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Evaluating the Long-Term Outcomes and Durability of Effect Following Treatment with Cladribine Tablets for Multiple Sclerosis

13/08/2025 11:01:37

Primary registry identifying number	Protocol number
LBCTR2020030215	MS700568_0026
MOH registration number	
Study registered at the country of origin	Study registered at the country of origin: Specify
Yes	
Type of registration	Type of registration: Justify
Prospective	N/A
Date of registration in national regulatory agency 25/07/2019	
Primary sponsor	Primary sponsor: Country of origin
Merck KGaA	Germany
Date of registration in primary registry	Date of registration in national regulatory agency
11/06/2021	25/07/2019
Public title	Acronym
Evaluating the Long-Term Outcomes and Durability of Effect Following Treatment with Cladribine Tablets for Multiple Sclerosis	
Scientific title	Acronym
An Exploratory Phase IV Ambispective Study of Patients Who Previously Participated in the CLARITY/CLARITY-EXT and ORACLE MS Clinical Trials	
Brief summary of the study: English	
The purpose of this study is to explore the long-term outcomes, durability of effect, and real-world treatment patterns in patients previously participating in the Phase III ORACLE MS and CLARITY/CLARITY-EXT clinical trials (i.e. parent studies). The results from this study may be of benefit to patients with multiple sclerosis (MS) and clinicians by helping to inform future treatment approaches and treatment decision-making.	
Brief summary of the study: Arabic	
تقييم النتائج طويلة الأمد و مدّة تأثير العلاج بأقراص Cladribine عند مرضى مصابين بتصلّب المتعدد	
Health conditions/problem studied: Specify	
Multiple Sclerosis	

*Optional blood sample: Patients willing to consent to provide an optional blood sample and who are seen at a site with available capabilities to store and ship samples, will have a blood draw taken at Study Visit 1 for pharmacogenetics testing.

Key inclusion and exclusion criteria: Inclusion criteria

1. Patients with MS randomised in CLARITY/CLARITY-EXT clinical trial(s) who have received ≥ 1 course of IMP (Cladribine Tablets or



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placebo). or Patients with their FCDE randomised in ORACLE MS clinical trial who hav 2. Informed Consent	e received≥ 1 course of IMP (Cladr	ibine Tablets or placebo).
Key inclusion and exclusion criteria: Gender	Key inclusion and exclusion of	criteria: Specify gender
Both		
Key inclusion and exclusion criteria: Age minimum	Key inclusion and exclusion	criteria: Age maximum
18	65	
Key inclusion and exclusion criteria: Exclusion criteria		
 Medical Conditions: Any condition, including any uncontrolled disease state other than MS, tha contraindication for participation in the study or that could interfere with the evaluation. 		titutes an inappropriate risk or a
For the MRI sub-study: 1. Female Participants Who are pregnant 2. Patient taking Cladribine Tablets as part of another study at the time of t	the start of this study	
Type of study		
Observational		
Type of intervention	Type of intervention: Specify	type
N/A	N/A	
Trial scope	Trial scope: Specify scope	
N/A	N/A	
Study design: Allocation	Study design: Masking	
N/A	N/A	
Study design: Control	Study phase	
N/A	N/A	
Study design: Purpose	Study design: Specify purpos	ie.
N/A	N/A	-
Study design: Assignment	Study design: Specify assign	mont
N/A	N/A	ment
		- On a sife
IMP has market authorization	IMP has market authorization	: Specify
Name of IMP	Year of authorization	Month of authorization
Type of IMP		
Pharmaceutical class		
Therapeutic indication		
Therapeutic benefit		

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Study model	Study model: Explain model
Cohort	NA
Study model: Specify model	
N/A	
Time perspective	Time perspective: Explain time perspective
Other	Ambispective: Retrospective and Prospective
Time perspective: Specify perspective	
Ambispective	
Target follow-up duration	Target follow-up duration: Unit
6	Weeks
Number of groups/cohorts	
4	
Biospecimen retention	Biospecimen description
Samples with DNA**	8ml blood sample will be collected for DNA analysis from consenting participants
	Actual approliment target size
Target sample size 8	Actual enrollment target size
Date of first enrollment: Type	Date of first enrollment: Date
Actual	15/08/2019
Date of study closure: Type	Date of study closure: Date
Actual	27/02/2021
Recruitment status	Recruitment status: Specify
Complete	
Date of completion	
27/02/2021	
IPD sharing statement plan	IPD sharing statement description
No	NO Individual Patient Data Sharing

Additional data URL

Admin comments





Trial status

Approved

Secondary Identifying Numbers		
Full name of issuing authority	Secondary identifying number	
European Clinical Trials Database	EudraCT number: 2019-000069-19	

Sources of Monetary or Material Support
Name
Merck KGaA Germany

Secondary Sponsors	
Name	
N/A	

Contact for Public/Scientific Queries						
Contact type	Contact full name	Address	Country	Telephone	Email	Affiliation
Public	Dr. Bassem Yamout	Hamra, Cairo Street	Lebanon	+9613221 222	-	American University of Beirut Medical Center
Scientific	Kristin Gabriel	EMD Serono, Inc. One Technology Place, Rockland MA 02370	United States of America	+1 781 427 1502	Kristin.Gabriel@ emdserono.com	EMD Serono, Inc.

Centers/Hospitals Involved in the Study				
Center/Hospital name	Name of principles investigator	Principles investigator speciality	Ethical approval	
Bellevue Medical Center	Dr. Souheil Gbeily	Neurology	Approved	
American University of Beirut Medical Center	Dr. Bassem Yamout	Neurology	Not approved	





Ethics Review				
Ethics approval obtained	Approval date	Contact name	Contact email	Contact phone
Bellevue Medical Center	10/09/2019	Prof. Souheil Jbeily	souheil.gebeily@gmail.com	9611682666

Countries of Recruitment
Name
Australia
Austria
Belgium
Bulgaria
Canada
Croatia
Czech Republic
Estonia
Finland
France
Georgia
Germany
Greece
Italy
Republic of Korea
Latvia
Lithuania
Morocco
Norway
Poland
Portugal



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Romania
Russian Federation
Republic of Serbia
Spain
Sweden
Switzerland
Tunisia
Ukraine
United Kingdom
United States of America
Lebanon

Health Conditions or Problems Studied			
Condition Code Keyword			
Multiple Sclerosis	Multiple sclerosis (G35)	MS, Multiple Sclerosis	

Interventions				
Intervention	Description	Keyword		
Optional Blood Sampling	Purpose: Pharmaco-genetics Testing	blood sampling		





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Primary Outcomes		
Name	Time Points	Measure
To evaluate long-term mobility after treatment with an investigational medicinal product (IMP; Cladribine Tablets or placebo) as part of the Phase III ORACLE MS and CLARITY/CLARITY-EXT clinical trials.	Proportion of study participants using a wheelchair (defined as unable to walk beyond approximately 5 meters even with aid, essentially restricted to wheelchair; wheels self in standard wheelchair and transfers alone; up and about in wheelchair some 12 hours a day) the majority of the time in the 3 months prior to Study Visit 1 for the CLARITY/CLARITY-EXT and ORACLE MS populations,	• Expanded Disability Status Scale (EDSS) score of 7.0 or higher (if available), or • Alternative clinical description data in medical records.





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Key Secondary Outcomes			
Name	Time Points	Measure	
To assess the long-term disability status after treatment with IMP as part of the Phase III ORACLE MS and CLARITY/CLARITY-EXT clinical trials for the CLARITY/CLARITY-EXT and ORACLE MS populations.	Proportion of study participants with 3-month sustained (i.e. ambulatory disability consistent with EDSS on at least 2 clinic visits no less than 3 months apart)	EDSS of 6.0 or higher in the last year prior to enrollment or corresponding clinical description in medical records	
To evaluate differences in clinical characteristics between long-term responders and study participants requiring alternate therapies following treatment with IMP for the CLARITY/CLARITY-EXT and ORACLE MS populations.	Clinical characteristics at Study Visit 1 of long-term responders (defined as study participants who did not demonstrate any evidence of disease reactivation based on Investigator assessment of clinical and imaging outcomes until Year 4 or later following their last doseb of IMP and who did not receive disease modifying treatment until Year 4 or later following their last doseb of IMP) compared to those of other study participants who started on alternate therapy less than 4 years following their last doseb of IMP for the CLARITY/CLARITY- EXT and ORACLE MS populations.		
To evaluate differences in magnetic resonance imaging (MRI) characteristics between long-term responders and study participants requiring alternate therapies following treatment with IMP for the CLARITY/CLARITY-EXT and ORACLE MS populations.	MRI characteristics at Study Visit 2 of long-term responders (defined as study participants who did not demonstrate any evidence of disease reactivation based on Investigator assessment of clinical and imaging outcomes until Year 4 or later following their last doseb of IMP and who did not receive disease modifying treatment until Year 4 or later following their last doseb of IMP) compared to those of other study participants who started on alternate therapy less than 4 years following their last doseb of IMP for the CLARITY/CLARITY- EXT and ORACLE MS populations.	-	



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Trial Results Summary results Study results globally Date of posting of results summaries Date of first journal publication of results **Results URL link Baseline characteristics** Participant flow Adverse events **Outcome measures** URL to protocol files