Study of investigational odronextamab versus investigator's choice in patients with previously untreated follicular lymphoma



Primary Objective: evaluate the efficacy and safety of odronextamab vs rituximab + investigator's choice of chemotherapy in patients with previously untreated FL



PATIENTS WITH PREVIOUSLY UNTREATED FL (N≈478)

OPEN LABEL INTERVENTION

Part 1: non-randomized

Odronextamab (IV) safety run-in

Part 2: randomized



Arm 1

Arm 2

Odronextamab (IV)

Rituximab (IV or SC) + investigator's choice of chemotherapy^a

Primary Endpoints

Part 1: Safety, tolerability, and DLT

Part 2: Complete response rate at 30 months (CR30)^b

Key Secondary Endpoints

Part 1:

Pharmacokinetics Progression-free survivalb,c Patient-reported outcomes **Immunogenicity** Event-free survivalb,c Overall survival

> CR30c Objective response rate^{b,c}



FIND OUT MORE

Scan here to find out more about this study at https://clinicaltrials.gov/ct2/show/NCT06091254

This information is intended for investigators interested in open clinical trials.

Odronextamab is an investigational agent and has not been evaluated by any regulatory authority.

Includes cyclophosphamide, doxorubicin, vincristine, prednisone (CHOP); and cyclophosphamide, vincristine, prednisone (CVP) or bendamustine. ^bBased on Independent Central Review (ICR). ^cBased on local investigator assessment. DLT, dose limiting toxicity; FL, follicular lymphoma; IV, intravenous; N, number of patients; R, randomized; SC, subcutaneous.



ODRONEXTAMAB

An Investigational CD20xCD3 Bispecific Antibody
Simultaneously engages CD20 on malignant B cells and CD3 on cytotoxic T cells¹

SELECTED INCLUSION CRITERIA^a

SELECTED EXCLUSION CRITERIA®



Previously untreated CD20+ FL (Grade 1-3a, stage II bulky, or stage III/IV)



CNS lymphoma or leptomeningeal lymphoma



Measurable disease on cross-sectional imaging documented by diagnostic imaging CT or MRI



Histological evidence of transformation to high-grade or diffuse large B-cell lymphoma



ECOG PS ≤2



Waldenström macroglobulinemia, Grade 3b follicular lymphoma, chronic lymphocytic leukemia, or small lymphocytic lymphoma



Adequate bone marrow and hepatic function



Treatment with any systemic anti-lymphoma therapy



For more information, visit <u>www.clinicaltrials.gov</u> or please call [+353 (0)61 533 400 or 844 REGN-MID]. NCT06091254

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alnclusion/exclusion criteria include a summary of selected criteria. Please review the complete study design on clinicaltrials.gov for complete details.

CD, cluster of differentiation; CNS, central nervous system; CT, computed tomography; ECOG PS, Eastern Cooperative Oncology Group performance status; FL, follicular lymphoma; MRI, magnetic resonance imaging.

Current per clinicaltrials.gov as of February 6, 2024.

1. Smith EJ et al. *Sci Rep.* 2015;5:17943. ODRO-EM-0002 v2.0 February 2024. ©2024 Regeneron Pharmaceuticals, Inc. All rights reserved

