

# CLINICAL TRIALS WATCH

ACCESSIBLE EASY READ INFORMATION ON:

## SERENADA STUDY

# SERENADA study

<b>1. Study Information</b>	
<b>Name of the study</b>	Safety and efficacy of EXV-802 and EXV-801 in the treatment of agitation in Alzheimer's disease dementia
<b>Study sponsor</b>	Exciva GmbH
<b>Disease</b>	Agitation in Alzheimer's disease
<b>Phase</b>	Phase III

<b>2. Information about the drug that will be tested in the study</b>	
<b>Name of drugs</b>	<p>EXV-802 (also called deraphan) + EXV-801 (also called deramciclane)</p> <p>EXV-802 is a fixed-dose combination of the approved drug dextromethorphan and the experimental anxiolytic EXV-801.</p>
<b>Administration</b>	Oral administration (capsules twice a day)
<b>Is the drugs already on the market for another medical condition?</b>	No, however dextromethorphan, one of the components of EXV-802, is a cough suppressant used in many cough and cold medicines. Additionally, the combination medicine dextromethorphan and bupropion is approved for major depressive disorder in US.
<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 administration of EXV-802</li><li>• An oral administration of EXV-801</li><li>• An oral administration of placebo (also called a dummy treatment which is an inactive substance identical in appearance to the drug being tested with no active therapeutic effect).</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 this study is to evaluate the efficacy and safety of EXV-802 and EXV-801 in treatment of agitation in people with Alzheimer's disease dementia.</li> </ul>
<b>How long will the treatment last?</b>	<ul style="list-style-type: none"> <li>6 weeks</li> </ul>
<b>What your involvement will entail?</b>	<ul style="list-style-type: none"> <li>During the study, participants will be asked to complete tests that will assess their agitation and aggression (e.g., CMAI-IPA, CGI-S, NPI-C).</li> </ul> <p>Further information on the procedures, tests and number of visits can be obtained from the study team.</p>

<b>4. Who can participate in this study?</b>	
<b>Who can participate in the study?</b>	<p>To take part in the study, participants must:</p> <ul style="list-style-type: none"> <li>Be between 55 and 90 years old</li> <li>Have a confirmed diagnosis of Alzheimer's disease dementia</li> <li>Have a clinically significant, moderate/severe agitation</li> <li>Have a diagnosis of agitation, according to the International Psychogeriatric Association (IPA) provisional definition of agitation</li> <li>Have a study partner who has a sufficient contact with the participant (at least on four individual days for at least two hours per day), is willing to participate in study procedures throughout the study duration.</li> </ul>
<b>Who cannot participate in the study?</b>	<p>Exclusion criteria include:</p> <ul style="list-style-type: none"> <li>Confirmed primary diagnosis of another form of dementia other than Alzheimer's disease dementia</li> </ul>

	<ul style="list-style-type: none"> <li>• Agitation symptoms that are primarily attributable to a condition other than Alzheimer’s disease dementia</li> <li>• History of uncontrolled seizures or a history of epilepsy</li> <li>• Major medical illness or unstable medical condition.</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>
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**5. Where and when will the study be conducted?**

<b>European country involved in the trial</b>	<ul style="list-style-type: none"> <li>• UK</li> </ul>
<b>Estimated start date of recruitment</b>	April 2026

**6. Information for your doctor**

<b>Clinicaltrials.gov identifier</b>	NCT07284472	<b>EU CT Number</b>	2024-517388-22-00
<b>Study contact information</b>	<a href="mailto:studies@exciva.com">studies@exciva.com</a>		
<b>Link to full text</b>	<a href="https://clinicaltrials.gov/study/NCT07284472">https://clinicaltrials.gov/study/NCT07284472</a>		

- ✓ The information contained in this document is based on information available on public registries (e.g. clinicaltrials.gov website) in April 2026.
- ✓ This document has been reviewed by a member of the European Dementia Carers Working Group.