Does urinary metabolite signature act as a biomarker of post-stroke depression?

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Background: It is difficult to conduct the precise diagnosis of post-stroke depression (PSD) in clinical practice due to the complex psychopathology of depressive disorder. Several studies showed that gas chromatography–mass spectrometry (GC-MS)-identified urinary metabolite biomarkers could significantly discriminate PSD from stroke survivors.

Methods: A systematic review was performed for the keywords of “urinary metabolite” and “PSD” using Medline, Cochrane Library, Embase, Web of Science, PsycINFO, Wanfang, CNKI, CBM, and VIP database from inception to 31 March 2022.

Results: Four related studies were included in the review. Differential urinary metabolites including lactic acid, palmitic acid, azelaic acid, and tyrosine were identified in all the included studies. As a significant deviation in the metabolite biomarker panel, glyceric acid, azelaic acid, phenylalanine, palmitic acid, pseudouridine, and tyrosine were found in at least 2 included studies, which indicated good potential for the differentiation of PSD.

Conclusion: The systematic review provided evidence that differential urinary metabolites analyzed by the GC-MS-based approach might be used as a biomarker for the diagnosis and prognosis of PSD.

KEYWORDS
metabolomics, urinary metabolite, biomarker, post stroke depression, systematic review

Highlights

- First systematic review was conducted to investigate the association between urinary metabolite and PSD.
- All the included studies identified differential urinary metabolites including lactic acid, azelaic acid, tyrosine, and palmitic acid.
- Azelaic acid, glyceric acid, palmitic acid, phenylalanine, pseudouridine, and tyrosine were found significantly differentiated in the metabolite biomarker panel, indicating good potential for the differentiation of PSD.
- The systematic review provided evidence that differential urinary metabolites analyzed by GC-MS-based might be used as a biomarker for the diagnosis and prognosis of PSD.

Introduction

Stroke is a serious cerebrovascular disease with high disability and mortality. Several complications were found in stroke survivors, including depression (1), anxiety (2), fatigue (3, 4), apathy (5), insomnia (6), psychosis (7), mania (8), dementia (9), cognitive disorder (10), and anosognosia (11). Of these, depression after stroke was studied by most researchers due to its high prevalence rate. It was indicated in a meta-analysis including 61 studies enrolling 25,488 patients that the morbidity of depression in stroke survivors was 31% (12). A number of representative studies (13–23) reported that the incidence rate of post-stroke depression (PSD) is from 11 to 41%, negatively impacting the life quality of patients with stroke and imposing heavy economic burdens on their families. Referring to previous studies, prior history of depression or severe stroke is a significant risk of PSD (24–27), whereas marriage (28), social support (29, 30), and levels of education (31) are ambiguous risk factors. However, the pathophysiological mechanisms of PSD are complex and remain to be clarified, which were found to be related to hypothalamic-pituitary-adrenal (HPA) axis dysfunction (32, 33), increased inflammation (34, 35), changes in neurotransmitters (36), and decreased neurotrophic factors (37–39).

Although the depressive disorder can be diagnosed by classical clinical nosologies (40), the diagnosis fundamentals of depression were under more disagreements (41, 42). Symptoms of depression were found to be a result of phenotypic, biological, genetic heterogeneity, and complex interactions between biological and environmental factors. Therefore, not fully understanding the complex psychopathology of depression leads to an ineffective treatment strategy (43). Fortunately, metabolomics is a quantifiable monitor of biochemical state to inform the molecular mechanisms of disorders. Hence, metabolomics acted as diagnostic, prognostic biomarkers (44) for psychiatric disorders in numerous studies. A urinary metabolomics study indicated that two sets of metabolites that could effectively discriminate ‘moderate’ and ‘severe’ patients with depression from healthy controls (HCs) were identified by nuclear magnetic resonance (NMR)- and gas chromatography–mass spectrometry (GC-MS)-based methods (45). Another study found that several specified metabolites could act as predictors of depression recovery (46, 47). Of note, recent studies (48–51) showed that GC-MS-identified urinary metabolite biomarkers could significantly distinguish PSD from stroke survivors, which could be used as an objective diagnostic tool for PSD.

The systematic review was conducted to clarify the specific role of urinary metabolites in PSD, evaluating whether they may act as a useful biomarker to early identify and diagnose depression after stroke as well as to predict the recovery of PSD.

Methods

Search strategy

This systematic review was conducted in accord with the guidelines recommended by Cochrane Collaboration (52) and the checklist of Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA). Relevant studies were searched by the following electronic databases from inception to 31 March 2022: Medline, Embase, Cochrane Library, Web of Science, PsycINFO database, China National Knowledge Infrastructure (CNKI), Wanfang Database, Chinese Scientific Journal Database (VIP database), and Chinese Biomedical Literature Database (CBM). The following keywords were searched: (“urinary metabolite” [Title/Abstract]) OR (“urinary biomarkers” [Title/Abstract]) OR (“urine metabolomics” [Title/Abstract]) AND (“post stroke depression” [Title/Abstract]) OR (“post stroke depressive disorder”[Title/Abstract]) OR (“post stroke depressive disorder” [Title/Abstract]) OR (“depression after stroke” [Title/Abstract]) OR (“depressive disorder after stroke” [Title/Abstract]).

Study selection

Inclusion criteria were as follows:

- Studies investigating urinary metabolite in patients with PSD;
- Studies assessing the depressive symptoms using diagnostic criteria or rating scales.

Exclusion criteria were as follows:

- Studies investigating other types of biomarkers from patients with PSD;
- Systematic reviews, meta-analyses, narrative reviews, book chapters, letters to the editor, case series, case reports, and animal studies.

Two researchers independently screened the titles and abstracts of searched articles and excluded unrelated studies, followed by the full-text review of the remaining articles by the
same researchers. Studies published in English or Chinese were searched. Disagreements were resolved after a discussion with the third author.

Data extraction

Data were extracted from all the included studies and recorded in an excel spreadsheet, including title, author, study design, participants’ characteristics, assessment of depression, and main findings. Any disagreement was resolved by a discussion with a third researcher.

Quality assessment

Two independent researchers used the modified versions of the Newcastle-Ottawa Quality Assessment Scale to assess the quality of included studies (53). The selection, the main outcomes and comparability of the studies were investigated by the scale. A score of 7 or above indicated a good quality. A score of 5–6 was considered a satisfactory quality. Scores less than 5 suggested unsatisfactory studies. With respect to selection, studies were considered to have a low, medium, or high risk of bias if they scored 3, 1–2, or 0 point, respectively. In terms of comparability, studies were considered to have a low, medium, or high risk of bias if they scored 2, 1, or 0 point, respectively. Referring to the outcome, studies were considered to have a low, medium, or high risk of bias if they scored 3, 2, or 1 point, respectively. The disagreement was resolved by a discussion with a third researcher.

Results

The initial literature search identified 301 results, of which only 4 were included in our analysis according to the inclusion criteria (Figure 1). The main characteristics of the included studies are summarized in Table 1. None of the studies mentioned that the recruited patients received any medication. Participants with alcohol abuse or illicit drug use were excluded in all the included studies.

Table 2 presents the results of the quality assessment using the modified versions of the Newcastle-Ottawa Quality Assessment Scale. All the included studies were of good quality.

Xie et al.’s study (50) included 89 stroke survivors without depression and 92 patients with PSD. A total of 12 differential urinary metabolites were identified between patients with PSD and non-depressed stroke survivors, including hydroxylamine, palmitic acid, lactic acid, azelaic acid, pyroglutamic acid, α-aminobutyric acid, uric acid, and tyrosine. Moreover, seven differential urinary metabolites in the panel were found to effectively disclose the difference between patients with PSD and non-depressed stroke survivors. The panel included hydroxylamine, myristic acid, palmitic acid, lactic acid, glyceric acid, azelaic acid, and tyrosine, demonstrating the differentiation of PSD. Interestingly, this study figured out four metabolic pathways, which included tyrosine, phenylalanine and tryptophan biosynthesis, glycerolipid metabolism, fatty acid biosynthesis, starch, and sucrose metabolism.

Liang et al.’s study (49) included 101 patients with PSD and 83 non-depressed patients after stroke. All the recruited patients were also diagnosed with type 2 diabetes mellitus (T2DM). The research identified 23 differential metabolites to discriminate the two groups. Patients in the PSD group were characterized by higher levels of sucrose, inositol, citric acid, lactic acid, methylsuccinic acid, vanillic acid, sorbitol, 2-methyl-3-hydroxybutyric acid, 3, 4-dihydroxybutyric acid, hydroxylamine, threitol, myristic acid, D-glucose, azelaic acid, palmitic acid, and fructose, along with lower levels of hypoxanthine, tyrosine, aminoethanol, malic acid, pseudouridine, indoxyl sulfate, and n-methylaminoisobutyric acid compared to non-depressed stroke survivors. Most of the differential metabolites were found to change significantly. Furthermore, the galactose metabolism was significantly affected in T2DM patients with PSD. The panel consisting of six differential metabolites including malic acid, pseudouridine, hypoxanthine, fructose, 3, 4-dihydroxybutyric acid, and inositol could effectively distinguish the two groups.

Zhang et al. (51) conducted a clinical study totally recruiting 130 patients with PSD, 128 non-depressed stroke survivors, and 127 HC candidates (28). A total of 17 differential urinary metabolites were identified to discriminate PSD from non-PSD. Higher levels of glycercic acid, azelaic acid, 5-hydroxyhexanoic acid, pseudouridine sucrose, lactic acid, and palmitic acid, along with lower levels of hippuric acid, tyrosine, phenylalanine, 3-hydroxyisobutryric acid, indoxyl sulfate, b-aminobutyric acid, leucine, hypoxanthine, ribose, and pyroglutamic acid were found in the PSD group compared with the non-PSD group. It was found that 6 metabolites (glyceric acid, azelaic acid, 5-hydroxyhexanoic acid, pseudouridine, phenylalanine, and tyrosine) significantly differentiated between PSD and non-PSD. The urinary biomarker panel could distinguish 72 patients with PSD from 146 non-PSD subjects.

Interestingly, Chen et al. (48) conducted a study investigating the elderly population including 124 patients with PSD, 78 HCs, and 122 non-depressed stroke survivors. Compared to elderly non-depressed subjects, the elderly subjects with PSD were characterized by higher levels of azelaic acid, palmitic acid, sucrose, α-aminobutyric acid, glycercic acid, fructose, and lactic acid, with lower levels of sorbitol, indoxyl sulfate, 3-hydroxyisobutyric acid, phenylalanine, tyrosine, and 3-hydroxyphenylacetic acid. Of these, 12 differential urinary metabolites remained significantly...
changed. The panel consisting of 8 differential urinary metabolites was the most significant deviations between elderly patients with PSD and elderly non-depressed subjects, including tyrosine, phenylalanine, 3-hydroxyphenylacetic acid, sucrose, palmitic acid, glycercic acid, α-aminobutyric acid, and azelaic acid. A significantly negative correlation was found between phenylalanine and age. Significantly negative correlations were also found between BMI and five differential urinary metabolites (sorbitol, tyrosine, phenylalanine, 3-hydroxyisobutyric acid, and azelaic acid). Furthermore, Hamilton Depression Rating Scale (HDRS) score and three differential urinary metabolites (3-hydroxyisobutyric acid, phenylalanine, and sucrose) were negatively correlated. In elderly subjects with PSD, three metabolic pathways, namely, phenylalanine metabolism, tyrosine, phenylalanine and tryptophan biosynthesis, and galactose metabolism were affected significantly.

According to all the included studies, the metabolites in the urine samples were profiled by a GC-MS-based metabolomics platform; the identification of differential urinary metabolites was analyzed by corresponding partial least-squares discriminant analysis (PLS-DA) loading plots. In addition, a simplified metabolite biomarker panel consisting of significantly different urinary metabolites was obtained to distinguish subjects with PSD from non-depressed stroke survivors via step-wise logistic-regression analysis. All of these are summarized in Table 3. Azelaic acid, lactic acid, palmitic acid, and tyrosine were identified in all the included studies. Most of the included studies (3 out of 4) identified these differential urinary metabolites including fructose, glycercic acid, indoxyl sulfate, and sucrose. Of all the significant deviations in the metabolite biomarker panel, glycercic acid, azelaic acid, phenylalanine, palmitic acid, tyrosine, and pseudouridine were found in at least 2 included studies, which indicated good potential for the differentiation of PSD. Moreover, of all the metabolic pathways detected, biosynthesis of phenylalanine, tyrosine, and tryptophan was significantly affected in 2 studies included.

**Discussion**

The incidence rates of stroke vary between 10 and 200 per 10,000 individuals between the age of 55–85 years and over 85 years. Referring to the latest statistics of the American Heart Association, 700,000 strokes occur every year in the United States (54). Neuropsychiatric disorders associated with stroke include depression, apathy, anxiety disorder, psychosis, mania, cognitive impairment, anosognosia, and fatigue. It was found in Folstein et al.'s study (55) that the depressive disorder was significantly more common in patients with stroke compared to patients undergoing other physical dysfunctions. With a high prevalence rate, PSD is a potential predictor of poor functional outcomes associated with imitations in daily activities (54), poor rehabilitation outcomes, cognitive problems (57, 58), social isolation, suicidal ideation (59), and higher all-cause mortality (60). However, the diagnosis of PSD is inconsistent and challenging. The neurological symptoms associated with stroke such as aphasia, abulia, or dementia might hinder the detection
**TABLE 1** Characteristics of included studies.

| Author    | Study design          | Sample size (N) | Age (Y) Gender | Stroke diagnosis | Depression assessment | Main findings |
|-----------|-----------------------|-----------------|----------------|-------------------|----------------------|--------------|
|           |                       |                 |                |                   |                      |              |
| Zhang et al. (51) | Cross-Sectional Studies | Training set HC:74, NSS:72, PSD:72 | Training set HC: 61.6 ± 5.8, NSS: 61.3 ± 8.7, PSD: 62.0 ± 6.3 | Training set HC: 32/42, NSS: 39/33, PSD: 31/44 | Previously diagnosed with unclear criteria | DSM-IV, HDRS |
|           |                       | Test set        | Test set       | Test set          |                      |              |
|           |                       | HC:53, NSS:56, PSD:58 | HC: 61.3 ± 6.1, NSS: 62.4 ± 8.0, PSD: 62.1 ± 8.3 | HC: 24/29, NSS: 21/35, PSD: 30/28 |                      |              |
|           |                       |                 |                |                   |                      |              |
| Xie at al. (50) | Cross-sectional studies | Training set NSS:59, PSD:58 | Training set NSS: 52.27 ± 5.25, PSD: 54.27 ± 4.19 | Training set NSS: 28/25, PSD: 25/23, Test set NSS: 16/14, Test set PSD: 19/15 | The criteria confirmed by the fourth national conference on cerebrovascular diseases | HDRS |
|           |                       | Test set        | Test set       | Test set          |                      |              |
|           |                       | NSS:30, PSD:34  | NSS: 51.77 ± 4.49, PSD: 54.14 ± 4.01 | NSS: 17/21, PSD: 19/15 |                      |              |
| Liang et al. (49) | Cross-sectional studies | Training set NSS:45, PSD:55 | Training set NSS: 59.55 ± 9.73, PSD: 60.83 ± 8.27 | Training set NSS: 20/25, PSD: 27/28, Test set NSS: 17/21, Test set PSD: 21/25 | The criteria confirmed by the fourth national conference on cerebrovascular diseases | HDRS |
|           |                       | Test set        | Test set       | Test set          |                      |              |
|           |                       | NSS:38, PSD:46  | NSS: 58.39 ± 9.3, PSD: 61.36 ± 7.87 | NSS: 15/21, PSD: 21/25 |                      |              |
| Chen et al. (48) | Cross-sectional studies | Training set HC:44, NSS:86, PSD:82 | Training set HC: 65.20 ± 3.71, NSS: 66.88 ± 4.84, PSD: 67.22 ± 4.69 | Training set HC: 18/26, NSS: 26/40, PSD: 39/43, Test set HC: 17/17, Test set NSS: 15/21, Test set PSD: 21/21 | The criteria confirmed by the fourth national conference on cerebrovascular diseases | DSM-IV, HDRS |
|           |                       | Test set        | Test set       | Test set          |                      |              |
|           |                       | HC:34, NSS:36, PSD:42 | HC: 65.41 ± 2.74, NSS: 67.19 ± 5.42, PSD: 66.17 ± 4.34 | HC: 17/17, NSS: 15/21, Test set PSD: 21/21 |                      |              |

DSM-IV, Diagnostic and Statistical Manual of Mental Disorders-IV; F, female; HC, healthy controls; HDRS, Hamilton Depression Rating Scale; M, male; N, number; NSS, non-depressed stroke survivors; PSD, post stroke depression; T2DM, type 2 diabetes mellitus; Y, year.
of depressive symptoms (61). Furthermore, if the patient refuses to participate in the therapy, it is not easy to early detect the signs of PSD.

Precision medicine is a promising strategy to overcome obstacles in the detection and diagnosis of psychiatric disorders. The rapid development of biotechnologies has been able to realize the target of precision medicine (62). As one of the high-throughput biotechnologies, metabolomics could play important roles in health and disorders (63). The data’s accuracy and comprehensive dimensionality are responsible for the realization of precision psychiatry. Analytical techniques for metabolomics include MS, NMR, and electrochemical detection. NMR is good at identifying novel metabolite structures. More sensitive than NMR, MS-based methods have an advantage in mass analysis capabilities. GC or liquid chromatography (LC) is usually used together with MS-based methods in metabolomics studies. LC-MS is able to detect the compounds of low molecular weight with high sensitivity and selectivity. Characterized by high sensitivity and resolution, GC-MS is able to identify unknown compounds including both the less polar and volatile metabolites and some polar compounds (64). Of note, GC-MS-based techniques were used to precisely characterize the urinary metabolic profiling in all the included studies, which enhance the credibility and scientificity of the results.

The pathophysiology of depressive disorder is closely related to the IgM-mediated immune responses. Azelaic acid, a naturally occurring acid found in grains, has anti-inflammatory properties. Conjugated azelaic acid was directed against by increased antibody titers, indicating enhanced oxidation of fatty acids (65), which could be observed in depressive disorder. In addition, as one of the most common saturated fatty acids, palmitic acid has been oxidatively modified in depression. Higher IgM antibody levels were found to direct against palmitoyl in depressed patients compared with normal subjects.

Considering the fact that azelaic acid, lactic acid, palmitic acid, and tyrosine were identified in all the included studies, a suitable statistical analysis technique can be used in future studies to calculate the area under the curve (AUC) value of the four specific metabolites to act as a significant biomarker to diagnose PSD. Lactic acid is an alpha-hydroxy acid (AHA) on account of the presence of a hydroxyl group adjacent to the carboxyl group. Lactate is the conjugate base of lactic acid. A few studies reported that lactate is metabolized by cerebral neurons of humans (66, 67). Glial cells could transform glucose into lactate, which was provided to the neurons (68, 69). As a result, extracellular fluid such as blood or cerebrospinal fluid is richer in lactate. However, studies found that both glial cells and neurons in the hippocampus of the brain decrease in depressed patients (70, 71), which could explain the identification of
TABLE 3 Summary of all the differential urinary metabolites identified in the included studies.

| Urinary metabolites        | Zhang et al. (51) | Xie et al. (50) | Liang et al. (49) | Chen et al. (48) |
|----------------------------|-------------------|----------------|------------------|-----------------|
| Aminoethanol               | /                 | /              | ▲                | /               |
| Azelaic acid               | ▲ ‡               | ▲ ‡            | ▲                | ▲ ‡             |
| Citric acid                | /                 | /              | ▲                | /               |
| Fructose                   | /                 | ▲              | ▲ ‡              | ▲               |
| Glucose                    | /                 | ▲              | ▲ ‡              | ▲               |
| Glyceric acid              | ▲ ‡               | ▲ ‡            | /                | ▲ ‡             |
| Hippuric acid              | ▲                 | /              | /                | /               |
| Hydroxyamine               | /                 | ▲              | ▲                | /               |
| Hypoxanthine               | ▲                 | /              | ▲ ‡              | /               |
| Indoxyl sulfate            | ▲                 | /              | ▲                | ▲               |
| Inositol                   | /                 | /              | ▲ ‡              | /               |
| Lactic acid                | ▲                 | ▲ ‡            | ▲                | ▲               |
| Leucine                    | ▲                 | /              | /                | /               |
| Malic acid                 | /                 | /              | ▲ ‡              | /               |
| Methylsuccinic acid        | /                 | /              | ▲                | /               |
| Myristic acid              | /                 | ▲ ‡            | ▲                | /               |
| n-Methyl nicotinimide      | /                 | /              | ▲                | /               |
| Palmitic acid              | ▲                 | ▲ ‡            | ▲                | ▲ ‡             |
| Phenylalanine              | ▲ ‡               | /              | /                | ▲ ‡             |
| Pseudouridine              | ▲ ‡               | /              | ▲ ‡              | /               |
| Pyroglutamic acid          | ▲                 | ▲              | /                | /               |
| Ribose                     | ▲                 | /              | /                | /               |
| Sorbitol                   | /                 | /              | ▲                | ▲               |
| Sucrose                    | ▲                 | /              | ▲                | ▲ ‡             |
| Threitol                   | /                 | /              | ▲                | /               |
| Tyrosine                   | ▲ ‡               | ▲ ‡            | ▲                | ▲ ‡             |
| Uric acid                  | /                 | ▲              | /                | /               |
| Vanillic acid              | /                 | /              | ▲                | /               |
| α-aminobutyric acid        | /                 | ▲              | /                | ▲               |
| β-aminoisobutyric acid     | ▲                 | /              | /                | /               |
| 2-methyl-3-hydroxybutyric acid | /              | /              | ▲                | /               |
| 3, 4-dihydroxybutyric acid | /                 | /              | ▲ ‡              | /               |
| 3-hydroxyisobutyric acid   | /                 | /              | ▲                | /               |
| 3-hydroxyphenylacetic acid | /                 | /              | ▲ ‡              | /               |
| 5-hydroxyhexanoic acid     | ▲ ‡               | /              | /                | /               |

▲, Identified; /, Not identified; ¶, significant deviation.

lower lactic acid in depressed subjects compared to those non-depressed subjects after stroke. Half of the included studies significantly identified the metabolic pathway of tyrosine, phenylalanine, and tryptophan. Closely involved in the context of depressive symptoms, indoleamine 2, 3-dioxygenase (IDO) is the crosstalk of the serotonergic system, the immune activation, and the kynurenic acid pathway. The breakdown of tryptophan which is the essential precursor of 5-HT is caused by IDO. Phenylalanine is the substrate for phenylalanine 4-hydroxylase (PAH) that forms another important amino acid called tyrosine. Tyrosine is converted into L-dopamine (L-DOPA) which is converted into dopamine (DA) in the tyrosine metabolic pathway. Furthermore, DA is converted into noradrenaline (NA) by DA hydroxylase. Numerous studies (72–83) confirmed that the dysfunction of monoamine neurotransmitters such as NE, DA, and 5-HT plays a crucial part in the pathogenesis of depressive disorder. Elevated DA, L-DOPA, and vanillylmandelic acid (VMA) levels were found in the hippocampus of CSDS-induced model mice (84, 85). The levels of VMA and L-DOPA
were reported to be related to depressive behaviors, which is consistent with the results of all the included studies that a higher level of tyrosine was identified in the PSD group than the non-depressed group. In contrast, patients with depression had reduced metabolites level of DA in the CSF (86) and in the hypothalamus of depression model mouse (87), results of which were consistent with the monoamine hypothesis. In summary, the different levels of metabolites might present the switch between depression and remission.

However, with regard to the plasma metabolites identified in PSD compared with non-depressed stroke survivors, the results of three previous studies were not consistent. Ding et al. (88) reported that subjects with PSD had elevated levels of oleic acid, palmitic acid, linoleic acid, pyroglutamate, proline, rhamnose, and decreased level of oxalate. In Wang et al.’s study (89), phenylacetylglutamine, p-chlorophenylalanine, and DHA levels were higher, while palmitic acid, betaine (trimethylglycine), and MHPG-SO4 were lower in the PSD group. Hu et al.’s study (90) found that the plasma of subjects with PSD was associated with higher levels of 3-methylhistidine, 1-methylhistidine, LDL and decreased level of oxalate. In Wang et al.’s study (89), phenylacetylglutamine, p-chlorophenylalanine, and DHA levels were higher, while palmitic acid, betaine (trimethylglycine), and MHPG-SO4 were lower in the PSD group. Hu et al.’s study (90) found that the plasma of subjects with PSD was associated with higher levels of 3-methylhistidine, 1-methylhistidine, LDL CH3-(CH2)n-, phenylalanine, and lower levels of tyrosine. The inconsistent results of these three studies might be related to different techniques to analyze the plasma sample. GC-MS analysis was used in Ding et al.’s study, Wang et al. applied LC-MS analysis, and the third study was based on a 1H NMR-based approach. Of note, the results of all the included studies in the systematic review were consistent due to the unified use of the GC-MS-based metabolomics method. In contrast, a GC-MS-based study (91) investigating fecal metabolites identified in PSD compared to HCs and non-depressed stroke survivors found that differential metabolites include amino acid metabolism (5-methoxytryptamine, l-kynurenine, glutamate, tyramine, cyanoalanine, maleic acid, and phenylacetic acid), lipid metabolism (lanosterol, squalene, lignoceric acid, and stigmasterol), carbohydrate metabolism (n-acetyl-d-mannosamine, arbutin, acetyl alanine, glutamate, and sucrose-6-phosphate), and nucleotide metabolism (cytosine), inconsistent with the results of included GC-MS-based research studying urinary metabolites. Therefore, the same metabonomic approach applied to analyze the same body fluid or tissue might be the reasonable strategy to precisely characterize the metabolic profiling as the diagnostic biomarker of PSD.

Nonetheless, it should be noted that only four related studies with a small sample size were included in the systematic review due to that very few studies have investigated this subject to date. Further studies investigating subgroups of different gender and age could better figure out biological differences among patients with PSD to guide more personalized treatment strategies. More studies of large sample size are also needed to investigate the essential mechanism through how urinary metabolites may interact in PSD. Furthermore, more randomized controlled trial (RCT) studies are needed to clarify the metabolomics-related role of antidepressants in alleviating depression after stroke.

Conclusion

The systematic review provided potential evidence that the GC-MS-based urinary metabolomics approach might be a useful diagnostic tool for PSD. Differential urinary metabolites significantly identified in those depressed compared to those non-depressed after stroke could be used as a precise biomarker for the diagnosis and prognosis of PSD.

Author contributions

WC and W-DS designed the study. WC, X-FWa, X-FWe, J-RZ, CH, WM, and W-DS performed the study. WC drafted the manuscript. All authors have read and approved the final manuscript.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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