Lower Hippocampal Volume in Patients with Schizophrenia and Bipolar Disorder: A Quantitative MRI Study

Jinya Sato 1, Yoji Hirano 1,2,* Noriaki Hirakawa 1, Junichi Takahashi 1, Naoya Oribe 1,3, Hironori Kuga 1,3, Itta Nakamura 1, Shogo Hirano 1, Takefumi Ueno 3, Osamu Togao 4, Akio Hiwatashi 5, Tomohiro Nakao 1 and Toshiaki Onitsuka 1,*

1 Department of Neuropsychiatry, Graduate School of Medical Sciences, Kyushu University, Fukuoka 812-8582, Japan; jinya.sato56@gmail.com (J.S.); hirakawa34@gmail.com (N.H.); yooni1977@gmail.com (J.T.); n.oribe@mac.com (N.O.); hirokuga.370@gmail.com (H.K.); nakamura.itta.352@m.kyushu-u.ac.jp (I.N.); hiranosho@gmail.com (S.H.); nakao.tomohiro.275@m.kyushu-u.ac.jp (T.N.)
2 Institute of Industrial Science, The University of Tokyo, Tokyo 153-8505, Japan
3 Hizen Psychiatric Medical Center, Division of Clinical Research, National Hospital Organization, Saga 842-0192, Japan; uenotk@gmail.com
4 Department of Molecular Imaging and Diagnosis, Graduate School of Medical Sciences, Kyushu University, Fukuoka 812-8582, Japan; toga@radiol.med.kyushu-u.ac.jp
5 Department of Clinical Radiology, Graduate School of Medical Sciences, Kyushu University, Fukuoka 812-8582, Japan; hiwatasi@radiol.med.kyushu-u.ac.jp
* Correspondence: yhouji@mac.com (Y.H.); onitsuka.toshiaki.939@m.kyushu-u.ac.jp (T.O.); Tel.: +81-92-642-5627 (Y.H. & T.O.)

Abstract: Since patients with schizophrenia (SZ) and bipolar disorder (BD) share many biological features, detecting biomarkers that differentiate SZ and BD patients is crucial for optimized treatments. High-resolution magnetic resonance imaging (MRI) is suitable for detecting subtle brain structural differences in patients with psychiatric disorders. In the present study, we adopted a neuroanatomically defined and manually delineated region of interest (ROI) method to evaluate the amygdalae, hippocampi, Heschl’s gyrus (HG), and planum temporale (PT), because these regions are crucial in the development of SZ and BD. ROI volumes were measured using high resolution MRI in 31 healthy subjects (HS), 23 SZ patients, and 21 BD patients. Right hippocampal volumes differed significantly among groups (HS > BD > SZ), whereas left hippocampal volumes were lower in SZ patients than in HS and BD patients (HS = BD > SZ). Volumes of the amygdalae, HG, and PT did not differ among the three groups. For clinical correlations, there were no significant associations between ROI volumes and demographics/clinical symptoms. Our study revealed significant lower hippocampal volume in patients with SZ and BD, and we suggest that the right hippocampal volume is a potential biomarker for differentiation between SZ and BD.

Keywords: schizophrenia; bipolar disorder; MRI; hippocampus; volume reduction

1. Introduction

Patients with schizophrenia (SZ) and bipolar disorder (BD) share many biological features, including genetic architecture (e.g., [1,2]) and structural (e.g., [3]) and functional brain abnormalities (e.g., [4–10]). Clinically, it is an urgent issue to accurately diagnose and distinguish between the two diseases for providing appropriate treatment and improving patient outcome, as they sometimes present similar pathologies. Thus, the identification of anatomical and functional biomarkers is crucial for differentiating SZ and BD patients to optimize pharmacological and psychosocial treatments for each condition [11]. Subcortical and cortical abnormalities in psychoses have been reported extensively, and high-resolution magnetic resonance imaging (MRI) is well-suited for detecting subtle brain structural differences in these populations [12–15]. A gold standard approach in MRI research is the
use of neuroanatomically defined and manually delineated regions of interest (ROIs). A voxel-based morphometric (VBM) method is useful to investigate the whole brain, but the anatomical warping and density-based methodology can lead to failure in detecting abnormalities that are identified using ROI analyses [16]. Furthermore, it has been suggested that the analytical flexibility and the complex and variable workflows have substantial effects on results [17]; therefore, simpler MRI analysis methods still carry value. In the present study, we adopted a neuroanatomically defined and manually delineated ROI method to evaluate the amygdalae, hippocampi, Heschl’s gyrus (HG), and planum temporale (PT) as regions that have been shown to be crucial in the development of SZ and BD [18–21].

Medial temporal lobe changes are one of the most consistent findings in MRI studies of SZ and BD. The hippocampus is involved in memory and spatial learning [22]. Hippocampal volume reductions have been reported repeatedly in patients with SZ and BD [23]. The amygdala is an almond-shaped nucleus located deep within the medial temporal lobes and plays a primary role in the processing and memory of emotional reactions [24,25]. Both increases and decreases in amygdala volumes have been reported in patients with BD, although a recent meta-analysis found no significant amygdala volume changes in BD [26]. Conversely, an international collaboration for the study of BD, the Enhancing Neuroimaging Genetics through Meta-Analysis (ENIGMA) Consortium, reported significantly smaller amygdaloid volumes in BD compared with healthy subjects (HS) [27].

The HG includes the primary auditory cortex [28], and the left PT plays an important role in language processing [29]. Structural abnormalities of HG and PT [18,30], alongside auditory and language-related deficits [31,32], are observed frequently in patients with SZ. For example, patients with first-episode SZ showed lower left PT and bilateral HG gray matter volumes compared with HS and patients with BD [30]. In addition, Kasai et al. reported progressive volume reductions in the left HG and PT gray matter in first-episode SZ patients, which were not observed in first-episode affective psychosis patients or HS [18]. More recently, Hirano et al. investigated the association between neurophysiological and structural deficits in SZ patients and found spontaneous gamma oscillation deficits [33] and cortical volume deficits in HG [34]. In BD patients, Reite et al. reported alterations in structural/functional asymmetry in HG [18]. Moreover, Takahashi et al. reported that BD patients showed significantly lower left PT volumes compared with HS [35].

In the present study, we adopted a neuroanatomically defined and manually delineated ROI method to evaluate the amygdalae, hippocampi, HG, and PT. As noted before, functional or structural abnormalities have been reported in both SZ and BD in these four ROIs. We hypothesized that medial temporal lobe abnormalities (mainly hippocampus) may be common to both SZ and BD, and lateral temporal lobe abnormalities may be specific to SZ.

2. Materials and Methods

2.1. Participants

We analyzed magnetic resonance (MR) images of 23 patients with SZ, 21 patients with BD, and 31 HS. All subjects had normal hearing, were aged 20–63 years, and were right-handed [36]. After a detailed description of the study, all participants provided informed consent, according to the regulations of the Ethics Committee of the Graduate School of Medical Sciences, Kyushu University. Experimental procedures were approved by the Kyushu University Institutional Review Board for Clinical Trials (approval number: 26018, date: June 12, 2014), and conformed to the Declaration of Helsinki. The exclusion criteria were (1) neurological illness or major head trauma, (2) having received electroconvulsive therapy, (3) alcohol or drug dependence, (4) alcohol or drug abuse within the past 5 years, or (5) a verbal intelligence quotient below 75. HS were screened using the Structured Clinical Interview nonpatient edition. None of the HS or their first-degree relatives had an Axis-I psychiatric disorder. The socioeconomic status (SES) of the subjects and their parental SES were measured using the Hollingshead two-factor index [37]. For SZ, the Positive and Negative Syndrome Scale (PANSS) [38] was administered to assess the severity of psychiatric symptoms. The Young Mania Rating Scale (YMRS) [39] and the
Hamilton Depression Rating Scale (HAM-D) [40] were administered to assess the severity of mood symptoms in BD. Table 1 shows the demographic and clinical characteristics of the participants.

Table 1. Demographic and clinical characteristics of the participants.

|                       | Healthy Subjects (HS) | Patients with Bipolar Disorder (BD) | Patients with Schizophrenia (SZ) | χ² or F or t | df | p   |
|-----------------------|-----------------------|-------------------------------------|----------------------------------|--------------|----|-----|
| Male/Female           | 11/20                 | 11/10                               | 8/15                             | 1.66         | 2  | 0.44|
| Age (years)           | 49.2 ± 9.8            | 48.9 ± 9.1                          | 44.3 ± 6.2                       | 2.50         | 2  | 0.73|
| Handedness a          | 92.0 ± 32.7           | 82.4 ± 51.0                         | 96.7 ± 8.7                       | 0.99         | 2  | 0.38|
| SES b                 | 2.4 ± 0.8             | 2.9 ± 1.0                           | 3.6 ± 1.0                        | 10.66        | 2  | <0.001|
| Parental SES          | 3.1 ± 0.9             | 2.9 ± 0.9                           | 2.8 ± 0.9                        | 1.10         | 2  | 0.34|
| Education (years)     | 14.4 ± 2.6            | 14.2 ± 2.3                          | 14.4 ± 2.0                       | 0.95         | 2  | 0.39|
| Symptom onset #       | 37.8 ± 11.2           | 26.7 ± 6.7                          | 31.95 ± 42.3                     | 5.60         | 42 | <0.001|
| Medication dose (CPZ equiv., mg) | 144.2 ± 183.9        | 599.8 ± 328.4                       | 5.60                             | 42 | <0.001|
| PANSS positive        |                       | 18.1 ± 7.9                          |                                  |              |    |     |
| PANSS negative        |                       | 22.5 ± 7.7                          |                                  |              |    |     |
| PANSS general         |                       | 42.9 ± 15.2                         |                                  |              |    |     |
| YMRS                  | 5.2 ± 5.7             |                                    |                                  |              |    |     |
| SIGH-D score          | 10.3 ± 5.9            |                                    |                                  |              |    |     |

Values are means ± SD unless otherwise noted. a Edinburgh Handedness Inventory, b Hollingshead two factor Index of Social Position. SES = socioeconomic status, YMRS = Young Mania Rating Scale, SIGH-D = Structured Interview Guide for the Hamilton Depression Rating Scale. # Symptom onset is the age at which the symptoms appeared.

2.2. MRI Data Acquisition

T1-weighted MR images were acquired using a three-dimensional (3D) turbo field echo sequence on a 3 Tesla scanner (Achieva TX, Philips Healthcare, Best, The Netherlands) at the Department of Radiology, Kyushu University Hospital. The imaging variables were as follows: repetition time = 8.2 ms, echo time = 3.8 ms, flip angle = 8°, field of view = 24 × 24 cm, number of echoes = 1, matrix = 240 × 240, inversion time = 1025.9 ms, number of slices = 190, and slice thickness = 1 mm. Images were aligned using the anterior and posterior commissure line and the sagittal sulcus to correct head tilt.

Total gray matter, white matter, and cerebrospinal fluid (CSF) volumes were calculated using Statistical Parametric Mapping 12 (SPM12, http://www.fil.ion.ucl.ac.uk/spm (accessed on 7 October 2020)), and VBM was performed using SPM12, running in MATLAB R2014a (The Math Works Inc., Natick, MA, USA). T1-weighted images were first segmented into gray matter, white matter, and CSF sections using tissue probability maps based on the International Consortium of Brain Mapping template for East Asian brains. Subsequently, we performed diffeomorphic anatomical registration through exponentiated lie algebra in SPM12 for intersubject registration of gray matter images [41]. The registered images were then smoothed with a Gaussian kernel of 8 mm full-width half-maximum and transformed into Montreal Neurological Institute stereotactic space using affine and nonlinear spatial normalization, implemented in SPM12. Total gray matter, white matter, and CSF volumes were generated from the VBM analysis. Total intracranial volume (ICV) was calculated as the sum of the gray matter, white matter, and CSF volumes.

2.3. Regions of Interest and Volume Measurement

The gray matter of the HG and PT, hippocampi, and amygdalae were outlined manually on a PC without knowledge of diagnosis (Figure 1). We used 3D information to provide reliable measures of the ROIs using a software package for medical image analysis [3D slicer, http://www.slicer.org (accessed on 7 October 2020)].
We used similar criteria to the previous works of Kwon et al. and Barta et al. for delineating HG and PT [42,43]. Briefly, HG was first identified in the most posterior coronal slice. Investigators manually drew the gray matter of HG from the most posterior to the most anterior slice. The most anterior slice of HG was defined as the last slice that HG could be identified anteriorly. In most cases, HG represented a single transverse convolution. In cases of more than one transverse convolution, we followed the literature’s definition [44–47]: when multiple convolutions originated medially from a common stem, these were all defined as HG; however, when they originated separately from the retrostriate region, only the most anterior gyrus was labeled as HG, and the more posterior gyri were identified as PT. The anterior border of PT was defined by the posterior border of HG. Posteriorly, investigators traced the gray matter of the PT on the coronal images to the end of the Sylvian fissure. The amygdalae and hippocampi were also outlined manually. For the amygdala and hippocampus, the most anterior slice used for the amygdala was the slice in which the white matter tract linking the temporal lobe with the rest of the brain (temporal stem) could be seen. The most posterior slice of the hippocampus was the last appearance of fibers of the crux of the fornix. Once drawn, ROIs could be viewed in any plane as 3D objects for further editing. All editing was performed blind to diagnosis. Interrater reliability of the ROIs was evaluated by three independent raters (J.S., N.H., and J.T.), who were also blind to diagnosis. Five cases were selected at random, and every slice was edited by the raters. The intraclass correlations for interrater reliability were calculated.
2.4. Statistical Analysis

To correct for brain size differences, we used relative volumes, which were calculated using the following formula: relative volume (%) = (absolute ROI volume/ICV) × 100. For the ROI analysis of the subcortical structures, we used a mixed model repeated measures analysis of variance (ANCOVA) with group (SZ, BD, and HS) as the between-subjects factor, hemisphere (left or right) and region (amygdala or hippocampus) as the within-subjects factors, and age as a covariate. When significant interactions related to group were observed, we evaluated group differences separately for each region to identify the source of the interactions. We used follow-up ANCOVAs with group as the between-subjects factor, hemisphere as the within-subjects factor, and age as a covariate for each region. We performed one-way analyses of variance (ANOVAs) of the average volume across both hemispheres for each ROI. For the ROI analysis of the cortical structures, we used a mixed model ANCOVA with group (SZ, BD, and HS) as the between-subjects factor, hemisphere (left or right) and region (HG or PT) as the within-subjects factors, and age as a covariate. Post hoc tests were the same as the subcortical structure analyses. All statistical tests in ANCOVA and ANOVA were 2-tailed with \( \alpha = 0.05 \). Exploratory analyses of the relationship between relative volumes of the ROIs and psychopathology scales were evaluated using Spearman’s \( \rho \). Due to our analyses involving multiple correlations, we set alpha levels at a conservative \( p < 0.001 \).

When significant group differences were observed in a particular region, optimal sensitivity and specificity of the ROI for the diagnosis of HS, SZ, or BD were determined via receiver operating characteristic (ROC) curve analysis using a nonparametric approach. We calculated the Youden index for each cutoff value as corresponding \((\text{sensitivity} + \text{specificity}) − 1\) to find the cutoff values that maximized discriminating power. We used IBM SPSS Statistics 26 for the analyses.

3. Results

3.1. Reliability

The intraclass correlations for interrater reliability were 0.95 for the left hippocampus, 0.92 for the right hippocampus, 0.91 for the left amygdala, 0.90 for the right amygdala, 0.91 for left HG, 0.93 for right HG, 0.91 for the left PT, and 0.95 for the right PT.

3.2. Relative Volumes of Each ROI

Table 2 shows the relative volumes of each ROI. For the amygdalae and hippocampi, the repeated measures ANCOVA showed significant hemisphere-by-region-by-group \((F[2,71] = 3.56, p = 0.03)\) and region-by-group interactions \((F[2,71] = 8.72, p < 0.0001)\) and significant main effects of group \((F[2,71] = 6.25, p = 0.003)\) and region \((F[1,71] = 102.23, p < 0.0001)\) but not hemisphere \((F[1,71] = 1.17, p = 0.28)\). There was no significant hemisphere-by-group \((F[2,71] = 1.38, p = 0.26)\) or hemisphere-by-region \((F[1,71] = 0.04, p = 0.84)\) interactions. To delineate the significant hemisphere-by-region-by-group interactions, post hoc tests were performed for each ROI.
To determine if schizophrenia (SZ) was significantly different from bipolar disorder (BD) and healthy subjects (HS) (Tukey’s honestly significant difference test, \( p < 0.05 \)).

| Table 2. Volume of each region of interest (ROI). | Healthy Subjects (HS) | Patients with Bipolar Disorder (BD) | Patients with Schizophrenia (SZ) | df | \( F \) | \( p \) |
|-------------------------------------------------|-----------------------|-------------------------------------|---------------------------------|----|-----|-----|
| **Intracranial Volume (mL)**                    | 1380 ± 128            | 1422 ± 136                          | 1418 ± 157                      | 2,72 | 0.79 | 0.46 |
| **Left Hippocampus**                            |                       |                                     |                                 |     |     |     |
| Absolute (mL)                                   | 2.91 ± 0.22           | 2.92 ± 0.24                         | 2.72 ± 0.24                     | 2,72 | 7.47 | 0.001* |
| Relative (%)                                    | 0.211 ± 0.015         | 0.206 ± 0.022                       | 0.193 ± 0.018                   |     |     |     |
| **Right Hippocampus**                           |                       |                                     |                                 |     |     |     |
| Absolute (mL)                                   | 2.95 ± 0.29           | 2.86 ± 0.20                         | 2.67 ± 0.28                     | 2,72 | 15.06 | <0.0001† |
| Relative (%)                                    | 0.214 ± 0.014         | 0.202 ± 0.020                       | 0.189 ± 0.017                   |     |     |     |
| **Left Amygdala**                               |                       |                                     |                                 |     |     |     |
| Absolute (mL)                                   | 1.26 ± 0.17           | 1.28 ± 0.14                         | 1.24 ± 0.16                     | 2,72 | 1.01 | 0.37 |
| Relative (%)                                    | 0.091 ± 0.009         | 0.091 ± 0.012                       | 0.087 ± 0.009                   |     |     |     |
| **Right Amygdala**                              |                       |                                     |                                 |     |     |     |
| Absolute (mL)                                   | 1.25 ± 0.20           | 1.31 ± 0.14                         | 1.26 ± 0.17                     | 2,72 | 0.78 | 0.46 |
| Relative (%)                                    | 0.091 ± 0.010         | 0.093 ± 0.012                       | 0.089 ± 0.008                   |     |     |     |
| **Left Heschl’s gyrus**                         |                       |                                     |                                 |     |     |     |
| Absolute (mL)                                   | 2.03 ± 0.50           | 2.02 ± 0.53                         | 1.95 ± 0.59                     | 2,72 | 0.71 | 0.97 |
| Relative (%)                                    | 0.147 ± 0.032         | 0.148 ± 0.039                       | 0.137 ± 0.035                   |     |     |     |
| **Right Heschl’s gyrus**                        |                       |                                     |                                 |     |     |     |
| Absolute (mL)                                   | 1.71 ± 0.48           | 1.77 ± 0.46                         | 1.75 ± 0.55                     | 2,72 | 0.03 | 0.97 |
| Relative (%)                                    | 0.123 ± 0.029         | 0.125 ± 0.034                       | 0.123 ± 0.034                   |     |     |     |
| **Left Planum temporale**                       |                       |                                     |                                 |     |     |     |
| Absolute (mL)                                   | 2.94 ± 0.60           | 2.88 ± 0.68                         | 2.79 ± 0.56                     | 2,72 | 1.20 | 0.31 |
| Relative (%)                                    | 0.213 ± 0.036         | 0.203 ± 0.046                       | 0.197 ± 0.032                   |     |     |     |
| **Right Planum temporale**                      |                       |                                     |                                 |     |     |     |
| Absolute (mL)                                   | 2.53 ± 0.76           | 2.56 ± 0.36                         | 2.63 ± 0.65                     | 2,72 | 0.06 | 0.94 |
| Relative (%)                                    | 0.183 ± 0.050         | 0.182 ± 0.032                       | 0.186 ± 0.044                   |     |     |     |

* Post hoc tests indicated that schizophrenia (SZ) was significantly different from bipolar disorder (BD) and healthy subjects (HS) (Tukey’s honestly significant difference test, \( p < 0.05 \)). † Post hoc tests indicated that volumes were significantly different as follows; HS > BD > SZ (Tukey Honestly Significant Difference, \( p < 0.05 \)).

For the amygdalae, the repeated-measures ANCOVA showed no significant main effects of group (\( F[2,71] = 0.53, p = 0.59 \)) or hemisphere (\( F[2,71] = 1.15, p = 0.29 \)), or a significant hemisphere-by-group interaction (\( F[2,71] = 0.39, p = 0.68 \)). These results indicated that there were no significant group differences among the three groups in either hemisphere.

For the hippocampus, the repeated-measures ANCOVA revealed a significant hemisphere-by-group interaction (\( F[2,71] = 3.56, p = 0.03 \)) and a main effect of group (\( F[2,71] = 9.21, p < 0.0001 \)) but not hemisphere (\( F[1,71] = 0.32, p = 0.57 \)). In the right hemisphere, the one-way ANOVA showed a significant group difference (\( F[2,72] = 15.06, p < 0.0001 \)), and post hoc Tukey’s honestly significant difference (HSD) tests revealed BD < HS (\( p = 0.04 \)), SZ < HS (\( p < 0.0001 \)) and SZ < BD (\( p = 0.03 \)) right hippocampal volumes. These results indicated that right hippocampal volumes were highest in HS, lower in the BD patients, and lowest in the SZ patients. In the left hemisphere, the one-way ANOVA showed a significant group difference (\( F[2,72] = 7.47, p = 0.001 \)), and post hoc HSD tests revealed SZ < HS (\( p = 0.001 \)) and SZ < BD (\( p = 0.04 \)) hippocampal volumes. No significant difference was found for the left hippocampus between HS and BD patients (\( p = 0.58 \)). These results revealed that SZ patients had a lower left hippocampal volume compared with HS and BD patients (see Figure 2).

For HG and PT, the repeated-measures ANCOVA showed a significant main effect of region (\( F[1,71] = 20.85, p < 0.001 \)). There were no other significant main effects or interactions, which indicated no significant group differences in the HG or PT (see Figure 3).

We also performed repeated-measures ANCOVAs using age and sex as covariates for ROI analysis; statistical results were the same using both methods.
Figure 2. Relative volumes of the left and right amygdalae and hippocampi in healthy subjects (HS) \((n = 31)\), patients with bipolar disorder (BD) \((n = 21)\), and patients with schizophrenia (SZ) \((n = 23)\). Horizontal lines indicate means. * \(p < 0.05\), ** \(p < 0.001\).

For HG and PT, the repeated-measures ANCOVA showed a significant main effect of region \((F[1,71] = 20.85, p < 0.001)\). There were no other significant main effects or interactions, which indicated no significant group differences in the HG or PT (see Figure 3).

We also performed repeated-measures ANCOVAs using age and sex as covariates for ROI analysis; statistical results were the same using both methods.

3.3. Correlations

Regarding the associations between ROI volume and demographic/clinical variables, we found no significant correlations in HS \((-0.34 \leq \rho \leq 0.38, 0.04 \leq p \leq 0.89)\), SZ patients \((-0.34 \leq \rho \leq 0.43, 0.04 \leq p \leq 0.98)\), or BD patients \((-0.40 \leq \rho \leq 0.48; 0.03 \leq p \leq 0.94)\).

3.4. ROC Analysis

Since significant differences among three groups were observed in the right hippocampus, we used ROC curve analysis to explore the discriminatory value of the relative volumes. Figure 4 shows the ROC curve of relative volumes of the right hippocampus between HS and BD, HS and SZ, and BD and SZ. The area under the curve (AUC) of the ROC analysis in BD vs. SZ was maximal for the right hippocampal relative volume \((AUC = 0.70, \text{Standard error} = 0.08, p = 0.03, 95\% \text{CI} = 0.54–0.85)\), indicating that the right hippocampal volume could be used to differentiate between BD and SZ subjects with moderate accuracy. The Youden index indicated a favorable cutoff point of 0.20%, which resulted in 62% sensitivity and 74% specificity.

4. Discussion

We investigated the volumes of the hippocampi, amygdalae, HG, and PT in SZ patients, BD patients, and HS. We found that right hippocampal volumes were lower in BD patients compared with controls, and further lower in SZ patients, whereas left hippocampal volumes were lower exclusively in SZ patients. There were no significant differences in either amygdala, HG, or PT among the three groups.

Our results are consistent with many previous studies (e.g., \([27,48]\)) although the recent study \([14]\) of the direct comparison found no significant volume differences between the two groups. Our findings suggest that the lateralized hippocampal volume abnormal...
3.3. Correlations

Regarding the associations between ROI volume and demographic/clinical variables, we found no significant correlations in HS ($-0.34 \leq \rho \leq 0.38$, $0.04 \leq p \leq 0.89$), SZ patients ($-0.34 \leq \rho \leq 0.43$, $0.04 \leq p \leq 0.98$), or BD patients ($-0.40 \leq \rho \leq 0.48$, $0.03 \leq p \leq 0.94$).

3.4. ROC Analysis

Since significant differences among three groups were observed in the right hippocampus, we used ROC curve analysis to explore the discriminatory value of the relative volumes. Figure 4 shows the ROC curve of relative volumes of the right hippocampus between HS and BD, HS and SZ, and BD and SZ. The area under the curve (AUC) of the ROC analysis in BD vs. SZ was maximal for the right hippocampal relative volume ($AUC = 0.70$, Standard error = 0.08, $p = 0.03$, 95% CI = 0.54–0.85), indicating that the right hippocampal volume could be used to differentiate between BD and SZ subjects with moderate accuracy. The Youden index indicated a favorable cutoff point of 0.20%, which resulted in 62% sensitivity and 74% specificity.

4. Discussion

We investigated the volumes of the hippocampi, amygdalae, HG, and PT in SZ patients, BD patients, and HS. We found that right hippocampal volumes were lower in BD patients compared with controls, and further lower in SZ patients, whereas left hippocampal volumes were lower exclusively in SZ patients. There were no significant differences in either amygdala, HG, or PT among the three groups.

Our results are consistent with many previous studies (e.g., [27,48]) although the recent study [14] of the direct comparison found no significant volume differences between the two groups. Our findings suggest that the lateralized hippocampal volume abnormality is related to the development of SZ and BD. We also found that the lower left hippocampal volume was not observed in BD patients and was specific to SZ, which is in line with numerous studies that have reported left hemisphere dominant abnormalities in SZ [49].

Although the degree of lowered volume in BD patients was not to that observed in SZ patients, the volume of the right hippocampus was found to be significantly lower in BD compared with HS. The ENIGMA Consortium reported significant lower volumes in the hippocampus, thalamus, and amygdala in patients with BD [27]; however, they did not evaluate laterality. We revealed that lower right hippocampal volume may be associated with pathology of BD. We highlight that in addition to large sample automated analysis methods, alternative approaches, such as the manual ROI drawing method, are important and valuable.

In the present study, there were no significant differences in either amygdala among the three groups. Moreover, contrary to our hypotheses and findings of previous studies (e.g., [18]), we did not find significant group differences in HG or PT. Bryant et al. reported no differences in the volumes of temporal lobe structures between female SZ patients and
In addition, Bora et al. performed voxelwise meta-analysis and reported that gray matter reductions of SZ and BD were less severe in sex-balanced samples [15]. Therefore, sex effects should be considered in future studies in larger sample sizes. We did not find any significant associations between the volumes of the ROIs and demographics/clinical symptoms. Although hippocampal volume reductions may increase the risk of psychosis, they may not be directly related to the severity of symptoms. Given the function of the hippocampus, it will be important to investigate relationships between hippocampal volumes and cognitive functional domains in future studies.

Because SZ and BD share the biological basis of psychoses [1–11,51–57] and they sometimes present similar pathologies, detection of anatomical and functional biomarkers that differentiate SZ and BD patients is crucial for the optimization of pharmacological and psychosocial treatments for each condition. The results of our ROC analyses suggest that the right hippocampal volume is a potential marker for differentiation between SZ and BD. To this end, multimodal functional neuroimaging studies [26,51–61] that combine structural MRI [3,16,18–21,23,26–28,42–44,51,61–63], functional MRI [26,53–61,64,65], magnetic resonance spectroscopy [66–69], positron emission tomography [69–72], electroencephalography, and magnetoencephalography [4–9,59,73–85] may lead to a better understanding of the biological bases of SZ and BD and enable the development of optimized treatments (e.g., [53,54,58,61,86,87]).

We must note that this study has some limitations. First, we had a relatively small sample size owing to our type of dataset and manual drawing approach, which requires considerable time and labor. Therefore, weaker correlations that were hypothesized may have gone undetected. Second, as mentioned earlier, sex differences were not explored in our study but warrant investigation in a larger sample, as suggested by Bryant et al. [50], Egloff et al. [88], and Bora et al. [15]. Third, the effect of the duration of illness should be considered. The current sample of BD had higher age and lower duration of illness compared to SZ. Thus, in the present study, we could not exclude the effect of earlier disease-specific differences and late unspecific processes, including disease progress or effects of medications. Importantly, metaregression analysis by Fuar-Poli et al. [89] reported that the higher the cumulative exposure to antipsychotic treatment, the greater the gray matter decreases in the SZ group over follow-up time, while no significant effects were observed for duration of illness. Although there were no significant correlations between any ROI volumes and duration of illness in our study, associations between the effect of disease duration and the hippocampal volumes should be examined in a future study. Finally, we could not exclude the medication effects on hippocampal volumes in clinical groups, as pointed out by a previous study [90].

In summary, we showed that patients with SZ and BD have significantly lower hippocampal volumes compared with HS. Furthermore, we suggest that the right hippocampal volume is a potential biomarker for differentiation between SZ and BD.

Author Contributions: Conceptualization, T.O. and J.S.; methodology, T.O., J.S., N.H., and Y.H.; software, J.S., H.K., T.U., S.H., and N.H.; validation, J.S., N.H., J.T., I.N., N.O., Y.H., and T.O.; formal analysis, T.O. and J.S.; investigation, J.S., Y.H., and T.O.; resources, A.H. and O.T.; data curation, J.S., T.O., and Y.H.; writing—original draft preparation, J.S.; writing—review and editing, Y.H., T.N., and T.O.; visualization, J.S. and Y.H.; supervision, Y.H.; and T.O.; project administration, Y.H., T.O., and T.N.; funding acquisition, Y.H. and T.O. All authors have read and agreed to the published version of the manuscript.

Funding: This research was supported, in part, by AMED (Japan Agency for Medical Research and Development) under Grant Number JP20dm0207069 (T.O.) and GAJJ020620 (JP19dm0107124h0004) (Y.H.); Grant-in-Aid for Young Scientists B JP22791129 (Y.H.), JP 15K19735 (N.O.), and JP 17K16385 (N.O.), a Grant-in-Aid for Scientific Research C: JP 16K10217 (T.O.), JP 19K08049 (N.O.), JP 15K09836 (Y.H.), JP 18K07604 (Y.H.), JP 19H03579 (Y.H.) and Fund for the Promotion of Joint International Research (Fostering Joint International Research B): JP20KK0193 (Y.H.) from the Japan Society for the Promotion of Science (JSPS); Medical Research Fund (Y.H.) from Takeda Science Foundation; SIRS Research Fund Award (Y.H.) from Schizophrenia International Research Society. The funding...
Sources had no further role in study design; in the collection, analysis, and interpretation of data; in the writing report; and in the decision to submit the paper for publication.

Institutional Review Board Statement: The study was conducted according to the guidelines of the Declaration of Helsinki, and approved by the Institutional Review Board of Kyushu University Institutional Review Board for Clinical Trials (approval number: 26018, date: 12 June 2014).

Informed Consent Statement: Informed consent was obtained from all subjects involved in the study.

Data Availability Statement: The data that support the findings of this study are available on request from the corresponding authors upon reasonable request. The data are not publicly available due to privacy or ethical restrictions.

Acknowledgments: We thank the participants who took part in the study. We thank Sarina Iwabuchi, for editing a draft of this manuscript.

Conflicts of Interest: The authors declare no conflict of interest.

References
1. Anttila, V.; Bulik-Sullivan, B.; Finucane, H.K.; Walters, R.K.; Bras, J.; Duncan, L.; Neale, B.M. Analysis of shared heritability in common disorders of the brain. Science 2018, 360, eaap8757. [PubMed]
2. Lee, S.H.; Ripke, S.; Neale, B.M.; Faraone, S.V.; Purcell, S.M.; Perlis, R.H.; Wray, N.R. Genetic relationship between five psychiatric disorders estimated from genome-wide SNPs. Nat. Genet. 2013, 45, 984–994. [PubMed]
3. Ivleva, E.I.; Clementz, B.A.; Dutcher, A.M.; Arnold, S.J.; Jon-Slaughter, H.; Aslan, S.; Tamminga, C.A. Brain structure biomarkers in the psychosis biotypes: Findings from the bipolar-schizophrenia network for intermediate phenotypes. Biol. Psychiatry 2017, 82, 26–39. [CrossRef]
4. Hirano, S.; Hirano, Y.; Maekawa, T.; Obayashi, C.; Oribe, N.; Kuroki, T.; Kanba, S.; Onitsuka, T. Abnormal neural oscillatory activity to speech sounds in schizophrenia: A magnetoencephalography study. J. Neurosci. 2008, 28, 4897–4903. [CrossRef] [PubMed]
5. Oribe, N.; Onitsuka, T.; Hirano, S.; Hirano, Y.; Maekawa, T.; Obayashi, C.; Ueno, T.; Kasai, K.; Kanba, S. Differentiation between bipolar disorder and schizophrenia revealed by neural oscillation to speech sounds: An MEG study. Bipolar Disord. 2010, 12, 804–812. [CrossRef]
6. Hirano, Y.; Hirano, S.; Maekawa, T.; Obayashi, C.; Oribe, N.; Monji, A.; Kasai, K.; Kanba, S.; Onitsuka, T. Auditory gating deficit to human voices in schizophrenia: A MEG study. Schizophr. Res. 2010, 117, 61–67. [CrossRef] [PubMed]
7. Tsuchimoto, R.; Kanba, S.; Hirano, S.; Oribe, N.; Ueno, T.; Hirano, Y.; Nakamura, I.; Oda, Y.; Miura, T.; Onitsuka, T. Reduced high and low frequency gamma synchronization in patients with chronic schizophrenia. Schizophr. Res. 2011, 133, 99–105. [CrossRef]
8. Isomura, S.; Onitsuka, T.; Tsuchimoto, R.; Nakamura, I.; Hirano, S.; Oda, Y.; Oribe, N.; Hirano, Y.; Ueno, T.; Kanba, S. Differentiation between major depressive disorder and bipolar disorder by auditory steady-state responses. J. Affect. Disord. 2016, 190, 800–806. [CrossRef]
9. Oribe, N.; Hirano, Y.; Del Re, E.; Mesholam-Gately, R.I.; Woodberry, K.A.; Ueno, T.; Kanba, S.; Onitsuka, T.; Shenton, M.E.; Spencer, K.M.; et al. Longitudinal evaluation of visual P300 amplitude in clinical high-risk subjects: An event-related potential study. Psychiatry Clin. Neurosci. 2020, 74, 527–534. [CrossRef]
10. Wolf, A.; Ueda, K.; Hirano, Y. Recent Updates of Eye-movement Abnormalities in Patients with Schizophrenia: A Scoping Review. Psychiatry Clin. Neurosci. 2020. [CrossRef]
11. Kato, T. Current understanding of bipolar disorder: Toward integration of biological basis and treatment strategies. Psychiatry Clin. Neurosci. 2019, 73, 526–540. [CrossRef] [PubMed]
12. Brugger, S.P.; Howes, O.D. Group heterogeneity and homogeneity of regional brain structure in schizophrenia: A comparative MRI study. JAMA Psychiatry 2017, 74, 1104–1111. [CrossRef]
13. Ellison-Wright, I.; Bullmore, E. Anatomy of bipolar disorder and schizophrenia: A meta-analysis. Schizophr. Res. 2010, 117, 1–12. [CrossRef]
14. Madeira, N.; Duarte, J.V.; Martins, R.; Costa, G.N.; Macedo, A.; Castelo-Branco, M. Morphometry and gyriﬁcation in bipolar disorder and schizophrenia: A comparative MRI study. NeuroImage 2020, 26, 10220. [CrossRef] [PubMed]
15. Bora, E.; Fornito, A.; Yucel, M.; Pantelis, C. The effects of gender on grey matter abnormalities in major psychoses: A comparative voxelwise meta-analysis of schizophrenia and bipolar disorder. Psychiatr. Med. 2012, 42, 295–307. [CrossRef]
16. Giulani, N.R.; Calhoun, V.D.; Pearlson, G.D.; Francis, A.; Buchanan, R.W. Voxel-based morphometry versus region of interest: A comparison of two methods for analyzing gray matter differences in schizophrenia. Schizophr. Res. 2005, 74, 135–147. [CrossRef]
17. Botvinik-Nezer, R.; Holzmeister, F.; Camerer, C.F.; Dreber, A.; Huber, J.; Johannesson, M.; Kirchler, M.; Iwanir, R.; Mumford, J.A.; Adcock, R.A.; et al. Variability in the analysis of a single neuroimaging dataset by many teams. Nature 2020, 582, 84–88. [CrossRef]
18. Kasai, K.; Shenton, M.E.; Salisbury, D.F.; Hirayasu, Y.; Onitsuka, T.; Spencer, M.H.; Yurgelun-Todd, D.A.; Kikinis, R.; Jolesz, F.A.; McCarley, R.W. Progressive decrease of left Heschl gyrus and planum temporale gray matter volume in first-episode schizophrenia: A longitudinal magnetic resonance imaging study. Arch. Gen. Psychiatry 2003, 60, 766–775. [CrossRef]
19. Reite, M.; Teale, P.; Rojas, D.C.; Reite, E.; Asherin, R.; Hernandez, O. MEG auditory evoked fields suggest altered structural/functional asymmetry in primary but not secondary auditory cortex in bipolar disorder. Bipolar Disord. 2009, 11, 371–381. [CrossRef]

20. Frey, A.; Nery, M.; Quevedo, J.; Soares, K. The role of hippocampus in the pathophysiology of bipolar disorder. Behav. Pharmacol. 2007, 18, 419–430. [CrossRef]

21. Lieberman, J.A.; Girgis, R.R.; Brucato, G.; Moore, H.; Provenzano, F.; Kegeles, L.; Javitt, D.; Kantrowitz, J.; Wall, M.M.; Corcoran, C.M.; et al. Hippocampal dysfunction in the pathophysiology of schizophrenia: A selective review and hypothesis for early detection and intervention. Mol. Psychiatry 2018, 23, 1764–1772. [CrossRef]

22. Eichenbaum, H. Hippocampus: Cognitive processes and neural representations that underlie declarative memory. Neuron 2004, 44, 109–120. [CrossRef]

23. Haukvik, U.K.; Tamnes, C.K.; Soderman, E.; Agartz, I. Neuroimaging hippocampal subfields in schizophrenia and bipolar disorder: A systematic review and meta-analysis. J. Psychiatr. Res. 2018, 104, 217–226. [CrossRef]

24. Morris, J.S.; Ohman, A.; Dolan, R.J. Conscious and unconscious emotional learning in the human amygdala. Nature 1998, 393, 467–470. [CrossRef]

25. LaBar, K.S.; Gatenby, J.C.; Gore, J.C.; LeDoux, J.E.; Phelps, E.A. Human amygdala activation during conditioned fear acquisition and extinction: A mixed-trial fMRI study. Neuron 1998, 20, 937–945. [CrossRef]

26. Ho, N.F.; Li Hui Chong, P.; Lee, D.R.; Chew, Q.H.; Chen, G.; Sim, K. The Amygdala in Schizophrenia and Bipolar Disorder: A Synthesis of Structural MRI, Diffusion Tensor Imaging, and Resting-State Functional Connectivity Findings. Harv. Rev. Psychiatry 2019, 27, 150–164. [CrossRef]

27. Hiba, D.P.; Westley, L.T.; van Erp, T.G.; Rasmussen, J.; Leonardo, C.D.; Faskowitz, J.; Haukvik, U.K.; Hartberg, C.B.; Doan, N.T.; Agartz, I.; et al. Subcortical volumetric abnormalities in bipolar disorder. Mol. Psychiatry 2016, 21, 1710–1716. [CrossRef]

28. Da Costa, S.; van der Zwaag, W.; Marques, J.P.; Frackowiak, R.S.; Clarke, S.; Saenz, M. Human primary auditory cortex follows the shape of Heschl’s gyrus. J. Neurosci. 2011, 31, 14067–14075. [CrossRef]

29. Nakada, T.; Fuji, Y.; Yoneoka, Y.; Kwee, I.L. Planum temporale: Where spoken and written language meet. Eur. Neurol. 2001, 46, 121–125. [CrossRef] [PubMed]

30. Hirayasu, Y.; McCarley, R.W.; Salisbury, D.F.; Tanaka, S.; Kwon, J.S.; Frumin, M.; Snyderman, D.; Yurgelun-Todd, D.; Kikinis, R.; Jolesz, F.A.; et al. Planum temporale and Heschl gyrus volume reduction in schizophrenia: A magnetic resonance imaging study of first-episode patients. Arch. Gen. Psychiatry 2000, 57, 692–699. [CrossRef] [PubMed]

31. Javitt, D.C.; Sweet, R.A. Auditory dysfunction in schizophrenia: Integrating clinical and basic features. Nat. Rev. Neurosci. 2015, 16, 535–550. [CrossRef]

32. Hirano, S.; Spencer, K.M.; Onitsuka, T.; Hirano, Y. Language-Related Neurophysiological Deficits in Schizophrenia. Clin. EEG Neurosci. 2020, 51, 222–233. [CrossRef]

33. Hirano, Y.; Oribe, N.; Kanba, S.; Onitsuka, T.; Nestor, P.G.; Spencer, K.M. Spontaneous Gamma Activity in Schizophrenia. JAMA Psychiatry 2015, 72, 813–821. [CrossRef]

34. Hirano, Y.; Oribe, N.; Onitsuka, T.; Kanba, S.; Nestor, P.G.; Hosokawa, T.; Levin, M.; Shenton, M.E.; McCarley, R.W.; Spencer, K.M. Auditory Cortex Volume and Gamma Oscillation Abnormalities in Schizophrenia. Clin. EEG Neurosci. 2020, 51, 244–251. [CrossRef] [PubMed]

35. Takahashi, T.; Malhi, G.S.; Wood, S.J.; Yucel, M.; Waltersfang, M.; Kawasaki, Y.; Suzuki, M.; Pantelis, C. Gray matter reduction of the superior temporal gyrus in patients with established bipolar I disorder. J. Affect. Disord. 2010, 123, 276–282. [CrossRef]

36. Oldfield, R.C. The assessment and analysis of handedness: The Edinburgh inventory. Neuropsychologia 1971, 9, 97–113. [CrossRef]

37. Hollingshead, A. Two Factor Index of Social Position; Yale University Press: New Haven, CT, USA, 1965.

38. Kay, S.R.; Fiszbein, A.; Opler, L.A. The positive and negative syndrome scale (PANSS) for schizophrenia. Schizophr. Bull. 1987, 13, 261–276. [CrossRef] [PubMed]

39. Young, R.C.; Biggs, J.T.; Ziegler, V.E.; Meyer, D.A. A rating scale for mania: Reliability, validity and sensitivity. Br. J. Psychiatry 1978, 133, 429–435. [CrossRef]

40. Williams, J.B. A structured interview guide for the Hamilton Depression Rating Scale. Arch. Gen. Psychiatry 1988, 45, 742–747. [CrossRef]

41. Ashburner, J.; Friston, K.J. Voxel-based morphometry—the methods. Neuroimage 2000, 11, 805–821. [CrossRef]

42. Kwon, J.S.; McCarley, R.W.; Hirayasu, Y.; Anderson, J.E.; Fischer, I.A.; Kikinis, R.; Jolesz, F.A.; Shenton, M.E. Left planum temporale volume reduction in schizophrenia. Arch. Gen. Psychiatry 1999, 56, 142–148. [CrossRef]

43. Bart, P.E.; Pearlson, G.D.; Brill, L.B.; Royall, R.; McGilchrist, I.K.; Pulver, A.E.; Powers, R.E.; Casanova, M.F.; Tien, A.Y.; Frangou, S.; et al. Planum temporale asymmetry reversal in schizophrenia: Replication and relationship to gray matter abnormalities. Am. J. Psychiatry 1997, 154, 661–667. [CrossRef]

44. Shapleske, J.; Rossell, S.L.; Simmons, A.; David, A.S.; Woodruff, P.W. Are auditory hallucinations the consequence of abnormal cerebral lateralization? A morphometric MRI study of the Sylvian fissure and planum temporale. Biol. Psychiatry 2001, 49, 685–693. [CrossRef]

45. Steinmetz, H.; Rademacher, J.; Huang, Y.X.; Heft, H.; Zilles, K.; Thron, A.; Freund, H.J. Cerebral asymmetry: MR planimetry of the human planum temporale. J. Comput. Assist. Tomogr. 1989, 13, 996–1005. [CrossRef]

46. Bart, P.E.; Petty, R.G.; McGilchrist, I.; Lewis, R.W.; Jeram, M.; Casanova, M.F.; Powers, R.E.; Brill, L.B.; Pearlson, G.D. Asymmetry of the planum temporale: Methodological considerations and clinical associations. Psychiatry Res. 1995, 61, 137–150. [CrossRef]
47. Honeycutt, N.A.; Musick, A.; Barta, P.E.; Pearlson, G.D. Measurement of the planum temporale (PT) on magnetic resonance imaging scans: Temporal PT alone and with parietal extension. Psychiatry Res. 2000, 98, 103–116. [CrossRef]

48. van Erp, T.G.; Hibar, D.P.; Rasmussen, J.M.; Glahn, D.C.; Pearlson, G.D.; Andreassen, O.A.; Agartz, I.; Westlye, L.T.; Haukvik, U.K.; Dale, A.M.; et al. Subcortical brain volume abnormalities in 2028 individuals with schizophrenia and 2540 healthy controls via the ENIGMA consortium. Mol. Psychiatry 2016, 21, 547–553. [CrossRef] [PubMed]

49. Seidman, L.J.; Faraone, S.V.; Goldstein, J.M.; Kremen, W.S.; Horton, N.J.; Makris, N.; Toomey, R.; Kennedy, D.; Caviness, V.S.; Tsuang, M.T. Left hippocampal volume as a vulnerability indicator for schizophrenia: A magnetic resonance imaging morphometric study of nonpsychotic first-degree relatives. Arch. Gen. Psychiatry 2002, 59, 839–849. [CrossRef]

50. Bryant, N.L.; Buchanan, R.W.; Vladar, K.; Breier, A.; Rothman, M. Gender differences in temporal lobe structures of patients with schizophrenia: A volumetric MRI study. Am. J. Psychiatry 1999, 156, 603–609. [PubMed]

51. Karantonis, J.A.; Rossell, S.L.; Carruthers, S.P.; Sumner, P.; Hughes, M.; Green, M.J.; Pantelis, C.; Burdick, K.E.; Cropley, V.; Van Rheenen, T.E. Brain morphology does not clearly map to cognition in individuals on the bipolar-schizophrenia-spectrum: A cross-diagnostic study of cognitive subgroups. J. Affect. Disord. 2021, 281, 776–785. [CrossRef] [PubMed]

52. Dezhina, Z.; Ranlund, S.; Kyriakopoulos, M.; Williams, S.C.R.; Dima, D. A systematic review of associations between functional MRI activity and polygenic risk for schizophrenia and bipolar disorder. Brain Imaging Behav. 2019, 13, 862–877. [CrossRef]

53. Goodkind, M.; Eckhoff, S.B.; Oathes, D.J.; Jiang, Y.; Chang, A.; Jones-Hagata, L.B.; Ortega, B.N.; Zaiko, Y.V.; Roach, E.L.; Korgaonkar, M.S.; et al. Identification of a common neurobiological substrate for mental illness. JAMA Psychiatry 2015, 72, 305–315. [CrossRef] [PubMed]

54. Le, B.D.; Stein, J.L. Mapping causal pathways from genetics to neuropsychiatric disorders using genome-wide imaging genetics: Current status and future directions. Psychiatry Clin. Neurosci. 2019, 73, 357–369. [CrossRef] [PubMed]

55. Appaji, A.; Nagendra, B.; Chako, D.M.; Padmanabha, A.; Jacob, A.; Hiremath, C.V.; Varambally, S.; Kesavan, M.; Venkatasubramanian, G.; Rao, S.V.; et al. Examination of retinal vascular trajectory in schizophrenia and bipolar disorder. Psychiatry Clin. Neurosci. 2019, 73, 738–744. [CrossRef]

56. Omori, W.; Itagaki, K.; Kajitani, N.; Abe, H.; Okada-Tsuchioka, M.; Okamoto, Y.; Takebayashi, M. Shared preventive factors associated with relapse after a response to electroconvulsive therapy in four major psychiatric disorders. Psychiatry Clin. Neurosci. 2019, 73, 494–500. [CrossRef] [PubMed]

57. Kubo, K. Increased densities of white matter neurons as a cross-disease feature of neuropsychiatric disorders. Psychiatry Clin. Neurosci. 2020, 74, 166–175. [CrossRef] [PubMed]

58. Smucny, J.; Cassady, J.; Carter, C.S. Comparing machine and deep learning-based algorithms for prediction of clinical improvement in depression using MRI. Hum. Brain Mapp. 2021, 42, 1197–1205. [CrossRef]

59. Knöchel, C.; Stäblein, M.; Storchak, H.; Reinke, B.; Jurcoane, A.; Prvulovic, D.; Linden, D.E.; van de Ven, V.; Ghinea, D.; Wenzel, S.; et al. Multimodal assessments of the hippocampal formation in schizophrenia and bipolar disorder: Evidences from neurobehavioral measures and functional and structural MRI. NeuroImage 2014, 6, 134–144. [CrossRef]

60. Hwang, H.C.; Kim, S.M.; Han, D.H. Different facial recognition patterns in schizophrenia and bipolar disorder assessed using a computerized emotional perception test and fMRI. J. Affect. Disord. 2021, 279, 83–88. [CrossRef]

61. Thompson, P.M.; Jahanshad, N.; Ching, C.R.; Salminen, L.E.; Thomopoulos, S.I.; Bright, J.; Baune, B.T.; Bertol, S.; Bralten, J.; Bruin, W.B.; et al. ENIGMA and global neuroscience: A decade of large-scale studies of the brain in health and disease across more than 40 countries. Transl. Psychiatry 2020, 10, 100. [CrossRef]

62. Nogovitsyn, N.; Souza, R.; Muller, M.; Srajer, A.; Metzak, P.D.; Hassel, S.; Ismail, Z.; Protzner, A.; Bray, S.L.; Lebel, C.; et al. Aberrant limbic brain structures in young individuals at risk for mental illness. Psychiatry Clin. Neurosci. 2020, 74, 294–302. [CrossRef] [PubMed]

63. Nemoto, K.; Shimokawa, T.; Fukunaga, M.; Yamashita, F.; Tamura, M.; Yamamori, H.; Yasuda, Y.; Azechi, H.; Kudo, N.; Watanabe, Y.; et al. Differentiation of schizophrenia using structural MRI with consideration of scanner differences: A real-world multisite cross-diagnostic study of cognitive subgroups. J. Affect. Disord. 2020, 281, 17–26. [CrossRef] [PubMed]

64. Roes, M.M.; Yin, J.; Taylor, L.; Metzak, P.D.; Lavigne, K.M.; Chinchani, A.; Tipper, C.M.; Woodward, T.S. Hallucination-Specific neurobehavioral measures and functional and structural MRI. NeuroImage 2014, 72, 547–553. [CrossRef] [PubMed]

65. van Dellen, E.; Börner, C.; Schutte, M.; van Montfort, S.; Abramovic, L.; Boks, M.P.; Cahn, W.; van Haren, N.; Mandl, R.; Stam, C.J.; et al. Functional brain networks in the schizophrenia spectrum and bipolar disorder with psychosis. NPJ Schizophr. Res. 2020, 6, 1–12. [CrossRef] [PubMed]

66. Reddy-Thootkur, M.; Kraguljac, N.V.; Lahey, A.C. The role of glutamate and GABA in cognitive dysfunction in schizophrenia and mood disorders—a systematic review of magnetic resonance spectroscopy studies. Schizophr. Res. 2020, 214, 862–877. [CrossRef]

67. Bustillo, J.R.; Jones, T.; Qualls, C.; Chavez, L.; Lin, D.; Lenroot, R.K.; Gasparovic, C. Proton magnetic resonance spectroscopic imaging of gray and white matter in bipolar-I and schizophrenia. J. Affect. Disord. 2019, 246, 745–753. [CrossRef] [PubMed]

68. Das, T.K.; Javadzadeh, A.; Dey, A.; Sabesan, P.; Theberge, J.; Radua, J.; Panagiotopoulou, L.; Barzdukas, A.; Fantini, I.; et al. Antioxidant defense in schizophrenia and bipolar disorder: A meta-analysis of MRS studies of anterior cingulate glutathione. Prog. Neuropsychopharmacol. Biol. Psychiatry 2019, 91, 94–102. [CrossRef] [PubMed]

69. Dogan, A.E.; Yuksel, C.; Du, F.; Chouinard, V.A.; Ongür, D. Brain lactate and pH in schizophrenia and bipolar disorder: A systematic review of findings from magnetic resonance studies. Neuropsychopharmacology 2018, 43, 1681–1690. [CrossRef]
