ORIGINAL RESEARCH

Mitochondrial Damage of Lymphocytes in Patients with Acute Relapse of Schizophrenia: A Correlational Study with Efficacy and Clinical Symptoms

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Objective: Accumulating evidence has demonstrated that schizophrenia is associated with mitochondrial and immune abnormalities. In this pilot case-control study, we investigated the level of mitochondrial impairment in lymphocytes in patients with acute relapse of schizophrenia and explored the correlation between the level of mitochondrial damage and symptoms or treatment response.

Methods: Lymphocytic mitochondrial damage was detected using mitochondrial fluorescence staining and flow cytometry in 37 patients (at admission and discharge) and 24 controls. Clinical symptoms were assessed using the Positive and Negative Syndrome Scale (PANSS) and Clinical Global Impression Scale (CGI-S).

Results: The levels of mitochondrial damage in CD3+ T, CD4+ T, and CD8+ T lymphocytes of the patients with schizophrenia at admission were significantly higher than those of the controls (p<0.05) and did not return to normal at discharge (p>0.05). The mitochondrial damage of T cells significantly improved at discharge for responsive patients only, as compared with that at admission (P<0.05). However, no significant difference was found in mitochondrial damage in CD19+ B cells between patients and healthy controls, or between admission and discharge (p>0.05). Furthermore, the reduction in mitochondrial damage of CD3, CD4, and CD8 lymphocytes was positively correlated with the reduction of the score of the PANSS positive scale at discharge (p<0.05), while no significant correlation was found between the level of mitochondrial damage in lymphocytes and the scores of PANSS and CGI-S.

Conclusion: Acute relapse of schizophrenia might be associated with higher levels of mitochondrial damage in peripheral blood T lymphocytes. The degree of recovery of mitochondrial impairment in the T cells may be used as a predictor of treatment response in schizophrenia. As this is a pilot study, the conclusion still needs further verification in large-scale studies.

Keywords: immune metabolism, mitochondrial impairment, lymphocyte, schizophrenia, biological marker

Introduction

Schizophrenia is a serious, complex, chronic, and disabling psychiatric disorder that causes severe cognitive, emotional, and social impairment.¹ Its typical clinical symptoms mainly include paranoid delusions, auditory hallucinations, disorganized behaviors, and disordered thoughts.^{1,2} According to statistics in 2016, approximately 21 million people were living with schizophrenia globally, which brought about an extremely high disease burden and accounted for 13.4 million years lived with disability (YLDS); the figures have been rising continuously as the population ages.³

The precise physiopathology of complicated psychiatric disorders has not been elucidated despite the extensive studies on neurotransmitters, neural connectivity, and synaptic plasticity in this field. A growing body of evidence has shown the role of immune-inflammatory responses or defective mitochondria in psychiatric illnesses.⁴⁻⁸

Studies have consistently demonstrated that schizophrenia is linked to chronic inflammation. Some studies using animal models of schizophrenia indicated immune-inflammatory reactions in the brain of animals.^{9,10} Postmortem and

positron emission tomography (PET) studies also found infiltration of T cells and B cells, as well as microglial activation in several brain regions, suggesting blood–brain barrier impairment and neuroinflammation, which may be involved in the pathophysiology of schizophrenia.^{11,12} Notably, cells of the nervous system and lymphocytes have similar receptors and signal transduction mechanisms, and in psychiatric disorders, the disturbances of transmitters in the central nervous system co-occur with changes in lymphocyte function and metabolism in peripheral blood. Thus, lymphocytes could be a peripheral model for metabolic alterations in the central nervous system of schizophrenia.¹³

Recent studies have linked mitochondrial dysfunction in schizophrenia to the activation of immune inflammation via the release of reactive oxygen species (ROS) and damage-related molecular markers.^{9,14} Mitochondria are complex organelles that respond to external and internal signals by regulating core functions of the cell, such as redox, energy metabolism, apoptosis, and calcium homeostasis. Perturbations such as oxidative stress can lead to misfolding of mitochondrial permeability transition (MPT) pores,^{15,16} which results in non-selective permeability of the inner mitochondrial membrane, membrane potential collapse, mitochondrial swelling, calcium overload, and cytochrome c release, thereby triggering the activation of caspase and apoptosis.^{17–19} MPT plays a key role in inducing ROS, Ca2+ toxicity, cell necrosis, and apoptosis.^{15,20} Mitochondria damage can also promote the release of mitochondrial danger-associated molecules (DAMPs) and ROS, which plays a key role in activating and maintaining inflammatory immune responses, and such inflammatory responses can further aggravate the mitochondrial damage.^{15,21,22}

Recent studies have found elevated levels of serum inflammatory factors and oxidative stress markers in patients with schizophrenia, which have important implications for cognitive impairment in patients with psychiatric disorders.^{23,24} In addition to its effects on the brain, inflammation is also believed to enhance the permeability of the blood–brain barrier and accelerate the entry of immune factors into the brain.²⁵ Microglia in the brain can be activated by the peripheral inflammatory immunological response, which causes them to release cytokines that affect synaptic plasticity and neurotransmitters, eventually resulting in cognitive, emotional, and behavioral alterations.²⁶ Other studies have also indicated that mitochondrial dysfunction plays an important role in immune response, and the dysfunction might be one of the causes of schizophrenia,²⁷ which is consistent with the role of mitochondria in neuronal growth, activity, and plasticity.²⁸

Circumstantial evidence also indicates that schizophrenia is associated with mitochondrial and immune abnormalities; however, most of these studies are focused on brain areas. For instance, some postmortem studies revealed that the number of mitochondria and volume of oligodendrocytes in brain regions such as the prefrontal cortex of schizophrenia is significantly reduced.^{29,30} Due to the limited availability of samples of brain areas, the discovery of peripheral blood biomarkers is crucial for the diagnosis and prognosis evaluation of schizophrenia. Thus, the present study aimed to do so by investigating the association between the level of mitochondrial damage in peripheral lymphocytes and schizophrenia.

Recently, a new immunofluorescence technology has emerged to reflect mitochondrial function by measuring singlecell mitochondrial mass (SCMM).^{31,32} Mitochondria are highly dynamic organelles with mitophagy mechanisms, and damaged mitochondria caused by mild stress can be eliminated by mitophagy. Intense stress leads to more MPT in mitochondria, which impairs mitophagy¹⁵ and results in abnormal mitochondria that cannot be eliminated, leading to increased mitochondrial mass.³³ Therefore, the parameter of mitochondrial mass may reflect mitochondrial disturbance.^{31,32} Flow cytometry assay in this study relies on a mitochondrial special dye, which is a green fluorescence that binds specifically to the mitochondria of living cells. The median fluorescence intensity (MFI) reflects the mitochondria mass (MM).^{31,34,35} At the same time, lymphocyte subsets were labeled with specific antibodies of CD3, CD4, CD8, and CD19 for the cell count of each subset. In the present study, the single-cell mitochondrial fluorescence intensity of each cell population, reflecting SCMM, was obtained by using the built-in algorithm based on the MFI and the counts of cell subsets, which was recorded as mitochondrial damage index (MDI).

Materials and Methods

Subjects

Patients who met the following criteria were included: meeting the diagnostic criteria for schizophrenia according to the Diagnostic and Statistical Manual of Mental Disorders Fourth Edition (DSM-IV); in an acute episode of recurrent schizophrenia; being 18–60 years of age; having a Clinical Global Impression Severity score (CGI-S) of \geq 4; having

complete clinical and biological data; having signed the informed consent form. Participants who met any of the following criteria were excluded: with current infections, allergies, autoimmune disorders, or use of anti-inflammatory, antioxidants and antiviral drugs; with a history of substance abuse; with other mental illnesses; with significant cognitive dysfunction; with cancer, hypertension, coronary heart disease, diabetes, or nervous system diseases. Healthy controls were recruited among healthy volunteers with no personal or family history of psychiatric disorders.

This study was conducted in accordance with the Declaration of Helsinki. It was approved by the Medical Research Ethics Committee of the Second Xiangya Hospital of Central South University, China (dated 02.01.2019, approval No. [2019] 2019–004). All the subjects provided written informed consent upon full explanation of the study procedure.

Measurements

All the recruited patients underwent a demographic interview, a systematic assessment of medical history, and an interview based on the Structured Clinical Interview for DSM-IV Axis I Disorders. The severity of psychotic symptoms was evaluated using PANSS and CGI-S. The test of lymphocyte mitochondria was performed only once for the control group, while performed at admission and discharge for the case group.

Venipuncture was performed between 6:00 a.m. and 7:00 a.m.; 3–5 mL of venous blood from each participant was collected using sterile EDTA anticoagulant tubes and stored at room temperature for 24–36 hours. Cells were labeled with a 96-well plate containing a mitochondria-specific dye (UB1024, UBBiotechnology Co., Ltd., Hangzhou, China) that can be activated by a 633-nm laser. The plate was wrapped with aluminum foil to keep it from light; the content was incubated at room temperature for 3–5 minutes and centrifuged for 1 minute at 250×g. Then, immunophenotypic antibodies were used to prepare the "mixed reagent" to label cell populations, including CD3-FITC (UB104411, UBBiotechnology Co., Ltd., Hangzhou, China), CD4-PE-Cy7 (UB105441), CD8-PE (UB106421), CD45-PerCP-Cy5.5 (UB109481), and CD19 (RUO: UBR26-200). Contents in the plate were added with 20 ul of "mixed reagent" and incubated for 15 minutes. Then, 100 µl of anticoagulated human peripheral blood was added and incubated for 15 minutes at room temperature, away from light. Hemolysin (NH Lysis Solution, 10×) 400 µl was processed in the same way.

Finally, flow cytometry (NovoCyte, Agilent Technologies, US) was used to detect the counts of T and B cell subsets (CD3, CD4, CD8, and CD19) in the peripheral blood (shown in Figures 1 and 2) and the fluorescence intensity of mitochondrial staining of each cell population (shown in Figure 3).

Statistical Analysis

The SPSS 25.0 software (IBM Corp) was used for statistical analyses. The results were tested for normality using the Shapiro–Wilk test. Student *t*-test (for normally distributed data) or Mann–Whitney *U*-test (for non-normally distributed data) were used to analyze the differences in continuous variables between patients and controls. A paired *t*-test (for normally distributed data) or Wilcoxon signed-rank test (for non-normally distributed data) was used to compare the mitochondrial damage before and after treatment. The Chi-square test was used to analyze categorical variables. Spearman-Rho correlation test was used to evaluate the possible relationship between the level of lymphocyte damage and the severity of the acute episode of relapsed schizophrenia (reflected by the score of PANSS). For all the tests, $P \leq 0.05$ was considered statistically significant.

Results

Demographic and Clinical Characteristics

The present case–control pilot study was conducted in the Second Xiangya Hospital, Hunan Province, China. A total of 61 participants were enrolled, including 37 patients and 24 healthy controls. The average age of onset in the case group was 20.41 ± 6.68 years, with a disease duration of 80.38 ± 85.95 months. The average length of hospital stay was 24.6 days (± 6.99 days), which was equivalent to the follow-up period. After treatment, the PANSS score of the patients with schizophrenia decreased significantly at discharge (p<0.001). The demographic and clinical characteristics of the patients and controls as well as the psychopathology features of the patients are presented in Table 1.



Figure I Flow cytometry analysis of T cell sub-populations.

Notes: (A) The gate was used to exclude beads and identify WBC cells. (B) The CD45-A and SSC-A plot was used to identify T cells as CD45+ SSC-A-. (C and D) T cell gate defined all T cells as CD3+CD4+, CD3+CD4+, CD3+CD8+, and CD3+CD8-.



Figure 2 Flow cytometry analysis of B cell subpopulations. Notes: (A) The gate was used to identify WBC cells. (B) The CD45-A and SSC-A plot was used to identify B cells as CD45+ SSC-A+. (C) CD19+ cells were distinguished by CD19-A+.

Differences in Mitochondrial Impairment in Lymphocytes Between Patients and Controls

MDI was used to reflect the degree of mitochondrial impairment in lymphocytes, as shown in Table 2. The mitochondrial damage in CD3, CD4, and CD8 T lymphocytes was significantly higher in the case group than in the control group (4.34



Figure 3 Representative histograms of mitochondria staining for each cell population.

Notes: (A–D) Characteristics of mitochondrial fluorescence of CD3+, CD4+, CD8+, and CD19+ cell populations, respectively, using the half-offset histogram of the APC channel. The abscissa Mito-A represents the fluorescence intensity of APC fluorescence of mitochondrial dye. APC, allophycocyanin;

[2.37-7.4] vs 1.76 [0.60-3.60], p<0.001; 8.78 [4.35-17.07] vs 4.48 [1.47-11.08], p=0.016; 11.16 [7.40-18.24] vs 3.43 [1.42-7.82], p<0.001) (Table 2). No significant difference in mitochondrial damage in CD19 B cells was found between patients and healthy controls (70.96 [26.94-122.63] vs 60.61 [18.72-136.40], p=0.626) (Table 2).

Table	Demographic an	d Clinical	Characteristics	of	Patients	and	Controls
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Characteristics	Schizophrenia (n=37)		Healthy Controls (n=24)	Р
Gender				
Male/female ^a	18/19		12/12	0.918
Age, mean±SD ^b	27.00±7.47		30.71±7.39	0.062
Age of onset, mean±SD ^b	20.41±6.68			
Time of disorder progression (Months), mean ± SD ^b	80.38±85.95			
PANSS score, mean±SD ^b	Admission	Discharge		
Positive score	20.24±6.05	11.22±4.44		<0.001
Negative score	29.16±6.11	18.84±6.99		<0.001
General score	49.43±6.57	30.89±6.52		<0.001
Total score	101.05±13.20	61.68±14.24		<0.001
CGI-S, mean ± SD ^b	5.95±0.88	3.27±0.990		<0.001

Notes: ^aChi-squared test. ^bStudent *t*-test, data are presented as mean \pm SD. **Abbreviation**: SD, standard deviation.

	Schizophrenia (n=37)	Controls (n=24)	
	Admission	Discharge	
MDI-CD19+ B cells, MDN (IQR) ^a	70.96 (26.94–122.63)	63.89 (38.56–110.97)	60.61 (18.72–136.40)
MDI-CD3+ T cells, MDN (IQR) ^a	4.34 (2.37–7.42)**	4.06 (2.85-6.04)**	1.76 (0.60–3.60)
MDI-CD4+ T cells, MDN (IQR) ^a	8.78 (4.35–17.07)*	8.65 (5.46-11.05)*	4.48 (1.47–11.08)
MDI-CD8+ T cells, MDN (IQR) ^a	11.16 (7.40–18.24)**	11.9 (7.99–18.68)**	3.43 (1.42–7.82)

 Table 2 Comparison of Parameters Between Patients and Controls

Notes: ^aMann–Whitney *U*-test or Wilcoxon signed-rank test; data are presented using median (interquartile range). *Compared with healthy controls, p < 0.05. **Compared with healthy controls, p < 0.01. No statistically significant differences were found in patients with schizophrenia between admission and discharge.

Abbreviations: IQR, interquartile range; MDN, median.

The Level of Mitochondrial Damage Was Mitigated in Responsive Patients

There were no significant differences in the level of mitochondrial damage between admission and discharge for the overall patient group, and both levels were higher than that of the controls (Table 2). However, when patients were divided into a responsive group (PANSS score reduction rate \geq 50%) and an unresponsive group (PANSS score reduction rate \leq 50%), which were matched for medication use based on the chlorpromazine-equivalent doses of their antipsychotics.³⁶ The result showed that the levels of mitochondrial damage in T cells, including CD3, CD4, and CD8 cells, were considerably lower at discharge than that at admission for the responsive group only (p<0.05) (Table 3). Furthermore, no significant change in the MDI level in CD19+ cells was noted before and after treatment, and no significant difference was found regarding CD19+ between patients and controls (Tables 2 and 3).

Associations Between the Severity of Symptoms and the Degree of Lymphocyte Mitochondrial Impairments

Spearman's rank correlation coefficient was used to analyze the correlation between the PANSS score and the level of mitochondrial damage, as the data were not normally distributed. No correlation of the degree of mitochondrial impairment was found with the total, positive, negative, and general scores of PANSS, or the GGI-S score (Table 4). However, the result showed that, after treatment, the improvement of mitochondrial damage in CD3, CD4, and CD8 lymphocytes were positively correlated with the reduction of PANSS scores, especially the positive score (Table 5). In addition, no significant association was found between the age of onset or disease duration and the mitochondrial damage (all P>0.05).

Discussion

To our knowledge, this pilot study was the first to explore the association of the severity of psychotic symptoms and patient response with the level of mitochondrial damage in lymphocytes among patients in an acute episode of recurrent

	The Responsive	e Group (n=20)	The Unresponsive Group (n=17)		
	Admission	Discharge	Admission	Discharge	
Dose of antipsychotics (mg/day) ^a	700.85±	±167.88	654.35±216.88		
MDI-CD19+ B cells (IQR) ^b MDI-CD3+ T cells (IQR) ^b MDI-CD4+ T cells (IQR) ^b MDI-CD8+ T cells (IQR) ^b	75.24 (35.57–139.67) 64.35 (49.61–11 5.96 (3.07–8.37) 3.84 (2.44–6.2 12.80 (5.26–18.78) 8.93 (5.55–11.8 14.38 (9.44–23.77) 10.24 (5.02–16.		63.35 (22.73–89.39) 3.78 (2.06–5.25) 6.24 (3.38–9.35) 10.06 (5.98–13.85)	59.12 (23.84–96.04) 4.41 (3.48–6.07) 7.94 (5.58–11.05) 13.04 (8.90–20.83)	

Table 3 Indicators of Responsive and Unresponsive Patients with Schizophrenia at Admission and Discharge

Notes: ^aChlorpromazine equivalents of antipsychotics, expressed as mean \pm standard deviation. ^bMann–Whitney *U*-test or Wilcoxon signed-rank test; data are presented using median (interquartile range). *p <0.05, admission vs discharge for the responsive group. No statistically significant difference was found between admission and discharge for the unresponsive group.

PANSS Scores		MDI-CD19+ B Cells	MDI-CD3+ T Cells	MDI-CD4+ T Cells	MDI-CD8+ T Cells
PANSS total score	r	-0.038	0.03	0.005	0.037
Positive score	r	-0.222	0.075	0.087	0.048
Negative score	r	0.072	0.017	-0.023	0.059
General score	r	-0.13	-0.018	-0.023	-0.039
CGI-S score	r	-0.085	0.087	0.085	0.065

Table 4 Correlations Between PANSS Scores and Mitochondrial Damage in Lymphocytes

Notes: Results were analyzed using the Spearman correlation test. *r*: Spearman correlation coefficient. No statistical difference was found in correlations presented in this table.

Table 5 Correlation Between the Reduction of PANSS Scores and Improvement of Lymphocyte Mitochondrial Impairment

 After Treatment

Clinical Symptoms		△MDI -CD19+ B Cells	△MDI -CD3+ T Cells	∆MDI-CD4+ T Cells	Δ MDI-CD8+ T Cells
Δ PANSS total score	r	0.105	0.333*	0.308	0.428**
Δ Positive score	r	0.21	0.382*	0.358*	0.461**
Δ Negative score	r	-0.05	0.14	0.202	0.183
Δ General score	r	0.035	0.255	0.221	0.323

Notes: Δ: changes in MDI or PANSS scores between admission and discharge. *p <0.05, **p <0.01. r: Spearman correlation coefficient.

schizophrenia. The results demonstrated that (1) mitochondrial damage in CD3+, CD4+, and CD8+ T lymphocytes was significantly higher in patients in an acute episode of recurrent schizophrenia, as compared to controls; (2) the levels of mitochondrial damage in CD3+, CD4+, and CD8+ T cells substantially improved in responsive patients only, which may partly reflect the therapeutic effect of schizophrenia; (3) the level of mitochondrial damage in CD19+ B cells did not change significantly through admission and discharge, as compared with the control group; (4) after treatment, alterations in the level of mitochondrial impairment in CD3, CD4, and CD8 lymphocytes were associated with the reduction of the PANSS positive score.

The most notable finding of this study was that the level of mitochondrial impairment in T lymphocytes was significantly higher in patients in an acute episode of relapsed schizophrenia, as compared to controls, which was in line with the results of many prior studies. Mitochondrial dysfunction can result in decreased energy generation, which substantially impacts neuronal differentiation and may be involved in abnormal cognitive and behavioral processes.^{37,38} It has been reported that mitochondrial dysfunction might be associated with many neuropsychiatric disorders,³⁹ such as autism,⁴⁰ Parkinson's disease, and Alzheimer's disease.⁴¹ According to prior studies, mitochondrial dysfunction was involved in the majority of mental diseases, and patients with mitochondrial diseases were more likely to present with psychiatric symptoms.^{42,43} Previous studies have suggested that the upstream abnormalities of genes might be associated with damaged mitochondria, leading to manifestations of diseases.⁸ For instance, the abnormality of genes associated with the risk of schizophrenia, such as the nuclear-encoded D-amino-acid oxidase activator (DAOA) and its splice variant LG72 that encodes mitochondrial proteins, might lead to mitochondrial damage.⁴⁴ Consistent with these findings, evidence from patients, cell lines, animal models, and postmortem studies also suggested that mitochondrial dysfunction might be involved in the pathogenesis of schizophrenia.^{38,45,46} A study found that oxygen consumption was reduced in the brains of schizophrenic patients,⁴⁷ and some other studies have shown decreased mitochondrial volume in the oligodendrocytes of the prefrontal lobe as well as a decreased number of mitochondria in the striatum.^{29,48,49}

Similar results have been observed in the peripheral blood of patients with psychiatric disorders.⁴⁶ Patients with schizophrenia were found to have fewer mitochondria in lymphocytes and lower mitochondrial density in the large activated lymphocytes, as compared with healthy controls.^{50,51} It was speculated that the reduction of mitochondria in lymphocytes might be an ultrastructural basis of immune dysfunction in patients with schizophrenia.^{4,45,52} Moreover, several recent studies found that the lymphocytes had lower oxygen consumption (mitochondrial respiration) and generated more mitochondrial oxidative stress-related products in patients with schizophrenia, as compared with healthy controls,^{38,53,54} implying that

patients with schizophrenia might have mitochondrial dysfunction in lymphocytes. Animal models of schizophrenia also showed mitochondrial impairment and up-regulation of immune-related genes, and the mitochondrial dysfunction was alleviated after treatment with clozapine.⁵⁵ The above findings are in line with the results of the present study, which showed mitochondrial damage in T lymphocytes, indicating that mitochondrial damage might be involved in the pathogenesis of schizophrenia. The present study found no mitochondrial damage in B lymphocytes, and a recent study also reported mitochondrial damage in T cells only in patients with schizophrenia.⁵⁶ A possible explanation is that cellular immunity may be more involved in the pathological process of schizophrenia than humoral immunity. To date, there is still insufficient evidence of mitochondrial damage in B lymphocytes in patients with schizophrenia. A meta-analysis suggested no significant difference in the number of B lymphocytes between patients with schizophrenia and healthy controls or before and after treatment, which was in line with the present study.⁵⁷ Conversely, some studies indicate that B cells are involved in the pathological process of schizophrenia. Set indicate that B cells are involved in the pathological process of schizophrenia.⁵⁸ The disparities might be attributed to the differences in medication use, disease duration, and other factors across different studies. Further studies on the role of B cells and their mitochondrial functions in patients with schizophrenia are warranted.

Mitochondrial dysfunction contributes to the development of inflammatory immunity, and inflammation and ROS can further promote mitochondrial damage.⁵⁹ Drugs that inhibit immune inflammation or oxidative stress may help stabilize mitochondrial function and reduce mitochondrial damage; thus, mitochondria might also be a therapeutic target for schizophrenia. Previous studies have shown that some psychotropic drugs have antioxidant components, such as N-acetylcysteine and coenzyme Q10, which can improve mitochondrial activity.⁶⁰ Several studies indicated that some antipsychotic drugs, such as olanzapine and clozapine, could protect mitochondrial activity through their anti-inflammatory and antioxidant effects or regulation of mitochondrial gene expression.^{53,60,61} However, some typical antipsychotics could negatively impact mitochondria.⁶⁰ These findings suggested that mitochondria might be a predictor of patient response and a target of treatment for schizophrenia.

In this pilot study, we found that patients who were responsive to treatment showed a considerable drop in the level of mitochondrial damage in T cells. As indicated by Bar-Yosef et al, patient response to treatment was associated with mitochondrial parameters, approximately half of the mitochondrial parameters (respiratory parameters) were improved in patients with schizophrenia who were responsive, but these parameters returned to normal only in those who received long-term treatment.⁶² Our results also indicated that parameters related to mitochondrial impairment in lymphocytes might be predictors of patient response. Nevertheless, no significant improvement was found in the mitochondrial damage in CD3, CD4, and CD8 lymphocytes after treatment, which might be due to the short follow-up duration (about 24 days) in this study.

Mitochondria are a major source of ROS, and their damage causes the increase of free radicals, resulting in cell dysfunction and neuronal loss.⁶³ Sultana et al demonstrated that the level of ROS produced by mitochondria in peripheral lymphocytes elevated in patients with Alzheimer's Disease, and the level was inversely correlated with the patients' cognitive level, ^{63,64} indicating that the level of ROS might reflect the brain impairment in Alzheimer's Disease and could be a potential biomarker for diagnosis and prognosis.⁶³ Furthermore, many studies suggested that immunological dysfunction and inflammation might contribute to the negative, cognitive, and positive symptoms of schizophrenia.⁷ In the present study, we attempted to explore the relationship between psychiatric symptoms and mitochondrial damage, but no correlation was found between the degree of mitochondrial damage before and after treatment and the positive, negative, or general symptoms of schizophrenia. This result was inconsistent with the findings of some previous studies. A recent study showed a linear relationship between mitochondrial damage and PANSS scores,⁵⁶ and Leirer et al found that the expression of mitochondrial genes in the peripheral blood was associated with negative symptoms of schizophrenia.⁶⁵ Some other investigations have revealed that the degree of mitochondrial deficit in the blood was positively correlated with the severity of paranoid symptoms or PANSS scores.^{46,50,56} The above studies have suggested an association between mitochondrial dysfunction and psychiatric symptoms. The disparities between our results and previous findings might be attributed to the insufficient sample size to detect a relationship, short treatment duration, different effects of drugs on mitochondria, or different stages of the patient's disease. Besides, abnormalities in proteins that regulate mitochondrial function were also found in patients with schizophrenia, and such abnormalities might be related to alterations in symptoms and treatment.⁶⁶ To some extent, mitochondrial damage in lymphocytes can also be used as a biomarker of treatment response.⁵⁶ This was supported by our findings, which suggested that the improvement in mitochondrial damage of CD3, CD4, and CD8

lymphocytes were positively correlated with the level of improvement of positive symptoms, but not the negative symptoms. Thus, our results indicated that the improvement of positive symptoms might be more associated with mitochondrial improvement, as compared with negative symptoms, which is generally consistent with the findings that the positive and negative symptoms of schizophrenia have different pathways of development, and that the positive symptoms are more associated with immune pathways.⁶⁵ It was also possible that the sample of our study was too small to detect a relationship between the improvement in negative symptoms and mitochondrial improvement. An open-label experiment showed that the improvement of mitochondrial function was conducive to the recovery of negative symptoms of schizophrenia.⁶⁷ Thus, future works with larger sample sizes, longer follow-up periods, and controlling for the types of drugs are needed to investigate the connection between mitochondrial damage and schizophrenia, to identify potential therapeutic targets.

Limitations

Some limitations in this study should be considered. The relatively small sample size might have reduced the statistical power to detect significant relationships. Thus, the negative findings might be interpreted as preliminary results, which need to be validated using larger cohorts. Furthermore, without an extended follow-up period and dynamic monitoring of mitochondrial damage in lymphocytes, our study focused primarily on the changes in the indicators before and after treatment. In addition, different antipsychotics might have varied effects on mitochondria, thus not controlling for the medication types might have led to certain inaccuracies. Future studies are warranted to explore the effect of different types of antipsychotic medications on mitochondria. Despite the above limitations, our pilot study is of great value, as it has provided preliminary data on mitochondrial impairment in lymphocytes in patients with schizophrenia. As the mitochondrial data in this study were obtained using peripheral blood, the mitochondrial damage in lymphocytes might only be an indirect indicator of the pathological changes in the central nervous system. Further studies are needed to identify more representative indicators by using more direct methods, such as in vivo quantitative neuroimaging.

Conclusion

In conclusion, this pilot study indicated that the level of mitochondrial damage in CD3, CD4, and CD8 T lymphocytes in the peripheral blood was significantly more severe in patients in an acute episode of relapsed schizophrenia, as compared to healthy controls. After treatment, the improvement of mitochondrial impairment was significant in responsive patients only and was positively associated with the improvement in positive symptoms. These findings suggested that mitochondrial impairment in lymphocytes might be present in patients with schizophrenia, and the degree of its improvement might be used as a predictor for patient response. However, the findings still need further verification in large-scale studies. Despite the limitations, this pilot study helped improve our understanding of the pathophysiology of schizophrenia as well as the role of mitochondria in the treatment of this disease.

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Disclosure

All authors have no conflicts of interest to declare in this work.

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