Multitasking Biomolecules in ME/CFS Pathogenesis Known Players on Their Unexpected Journey

Alain Moreau PhD

Full Professor

Department of Stomatology, Faculty of Dentistry / Department of Biochemistry & Molecular Medicine, Faculty of Medicine Université de Montréal

Scientific Director

Viscogliosi Laboratory in Molecular Genetics of Musculoskeletal Diseases, CHU Sainte-Justine Research Center Open Medicine Foundation Collaborative ME/CFS Research Center at CHU Sainte-Justine/Université de Montréal

Director

Interdisciplinary Canadian Collaborative Myalgic Encephalomyelitis – ICanCME – Research Network

alain.moreau@umontreal.ca

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Institute of Musculoskeletal Health and Arthritis









Research Centre CHU sity Hospital Centre

e Montréa



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ograms are funded by:

Research program on ME/CFS

Pr. Moreau ME/CFS research program has been approved by Sainte-Justine University Hospital Ethic Review Board (protocol #2015-829)

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Member of the Institute of Musculoskeletal Health & Arthritis (CIHR), Institute Advisory Board Ο

Health Research

- Member of Open Medicine Foundation Scientific Advisory Board (USA) Ο
- Senior Editorial Board Member, Scientific Reports, Nature Co (UK) Ο
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en santé du Canada

? OPEN MEDICINE FOUNDATION

> ME/CFS Collaborative Research Center at CHU Sainte-Justine/Université de Montréal

https://www.omf.ngo/collaborative-research-center-montreal/



Alain Moreau PhD, Scientific Director Anita Franco, MSc, Lab coordinator Yaned Gaitan MSc, Research assistant Viorica Lascau MSc, Manager – certification Mohamed Elbakry PhD, Research Associate Dashen Wang MD, PhD, Research Associate Sophie Perreault, RN Clinical research nurse Valerie Tremblay, RN Clinical research nurse Iurie Caraus PhD, Postdoctoral Fellow Wesam Elremaly PhD, Postdoctoral Fellow Corinne Leveau BSc, PhD student Evguenia Nepotchatykh BSc, PhD student Lynda Chalder MSc, PhD student Bita Rostmani, undergraduate student Marie-Yvonne Akoume PhD, Professor (Université de Libraville) Dawei Li, PhD, Associate Professor (University of Vermont)





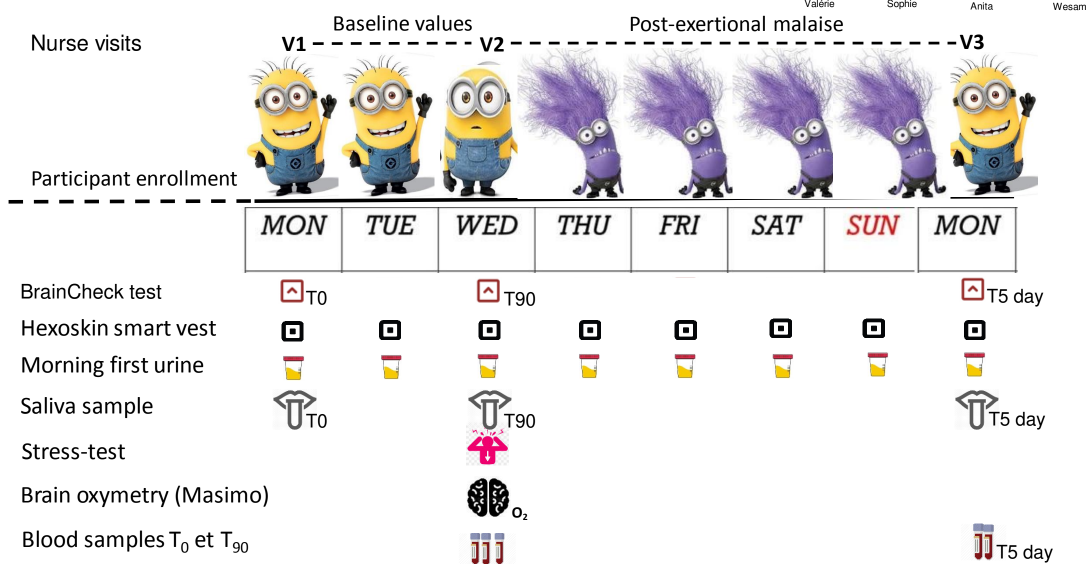


PROVOCATION STUDY: A NEW APPROACH Development of a stress challenge inducing post-exertional malaise (PEM) Compression at 0,006 Hz, HEXO varying from 0-4 psi stimulation 120 Healthy Subject Patient Blood sampling T_0 T_{90 min} Hexoskin during the stress-test smart vest Brain oxymetry Valérie Sophie Anita Wesam Viorica

XPERIMENTAL APPROACH Stress-test version 2.0



Anita



6 SLEEP DISTURBANCES IN ME/CFS

Longitudinal sleep assessment with Hexoskin smart biometric vest

Sleep Data – EM-169 (F, 42 y)	Pre-Stress Test (night 2)	Post-Stress Test (night 3)	Post-Stress Test (night 7)	Post-Stress Test Worse Night (#3)
Sleep Position Changes (#)	47	105	46	105
Total Sleep Time (hh:mm:ss)	07:08:40	07:13:20	08:28:20	07:13:20
REM Sleep Time (hh:mm:ss)	02:04:20	01:52:20	03:33:20	01:52:20
Non-REM Sleep Time (hh:mm:ss)	05:04:20	05:21:00	05:55:00	5:21:00
Time Awake (hh:mm:ss)	00:22:40	00:47:00	00:47:20	00:47:00

Sleep Data – EM-170 (F, 50 y)	Pre-Stress Test (night 2)	Post-Stress Test (night 3)	Post-Stress Test (night 7)	Post-Stress Test Worse Night (#5)
Sleep Position Changes (#)	61	65	64	59
Total Sleep Time (hh:mm:ss)	08:47:40	10:53:00	07:15:20	6:13:40
REM Sleep Time (hh:mm:ss)	1:51:00	3:23:20	01:08:00	00:55:00
Non-REM Sleep Time (hh:mm:ss)	6:56:40	7:29:40	6:07:20	5:18:00
Time Awake (hh:mm:ss)	00:42:00	1:02:20	00:33:00	1:23:20
Sleep Data – EM-171 (M, 40 y)	Pre-Stress Test (night 2)	Post-Stress Test (night 3)	Post-Stress Test (night 7)	Post-Stress Test Worse Night (#6)
Sleep Data – EM-171 (M, 40 y) Sleep Position Changes (#)	Pre-Stress Test (night 2) 16	Post-Stress Test (night 3) 22	Post-Stress Test (night 7) 21	Post-Stress Test Worse Night (#6) 76
Sleep Position Changes (#)	16	22	21	76
Sleep Position Changes (#) Total Sleep Time (hh:mm:ss)	16 07:54:40	22 08:27:40	21 06:18:00	76 05:57:00

7 MULTITASKING BIOMOLECULES (1) Study of ThrombOsPondin-1 in ME/CFS – STOP-ME







Dr. Wesam Elremaly



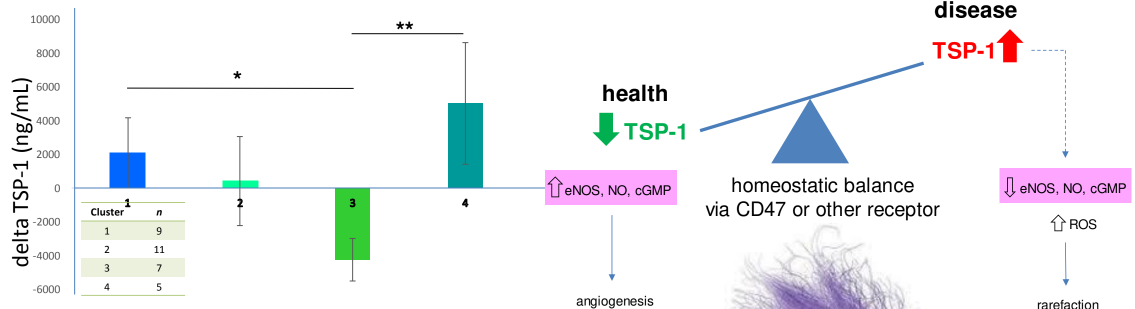
THE PROBLEM: Little is known about the mechanisms causing brain fog, orthostatic intolerance as well as postural orthostatic tachycardia (POTS) in ME/CFS.



OUR HYPOTHESIS: We propose that elevation of circulating thrombospondin-1 (TSP-1) levels could induce a brain fog and PEM in ME/CFS by reducing brain-blood flow. Conversely, a rapid decrease in blood TSP-1 levels could induce an orthostatic intolerance or even POTS.

ROLE OF THROMBOSPONDIN-1 IN ME/CFS?

Changes in plasma TSP-1 levels could be involved in ME/CFS pathogenesis



vasodilation

blood flow

- Cluster 3 encompasses ME/CFS patients showing a strong elevation of TSP-1 blood levels after the application of the stresstest. This subgroup including all ME/CFS patients exhibiting an orthostatic intolerance.
- Cluster 4 encompasses ME/CFS patients at-risk of developing brain fog.

what brain fog?

vasoconstriction

ischemia

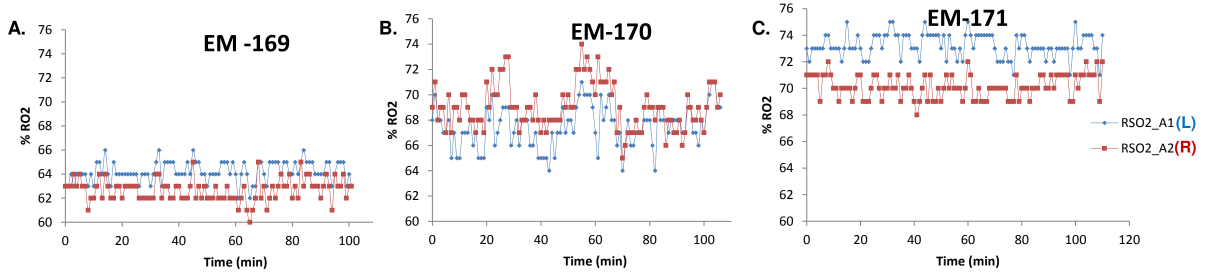
PRELIMINARY DATA

Changes in plasma TSP-1 levels and brain oxygen levels during the stress-test

 Table 3. Changes in plasma TSP-1 levels at different time points

Patient ID	Sex	Age (year)	TSP-1 at baseline (T0 min)	TSP-1 post-stress test (T90 min)	TSP-1 post-stress test (T+ 5 days)	PEM score (DSQ)	Medication
EM 169	F	42	25 665 ng/mL	16 956 ng/mL	13 993 ng/mL	92	
EM 170	F	50	18 926 ng/mL	18 602 ng/mL	10 038 ng/mL	65	Pregabalin
EM 171	М	40	8 054 ng/mL	17 325 ng/mL	7 718 ng/mL	86	Pregabalin (25mg+125mg) + Vit D3

Changes in brain oxygen levels during the stress-test



10 LONGITUDINAL NEUROCOGNITIVE ASSESSMENT Effects of plasma TSP-1 levels on neurocognitive functions

21/2020	Bra	InCheck	
	CLINICAL R ASSESSMENT DATE		BrainCheck
IDENTIFYING INFORMATION			T
Name: EM169c EM169c	DOB: 10/07/1977	Age: 42	Sex: F
NEUROCOGNITIVE ASSESSMENT			
BRAINCHECK COMBINED TEST RESULT	rs:		
\sim	14th Population Percer	ntile, LOW AVG	
84	Validity Test: PASS	N	Aalingering Test: PASS
STANDARD SCORE RANGE: 0-200		mpairment: POSSIBLE Cinica	al correlation warranted
			E
BRAINCHECK INDIVIDUAL TEST RESULT	5:		-
TTENTION			——————————————————————————————————————
Trails B	0	71st Population Percer	ntile, AVG
atients with impairment may struggle with novigating (s blowing a map, paying bills correctly, playing familiar ga brangly predict a decline in mobility and the inability to d	mes. Lower scores	and the second se	r indication of dysfunction
EXECUTIVE FUNCTION			
Digit Symbol Substitution	•	40th Population Perce	ntie, AVG
Patients with impoinment may struggle with paying atten eading, basic anthmetic. Lower scores have been associa		Impression: UNLIKELY	indication of dysfunction
need, ansiety, and substance use	LOW AN	C HICH Validity Test: Pass	
Stroop		83rd Population Perce	ntile , ABOVE AVG
blients with impointent may struggle with following co			r indication of dysfunction
ectsion making, poor judgment, socially inappropriate be ethdrawol, maintaining a healthy diet.	LOW AV	C HICH Validity Test. Pass	
MEMORY			
Immediate Recognition	-	1st Population Percent	tie, VERY LOW
Patients with imporment may struggle with repeating the one question repeatedly within a few minutes of each a		Impression: LIKELY in	ndication of dysfunction
hey were going to do, forgetting where they placed some themion to the TV		C HIGH Validity Test: Pass	
Delayed Recognition	0	5th Population Percent	tie, LOW
blents with impairment may struggle with repeating the		Impression: POSSIBLE	E indication of dysfunction
day or next day, forgetting the content of a conversation,	ar new ang to hely on a	11.0.0	



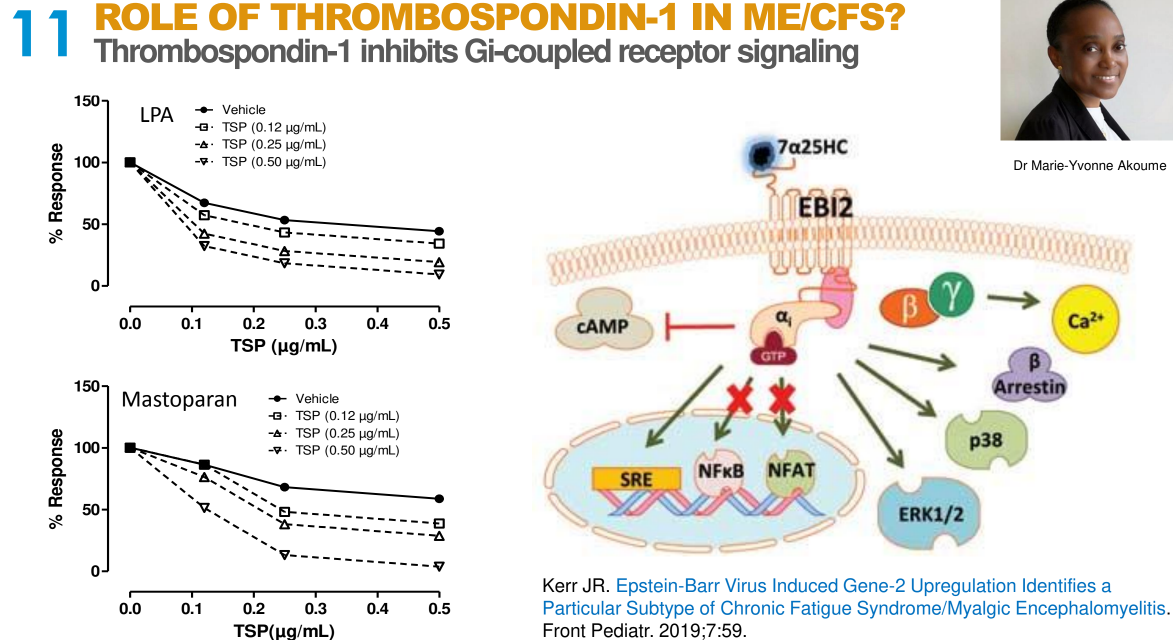
Dr. Wesam Elremaly

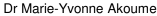
Table 1. Clinical and demographic data of participants

Patient ID	Sex	Age (year)	Illness duration (years)	Sleep score	Cognitive score	PEM score	ANI score
EM 169	F	42	6	28	61	92	33
EM 170	F	50	5	38	69	65	34
EM 171	М	40	3	53	60	86	41

Table 2. Clinical results with BrainCheck

Patient ID	At baseline (T0 min)	Post-stress test (T90 min)	Post-stress test (T+5 days)	Neurocognitive effects
EM 169	98	94	84	Likely a memory dysfunction
EM 170	83	96	101	Likely a memory dysfunction
EM 171	101	85	111	Possible executive function dysfunction





12 THERAPEUTIC OPTIONS FOR ME/CFS PATIENTS How to decrease plasma TSP-1 levels or block its signaling action?







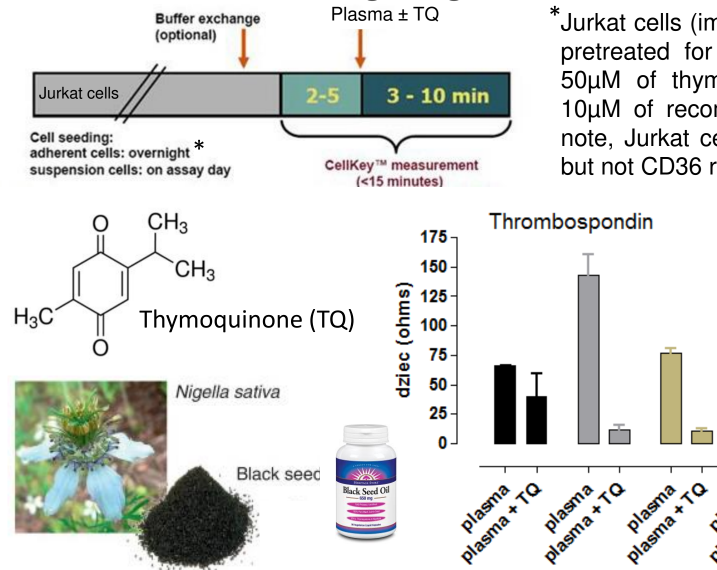
- $_{\odot}\,$ Interestingly, $\alpha 2\delta$ -1 is the high affinity receptor for TSP-1 in the brain.
- Two commonly prescribed anti-epileptic, anti-neuropathic pain medications, gabapentin (Neurontin[™]) and pregabalin (Lyrica[™]) are targeting α2δ-1 receptor. Both drugs are being used off-label for ME/CFS and fibromyalgia patients.
- Vitamin D3 supplementation for 12 weeks markedly reduced TSP-1 levels by almost 2.5 fold (522.7 ± 379.8 ng/mL vs 206.7 ± 204.5 ng/mL, p<0.001.¹
- Low-dose of cyclophosphamide.²
- Hyperbaric oxygenation therapy could be effective to decrease blood TSP-1 levels but it remains to be tested by a clinical trial. A direct link between TSP-1 activity and hyperoxic condition has not been made yet.³

 ¹ Amarasekera AT, et al. Vitamin D supplementation lowers thrombospondin-1 levels and blood pressure in healthy adults. *PLoS One*. 2017;12(5):e0174435.
 ² Lansiaux, A. et al. Circulating thrombospondin 1 level as a surrogate marker in patients receiving cyclophosphamide-based metronomic chemotherapy. Invest New Drugs 30, 403–404 (2012).

³ Asadamongkol B, Zhang JH. The development of hyperbaric oxygen therapy for skin rejuvenation and treatment of photoaging. Med Gas Res. 2014;4(1):7

ERAPEUTIC OPTIONS FOR ME/CFS PATI





Jurkat cells (immortalized human lymphocytes T) were pretreated for 2 hours with plasma with or without 50µM of thymoquinone (TQ). Then stimulated with 10µM of recombinant thrombospondin-1 proteins. Of note, Jurkat cells express $\alpha 2\delta$ -1 and CD47 receptors but not CD36 receptor.

plasma TO

125/12 * 10

plasma

plasma TQ

Control

G1 (EM-113)

G3 (EM-103)

G4 (EM-12)

G5 (EM-117)

G2 (EM-98)

4 **MULTITASKING BIOMOLECULES (2)** Role of SMPDL3B in ME/CFS pathophysiology



Bita Rostami



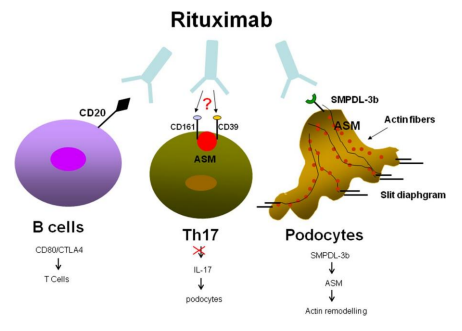
Dr. Wesam Elremaly



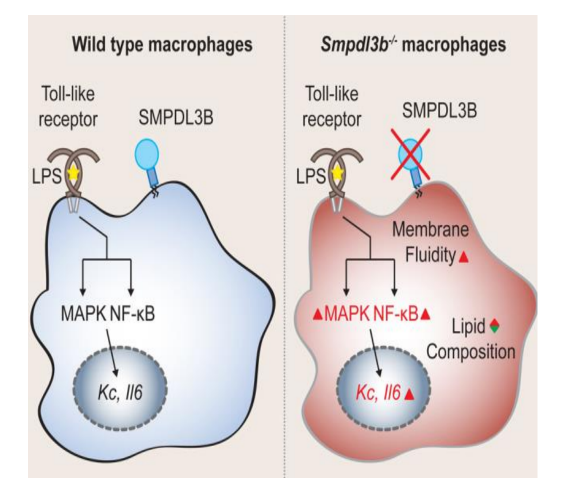
THE PROBLEM: Little is known about the mechanism underlying lipid metabolism alteration occurring in ME/CFS.



OUR HYPOTHESIS: We propose that sphingomyelin phosphodiesterase acid-like 3b (SMPDL3B) is involved in ME/CFS pathogenesis by modulating innate immunity and lipid metabolism. We have identified SMPDL3B as a possible alternative target of Rituximab in ME/CFS pathogenesis.



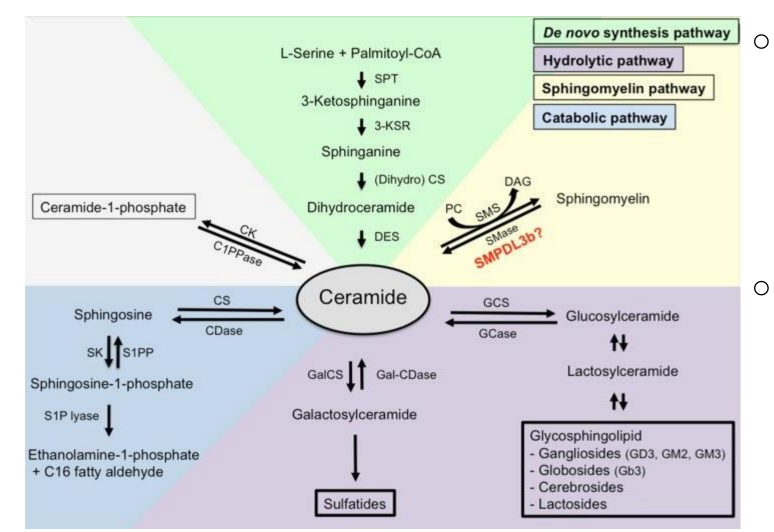
15 LIPID-MODIFYING ENZYME SMPDL3B Possible role of in the regulation of innate immunity in ME/CFS



- SMPDL3B expression is prominently observed in macrophages and DCs.
- Consistent with a possible role for this enzyme in the course of inflammatory processes.
- SmpdI3b transcription in bone marrow-derived macrophages (BMDMs) and DCs (BMDCs) is robustly induced upon TLR stimulation

Heinz LX, Baumann CL, Köberlin MS, et al. The Lipid-Modifying Enzyme SMPDL3B Negatively Regulates Innate Immunity. Cell Rep. 2015;11(12):1919-1928.

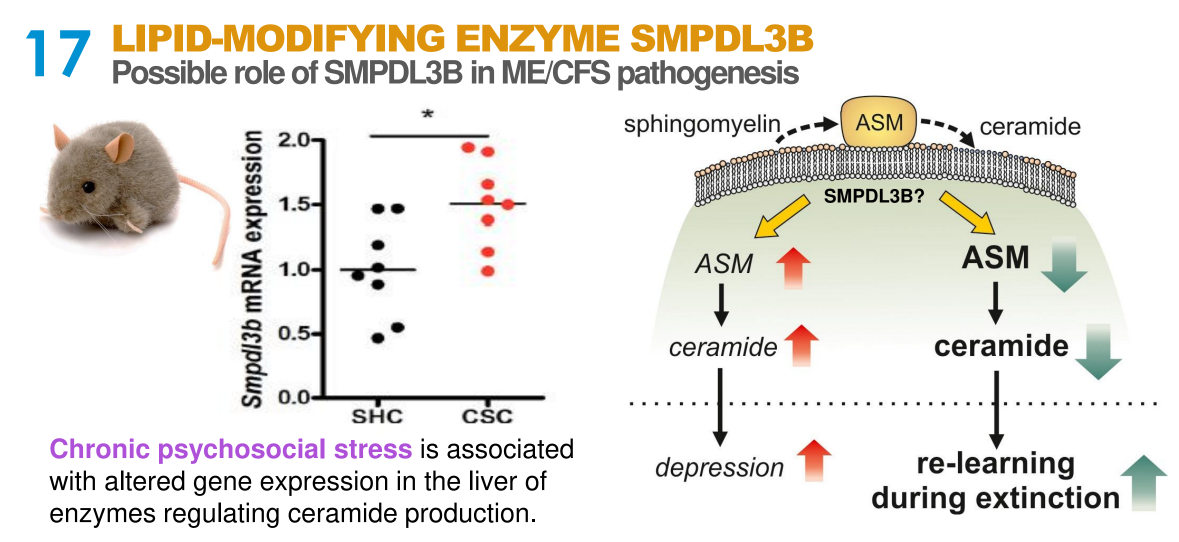
16 LIPID-MODIFYING ENZYME SMPDL3B SMPDL3B is a relevant molecule if ME/CFS pathogenesis



Merscher S, Fornoni A. Podocyte pathology and nephropathy - sphingolipids in glomerular diseases. Front Endocrinol (Lausanne). 2014;5:127.

In males, over 50% (16/30) of the sphingolipids that were decreased were ceramides, and 47% (14/30) were sphingomyelin species. In females, 86% (18/21) were ceramides and 14% (3/21) were sphingomyelins in females.

Naviaux R.K. et al. "Metabolic features of chronic fatigue syndrome." PNAS 2016; 113(37): E5472–E5480



Reichel M, Rhein C, Hofmann LM, et al. Chronic Psychosocial Stress in Mice Is Associated With Increased Acid Sphingomyelinase Activity in Liver and Serum and With Hepatic C16:0-Ceramide Accumulation. Front Psychiatry. 2018;9:496. Huston JP, Kornhuber J, Mühle C, et al. A sphingolipid mechanism for behavioral extinction. J Neurochem. 2016;137(4):589-603.

18 THERAPEUTIC OPTION FOR ME/CFS PATIENTS How to increase Smpdl3b gene expression?

List of top 5 up-regulated genes in Lateral Entorhinal Cortex after 7 days of AICAR administration (ACR7) and exercise (RUN7)

GENE	FOLD		
	ACR7	RUN7	
lgsf1	3.21	2.29	
Nr2f2	2.40	2.37	
Smpdl3b	2.35	2.10	
Tmie	2.19	1.93	
Gm4983	1.90	1.96	



AMPK agonist



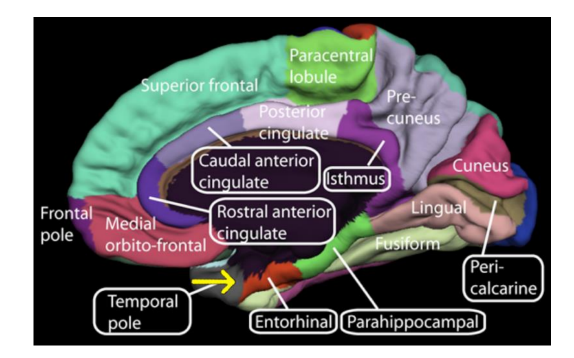
5-Aminoimidazole-4-carboxamide ribonucleotide

The entorhinal cortex (EC) is an area of the brain located in the medial temporal lobe and functions as a hub in a widespread network for memory, navigation and the perception of time. The EC is the main interface between the hippocampus and neocortex.

19 IMPORTANCE OF ENTORHINAL CORTEX Possible link between memory, bodily sensation and fatigue rating in ME/CFS

- The parahippocampal gyrus (PaHcG), which includes the entorhinal cortex, is involved in aspects of limbic function as well as memory retrieval and storage.
- Reduced connectivity in ME/CFS participants between PaHcG and regions that encompassed left postcentral gyrus (i.e., primary sensory cortex) and supramarginal gyrus suggests abnormality in the link between memory and bodily sensation.¹
- Such lower connectivity was strongly correlated to higher fatigue ratings of ME/CFS participants.¹

¹ Boissoneault J, Letzen J, Lai S, et al. Abnormal resting state functional connectivity in patients with chronic fatigue syndrome: an arterial spin-labeling fMRI study. *Magn Reson Imaging*. 2016;34(4):603-608.



By Hagmann P, Cammoun L, Gigandet X, Meuli R, Honey CJ, et al. https://commons.wikimedia.org/w/index.php?curid=8636113

20 PRELIMINARY DATA Possible role of SMPDL3B in ME/CFS pathogenesis

Hypermethylation of SMPDL3B gene occurs in ME/CFS



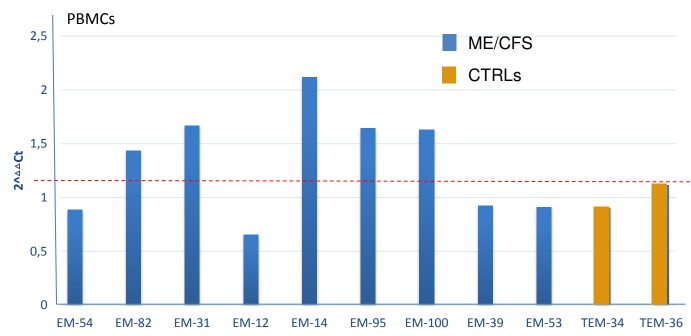


Lynda Chalder Dr. Dawei Li

Family #14

F14	bValDiff	absbValDiff	F14-01 ME	F14-02 Healthy	gene
cg05320933	0,3630	0,3630	0,6202	0,2572	SMPDL3B
cg23448720	0,3148	0,3148	0,4982	0,1834	SMPDL3B

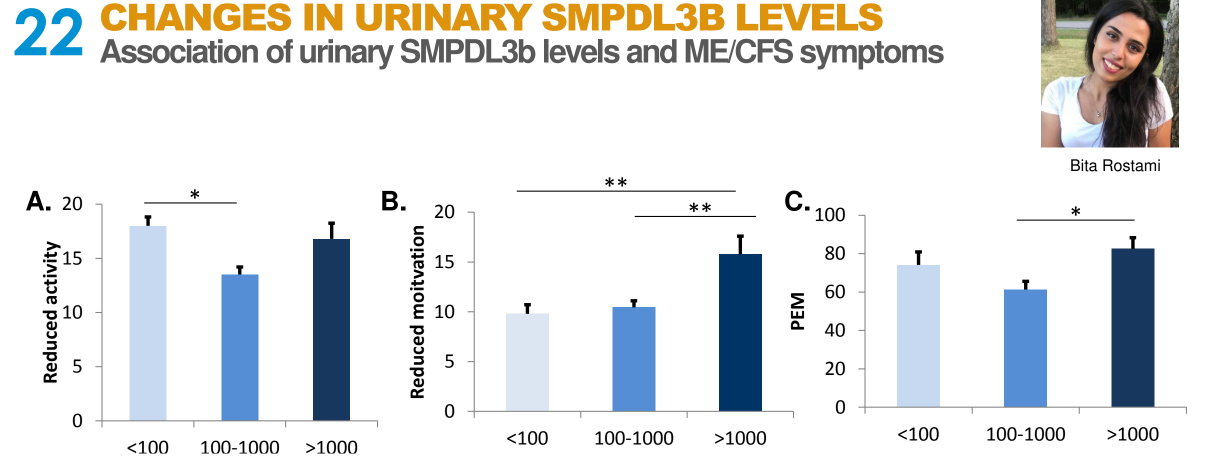
SMPDL3B² expression





Bita Rostami

MAB EFFECTS IN ME/CFS? 21 MiR-140-5p expression could influence Rituximab bioavailability Α. Β. Mixed connective tissue disease miR-140-5p Mean Relative Expression Evguenia Nepotchatykh Mature T-cell neoplasm Chronic fatigue syndrome p = 0.008p = 0.0308 CO (1) Stage II monocytoid B-cell lymphoma Relative Expression 6 CO (10) CO (1) Advanced stage diffuse large B-cell lymphoma CO (2) CO (1) KRT20 (CD20) 0 ME baseline **CTRL** Baseline ME 90min CTRL 90min -1 Baseline 90min miR-140-5p Rituximab С. 8,1 8.0 5,6 6,0 4,4 4,3 4,0 4,0 2,7 2,3^{2,5} 2,4 2,3 2,0 2,0 1,8 1,6 2,0 1.3 1,2 0,6^{0,9} 1,0 8,0 0,90.8 0,8 07 0,4_{0,3} 0,20,4 0,0^{0,2}).4 0.0 EM-12-base EM-12-90min EM-14-BASE EM-14-90MIN EM-17-base EM-17-90MIN EM-19-base EM-19-90MIN EM-31-BASE EM-31-90MIN EM-32-base EM-32-90min EM-34-BASE EM-34-90min EM-36-base EM-36-90min EM-39-base M-39-90min EM-41-base EM-41-90min EM-53-base EM-53-90min EM-54-base EM-54-90min EM-61-base EM-61-90min EM-71-base EM-71-90min EM-74-base EM-74-90min EM-77-base EM-77-90min EM-85-base EM-85-90min EM-90-base EM90-90min EM-92-base EM-92-90min EM-95-base EM-95-90min EM-100-base EM-100-90min EM-109-base EM-109-90min EM-16-base EM-16-90min EM-23-base EM-23-90min EM-56-base EM-56-90min EM-58-base EM-58-90min EM-59-base EM-59-90min EM-70-base EM-70-90min EM-82-base EM-82-90min EM-98-base EM-98-90min



Classification of urinary SMPDL3B into <100, 100-1000 and > 1000 ng/ml/mg creatinine and DSQ questionnaires

23 ACKNOWLEDGEMENTS

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Anita Franco



Lynda Chalder



Evguenia Nepotchatykh



Viorica Lascau



Dr. Marie-Yvonne Akoume



Dr. Iurie Caraus





Bita Rostami





Sophie Perreault



Corinne Leveau

Dr. Wesam Elremaly

Dr. Dawei Li