.63%) Females:9 (47.37%)	Males: 10 (66.67%) Females: 5 (33.33%)	0.376 ^a
Right-handed	27 (90 %)	7 (36.84 %)	14 (93.33 %)	0.180 ^a

Lesion size (ml)	_____	32.45 (\pm 33.21)	38.16 (\pm 46.94)	0.859
TIV	1395.4 (\pm 110.01)	1307.0 (\pm 157.71)	1277.7 (\pm 98.30)	0.127

HC: Healthy Controls, **LLP:** Left Lesioned Patients, **RLP:** Right Lesioned Patients,

TIV: Total intracranial volume

* p-values are obtained by one-way Kruskal Wallis non-parametric ANOVA

^a Chi squared test

Table 2: Significant clusters for the LH and RH resulting from the vertex wise comparison of cortical thickness, surface area, cortical volume and LGI between LLP and HC groups.

LLP < HC clusters are presented in white cells and LLP > HC are presented in grey cells.

LH/RH	Cortical index	Brain area	Cluster size (mm^2)	MNI x, y, z	p-value
RH	Cortical thickness	_____	_____	_____	_____
	Surface area	Occipital pole	1538.71	9.4, -97.3, 13.2	0.0076
	Cortical volume	Superior occipital gyrus	969.34	14.7, -94.0, 12.4	0.0015
	LGI	Superior occipital gyrus	3065.30	26.1, -71.0, 32.5	0.0000
		Superior frontal gyrus	2703.80	7.4, 24.0, 57.1	0.0000
		Superior part of the precentral sulcus	753.30	22.6, -6.8, 48.7	0.0000
LH	Cortical thickness	Postcentral sulcus	490.93	-31.1, -40.6, 44.4	0.0001

		Superior segment of the circular sulcus of the insula	470.17	-35.8, -9.7, 16.9	0.0001
		Inferior part of the precentral sulcus	453.18	-42.2, -1.3, 37.7	0.0001
		Inferior frontal sulcus	250.97	-37.4, 14.4, 23.0	0.0001
		Angular gyrus	145.55	-52.7, -52.4, 18.5	0.0001
		Orbital sulci	135.10	-20.0, 24.7, -16.3	0.0001
		Long insular gyrus and central sulcus of the insula	134.65	-36.7, 1.5, -9.8	0.0001
		Lateral aspect of the superior temporal gyrus	124.22	61.4, -48.1, 15.5	0.0003
		Intraparietal sulcus and transverse parietal sulci	106.94	-29.8, -54.3, 35.9	0.0010
		Planum polar of the superior temporal gyrus	105.27	-44.5, 8.7, -24.1	0.0011
	Surface area	Transverse temporal sulcus	3241.29	-44.6, -30.5, 2.4	0.0001
		Inferior segment of the circular sulcus of the insula	3009.35	-45.9, -20.5, -4.6	0.0001
		Precentral gyrus	858.32	-49.8, -8.4, 41.3	0.0001
		Intraparietal sulcus and transverse parietal sulci	725.58	-34.1, -58.4, 37.1	0.0001

		Subcentral gyrus and sulci	117.20	-62.0, -8.9, 26.9	0.0005
		Superior part of the precentral sulcus	93.23	-26.7, -13.2, 53.9	0.0044
	Cortical volume	Inferior segment of the circular sulcus of the insula	2610.56	-44.0, -23.7, -1.7	0.0001
		Transverse temporal sulcus	1058.67	-49.7, -30.4, 3.4	0.0001
		Angular gyrus	890.85	-41.3, -65.4, 39.6	0.0001
		Inferior part of the precentral sulcus	507.72	-48.5, -1.9, 40.0	0.0001
		Lingual gyrus	365.83	-12.9, -82.2, -12.6	0.0001
		Inferior temporal gyrus	143.56	-55.7, -53.8, -13.0	0.0001
		Middle temporal gyrus	129.62	-59.2, -58.2, 2.3	0.0001
		Lateral occipito-temporal gyrus	128.40	-32.5, -34.8, -24.6	0.0001
		Subcentral gyrus and sulci	107.54	-62.0, -8.9, 26.9	0.0010
	LGI	Subcentral gyrus and sulci	4447.37	-57.2, -0.1, 10.7	0.0001
		Anterior occipital sulcus and preoccipital notch	619.36	-39.8, -68.1, -0.0	0.0001

Table 3: Significant clusters for the LH and RH resulting from the vertex wise comparison of cortical thickness, surface area, cortical volume and LGI between RLP and HC groups.
RLP < HC clusters are presented in white cells and RLP > HC are presented in grey cells.

LH/RH	Cortical index	Brain area	Cluster size (mm ²)	MNI x, y, z	p-value
RH	Cortical thickness	Postcentral gyrus	100.64	46.6, -24.8, 51.3	0.0015
		Pericallosal sulcus	71.81	3.6, 2.0, 28.7	0.0394
	Surface area	Posterior ramus of the lateral sulcus	1226.49	33.1, -28.7, 19.9	0.0001
		Angular gyrus	1079.26	43.9, -62.8, 36.1	0.0001
		Middle frontal sulcus	430.07	37.9, 24.2, 38.1	0.0001
		Middle occipital gyrus	277.33	47.1, -74.8, 4.6	0.0001
		Middle temporal gyrus	265.06	56.8, -0.1, -28.8	0.0001
		Suborbital sulcus	168.70	8.5, 45.8, -14.8	0.0001
		Supramarginal gyrus	141.42	60.4, -40.2, 27.5	0.0001
		Postcentral gyrus	122.57	43.6, -29.0, 51.5	0.0002
	Cortical volume	Posterior ramus of the lateral sulcus	775.09	33.0, -29.4, 19.4	0.0001
		Angular gyrus	192.83	40.9, -66.7, 37.9	0.0001
		Postcentral gyrus	166.21	47.8, -23.9, 52.9	0.0001
		Anterior occipital sulcus and middle occipital gyrus	103.97	44.0, -67.7, 7.0	0.0008
		Middle occipital gyrus	87.51	38.4, -81.8, 26.5	0.0078
		Inferior temporal gyrus	80.79	51.7, -44.0, -19.5	0.0163
		Subcentral gyrus and sulci	75.60	63.2, -13.0, 18.7	0.0268
	LGI	—	—	—	—
LH	Cortical thickness	—	—	—	—
	Surface area	—	—	—	—
	Cortical volume	—	—	—	—

	LGI	Middle occipital gyrus	2228.28	-40.3, -81.8, 17.3	0.0001
		Precuneus	1211.78	-6.4, -69.5, 46.0	0.0001