Cerebrospinal fluid metabolomics in West Syndrome: central role of the serine metabolic pathway

Emmanuelle Lagrue1,2, Blandine Madji Hounoum1,3, Clémentine Rullier2, Christian R Andres1,3, Patrick Emond1,3,4, Cinzia Bocca4, Pierre Castelnau1,2* and Hélène Blasco1,3

1UMR 1253, iBrain, Université de Tours, Inserm, Tours, France
2CHRU de Tours, Hôpital Clocheville, Service de Neuropédiatrie et Handicaps, 49 Boulevard Béranger, Tours, France
3CHRU de Tours, Hôpital Bretonneau, Laboratoire de Biochimie et Biologie Moléculaire, 10 Boulevard Tonnellé, Tours, France
4PPF, CHRU de Tours, Hôpital Bretonneau, 10 Boulevard Tonnellé, Tours, France

Abstract

Purpose: West Syndrome (WS) is a rare epileptic condition which specifically affects young infants, with a potentially severe outcome. Its pathophysiology remains unclear, which hinders progress in developing targeted medications. Here, we sought to determine whether WS patients have a specific cerebrospinal fluid (CSF) metabolic profile which could help to characterize the alterations involved.

Method: CSF samples were collected during the 2010–2016 period from WS patients (n=9). The control group (n=9) included normally developing and seizure-free children who underwent a lumbar puncture (LP). Targeted and untargeted CSF metabolomics analyses were performed by Liquid Chromatography coupled to High Resolution Mass Spectrometry (LC-HRMS). The metabolic patterns were analyzed by multivariate analysis based on multiple machine-learning methods (i.e. biosigner algorithm including the use of training, test sets and bootstraps) and univariate statistical analysis.

Results: Biosigner strategy revealed a significant model discriminating WS and controls from 2 metabolites including serine. The model correctly predicted diagnosis for 83% of subjects. Serine levels were also statistically different between the two groups by univariate analysis (p=0.0023).

Conclusion: This is the first metabolomics study in WS. Our results suggest a pivotal need of serine in this disorder. The symptoms observed in WS are consistent with alterations of the serine metabolism pathway. We provide new data concerning this severe epileptic syndrome and highlight the potential value of metabolomics studies in pediatric neurological disorders, even when patients are scarce.

Introduction

West Syndrome (WS) refers to a subset of infantile spasm that can either be secondary to a neurological disorder or of unknown cause. Though it was identified in the 19th century, our understanding of its pathophysiology remains relatively poor and robust biomarkers are needed. Using proteomics approach, Greene et al. highlighted some potential biomarkers in animal models. However, this proteomics profile is not specific and such a technically challenging approach precludes its use in routine practice. Metabolomics is a powerful strategy to ascertain metabolic signatures from small molecules in biological fluids. In this study, we used for the first time a metabolomics approach, based on mass spectrometry to explore WS. This may provide an opportunity to improve the management and therapy of this disease.

Methods

Patient recruitment and data selection

The patients were recruited in the pediatric neurology unit of the Tours University Hospital (July 2010 - May 2016). The inclusion criterion was a diagnosis of WS, based on the ILAE definition [1]. CSF samples were collected before introduction of antiepileptic drugs. The control group included normally developing and seizure-free children, who initially presented with various symptoms, requiring a lumbar puncture (LP) during their investigation. The following elements were collected: sex, age at epilepsy onset, psychomotor development, developmental regression, WS etiology, ongoing treatment, age at LP, fasting or non-fasting state at LP. Ethical consent was obtained from both parents.

Sample preparation and LC-HRMS analysis

We prepared the samples and quality control samples (QC) for metabolomics by standard methods using LC-HRMS. Analysis of LC-HRMS was performed using an UPLC Ultimate 3000 system (Dionex), coupled to a Q-Exactive Mass Spectrometer (Thermo Fisher Scientific, Bremen, Germany), and operated in the positive (ESI+) and negative (ESI-) electrospray ionization modes, as previously published [2].

Data preprocessing

We performed both untargeted and targeted analysis. SIEVE® software (Thermo Fisher Scientific) was used to process raw data.
for untargeted analysis and metabolites were identified from our own library for targeted analysis. We only kept metabolites having a coefficient of variation < 30% in the QC samples in the final dataset.

#### Multivariate data analysis

Clustering of QC and patients’ samples was assessed using principal component analysis (PCA) to evaluate outliers. OPLS-DA was first performed using SIMCA® version 13.0 (Umetrics, Umeå, Sweden) to provide the Score Scatter Plot and the Loading Score Plot to visualize both, the discrimination of samples and the most discriminant variables.

We also used the biosigner algorithm in R [3] to assess as a new strategy for discovering significant molecular signatures. This data-mining algorithm is independently wrapped around different machine-learning approaches, i.e. partial least-squares discriminant analysis (PLS-DA), random forest (RF), and support vector machines (SVM). This strategy aims at finding the smallest pattern that provides a significant model after the combination of sampling (bootstrap, n=1000), ranking of relevant metabolites and the evaluation of performance after permutation within the test set and the half-interval search. The final training of the model is based on all samples from the dataset and the selected features. First, the dataset is split into training and testing datasets (by boot-strapping, controlling class proportions), then a model is built on the training set and performance of prediction is evaluated on the test set.

#### Univariate analysis

The univariate analysis of metabolites was based on fold-change (FC) values and the threshold of significance with the volcano plot and the non-parametric Wilcoxon test using Metaboanalyst, version 2.1. We highlighted only metabolites with \( p < 0.05 \) and FC > 1.5.

#### Results

##### CSF collection

Nine WS patients (six girls/three boys) were included. The clinical data are summarized in Table 1. The clinical features of the control group were as follows:

- patient 1: 14 years and 1-month-old girl, limb pain;
- patient 2: 4 years and 11-month-old girl, ataxia
- patient 3: 10 years and 4-month-old boy, headache and dizziness
- patient 4: 14 years and 6-month-old girl, severe chronic back pain
- patient 5: 9 years and 1-month-old girl, headache and ataxia
- patient 6: 9 years and 4-month-old girl paresthesia
- patient 7: 15 years and 6-month-old girl, right hemiparesis
- patient 8: 15-year-old girl, fainting
- patient 9: 12 years and 10 month-old-girl, diplopia

##### LC-HRMS analysis

Untargeted analysis revealed 291 and 82 frames for ESI+ and ESI- mode acquisitions, respectively and targeted analysis led to the identification of 62 metabolites.

##### Multivariate analysis

PCA showed a tight cluster for the QC and for subjects within each group (data not shown). The Figure 1A shows the Score Scatter Plot built from OPLS-DA \( (p=0.0001) \) performed using the 10 most relevant metabolites (Figure 1B) such as serine, tryptophan, histidine, 3-methylhistidine, glutamic acid, citrate, vitamin B5 and unidentified metabolites. Biosigner analysis revealed a significant signature for WS patients from 2 metabolites: serine and an unknown metabolite X363 from ESI –. The percentage of correctly predicted patients was 83% in the test set after 1000 bootstraps.

##### Univariate analysis

Volcano plot revealed 8 metabolites with \( p\text{-value} < 0.05 \) and FC > 1.5:

- serine \( (p=0.0023) \)
- 3-methylhistidine \( (p=0.024) \)
- histidine \( (p=0.0152) \)
- vitamin B5 \( (p=0.0152) \)
- succinate \( (p=0.03) \)
- 3 unidentified metabolites including X363 \( (p=0.005, \text{data not shown}) \).

#### Discussion

This is the first CSF metabolomics study in WS patients. One of its strengths lies in the combination of several statistical approaches, i.e. the biosigner machine-learning strategy and univariate analysis, based on the volcano plot. Despite the small cohort size, PCA findings within each group of subjects revealed no bias due to fasting or treatment, and robust multivariate analysis provided an excellent model to discriminate WS and controls. We could not select matched controls due to ethical issues, but this preliminary study enabled to show that metabolomics strategy may be used in epilepsy.

##### A specific CSF metabolome in WS patients

OPLS-DA revealed an excellent model to discriminate WS and controls from only 10 metabolites. Surprisingly, the highly restrictive biosigner analysis also provided a significant model with an accuracy superior to 80%, from only 2 metabolites. This new method, based on different types of recent machine-learning strategies, including bootstrap resampling, is considered the most robust for highlighting discriminant compounds and judging the fitness of a model.

##### Discriminant role of serine

The most prominently affected metabolite was serine. Elevated serine levels have already been observed in animal models. In humans, in situ elevation of serine levels is associated with intractable seizures. Thus, serine homeostasis appears to play an important role in cerebral neurotransmission and epileptogenesis. Both glutamate and serine elevation are required to activate NMDARs, representing a key factor in synaptic plasticity and excitatory transmission [4]. It has been shown in a rat model of chronic epilepsy that altered D-serine levels modify NMDAR subunit composition, altering long-term potentiation and subsequent memory functions [5]. Serine also participates in the regulation of the mTOR (mammalian target of rapamycin) pathway, implicated in synaptic plasticity, neuronal repair mechanisms, and memory [6,7]. Inhibition of the mTOR pathway is considered to be a promising anticonvulsant strategy [7]. The ketogenic diet, which increases serine levels in CSF, also has a beneficial effect in severe epileptic patients [8].

We also observed that the concentration of tryptophan tended to be higher in WS patients. This is consistent with the findings of serine, as tryptophan results from the irreversible condensation of indole and serine. Tryptophan degradation secondarily leads to the production of nicotinamide adenine dinucleotide (NAD+), known as the kynurenine pathway. Our results further emphasize the involvement of this pathway in epilepsy [9,10]. However, its involvement in the pathophysiology of WS is still hypothetical and requires further specific studies.

Other biochemical pathways involved in WS

Citrate and succinate involved in the tricarboxylic cycle were also identified. Citrate is highly linked to epileptogenic process as its transport is altered in patients carrying the SLC13A5 gene mutation, in which epilepsy is a key feature.

Modulation of the histidine-carnosine pathway was also highlighted. This data is consistent with a previous report showing that histidine and carnosine both exhibit anticonvulsant effects in the rat PTZ kindling seizures model and penicillin-induced epileptiform activity in rats.

In conclusion, this is the first study to address the potential use of metabolomics in WS. We showed a discriminant profile and our data shed light on the pathways involved in WS pathophysiology, especially concerning serine. The next step will be to validate these results on an independent cohort, comparing unknown cause WS patients to other epileptic children.

Declarations

Authorship / Contributorship / Acknowledgements: All authors listed have contributed sufficiently to the project to be included as authors, have approved the manuscript and this submission.

Funding information: To the best of our knowledge, no conflict of interest, financial or other, exists.

Competing interest: We don’t have any competing interest to declare.

References
