



*Making dementia a priority:  
changing perceptions, practice and policy.*

# CLINICAL TRIALS WATCH

ACCESSIBLE EASY READ INFORMATION ON:

**T-817MA STUDY**

# T-817MA study

<b>1. Study Information</b>	
<b>Name of the study</b>	Efficacy and Safety of T-817MA in patients with mild cognitive impairment due to Alzheimer's disease or mild Alzheimer's disease
<b>Study sponsor</b>	FUJIFILM Toyama Chemical Co., Ltd.
<b>Disease</b>	Alzheimer's disease
<b>Phase</b>	Phase II

<b>2. Information about the drug that will be tested in the study</b>	
<b>Name of drug</b>	T-817MA (also named Edonerpic)
<b>Administration</b>	Tablet
<b>Is the drug already on the market for another medical condition?</b>	No
<b>Will all participants receive the same drug?</b>	<p>Participants will be selected by chance to receive one of the following options:</p> <ul style="list-style-type: none"><li>• An oral tablet of T-817MA (224mg) once daily for first 4 weeks and then two oral tablets of T-817MA (448mg) for the following weeks</li><li>• An oral tablet of placebo (inactive substance identical in appearance to the drug being tested).</li></ul> <p>Neither the participant nor their doctor will know if the person is receiving the investigational drug or the placebo.</p>

<b>3. Information about participating in the trial</b>	
<b>What are the researchers trying to find out?</b>	<ul style="list-style-type: none"><li>• The purpose of the study is to evaluate the efficacy and safety of T-817MA in people with mild cognitive impairment due to Alzheimer's disease or mild Alzheimer's disease.</li></ul>
<b>How long will the treatment last?</b>	<ul style="list-style-type: none"><li>• Around 1.5 year (5 visits)</li></ul>
<b>What your involvement will entail?</b>	<ul style="list-style-type: none"><li>• During the study, participants will have to undergo brain scan (MRI) and CSF examination (spinal tap) to see if they have amyloid pathology in their brain</li></ul>

	<ul style="list-style-type: none"> <li>• Complete a memory test (MMSE) to assess cognitive function</li> <li>• Complete a test that will assess memory, orientation, judgment and problem solving, personal care and community affairs (this is a test called CDR)</li> <li>• Perform an electroencephalogram (this is a test or record of brain electrical activity using electrodes placed along the scalp)</li> <li>• Complete some laboratory tests to evaluate the emergent adverse effects (unfavourable signs, symptoms or diseases temporally associated with the use of the drug tested in the study).</li> </ul> <p>Further information on the number of visits can be obtained from the study team.</p>
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<h4 style="color: red; margin: 0;">4. Who can participate in this study?</h4>	
<p><b>Who can participate in the study?</b></p>	<p>To take part in the study, participants must:</p> <ul style="list-style-type: none"> <li>• Be 50 to 80 years old</li> <li>• Have a diagnosis of mild cognitive impairment due to Alzheimer’s disease or mild Alzheimer’s disease according to the National Institute on Aging/Alzheimer’s Association core clinical criteria</li> <li>• Have evidence of abnormal accumulation of amyloid and p-tau181 in their brain (as per results of CSF examination (spinal tap))</li> <li>• Have a score between 24 and 30 points in the MMSE test (a test about your memory). This would suggest that the person has an impairment in their memory that is at a very mild stage</li> <li>• Be post-menopausal or permanently sterilized for female participants</li> </ul>

	<ul style="list-style-type: none"> <li>• Be required to use highly effective methods of contraception during the study and until 104 days after the last dose for male participants</li> <li>• If the person is taking acetylcholinesterase inhibitors as an approved anti-dementia medication (i.e. donepezil, rivastigmine, galantamine) the dosing regimen must have been stable for at least 3 months prior to the screening visit</li> <li>• Have a study partner who has a sufficient contact with the participant, is willing to participate in study procedures throughout the study duration.</li> </ul>
<p><b>Who cannot participate in the study?</b></p>	<p>Exclusion criteria include:</p> <ul style="list-style-type: none"> <li>• Contraindication to PET imaging, MRI procedures and lumbar puncture</li> <li>• Any results within the previous two years that showed a pathology inconsistent with a diagnosis of Alzheimer's disease</li> <li>• A current treatment of memantine.</li> </ul> <p>The above list is not exhaustive. It includes the most common conditions and diseases that might exclude people from the study.</p>

**5. Where and when will the study be conducted?**

<p><b>European countries involved in the trial</b></p>	<ul style="list-style-type: none"> <li>• Czech Republic</li> <li>• Germany</li> <li>• Hungary</li> <li>• Ireland</li> <li>• Netherlands</li> <li>• Spain</li> <li>• UK</li> </ul>
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<b>Estimated start date of recruitment</b>	December 2019
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<b>6. Information for your doctor</b>			
<b>EudraCT Number:</b>	2018-003567-66	<b>Clinicaltrials.gov identifier</b>	NCT04191486
<b>Study contact information</b>	Larah de Groot +31 30 656 9186 <a href="mailto:larah.degroot@juliusclinical.com">larah.degroot@juliusclinical.com</a>		
<b>Link to full text</b>	<a href="https://clinicaltrials.gov/ct2/show/NCT04191486">https://clinicaltrials.gov/ct2/show/NCT04191486</a>		

- ✓ The information contained in this document is based on information available on public registries (e.g. clinicaltrials.gov website) on February 2021.
- ✓ This document has been reviewed by the pharmaceutical company running this trial.
- ✓ This document has been reviewed by a member of the European Working Group of People with Dementia.