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Magnetic resonance spectroscopy metabolomics of cerebrospinal fluid in patients with multiple sclerosis, clinically isolated syndrome, other inflammatory brain diseases and controls

Metabolomika cerebrospinálneho likvoru pomocou magnetickej rezonančnej spektroskopie u pacientov so sclerosis multiplex, s klinicky izolovaným syndrómom, inými zápalovými ochoreniami mozgu a u zdravých kontrol

Abstract

Aim: Early recognition of multiple sclerosis (MS) allows patients to begin treatment earlier and delay disease progression. We performed an analysis of cerebrospinal fluid (CSF) metabolites to find early predictors of MS. Methods: We included 56 participants with suspected MS before any treatment. Out of those, 28 patients were diagnosed with definite MS, 17 with clinically isolated syndrome (CIS) according to McDonald 2010 criteria, and 11 with other demyelinating diseases (DEM) of the CNS. The control group (CON) included 29 participants without any confirmed CNS disease. Proton nuclear magnetic resonance spectroscopy was used to measure CSF metabolites. Results: Glutamine, correlating with Expanded Disability Status Scale (EDSS), was the only metabolite capable to distinguish between CIS and MS, DEM, and CON. Valine, leucine, isoleucine, decreased in CIS and MS when compared with CON, did not differ from DEM. Citrate CSF levels specified MS and CIS against DEM but did not help to distinguish between CIS and MS. Citrate showed significant correlations with age, disease duration, and EDSS in MS patients. Acetate, acetone, pyruvate, formate and histidine CSF levels were not significant predictors of MS or CIS, although they correlated with selective variables. Conclusion: This work shows the predictive role of CSF glutamine in diagnosing MS since its early stages, pinpointing an important role of the glutamate/glutamine cycle in MS pathogenesis. Another potential predictor of MS was citrate. Other metabolites were not identified as sensitive CSF markers of MS.

Key words

metabolomics – cerebrospinal fluid – proton nuclear magnetic resonance spectroscopy (1H-NMRS) – multiple sclerosis – clinically isolated syndrome – inflammatory demyelinating brain diseases

Klíčová slova

cerebro-spinálny likvor – metabolomika – jednoprotonová nukleárna magnetická rezonančná spektroskopia (1H-NMRS) – sclerosis multiplex – klinicky izolovaný syndróm – zápalové demyelizačné ochorenia mozgu

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Súhrn

Ciel: Včasné rozpoznanie sclerosis multiplex (SM) pomáha začať liečbu pacientov skôr, a tak oddialiť progresiu ochorenia. Urobili sme analýzu metabolitov cerebro-spinálneho likvoru (cerebrospinal fluid; CSF), s cieľom zistiť prediktory včas, a tak oddialiť SM. Metódy: Do štúdie bolo zaradených 56 jedincov s podozrením na SM, pred začatím akejkoľvek liečby. Z nich bolo 28 diagnostikovaných ako definitívna SM, u 17 pacientov sme zistili klinicky izolovaný syndróm (clinically isolated syndrome; CIS) podľa McDonaldových kritérií z roku 2010, v 11 prípadoch sa jednalo o iné demyelinizačné ochorenie CNS (DEM). Kontrolnú skupinu (CON) tvorili 29 jedinci, ktorí nemali dokázané žiadne ochorenie CNS. Na meranie metabolitov CSF bola použitá protonová nukleárna magnetická rezonančná spektroskopia. Výsledky: Glutamín, ktorý koreloval s Expanded Disability Status Scale (EDSS), bol jediným metabolitom, ktorý dokázal odlíšiť CIS, SM, DEM a CON. Valín, leucín, isoleucín, znížené u CIS a SM v porovnaní s CON, sa neodlišovali od DEM. Hladiny citrátu v CSF špecifikovali SM a CIS oproti DEM, ale nepomohli v rozlíšení CIS a SM. Citrát ukazoval signifikantné korelácie s vekom, dľžkou trvania ochorenia a EDSS u SM pacientov. Acetát, aceton, pyruvát, formát, histidin v CSF neboli signifikantnými prediktormi SM alebo CIS, hoci korelovali s niektorými vybranými premennými. Záver: Táto práca ukazuje prediktívnu úlohu glutamínu v CSF v stanovení diagnózy SM od jej včasných štádií, vypichujúc tak dôležitú úlohu glutamát/glutamínového cyklu v patogenéze SM. Ďalší potenciálny prediktor SM bol citrát. Ďalšie metabolity neboli identifikované ako senzitívne CSF markery SM.

Introduction

Multiple sclerosis (MS) is a heterogeneous disease with an unpredictable disease course. It is known that accumulating disability reflects the progression of neuronal damage, which can occur during clinically silent inflammatory episodes. Neurodegeneration was traditionally considered as a secondary phenomenon to inflammation and demyelination. However, recent data indicate that neurodegeneration develops along with inflammation and demyelination [1]. While both inflammation and demyelination are well described and understood cellular processes, neurodegeneration is still mattered of debate.

In MS, our current treatments reduce inflammation and, therefore relapses and shortterm disability, and we hope this will slowdown the rate of neurodegeneration [2]. Studies that have analyzed early treatment in patients highly likely to have MS (clinically isolated events with the evidence of lesions on brain MRI) provided significant benefits in delaying further changes in MRI and attacks [2,3]. Early recognition of the inflammatory process enables patients to begin treatment with an immunomodulatory agent but may also increase the rate of false--positive MS diagnoses [3]. However, in MS, there is no such specific test available which is why one needs to rely on "circumstantial" evidence". The diagnosis is based on typical, yet not limited to, clinical findings, MRI, and cerebrospinal fluid (CSF) examination as well as other investigations [4].

In our previous work, we found superiority of neurodegenerative metabolic pathways in brain tissue of MS patients using the *in vivo* proton nuclear magnetic resonance spectroscopy (1H-NMRS) method [5]. Now we decided to test CSF metabolites as

we believe that additional molecules of CSF could help to improve the differential diagnosis of MS and its subtypes and predict the course of the disease, thus selecting optimal therapy for each patient.

Both inflammatory and neurodegenerative markers of CSF have yet been described [6]. However, so far, only several groups of researchers studied concentrations of metabolomics in CSF using 1H-NMRS [7–13]. Nevertheless, the results are partly inconsistent.

According to our knowledge, our work is the first one presenting comparative analysis of CSF metabolites to find early predictors of MS

Materials and methods Patients and controls

Informed written consent was obtained from all study participants after approving the study protocol by the local ethics committee. Only participants with suspected MS before any treatment were included. The patients underwent clinical and paraclinical examinations to prove or exclude MS. Clinical disability was evaluated by 2 neurologists specialized in MS using Expanded Disability Status Scale (EDSS). The neurologists have been Neurostatus certified in EDSS evaluation. MRI examination of the brain and cervical spinal cord performed according to the approved protocol (MAGNIMS) provided evidence of pathological lesions and their activity. Evoked potentials (visual, acoustic, somatosensory tests) informed about potential demyelinating and axonal changes of the tracts.

CSF and blood samples were used for biochemical and immunological tests. In CSF, we tested oligoclonal bands (OCB), flow cytometry, immunoglobulins, and selectively

antibodies against autoimmune encephalitis. MS and CIS diagnosis were based on the McDonald criteria 2010 [3]. Based on the results, 28 patients were diagnosed with definite MS, 17 with CIS (patients with MS after the first clinical relapse), and 11 with other inflammatory brain diseases of the CNS (demyelinating diseases of the CNS; DEM). These patients presented with either isolated monophasic optic neuritis or with idiopathic myelitis, vasculitis or monophasic forms of the CNS demyelination. MRI lesions were different from MS and did not comply with the Barkhof criteria for MS [14]. In this group, OCB were negative (type 1). The patients also did not fulfill MRI criteria for neuromyelitis optica, and IgG antibodies against aquaporin-4 were negative. Three out of the patients were tested to anti-myelin oligodendrocyte glycoprotein (anti-MOG) with negative results.

Due to ethical concerns surrounding the collection of CSF from healthy individuals, healthy controls were not recruited for this study. The control group (CON) included 29 participants without any confirmed CNS disease. These were patients with episodic severe headache, lower back pain or peripheral neuropathy, and normal MRI scans of the brain and spinal cord. In these patients, lumbar punction and evaluation of CSF samples were indicated to exclude intracranial bleeding, neuroinfections, or a disorder of lower motoneuron, e.g., peripheral neuropathy. Biochemical tests were indicated in all samples, and immunological tests were applied in selected CFS samples accordingly.

Sample preparation

The stock solution consisted of phosphate buffer 250 mL and 0.28 mMTMSP-d4 (trimethylsilylpropionic acid -d4) as a chemical

shift reference in deuterated water. CSF was immediately centrifuged at 2,000 rpm at 4° C for 20 min and then frozen to -80 °C until examined. After thawing, 500 µL of CSF was softly mixed with a 100 µL stock solution and transferred into a 5- mm 1H-NMR tube. Samples were randomized for acquisition.

1H-NMRS data acquisition

1H-NMRS data were acquired on 600 MHz 1H-NMRS spectrometer Avance III (Bruker, Billerica, MA, USA) equipped with cryoprobe at T = 310 K. Initial settings (basal shimming, receiver gain, water suppression frequency) was done on an independent sample and adopted for measurements. After the preparation, samples were stored in a Sample Jet, cooled at 5 °C. Before the measurement, each sample was preheated on the 310 K for 5 min. An exponential noise filter was used to introduce 0.3-Hz line broadening before Fourier transform. The TMSP-d4 signal was assigned a chemical shift of 0.000 ppm.

We modified the standard profiling protocols from Bruker as follows: NOESY with presaturation: FID size 64k, dummy scans 4, number of scans 128, spectral width 20.4750 ppm; COSY with presaturation: FID size 4k, dummy scans 8, number of scans 1, spectral width 16.0125 ppm; homonuclear J-resolved: FID size 8k, dummy scans 16, number of scans 4; CPMG with pre-saturation: FID size 64k, dummy scans 4, number of scans 128, spectral width 20.0156 ppm. All experiments were conducted with a relaxation delay of 5 s.

Data analysis

All spectra were binned to bins of the size of 0.001 ppm, starting from 0.500 ppm to 9.000 ppm, with an excluded water region of 4.6-4.9 ppm. Spectra were solved with the help of the human metabolomics database, chenomics software and by researching in metabolomics literature. For all compounds, the multiplicity of peaks was confirmed in j-resolved spectra and homonuclear cross peaks were confirmed in COSY spectra. After the metabolites were identified (Fig. 1), we chose spectra subregions with only a single metabolite assigned, or minimally affected by other cometabolites. In 0.001 ppm binned spectra, we summed integrals of selected signals. Metabolites showing weak intensive peaks or strong peak overlap were excluded from the evaluation. Statistical tests were performed in Matlab R2015a (MathWorks, Naticks, MA, USA).

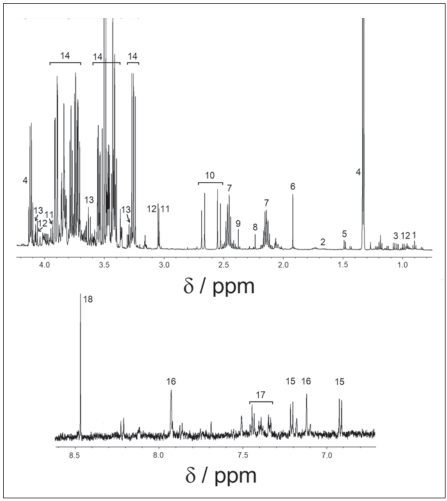


Fig. 1. 1H-NMR spectrum of cerebrospinal fluid, cpmg acquisition.

1 – isoleucine; 2 – leucine; 3 – valine; 4 – lactate; 5 – alanine; 6 – acetate; 7 – glutamine; 8 – acetone; 9 – pyruvate; 10 – citrate; 11 – creatine; 12 – creatinine; 13 – myo-inositol; 14 – glucose; 15 – tyrosine; 16 – histidine; 17 – phenylalanine; 18 – formate

Obr. 1. 1H-NMR spektrum v cerebrospinálnom likvore, nábor v pulzovej sekvencii 1H-NMR.

1 – izoleucín; 2 – leucín; 3 – valín; 4 – laktát; 5 – alanín; 6 – acetát; 7 – glutamín; 8 – acetón; 9 – pyruvát; 10 – citrát; 11 – kreatin; 12 – kreatinín; 13 – myo-inozitol; 14 – glukóza; 15 – tyro-zín; 16 – histidín; 17 – fenylalanín; 18 – formát

Results

Finally, 18 metabolites: valine, isoleucine, leucine (branched-chain amino acids [BCAA]), lactate, alanine, formate, tyrosine, phenylalanine, glutamine, histidine, citrate, acetate, acetone, pyruvate, myoinositol, glucose, creatine and creatinine were identified in 1H-NMRS spectra.

Glutamine was the only metabolite which levels differed between CIS, MS and CON. Glutamine also correlated with EDSS in MS and CIS. In CSF, BCCA (valine, leucine, isoleucine) were decreased in CIS/MS patients after the first clinical relapse and MS compared with CON and also correlated with age and disease duration (DD) of MS patients. Citrate CSF levels showed signifi-cant correlations

with age, DD and EDSS in MS patients (Fig. 2). The acetate CSF levels were significantly increased in MS against all other groups. Other CSF metabolites significantly different between groups were formate, histidine, acetone, and pyruvate.

Correlations of relative metabolites CSF levels between subgroups are summarized in Tab. 1. Results from statistic calculation related to patients parameters are summarized in Tab. 2.

Discussion Glutamine

Glutamine is an ubiquitous amino acid in mammalian tissues and in the blood, where

Tab. 1. Statistic evaluation of the relative concentration of metabolites in cerebrospinal fluid from patients with MS, CIS/MS after a first clinical result, DEM, and CON.

	MS/CON	MS/DEM	MS/CIS	CIS/DEM	CIS/CON	DEM/CON
glutamine	0.0004*	0.01*	0.02*	_	0.04*	_
	15%	10%	9%		7%	
valine	0.09	-	-	_	0.04*	_
	-6%				-11%	
isoleucine	0.09	-	-	-	0.02*	-
	-11%				-19%	
leucine	0.08	-	-	_	0.07	_
	-14%				-13%	
formate	0.04*	-	0.09	_	_	_
	18%		18%			
histidine	0.04*	-	_	_	_	_
nistiaine	-39%					
citrate	_	0.07	-	0.06	_	0.08
		10%		13%		-10%
acetate	0.03*	0.05	0.03*	_	_	_
	14%	16%	13%			
acetone	0.04*	-	0.03*	0.07	-	0.05
	-37%		-35%	42%		-32%
pyruvate	0.02*	-	0.04*	-	_	-
	-16%		-22%			

^{*}statistical significance P < 0.05, Mann-Whitney U test

CON – controls; CIS/MS – patients after the first clinical relapse (clinically isolated syndrome vs. multiple sclerosis); DEM – other inflammatory brain diseases; MS – multiple sclerosis

Tab. 2. Correlation of cerebrospinal fluid metabolite levels with MS and CIS/MS after the first clinical relaps patients' EDSS, age and DD.

	age/years	EDSS	DD/years	
alanine	pval (MS) < 0.05	pval (MS + CIS) < 0.05	pval (MS) < 0.005	
	pval (MS + CIS) < 0.005		pval (MS + CIS) < 0.005	
valine	pval (MS) < 0.05		pval (MS) <0.05	
	pval (MS + CIS) < 0.01		pval (MS + CIS) < 0.05	
isoleucine	pval (MS) < 0.05		pval (MS) < 0.05	
leucine	pval (MS) < 0.05		pval (MS) < 0.05	
glutamine		pval (MS) < 0.05		
		pval (MS + CIS) < 0.05		
citrate	pval (MS) < 0.0005	pval (MS) < 0.05	pval (MS) < 0.05	
	pval (MS + CIS) < 0.0001	pval (MS + CIS) < 0.005	pval (MS) < 0.05	
acetate	pval (MS + CIS) < 0.05		pval (MS) < 0.01	
			pval (MS + CIS) < 0.001	

statistical significance P < 0.05, all relative CSF metabolite levels were increasing with evaluated patients' parameter

CIS – clinically isolated syndrome (patients after the first clinical relapse); DD – disease duration in years; EDSS – Expanded Disability Status Scale; MS – multiple sclerosis

it is a precursor or product in multiple metabolic pathways in both the CNS and in the peripheral tissues. Glutamine participates in a glutamine-glutamate cycle that plays a role in neuroglia communication in the synapse. After the release of neurotransmitter glutamate into the synapse, this is rapidly up-taken by astrocytes and metabolized to glutamine, which is delivered into neurons, metabolized back to glutamate and repeatedly used for neurotransmission [15]. Glutamate, the principal CNS excitatory neurotransmitter, is the most abundant amino acid in the brain with the extracellular concentration much lower than the intracellular one [16]. There is growing evidence that glutamate plays a role in the pathology of MS. Increased glutamate CSF levels were observed in MS patients by Bakhartova et al [17] and Sarchielli et al [18] and correlated with disease severity and course [18].

Glutamine crosses the blood-brain barrier (BBB) via facilitating transport systems [19], using its carrier [20]. The concentration of glutamine in the CSF is very close to its concentrations in the arterial blood or extracellular fluid [21]. In our work, we observed increased CSF levels of glutamine, co-metabolite of glutamate, in patients with MS and in CIS/MS patients after the first clinical relapse when compared to CON and DEM. In our patients, the transmission of glutamine from the arterial blood to the CSF via hyperpermeable BBB cannot be rejected, but this mechanism does not explain a selective increase of glutamine only in MS and CIS/MS patients after the first clinical relapse over other forms of brain diseases, where disruption of the BBB is also found [22]. We supposed the active transformation of glutamate to glutamine inside the brain of MS and CIS/MS patients after the first clinical relanse

MS is considered to be the mostly T-cell mediated disease, and T-cell activation is associated with abnormal glutamate/glutamine functioning. T-cell activation is dependent on extracellular glutamine and T-cell activation induces expression of glutamine transporters [23], which has also been proven in the peripheral blood [24]. The depletion of glutamine blocks proliferation and cytokine production and this cannot be rescued by supplying biosynthetic precursors of glutamine. T-cells are located at the active edge of MS lesions, and the presence of perivascular T-cell infiltrates throughout the CNS is a consistent feature

in the early stages of MS [25]. Although various processes involving glutamine are well described, it is unknown, if increased glutamine levels represent rather the cause or the consequence in developing CIS and further MS. Based on our results, we assume its participation in disease development, since the glutamine CSF level was increased in MS patients over CIS (MS patients after a first clinical relapse), in MS and CIS (MS patients after a first clinical relaps) over CON, and it correlated with the degree of disability.

Contrary to our results, Ashley et al [10] reported a decreased CSF glutamine level in MS against controls. However, the investigation in highly homogeneous CIS patient cohorts produced results with increased CSF glutamine levels [7]. Increased levels of glutamine in the CSF were also documented in different forms of neurodegeneration such as in Alzheimer disease [26] or amyotrophic lateral sclerosis [27]. Further examination in this field is required to elucidate links between glutamate and different types of neurodegenerative disorders.

Leucine, isoleucine, valine (BCAA)

The branched-chain amino acids leucine, isoleucine and valine cross the BBB swiftly. It seems probable that BCAA participate in a "glutamate-BCAA cycle", which involves shuttling BCAA between astrocytes and neurons [28]. The metabolism of BCAA in glial and neural cells was studied in detail by Murin et al [29–31]. Besides function as "buffering" of an internal pool of glutamate, BCAA influence the immune properties of microglial cells and their responsiveness to pro-inflammatory signals in vitro [32].

Valine, leucine and isoleucine CSF levels were found decreased in MS, CIS (MS patients after the first clinical relapse) and DEM against the CON group (Tab. 1) and correlated positively with DD and age of MS patients (Tab. 2). Since these essential amino acids are transported into the brain by crossing BBB via controlled, facilitated transport and because they are used for accelerated glutamate synthesis, their reduced CSF levels may indicate their partial depletion due to this process.

Citrate

There is insufficient knowledge about specific CNS processes in which citrate may play a prominent role. In general, citrate is the key

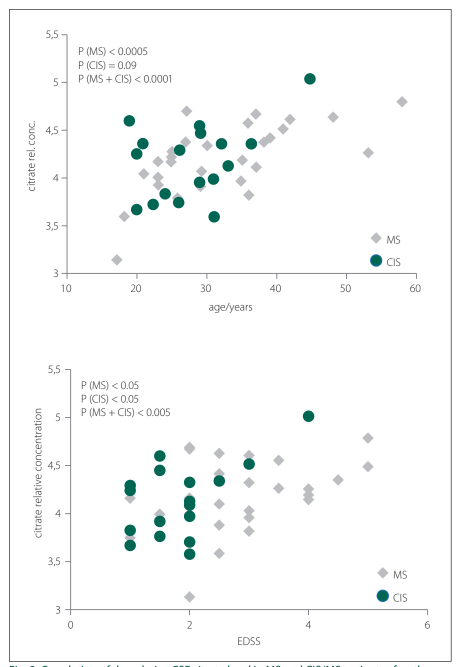


Fig. 2. Correlation of the relative CSF citrate level in MS and CIS/MS patients after the first clinical relapse with age and EDSS score.

 ${\it CIS-clinically isolated syndrome; CSF-cerebrospinal fluid; EDSS-Expanded Disability Status Score; MS-multiple sclerosis}$

Obr. 2. Korelácia relatívnych hladín citrátu v cerebrospinálnom likvore s vekom a skóre EDSS pacientov s MS a CIS.

CIS – klinicky izolovaný syndróm; CSF – cerebrospinálny likvor; EDSS – Expanded Disability Status Scale; MS – sclerosis multiplex

metabolite of Krebs cycle. Besides this, citrate plays an important role in the fatty-acid synthesis, the main component of brain lipids [33]. Citrate is released from astrocytes in large amounts, which is in keeping with

high concentrations found in the CSF [34]. However, the functional importance of high concentrations of citrate in the CSF and the large release of citrate from astrocytes are not fully understood. Rather than cit-

rate serving as a precursor for the synthesis of transmitter glutamate, citrate released from astrocytes may regulate the extracellular concentrations of Ca²⁺, Mg²⁺ and Zn²⁺ by chelation [35], thereby modulating neuronal excitability as an endogenous modulator of glutamate receptors, in particular, the N-methyl-D-aspartate (NMDA) receptor subtype. Citrate has a preference for Zn²⁺ over Ca²⁺ and Mg²⁺ [35] and is potentially able to abolish the inhibitory action of Zn²⁺ of the NMDA receptors in a reverse manner [36]. It is therefore attractive to suggest a hitherto unknown regulatory function of citrate in particular astrocytes [37].

In our study, we did not observe significant differences in citrate CSF levels between patient and control groups, contrary to other results, where decreased CSF citrate level was found in MS patients [7,38]. However, we observed the relationship of citrate CSF level to several clinical parameters of MS and CIS (Tab. 2). The CSF levels of citrate increased with EDSS score, DD and age in MS and with EDSS score and age in CIS (MS patients after the first clinical relapse) (Fig. 2).

Glucose, lactate, and pyruvate

Astrocytes have the highest concentration of the primary glucose transporter. Glucose, transported into astrocyte, is entering glycolysis with pyruvate as a product. Pyruvate has two fates: it can enter Krebs cycle in mitochondria or it is converted to lactate, which is then transported to neurons. Once lactate enters neurons, it is converted back to pyruvate. Neuronal pyruvate can be shuttled to mitochondria and funneled into Krebs cycle to gain energy or convert to citrate for lipid synthesis. Neurons show a relative preference for lactate over glucose when both substrates are present. However, we do not yet know the relative contributions of glycolysis and oxidative phosphorylation to the immediate energy demands of the neurons

In our study, CSF levels of glucose and lactate were not changed in any of the groups. Based on our results, it seems that neither "lactate shuttle" nor glycolysis was affected in the evaluated groups. Other studies published conflicting results in the CSF in MS: decreased glucose [9], increased glucose [39], lactate without change [9], increased lactate [7] and decreased lactate [40]. We found decreased CSF pyruvate levels in MS patients against CIS (MS patients after the first clinical relapse) and CON. Current research revealed

many mitochondrial abnormalities involved in the development and progression of MS [41] that are also supposed in our MS patients. Decreased pyruvate may also balance mitochondrial citrate overproduction in an advanced disease.

Histidine

Histidine is a precursor of histamine, a neurotransmitter for the brain and spinal cord. Elevated CSF histamine levels in MS patients were already reported [42,43]. Histamine can be found in large amounts in granules of mast cells, a type of white blood cells, and in granulocytes, derived from myeloid stem cells. Mast cells participate in innate and adaptive immunity, inflammation and autoimmunity [44]. We observed decreased histidine CSF levels in MS patients when compared to CON. Considering that histamine is produced from histidine by irreversible decarboxylation and histidine is an essential amino acid for the brain, entering the brain via BBB transporter systems, the increased histamine can be manifested by the exhaustion of histidine

Acetate and formate

Previous data showed that levels of acetate and formate in MS patients were altered [9,39,45]. Our analysis revealed statistically significantly increased levels of both acetate and formate in CSF of MS patients against CON, CIS patients, and DEM (Tab. 1).

In general, the majority of acetate is generated by the enzymatic hydrolysis of acetyl-CoA. Acetyl-CoA is a co-substrate for the synthesis of citrate, and therefore sustaining the reactions of the citric acid cycle has a profound impact on the energy metabolism and biosynthetic reactions of fatty acids, cholesterol, ketone bodies and non-essential amino acids and their derivates including neurotransmitters glutamate, gamma-aminobutyric acid (GABA) and acetylcholine. Furthermore, acetyl-CoA is a common donor of acetyl residue for the synthesis of N-acetylaspartate or N-acetylaspartylglutamate; and also is essential for the sustaining of protein acetylation [46]. Reactivation of acetate into acetyl-CoA may take place in cytosolic, mitochondrial and nuclear compartments of glial cells and neurons. The cytosolic expression of acetyl-CoA synthase in oligodendrocytes [47] suggests their capability to utilize acetate for lipid synthesis and myelination [46].

Formate may be generated either by peroxisomal α -oxidation of fatty acids [48] or disturbed mitochondrial metabolism [49]. Since brain lipids consist of a considerable proportion of fatty acids [33], the demyelination accompanying lipolysis can elevate the level of free fatty acids, and then formate. In the brain, acetate and formate are further metabolized in glia and neurons [50,51]. We assume that increased production of free fatty acids by demyelination leads to their elevated mitochondrial and peroxisomal degradation, followed by the increased formation of acetate and formate. We suggest that the quantities of newly generated acetate and formate bodies exceed the catabolic capacity of the brain cells and therefore lead to their accumulation in the CSF of our MS patients.

Alanine

A variety of data suggest that beta-alanine should be considered a small-molecule neurotransmitter that may function as an important carrier for ammonia transfer to complete brain glutamate-glutamine cycling. Alanine is a non-essential amino acid, crossing BBB by transport systems [19]. In vitro, both cortical and cerebral neurons show several-fold-higher de novo synthesis and release of alanine [53,54] confirming neuronal bonding of nitrogen. We found a significant increase in alanine CSF level in MS patients correlating with age, DD and disability score, which may show alanine participation in disease progress and duration. However, alanine may be a part of pyruvate recycling, where the changes of alanine with the age should be expected [55]. Therefore, a link between alanine and MS is only disputable.

Other unchanged metabolites

Creatine and creatinine serve as indices of one aspect of brain energy metabolism, the creatine-creatine phosphate shuttle. No changes in CSF levels of both metabolites in our study indicate that this part of energy metabolism is not affected in CIS (MS patients after the first clinical relapse) and in MS, or DEM patients.

Furthermore, we did not find an alteration in CSF levels of tyrosine and phenylalanine. In the brain tyrosine functions as a precursor of catecholamines (dopamine, norepinephrine, epinephrine). It was reported that the activity of neurotransmitter dopamine decreases in the CSF in MS [56]. However, we did not prove that.

Limitations

Although we have identified disease unique metabolite profiles, the future use of metabolite biomarkers in the clinical environment will require optimisation to improve accuracy. In our study, the accuracy of diseaseunique metabolites profiles may be affected by a lower number of participants in the groups and heterogeneity of a disease course in both MS and non-MS diseases. Other limitation of this study could be the fact that we tested CSF in isolation, the samples were not coupled with serum of study participants. We did not test anti-MOG in all study participants due to technical problems. The test was not widely available in our country at the time of study beginning. Further studies may improve our knowledge about CSF biomarkers in demyelinating disorders.

Conclusion

In our work, glutamine, which correlated with EDSS, was the only metabolite capable of distinguishing between MS after the first clinical relapse or CIS and MS concerning other forms of the demyelinating brain diseases and controls. This work is the first one to show the predictive role of CSF glutamine in the early stages of MS, pinpointing an important role of the glutamate/glutamine cycle in MS pathogenesis. Another potential predictor of MS was citrate, as citrate CSF levels specified MS and CIS against DEM. However, it did not help to distinguish between early and later stages of MS. Citrate showed significant correlations with age, DD and EDSS in MS patients. The BCAA (acetate, acetone, pyruvate, formate, histidine) CSF levels are not significant predictors of MS

Ethical statement

This study was performed following the ethical standards laid down in the 1964 Declaration of Helsinki and its later amendments. Ethics approval was performed by the Ethical committee of the Jessenius Faculty of Medicine in Martin, Comenius University (EK 1678/2015). Informed written consent was obtained from all study participants.

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Disclosures

The authors declare they have no potential conflicts of interest concerning drugs, products or services used in the study.

Authors contribution

EB, LP, SS, EK – research, data collection, clinical assessment; EB, EK – study design, writing of the manuscript; EB – performance of 1H-NMRS, statistical analyses, collection of the data; DC, PH, SS, JL, EK – critical reading and correction of the manuscript.

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MAGNETIC RESONANCE SPECTROSCOPY METABOLOMICS OF CEREBROSPINAL FLUID IN PATIENTS

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