

c	THURSDAY, MARCH 21, 2024
08:10-10:50	Neuroimmunology 1 HALL A
Chairs:	Bruno Gran, UK; Olaf Stuve, USA
08:10-09:00	Is Hashimoto's encephalitis/encephalopathy a valid construct in 2024?
	Capsule: Hashimoto's encephalopathy is a rare condition manifesting with variety of symptoms ranging from disturbances of consciousness, seizures, myoclonus
	to rapidly progressive cognitive decline observed in euthyroid patients with anti-thyroid antibodies. It is a steroid-responsive disorder. However, majority of
	described cases are from the period before tests for novel antibodies were available. Anti-thyroid antibodies in a patient with encephalopathy could be an
	incidental finding. Is therefore Hashimoto's encephalitis/encephalopathy a valid construct in 2024? Moderator: Uros Rot , Slovenia
08:10-08:20	Introduction and Pre-Debate Voting
08:20-08:35	Yes: Alasdair Coles, UK
08:35-08:50	No: Divyanshu Dubey, USA
08:50-09:00	Discussion, Rebuttals and Post-Debate Voting
09:00-09:50	Is long COVID an autoimmune disease?
	Capsule: Long COVID refers to diverse symptoms, neurological and otherwise, that follow COVID-19 infection. The existence of this condition as a unique syndrome and its cause(s) remain uncertain. Is there reason to believe that long COVID is an autoimmune disease?
09:00-09:10	Moderator: Thomas Pollak, UK
09.00-09.10	Introduction and Pre-Debate Voting
09:10-09:25	Yes: <u>Michael D. Geschwind,</u> USA
09:25-09:40	No: <u>Hans-Peter Hartung</u> , Germany
09:40-09:50	Discussion, Rebuttals and Post-Debate Voting
09:50-10:50	A prolonged (1 year) corticosteroid taper is sufficient to prevent relapses in patients with MOGAD
	Capsule: MOGAD is a monophasic or relapsing disease associated with MOG-IgG autoantibodies, manifesting primarily as optic neuritis or as acute disseminated
	encephalomyelitis in children. It is controversial whether to start relapse prevention treatments following a first episode of MOGAD. Some experts have suggested
	that a prolonged taper of corticosteroids over 1 year is needed tol reduce the risk of relapses in MOGAD. How strong is the evidence and is this a recommendation that should be endorsed?
	Moderator: Joab Chapman, Israel
09:50-10:00	Introduction and Pre-Debate Voting
10:00-10:15	Yes: Michael Levy, USA
10:15-10:30	No: Brian G. Weinshenker, USA
10:30-10:50	Discussion, Rebuttals and Post-Debate Voting
10:50-11: 20	Coffee Break, Exhibition & ePosters Visits



THURSDAY, MARCH 21, 2024			
11: 20-12:00	Opening Ceremony Plenary Hall		
11:20-11:25	Natan Bornstein, Israel		
11:25-11:30	Amos Korczyn, Israel		
11:30-11:35	Anthony Schapira, UK		
11:35-11:40	Richard J Davenport, UK		
11:40-12:00	Best e Posters award		
12:00 -13:00	Plenary Session Plenary Hall		
Chairs:	Brian G. Weinshenker, USA; John Hardy, UK		
12:00-12:10	CONy Excellence in Neurology Award to Prof. Angela Vincent.		
12:10-13:00	Antibody-mediated diseases: past, present and questions for the future <u>Angela Vincent</u> , UK		
13:00-14:00	Industry Sponsored Symposium Plenary Hall		
14:10-15:00	MTE		
14:00-15:00	Lunch Break, Exhibition & ePosters Visits		



THURSDAY, MARCH 21, 2024			
15:00-16:40	Antibody testing HALL A		
Chair:	Brian G. Weinshenker, USA		
15:00-15:50	All patients with inflammatory optic neuritis should be screened for AQP4-IgG and MOG-IgG antibodies		
	Capsule: Optic neuritis has discriminating clinical and paraclinical characteristics, different responses to treatment and prognosis in patients with MS, NMOSD of		
	MOGAD but there is a significant clinical overlap between the entities. Immune treatment differs markedly between them. Should we therefore screen all		
	patients with inflammatory optic neuritis for AQP4-IgG and MOG-IgG antibodies?		
15:00-15:10	Moderator: Ruth Geraldes, UK		
15.10 15.25	Introduction and Pre-Debate Voting		
15:10-15:25	Yes: <u>Uros Rot</u> , Slovenia		
15:25-15:40	No: Saif Huda, UK		
15:40-15:50	Discussion, Rebuttals and Post-Debate Voting		
15:50-16:40	Is it sufficient to send focused antibody testing on patients with suspected autoimmune encephalitis, or should all patients be screened with a panel of antibody		
	tests?		
	Capsule: Autoimmune encephalitis (AE) is frequently associated with antibodies against neuronal, synaptic or glial proteins. Several clinical syndromes of AE		
	have been reported to date. Correct diagnosis of AE depends on the disease phenotype, exclusion of alternative cause (infection, metabolic and neuropsychiatric)		
	and identification of a specific antibody in serum and CSF. Both missed diagnosis and misdiagnosis of AE are recognised, and seronegative AE has been reported.		
	Detection of serum antibody alone may be of uncertain significance without the clinical phenotype. Should antibody testing in suspected AE be focused on the		
	specific syndrome or routinely carried out on an extensive panel of autoimmune and paraneoplastic antibodies?		
15:50-16:00	Moderator: <u>Abhijit Chaudhuri</u> , UK		
	Introduction and Pre-Debate Voting		
16:00-16:15	Yes: <u>Eoin Flanagan</u> , USA		
16:15-16:30	No: <u>Ruth Geraldes,</u> UK		
16:30-16:40	Discussion, Rebuttals and Post-Debate Voting		
16:40- 17:00	Coffee Break, Exhibition & ePosters Visits		



THURSDAY, MARCH 21, 2024		
17:00-18:40	Corticosteroid treatment	HALL A
Chair:	Boleslav Lichterman, Russia	
17:00-17:50	Diabetes mellitus-associated plexopathy is an inflammatory vasculitis and should be treated with high dose corticosteroids	
	Capsule: An infrequent but very disabling complication of diabetes mellitus is lumbosacral plexopathy (also known as diabetic asymmetrical accompanied by prominent pain and proximal weakness. The pathogenesis is controversial, but nerve biopsies microvasculitis with ischemic and inflammatory changes. Corticosteroids can be very effective in alleviating pain and many ex advocate this treatment in spite of the lack of definite data	have demonstrated evidence of
17:00-17:10	Moderator: Divyanshu Dubey , USA	
17.10 17.25	Introduction and Pre-Debate Voting	
17:10-17:25	Yes: <u>P. James B. Dyck</u> , USA	
17:25-17:40	No: <u>Alasdair Coles</u> ,UK	
17:40-17:50	Discussion, Rebuttals and Post-Debate Voting	
17:50-18:40	Cerebral amyloid angiopathy (CAA) may lead to inflammatory vasculopathy; patients with cerebral amyloid angiitis should re diagnosis.	ceive corticosteroids on
	Capsule: CAA is a vasculopathy characterised by amyloid beta (AB) deposition in cortical and meningeal blood vessels. Cerebra an inflammatory response, leading to perivascular inflammation and vasculitis. Acute, subacute, as well as chronic or progress neurological dysfunction are recognised in CAA and often attributed to the inflammatory response. Early corticosteroid there beneficial in cerebral amyloid angiitis.	sive focal and multifocal
17:50-18:00	Moderator: Angela Vincent, UK	
18:00-18:15	Introduction and Pre-Debate Voting Yes: Joab Chapman, Israel	
18:15-18:30	No: <u>Abhijit Chaudhuri</u> , UK	
18:30-18:40	Discussion, Rebuttals and Post-Debate Voting	
18:45	Networking Reception (Exhibition Area)	



08:10 - 10:50	Alzheimer's disease (AD): Biomarkers HALL B
Chairs:	Marina Janelidze, Georgia; Claire Sexton ,USA
08:10-09:00	Biomarkers are useful in subjective cognitive complaints and should be tested in each patient
	<i>Capsule:</i> Patients with SCC are at increased risk to develop dementia. it is important to identify who is at risk. Are there any biomarkers which van help?
08:10-08:20	Moderator: <u>Thomas C. Neylan</u> , USA Introduction and Pre-Debate Voting
08:20-08:35	Yes: Paul Edison, UK
08:35-08:50	No <u>: Zvezdan Pirtosek</u> , Slovenia
08:50-09:00	Discussion, Rebuttals and Post-Debate Voting
09:00-09:50	Are serum markers such as phospho-tau useful in diagnosing AD ?
	Capsule: In a chronic medical condition, early diagnosis becomes important when treatment is available that can alter its course. Regarding AD, there is hope that drugs or prevention strategies will have the capacity of slowing down the neurodegeneration. Such treatments may provide greatest benefit to early stage since higher levels of functioning, independence, and quality of life will be maintained. Blood-based biomarkers would be critical in making early diagnosis accessible in routine clinical care. This debate will focus on the central question whethe
	AD can (and should) be diagnosed early based on biomarkers measured in blood.
09:00-09:10	Moderator: <u>Xiaoping Wang</u> , People's Republic of China Introduction and Pre-Debate Voting
09:10-09:25	Yes: Robert Perneczky, Germany
09:25-09:40	No: Arfan Ikram, The Netherlands
09:40-09:50	Discussion, Rebuttals and Post-Debate Voting
09:50-10:50	Sleep, Alzheimer's and Dementia – session in cooperation with Alzheimer's Association
09:50-09:55	Moderators: <u>Claire Sexton</u> , USA; <u>Lea Grinberg,</u> USA
09:55-10:10	Sleep in clinical populations: Thomas C. Neylan, USA
10:10-10:25	Neuropathology and neuroimaging of sleep: <u>Neus Falgas</u> , Spain
10:25-10:40	Sleep as risk factor - evidence for interventions: Sharon Naismith, Australia
10:40-10:50	Panel Discussion
10:50 -11:20	Coffee Break, Exhibition & ePosters Visits



	THURSDAY, MARCH 21, 2024	
11:20-12:00	Opening ceremony (Plenary Hall)	
12:00-13:00	Plenary Session (Plenary Hall)	
13:00-14:00	Industry Sponsored Symposium (Plenary Hall)	
14:10-15:00	MTE	
14:00-15:00	Lunch Break, Exhibition & ePosters Visits	
15:00-16:40	AD: Therapy HALL B	
Chairs:	<u>Panteleimon Giannakopoulos</u> , Switzerland ; <u>Yvonne Freund-Levi</u> , Sweden	
15:00-15:50	Obstructive sleep apnea is detrimental in patients with dementia and should always be treated	
	Capsule: An overwhelming body of work suggests that obstructive sleep apnea is more prevalent in patients with dementia and may be one of the risks for development of dementia. Whilst the exact mechanics of this bidirectional relationship are not fully understood, several studies advocate that early diagnosis, and early treatment of sleep apnea in patients with dementia may improve their quality of life, and possibly also decelerate the neurodegenerative process. In this debate the major limitations and/or potential contraindications, as well as the most promising aspects of OSA-treatment approach will be discussed.	
15:00-15:10	Moderator: <u>Michael D. Geschwind</u> , USA Introduction and Pre-Debate Voting	
15:10-15:25	Yes: Ivana Rosenzweig, UK	
15:25-15:40	No: Sharon Naismith, Australia	
15:40-15:50	Discussion, Rebuttals and Post-Debate Voting	
15:50-16:40	Is cognitive reserve a useful term?	
	Capsule: The concept of reserve was established to account for the observation that a given degree of neurodegenerative pathology may result in varying severities in different individuals. There is a large amount of evidence on epidemiological risk and protective factors for neurodegenerative diseases and dementia, yet the biological mechanisms that underpin the protective effects of certain lifestyle and physiological variables remain poorly understood, limiting the development of more effective preventive and treatment strategies. Additionally, different definitions and concepts of reserve exist, which hampers the coordination of research and comparison of results across studies. Is cognitive reserve just another buzz word or is the phenomenon supported by enough scientific evidence?	
15:50-16:00	Moderator: <u>Robert Perneczky</u> , Germany Introduction and Pre-Debate Voting	
16:00-16:15	Yes: <u>Yaakov Stern</u> , USA	
16:15-16:30	No: <u>Amos Korczyn</u> , Israel	
16:30-16:40	Discussion, Rebuttals and Post-Debate Voting	
16:40- 17:00	Coffee Break, Exhibition & ePosters Visits	



THURSDAY, MARCH 21, 2024		
17:00-18:40	AD 3	HALL B
Chairs:	Judith Aharon Peretz, Israel; Milica G. Kramberger, Slovenia	
17:00-17:50	Antiamyloid drugs have only a very limited effect and will not be clinically useful for most patients	
	Capsule: Several large clinical trials have demonstrated potential utility of amyloid-targeting approaches in slowing the pr change the course of the disease in some people in the early stages, giving them more time to participate in daily life. How treatments have also been shown to have significant side effects and high cost. In this debate the major limitations as we amyloid-targeting approach will be discussed.	vever, while promising, these
17:00-17:10	Moderator: John Hardy, UK	
17:10-17:25	Introduction and Pre-Debate Voting Yes: Dorota Religa , Sweden	
17:25-17:40	No: <u>Paul Edison</u> , UK	
17:40-17:50	Discussion, Rebuttals and Post-Debate Voting	
17:50-18:40	Should lecanemab use be extended beyond 18 months?	
	Capsule: There is only one phase 3 randomized trial of lecanemab and one of donanemab. Both were undertaken to support have uncontrolled, long-term extended treatment options provided for participants who completed the 18 month trials and clinical effects of these antibodies are small and dropouts and adverse events fairly common a question arises about whet beyond the length of the trials and whether any clinical benefit might become apparent over the long-term. Only a few hur exposed to these antibodies beyond 18 months; and no regular clinic patient in the USA or Japan could have been exposed months. Thus this issue is ripe for debate as evidence is sparse or absent. This debate might highlight what needs to be contreatment.	d wished to continue treatment. As the her treatment should be continued ndred clinical trials patients have been I to lecanemab for more than 8
17:50-18:00	Moderator: Zvezdan Pirtosek, Slovenia	
18:00-18:15	Yes: Dorota Religa , Sweden	
18:15-18:30	No: Lon Schneider , USA	
18:30-18:40	Discussion, Rebuttals and Post-Debate Voting	



THURSDAY, MARCH 21, 2024			
08:10 - 10:50	Parkinson's disease (PD)	HALL C	
Chairs:	<u>Cristian Falup-Pecurariu</u> , Romania; <u>Daniel Weintraub</u> , USA		
08:10-09:00	Has the DLB vs. PD(D) distinction outlived its usefulness?		
	Capsule: The timing of dementia relative to parkinsonism is the major clinical distinction between DLB and PD. There is evid		
	neuropsychological, and neuropathological features of DLB and PD, remarkably convergent neuropathologic changes at au		
	believe that PD and DLB are different phenotypic expressions of the same underlying process and therefore, maybe it is time to consider that the DLB vs. PD(D)		
	distinction outlived its usefulness?		
08:10-08:20	Moderator: Nestor Galvez-Jimenez, USA		
08:20-08:35	Introduction and Pre-Debate Voting		
	Yes: <u>Lea Grinberg</u> , USA No: Vladimira Vuletic, Croatia		
08:35-08:50			
08:50-09:00	Discussion, Rebuttals and Post-Debate Voting		
09:00-09:50	With the limitation of dopamine-based therapy, should therapy in PD be directed in other directions?		
	Capsule: PD is not only a disease with a shortage of dopamine but is a multi-transmitter disease. Should other options than	simply dopamine replacement be	
	discussed, or does levodopa still have untapped potential?		
09:00-09:10	Moderator: Amos Korczyn, Israel		
00.40.00.05	Introduction and Pre-Debate Voting		
09:10-09:25	Yes: Jaroslaw Slawek, Poland		
09:25-09:40	No: <u>Heinz Reichmann</u> , Germany		
09:40-09:50	Discussion, Rebuttals and Post-Debate Voting		
09:50-10:50	Is it possible to modify the disease course in PD?		
	Capsule: As we may soon get options to diagnose PD in its prodromal phase, disease-modifying therapies may get a second	l chance.	
09:50-10:00	Moderator: Avner Thaler, Israel		
09.30-10.00	Introduction and Pre-Debate Voting		
10:00-10:15	Yes: <u>Vladimira Vuletic</u> , Croatia		
10:15-10:30	No: Jaroslaw Slawek ,Poland		
10:30-10:50	Discussion, Rebuttals and Post-Debate Voting		
10:50 -11:20	Coffee Break, Exhibition & ePosters Visits		



THURSDAY, MARCH 21, 2024		
11:20-12:00	Opening Ceremony (Plenary Hall)	
12:00-13:00	Plenary Session (Plenary Hall)	
13:00-14:00	Industry Sponsored Symposium (Plenary Hall)	
14:10-15:00	MTE	
14:00-15:00	Lunch Break, Exhibition & ePosters Visits	
15:00-16:40	Precision therapies in PD HALL C	
Chairs:	Gilad Yahalom, Israel; Weidong Le, China	
15:00-15:50	Are we ready for precision medicine in PD?	
	Capsule: Developments in -omic technologies as well as deep phenotyping support the heterogeneity of PD risk and progression. However, this has not yet translated into "personalized medicine" outside of the research environment. Are we ready yet to tailor disease prevention and/or treatment according to different disease profiles?	
15:00-15:10	Moderator: <u>Yoav Ben-Shlomo</u> , UK Introduction and Pre-Debate Voting	
15:10-15:25	Yes: <u>K. Ray Chaudhuri,</u> UK	
15:25-15:40	No: <u>Evzen Ruzicka</u> , Czech Republic	
15:40-15:50	Discussion, Rebuttals and Post-Debate Voting	
15:50-16:40	GBA targeted therapies are a waste of money	
	Capsule: Genetic variants of the GBA1 gene are the commonest genetic risk factor for PD. They result in a number of specific biochemical alterations including lysosomal and mitochondrial dysfunction and accumulation of alpha-synuclein. Is the GBA1 gene and its product (glucocerebrosidase) a reasonable target for therapies to modify the course of PD?	
15:50-16:00	Moderator: <u>Ziv Gan-Or</u> , Canada Introduction and Pre-Debate Voting	
16:00-16:15	Yes: <u>Avner Thaler</u> , Israel	
16:15-16:30	No : <u>Anthony Schapira</u> , UK	
16:30-16:40	Discussion, Rebuttals and Post-Debate Voting	
16:40- 17:00	Coffee Break, Exhibition & ePosters Visits	



THURSDAY, MARCH 21, 2024		
17:00-18:40	PD 3	HALL C
Chairs:	Yehonatan Sharabi, Israel; Ilana Schlesinger, Israel	
17:00-17:50	Telemedicine is valuable for PD patients care and will become the predominant method	
	Capsule: The COVID-19 pandemia forced many of us to start applying telemedicine. It is also known for many years that whole day gives a better insight than taking the history from the patient. Thus, telemedicine may be an option for many	
17:00-17:10	Moderator: <u>Yoav Ben-Shlomo</u> , UK Introduction and Pre-Debate Voting	
17:10-17:25	Yes: <u>Heinz Reichmann</u> , Germany	
17:25-17:40	No: Anthony Schapira, UK	
17:40-17:50	Discussion, Rebuttals and Post-Debate Voting	
17:50-18:40	Should idiopathic REM-sleep behavior disorder (iRBD) patients be informed about potential PD prognosis as long as ther ?	e is no disease modifying therapy (DMT)
	Capsule: iRBD is linked with an increased risk of PD and other alpha-synucleinopathies, but presently there is no consense patients However, presently, there is no proven neuroprotective strategy, or DMT, to prevent the development of neur limited data concerning counselling of iRBD patients. What are the potential ethical and clinical conundrums in prognost	ological deficits and there are only very
17:50-18:00	Moderator: K. Ray Chaudhuri, UK	
18:00-18:15	Yes: Ivana Rosenzweig, UK	
18:15-18:30	No: Danielle Wasserman Berkovitz, Israel	
18:30-18:40	Discussion, Rebuttals and Post-Debate Voting	



FRIDAY, MARCH 22,2024			
07:40-08:30	ePoster Guided Tour		
08:30-11:00	STROKE: Acute stroke	HALL B	
Chairs:	<u>Vida Demarin</u> , Croatia ; <u>Natan Bornstein</u> , Israel		
08:30-09:20	In people with intracerebral hemorrhage (ICH) minimally invasive neurosurgery should be routinely discussed		
	Capsule: Acute stroke due to supratentorial intracerebral hemorrhage is associated with high morbidity and mortality. Oper		
	evacuation has not been found to have any benefit in large, randomized trials. Recently minimally invasive catheter evacuation		
	with the aim of decreasing clot size and iron toxicity showed promising results . Whether minimally invasive surgery should be routinely discussed in patients with ICU is the storie of this debate		
08:30-08:40	with ICH is the topic of this debate.		
08.30-08.40	Moderator: <u>Robert Mikulik,</u> Czech Republic Introduction and Pre-Debate Voting		
08:40-08:55	Yes: Marina Roje Bedekovic, Croatia		
08:55-09:10	No: Laszlo Csiba, Hungary		
09:10-09:20	Discussion, Rebuttals and Post-Debate Voting		
09:20-10:10	Mecanical Thrombectomy (MT) for large core infarcts is a worthwhile use of health care resources.		
	Capsule: Patients with large infarct core on baseline imaging were excluded from MT studies due to their assumed poor ou		
	reperfusion therapies are safe and beneficial in this group of patients. The debate will try to answer the open questions: What is the best imaging modality		
00.00.00.00	to diagnose the large infarct core? Can analysis of risk-benefit balance justify MT for this group of patients? How can the risk of the second s	sk of complications be reduced?	
09:20-09:30	Moderator: <u>Laszlo Csiba,</u> Hungary Introduction and Pre-Debate Voting		
09:30-09:45	Yes: <u>Ashfaq Shuaib</u> ,Canada		
09:45-10:00	No: Amith Sitaram, UK		
10:00-10:10	Discussion, Rebuttals and Post-Debate Voting		
10:10-11:00	Should patients with mild non-disabling stroke admitted within 4.5h be considered for thrombolysis?		
	Capsule: The role of thrombolysis in treatment of patients with AIS with low National Institutes of Health Stroke Scale (NIHS	SS) scores is not well understood.	
	However, in the absence of definitive evidence, practice pattern is widely variable. Here we debate the benefit of mild through through the benefit of mild the benefit of		
	most recent evidence.	·	
10:00-10:20	Moderator: Robert Mikulik, Czech Republic		
10:20-10:35	Yes: <u>Ashfaq Shuaib,</u> Canada		
10:35-10:50	No: Jesse Dawson, UK		
10:50-11:00	Discussion, Rebuttals and Post-Debate Voting		
11:00-11:30	Coffee Break, Exhibition & e-Posters Visit		



FRIDAY, MARCH 22,2024			
11:30-13:30	Plenary Session	Plenary Hall	
Chairs:	Lea Grinberg USA/Brazil ; Ivana Rosenzweig, UK		
11:30-12:00	Sleep disorders and brain health Claudio Bassetti, Switzerland		
12:00-12:30	The use of AI in neurology . <u>Sanjay Budheo</u> , UK		
12:30-13:00	Functional disorders, with a focus on Functional Cognitive Disorder. Narinder Kapur, UK		
13:00-13:30	The problem with Alzheimer's disease. Amos Korczyn, Israel		
13:40-14:30	MTE		
13:30-14:30	Lunch Break, Exhibition & e-Posters Visit		
14:30-16:10	STROKE: Prevention and recovery	HALL B	
Chair:	Marina Roje Bedekovic, Croatia		
14:30-15:20	Is there an added value to use virtual reality in rehabilitation after stroke?		
	Capsule: Within the last 10 years there is growing interest in possible applications of advanced technolgies such as virtu have been made to use these techniques in motor, cognitive and emotional therapies in rehabilitation after stroke. It is offer additional value for patients or may even be counterproductive taking away resources which could be used better.	however controversial if this will	
14:30-14:40	Moderator: <u>Nirmal Surya</u> , India Introduction and Pre-Debate Voting		
14:40-14:55	Yes: Volker Hoemberg, Germany		
14:55-15:10	No: Dafin Fior Muresanu , Romania		
15:10-15:20	Discussion, Rebuttals and Post-Debate Voting		
15:20-16:10	Should GLP-1 agonists be used to reduce stroke risk in non-diabetic obese individuals?		
	Capsule: There is evidence from studies in diabetes that GLT-1 receptor agonist treatment may reduce cardiovascular ev study in obese people with no evidence of diabetes reveals that GLP-1 receptor agonists may also be effective in the prev		
	obese individuals. Should GLP-1 receptor agonists be used in obese non-diabetic individuals for stroke prevention?		
15:20-15:30	Moderator: <u>Vida Demarin</u> , Croatia		
15:30-15:45	Introduction and Pre-Debate Voting		
	Yes: <u>Natan Bornstein</u> , Israel		
15:45-16:00	No: <u>Jesse Dawson</u> , UK		
16:00-16:10	Discussion, Rebuttals and Post-Debate Voting		
16:10-16:30	Coffee Break, Exhibition & e-Posters Visit		



	FRIDAY, MARCH 22,2024	
16:30-18:10	STROKE: Prevention and recovery HALL B	
Chair:	Zrinka Hrgovic, Croatia	
16:30-17:20	Should we aggressively treat asymptomatic small vessels disease (SVD) with drugs?	
	Capsule: Asymptomatic SVD is perhaps the most common abnormality noted on CT and MR imaging in the elderly population. The presence of SVD incr the risk of symptomatic stroke, dementia, falls, and early mortality. Thus far no definite treatments have been shown to slow the progression of SVD. SI patients with SVD be treated aggressively with antiplatelet medications or is risk factor management sufficient to manage patients in whom SVD is pre on brain imaging?	hould
16:30-16:40	Moderator: <u>Ashfaq Shuaib</u> , Canada Introduction and Pre-Debate Voting	
16:40-16:55	Yes: Natan Bornstein, Israel	
16:55-17:10	No: <u>Alan Cameron</u> ,UK	
17:10-17:20	Discussion, Rebuttals and Post-Debate Voting	
17:20-18:10	Can authorized alternative nicotine delivery systems be conducive in the effort of risk mitigation for smokers in high-risk patients?	
	Capsule: Capsule: Smoking is an important risk factor for multiple pathologies including atherosclerosis and stroke,. Whether heavy smokers can benefi switching from combustible tobacco smoking to alternative nicotine consumption is a promising approach for such patients.	t by
17:20-17:30	Moderator: Natan Bornstein, Israel	
17:30-17:45	Yes: <u>Yehya Orsan</u> , Israel	
17:45-18:00	No: Laszlo Csiba, Hungary	
18:00-18:10	Discussion, Rebuttals and Post-Debate Voting	



	FRIDAY, MARCH 22, 2024
07:40-08:30	ePoster Guided Tour
08:30-11:00	HEADACHE : Migraine and cluster headache HALL C
Chairs:	Vlasta Vukovic Cvetkovic, Denmark ;Ruta Mameniskiene, Lithuania
08:30-09:20	Are anti-CGRP mAbs effective in prevention of cluster headache?
	Capsule: Cluster headache is a serious medical condition that lacks disease-specific and mechanism-based treatments. There is some evidence that galcanezumab may be effective in reducing the frequency of cluster headache attacks, but the evidence is weak. Should CGRP mAbs be used in people with episodic cluster headache?
08:30-08:40	Moderator: <u>Dimos Mitsikostas</u> , Greece Introduction and Pre-Debate Voting
08:40-08:55	YES: <u>Giorgio Lambru</u> , UK
08:55-09:10	NO: Hakan Ashina, Denmark
09:10-09:20	Discussion, Rebuttals and Post-Debate Voting
09:20-10:10	Premonitory symptoms in migraine and cluster headache are important for treatment
	Capsule: Premonitory symptoms in migraine and cluster headache involve activation of the central parts of the trigeminovascular system (TVS). Whether preventive migraine treatments acting on the peripheral parts of the TVS can reduce the incidence of premonitory symptoms, not just the incidence of headache attacks, in people with migraine and/or cluster headache, remains an attractive hypothesis.
09:20-09:30	Moderator: <u>Cristina Tassorelli</u> , Italy Introduction and Pre-Debate Voting
09:30-09:45	YES: <u>Anna Andreou,</u> UK
09:45-10:00	NO: Hakan Ashina, Denmark
10:00-10:10	Discussion, Rebuttals and Post-Debate Voting
10:10-11:00	Is central 5-HT1F agonism essential for ditans to be effective?
	Capsule: 5-HT1F receptors have been identified in both peripheral and central parts of the TVS. Ditans penetrating the blood brain barrier and activating 5- HT1F receptors in both parts of the TVS, also induce adverse effects that limit their use. Is central 5-HT1F agonism essential for ditans to be effective, or is the peripheral action enough, like with all other migraine-specific and mechanism based symptomatic treatments?
10:10-10:20	Moderator: <u>Dimos Mitsikostas</u> , Greece Introduction and Pre-Debate Voting
10:20-10:35	Yes: <u>Anna Andreou</u> , UK
10:35-10:50	No: <u>Antoinette Maassen van den Brink</u> , The Netherlands
10:50-11:00	Discussion, Rebuttals and Post-Debate Voting
11:00-11:30	Coffee Break, Exhibition & e-Posters Visit



FRIDAY, MARCH 22, 2024		
11:30-13:30	Plenary Sessions (Plenary Hall)	
13:40-14:30	MTE	
13:30-14:30	Lunch Break, Exhibition & e-Posters Visit	
14:30-16:10	HEADACHE : Ditans and Gepants	HALL C
Chair:	Tomas Nezadal, Czech Republic	
14:30-15:20	Addition of a gepant for the acute care of migraine attacks is safe and effective in patients on anti-CGRP mAbs	
	Capsule: The consideration of using gepants for aborting migraine attacks remains a topic of debate when managing pati anti-CGRP mAbs as preventive therapy. Moreover, considering the safety aspects of combining two drugs that target CGR adding a gepant for immediate migraine relief in patients on anti-CGRP mAbs preventive treatment	
14:30-14:40	Moderator: <u>Messoud Ashina,</u> Denmark Introduction and Pre-Debate Voting	
14:40-14:55	Yes:Cristina Tassorelli, Italy	
14:55-15:10	No: Gisela Terwindt, The Netherlands	
15:10-15:20	Discussion, Rebuttals and Post-Debate Voting	
15:20-16:10	Are gepants and ditans efficacious and safe for people with vascular risk factors?	
	Capsule: Preclinical studies showed that CGRP is a potent vasoactive neuropeptide, yet activation of 5-HT1F receptors doe arteries. Is there evidence supporting the use of treatments targeting either the CGRP neuropeptide, e.g., the gepants, or in people with migraine and vascular risk factors?	-
15:20-15:30	Moderator: <u>Messoud Ashina</u> , Denmark Introduction and Pre-Debate Voting	
15:30-15:45	Yes: Jose Miguel Lainez, Spain	
15:45-16:00	No: Gisela Terwindt, The Netherlands	
16:00-16:10	Discussion, Rebuttals and Post-Debate Voting	
16:10-16:30	Coffee Break, Exhibition & ePosters Visit	



	FRIDAY, MARCH 22, 2024	
16:30-18:10	Medication overuse headache and artificial intelligence	HALL C
Chair:	Licia Grazzi, Italy	
16:30-17:20	Is Artificial Intelligence (AI) ready for inclusion in headache management?	
	Capsule: AI systems have gotten our attention as they have emerged in all aspects of the working and education worlds. as a way to improve the quality of diagnosis and treatment planning, as well as a way to deliver care more efficiently. Bu to its promise. Recently however, with increased power of operating systems and new designs for analyzing data, this m revolutionize medicine including the care of patients with headache disorders. It the time now?	t most would agree it has not lived up
16:30-16:40	Moderator: Alan Rapoport, USA	
	Introduction and Pre-Debate Voting	
16:40-16:55	Yes: Robert Cowan, USA	
16:55-17:10	No: <u>Morris Levin</u> , USA	
17:10-17:20	Discussion, Rebuttals and Post-Debate Voting	
17:20-18:10	Is it necessary to detoxify most patients with Medication Overuse Headache (MOH) in order to achieve success?	
	Capsule: Ever since the concept for MOH was promoted, a key tenet was that patients with this syndrome could not impr was removed. Only then, would preventive medications be effective in reversing the chronification of the primary headac dogma into question, but there is still disagreement about the best way(s) to help patients.	
17:20-17:30	Moderator: <u>Cristina Tassorelli</u> , Italy Introduction and Pre-Debate Voting	
17:30-17:45	Yes: <u>Morris Levin</u> , USA	
17:45-18:00	No: <u>Alan Rapoport</u> , USA	
18:00-18:10	Discussion, Rebuttals and Post-Debate Voting	



	SATURDAY,MARCH 23,2024	
<u>07:30-08:20</u>	ePoster Guided Tour	
08:20-11:00	Amyotrophic lateral sclerosis/Motorneuron disease (ALS/MND) HALL A	
Chairs:	Pam Shaw, UK; Albert Ludolph, Germany	
08:20-09:10	Human cell models are better than mouse models in therapy development for ALS/MND	
	Capsule: Given the limitations of currently available mouse models of ALS/MND and the lack of a mouse model for sporadic disease, human cellulo	ar models
00.00.00.00	have key advantages in the evaluation of potential neuroprotective therapies.	
08:20-08:30	Moderator: <u>Amir Dori</u> , Israel	
00.20 00.45	Introduction and Pre-Debate Voting	
08:30-08:45 08:45-09:00	Yes : Pam Shaw, UK	
	No: Linda Greensmith ,UK	
09:00-09:10	Discussion, Rebuttals and Post-Debate Voting	
09:10-10:00	ALS/MND treatments have to demonstrate efficacy on both prolonging survival and slowing disease progression	
	Capsule: In the absence of improved function, prolonged survival by a few months may not necessarily indicate attenuation of disease progression.	. Therefore,
09:10-09:20	evidence for slowed disease progression is required Moderator: Amir Dori, Israel	
09.10-09.20	Introduction and Pre-Debate Voting	
09:20-09:35	Yes: <u>Albert Ludolph</u> , Germany	
09:35-09:50	No: Osman Sinanovic, Bosnia and Herzegovina	
09:50-10:00	Discussion, Rebuttals and Post-Debate Voting	
10:00-11:00	The euphoria on personalized ALS/MND treatment with Antisense oligonucleotides (ASO) is premature	
	Capsule: Treatment with ASOs for specific genetic subtypes of ALS/MND has shown some promising results, but euphoria is premature given the r these subtypes and the failure of some ASO approaches.	rarity of
10:00-10:10	Moderator: Osman Sinanovic, Bosnia and Herzegovina	
10.10 10.25	Introduction and Pre-Debate Voting	
10:10-10:25	Yes: <u>Pam Shaw,</u> UK	
10:25-10:40	No: Giancarlo Logroscino, Italy	
10:40- 11:00	Discussion, Rebuttals and Post-Debate Voting	
11:00-11:30	Coffee Break, Exhibition & e-Posters Visit	



	SATURDAY,MARCH 23,2024	
11:30-12:30	Plenary Session Plenary Hal	I
Chairs:	George Chakhava, Georgia; Odelia Elkana, Israel	
11:30-12:00	Aphantasia – when all is dark in the mind's eye	
	Adam Zeman, UK	
12:00-12:30	The impossible pricing of anti-amyloid medication. A global challenge.	
12:30-13:30	Paola Barbarino, UK	
12:30-13:30	Industry Sponsored Symposium (Plenary Hall) Plenary Ha MTE	<u>11</u>
13:30-14:30	Lunch Break, Exhibition & e-Posters Visit	
14:30-16:10	Amyotrophic lateral sclerosis/ motorneuron disease (ALS/MND) 2 HALL	Δ
Chairs:	Ervin Jancic, Croatia; Nana Kvirkvelia, Georgia	
14:30-15:20	Tracheostomy ventilation in ALS/MND should be offered to all patients	
	Capsule: Neuromuscular respiratory failure is the cause of death in most patients with ALS/MND. Invasive tracheostomy ventilation adds	a burden of care
	while non-invasive respiratory support is readily available.	
14:30-14:40	Moderator: David Oliver, UK	
	Introduction and Pre-Debate Voting	
14:40-14:55	Yes: Giancarlo Logroscino, Italy	
14:55-15:10	No: <u>Albert Ludolph</u> , Germany	
15:10-15:20	Discussion, Rebuttals and Post-Debate Voting	
15:20-16:10	Generalized myasthenia gravis (gMG) patients with highly active disease should be treated with innovative treatments earlier	
	Capsule : The development of immunological therapies against myasthenia gravis(MG) pose a practical question on when to treat and whe Previous drugs, including azathioprine, steroids and others, are quite useful, and lacking head to head comparison, the problem is maintain	•
15:20-15:30	Moderator: Hans-Peter Hartung, Germany	
15:30-15:45	Yes: Hans-Peter Hartung, Germany	
15:45-16:00	No: Jacek Losy, Poland	
16:00-16:10	Discussion, Rebuttals and Post-Debate Voting	
16:10-16:30	Coffee Break, Exhibition & e-Posters Visit	



SATURDAY,MARCH 23,2024		
16:30-18:10	Neurodegenerative diseases	HALL A
Chairs:	Johannes Attems, UK <u>; Bogdan Popescu,</u> Romania	
16:30-17:20	Will anti-tau drugs be useful in fighting tau-driven diseases?	
	Capsule: Tauopathies are very common pathology changes in neurodegenerative diseases, particularly AD, in which the hyperphosphorylated tau proteins. The large number of individuals affected by tauopathies and the theory that this lead search for disease-modifying therapies targeting tau pathology. None has proven useful so far. Is the search worthwhile	ds to neurodegeneration have led to a
16:30-16:40	Moderator: Bogdan Popescu , Romania Introduction and Pre-Debate Voting	
16:40-16:55	Yes: Lea Grinberg, USA	
16:55-17:10	No : <u>Jesse Cedarbaum</u> , USA	
17:10-17:20	Discussion, Rebuttals and Post-Debate Voting	
17:20-18:10	Pathological changes in microglia were linked to pathology in different neurodegenerative diseases. Can we target mice approach?	roglia to develop a therapeutic
	Capsule: Activation of microglia is considered a pathological hallmark in many neurodegenerative diseases. While micro activity, impairment in their activity may exacerbate neuroinflammation and result in neuronal death. Should microglia intervention to ameliorate disease pathology in neurodegenerative diseases?	
17:20-17:30	Moderator: <u>Jesse Cedarbaum</u> , USA Introduction and Pre-Debate Voting	
17:30-17:45	Yes: Dan Frenkel, Israel	
17:45-18:00	No: Bogdan Popescu, Romania	
18:00-18:10	Discussion, Rebuttals and Post-Debate Voting	



	SATURDAY, MARCH 23, 2024	
<u>07:30-08:20</u>	ePoster Guided Tour	
08:20-11:00	Epilepsy: Drug treatment	HALL B
Chairs:	Matthias Koepp, UK ; Eva Andermann, Canada	
08:20-09:10	8:20-09:10 Should we routinely utilize therapeutic drug monitoring when prescribing drug therapy for people with epilepsy?	
	Capsule: It is debated whether monitoring of serum levels improves care for people with epilepsy, and the literature contain. What should we do?	s conflicting recommendations.
08:20-08:30	Moderator: <u>Michael Sperling</u> , USA Introduction and Pre-Debate Voting	
08:30-08:45	Yes: <u>William Theodore,</u> USA	
08:45-09:00	No: <u>Elinor Ben-Menachem</u> , Sweden	
09:00-09:10	Discussion, Rebuttals and Post-Debate Voting	
09:10-10:00	Should women who wish to become pregnant be prescribed supplemental folic acid?	
	Capsule: A recent study suggests that cancer risk is increased in children born to women who took folic acid during pregnant clinical practice?	cy. Is this study sufficient to alter
09:10-09:20	Moderator: <u>Martin Holtkamp</u> , Germany Introduction and Pre-Debate Voting	
09:20-09:35	Yes: Jacqueline French,USA	
09:35-09:50	No: <u>Alla Guekht,</u> Russia	
09:50-10:00	Discussion, Rebuttals and Post-Debate Voting	
10:00-11:00	Should polytherapy be used early, after one drug has failed to control seizures?	
	Capsule: Is rational polytherapy a sensible practice after failure of one antiseizure medication, given the low probability of rolls there evidence that polytherapy is superior to monotherapy?	esponse to another monotherapy
10:00-10:10	Moderator: Zeljka Petelin-Gadze, Croatia Introduction and Pre-Debate Voting	
10:10-10:25	Yes: <u>Manjari Tripathi</u> , India	
10:25-10:40	No: Elinor Ben-Menachem, Sweden	
10:40-11:00	Discussion, Rebuttals and Post-Debate Voting	
11:00-11:30	Coffee Break, Exhibition & e-Posters Visit	



SATURDAY, MARCH 23, 2024	
11:30-12:30	Plenary Sessions (Plenary Hall)
12:30-13:30	Industry Sponsored Symposium (Plenary Hall)
13:30-14:30	MTE
13:30-14:30	Lunch Break, Exhibition & e-Posters Visit
14:30-16:10	The unusual cases HALL B
Chairs:	Nandan Yardi , India <u>; Natasa Pejanovic Skobic</u> , Bosnia and Herzegovina
14:30-15:20	Mix of interesting cases related to diagnosis and management challenges in epilepsy, discussing advanced drug therapy and a modes of administration. Case Studies: Michael Sperling, USA
15:20-16:10	Should we prefer one method of neurostimulation over others?
	Capsule: Vagus nerve stimulation, deep brain stimulation, responsive neurostimulation, and extracranial stimulation are used to treat people with uncontrolled seizures. Is there evidence that some method is superior or inferior to others?
15:20-15:30	Moderator: Alla Guekht, Russia Introduction and Pre-Debate Voting
15:30-15:45	Yes: Martin Holtkamp, Germany
15:45-16:00	No: Ivan Rektor, Czech Republic
16:00-16:10	Discussion, Rebuttals and Post-Debate Voting
16:10-16:30	Coffee Break, Exhibition & e-Posters Visit



SATURDAY, MARCH 23, 2024		
16:30-18:10	Epilepsy 3 HALL B	
Chairs:	Man Mohan Mehndiratta, India; Andreja Bujan Kovac, Croatia	
16:30-17:20	Should laser interstitial thermal ablation therapy (LITT) be preferred over open surgery, either anterior temporal lobectomy or amygdalohippocampectomy for mesial temporal sclerosis?	
	Capsule: Thermal ablation has been used for over 11 years and offers clear advantages with regard to perioperative pain, morbidity, and period of disability. Are these advantages sufficient to recommend this procedure as the first preferred therapy for people with seizures due to mesial temporal sclerosis?	
16:30-16:40	Moderator: <u>Lilach Goldstein,</u> Israel Introduction and Pre-Debate Voting	
16:40-16:55	Yes: Michael Sperling, USA	
16:55-17:10	No: Matthias Koepp, UK	
17:10-17:20	Discussion, Rebuttals and Post-Debate Voting	
17:20-18:10	Should intracranial EEG be largely restricted to patients with non-lesional epilepsy unless there is discordance between the lesion and the scalp EEG?	
	Capsule: Some would argue that intracranial EEG is vastly overused without adequate evidence that it improves outcome in patients with obvious epileptogenic lesions. Is there evidence to support its use or is excision of a lesion with adequate margin sufficient to achieve a good post-surgical outcome?	
17:20-17:30	Moderator: <u>Matthias Koepp</u> , UK Introduction and Pre-Debate Voting	
17:30-17:45	Yes: <u>William Theodore</u> , USA	
17:45-18:00	No: Ivan Rektor, Czech Republic	
18:00-18:10	Discussion, Rebuttals and Post-Debate Voting	



SATURDAY, MARCH 23, 2024		
07:30-08:20	ePoster Guided Tour	
08:20-11:00	Multiple sclerosis: drug therapy	HALL C
Chairs:	Jacek Losy, Poland <u>; Dimitrios Karussis</u> , Israel	
08:20-09:10	Radiologically isolated syndrome (RIS): Disease modifying therapies (DMT) should only be started when symptoms have occ	
	Capsule: RIS is considered a possible early indicator of MS or "pre-MS", but since no clinical symptoms have occurred, it is unl	-
	without actually ever developing MS. With studies showing that more than half of RIS patients still remain symptom-free for	years+, there may not be a clear
	indication for starting therapy	
08:20-08:30	Moderator: <u>Alicia Kalinowska,</u> Poland	
00.20 00.45	Introduction and Pre-Debate Voting	
08:30-08:45 08:45-09:00	Yes: <u>Agne Straukiene</u> , UK No: Daniel Salo Reich , USA	
09:00-09:10	Discussion, Rebuttals and Post-Debate Voting	
09:10-10:00	DMT should be discontinued after prolonged stability	
	Capsule: There is a sense that in some patients, MS "burns out" and patients no longer have relapses or new MRI lesions. Since	-
	MS activity (relapse and MRI), it is perhaps reasonable to stop the therapy after a period of time when no MS activity has been been been as the stop of the therapy after a period of time when no MS activity has been been as the stop of the therapy after a period of time when no MS activity has been as the stop of the therapy after a period of time when no MS activity has been as the stop of the therapy after a period of time when no MS activity has been as the stop of the therapy after a period of time when no MS activity has been as the stop of the therapy after a period of time when no MS activity has been as the stop of the therapy after a period of time when no MS activity has been as the stop of the therapy after a period of time when no MS activity has been as the stop of the therapy after a period of time when no MS activity has been as the stop of the therapy after a period of time when no MS activity has been as the stop of the therapy after a period of time when no MS activity has been as the stop of the therapy after a period of time when no MS activity has been as the stop of the therapy after a period of time when no MS activity has been as the stop of the therapy after a period of therapy after a period of the therapy after a period of therapy after a period of the therapy after a period of the therapy after a period of therapy after	en perceived.
09:10-09:20	Moderator: <u>Konrad Rejdak</u> , Poland	
	Introduction and Pre-Debate Voting	
09:20-09:35	Yes: Per Soelberg Sørensen, Denmark	
09:35-09:50	No: <u>Klaus Schmierer</u> , UK	
09:50-10:00	Discussion, Rebuttals and Post-Debate Voting	
10:00-11:00	The central vein sign should be used as a diagnostic criterion	
	Capsule: The central vein sign (CVS) adds an "in vivo histology" marker to the pattern approach of the McDonald criteria, lar	
	space and time. Through its apparent specificity for MS lesions, the CVS may enable a more accurate diagnosis, departing fro	
	better explanation". However, is the evidence sufficiently strong, and MRI-technology available widely enough, to include the	e CVS in the standard set of criteria?
10:00-10:10	Moderator: Konrad Rejdak, Poland	
10.10 11.25	Introduction and Pre-Debate Voting	
10:10-11:25	Yes: Declan Chard, UK	
10:25-10:40	No : Daniel Salo Reich, USA	
10:40-11:00	Discussion, Rebuttals and Post-Debate Voting	
11:00-11:30	Coffee Break, Exhibition & e-Posters Visit	



SATURDAY, MARCH 23, 2024		
11:30-12:30	Plenary Session (Plenary Hall)	
12:30-13:30	Industry Sponsored Symposium (Plenary Hall)	
13:30-14:30	MTE	
13:30-14:30	Lunch Break, Exhibition & e-Posters Visit	
14:30-16:10	Multiple sclerosis 2 HALL C	
Chair:	Homa Ebrahimi, Iran	
14:30-15:20	Epstein-Barr virus (EBV) is a driver of ongoing MS disease activity	
	Capsule: EBV is a confirmed risk factor for the development of MS. However, does EBV re-activation play an important role even after the disease has been triggered? Recent evidence suggest that MS patients display aberrant EBV gene expression and regulation of both viral and cellular genes in B cells and dysregulated EBV latency. Antiviral treatments can ameliorate EBV replication, viral loads, lytic gene expression, and EBV-mediated inflammation suggesting that dysregulation of EBV latency drives MS activity. But is the evidence robust enough, and what are the counter arguments?	
14:30-14:40	Moderator: <u>Floriana De Angelis</u> , UK Introduction and Pre-Debate Voting	
14:40-14:55	Yes: <u>Francesca Aloisi</u> , Italy	
14:55-15:10	No : <u>Gavin Giovannoni</u> , UK	
15:10-15:20	Discussion, Rebuttals and Post-Debate Voting	
15:20-16:10	Multiple Sclerosis is progressive from onset	
	Capsule: The classical vision of MS characterized in more than 90% of cases by two sequential phases, an initial relapsing remitting course followed by a secondary progressive course is now challenged by the recent observation that a progression independent from relapse activity (PIRA) can be seen very early in the disease. So is MS always progressive from the beginning?	
15:20-15:30	Moderator: <u>Per Soelberg Sorensen,</u> Denmark Introduction and Pre-Debate Voting	
15:30-15:45	Yes: <u>Floriana De Angelis</u> , UK	
15:45-16:00	No: <u>Giancarlo Comi</u> , Italy	
16:00-16:10	Discussion, Rebuttals and Post-Debate Voting	
16:10-16:30	Coffee Break, Exhibition & e-Posters Visit	



SATURDAY, MARCH 23, 2024		
16:30- 18:10	MS : Diagnosis and pathogenesis	HALL C
Chair:	Alicja Kalinowska, Poland ; Peter Feys, Belgium	
16:30-17:20	Microglia are the main drivers of progressive MS	
	Capsule: Microglia are known for their involvement in the immune response within the central nervous system, and their activation observed in MS. Activated microglia can release neurotoxic substances, potentially damaging surrounding neurons and exacerb the other hand, MS is a complex and multifactorial disease with contributions from various immune cells, genetics, and environmer are a heterogeneous cell population and their functions can vary based on their activation state and include also neuroprote microglia may overlook other crucial elements of MS pathogenesis and might oversimplify the complex nature of the disease. Are not progressive MS?	nating the progression of MS. On natal factors. In addition, microglia active effects. Focusing solely on
16:30-16:40	Moderator: <u>Ron Milo</u> , Israel Introduction and Pre-Debate Voting	
16:40-16:55	Yes: <u>Alicja Kalinowska,</u> Poland	
16:55-17:10	No: Sharmilee Gnanapavan, UK	
17:10-17:20	Discussion, Rebuttals and Post-Debate Voting	
17:20-18:10	Enhancing recovery in MS – in cooperation with European Charcot Foundation (ECF)	
Chairman:	<u>Giancarlo Comi</u> , Italy Strategies to promote remyelination: <u>Hans Peter Hartung</u> , Germany Advances in neurorehabilitation: <u>Peter Feys</u> , Belgium Role of neuromodulation in enhancing recovery: <u>Letizia Leocani</u> , Italy	
18:10	Closing ceremony Plenar	y Hall
	Ivan Rektor, Czech Republic	
	<u>Natan Bornstein</u> , Israel	