Amgen United States

Retrospective Cohort Study Evaluating Clinical Outcomes in Patients with Granulomatosis with Polyangiitis (GPA)/Microscopic Polyangiitis (MPA) Treated with Avacopan (20230026)

First published: 09/04/2024 Last updated 19/10/2024 Study Ongoing Administrative details **PURI** https://redirect.ema.europa.eu/resource/100000065 EU PAS number EUPAS1000000065 Study ID 1000000065 DARWIN EU® study No Study countries **United States** Study status Ongoing Research institutions and networks **Institutions**

First published: 01/02/2024

Last updated 21/02/2024

Institution

Mass General Brigham

Cleveland Clinic Foundation's Center for Vasculitis Care and Research

Contact details

Study institution contact

Global Development Leader Amgen Inc.

Study contact

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Primary lead investigator

Global Development Leader Amgen Inc.

Primary lead investigator

Study timelines

Date when funding contract was signed

Planned:

29/09/2023

Actual:

27/10/2023

Study start date

Planned:

02/04/2024

Actual:

02/04/2024

Data analysis start date

Planned:

01/04/2024

Actual:

02/04/2024

Date of final study report

Planned:

03/04/2026

Sources of funding

Pharmaceutical company and other private sector

Amgen Inc.	
Study protocol =	ol-Published Original avacopan 20230026 .pdf(603.86 KB)
	u-rublished Original avacopan 20230020 .pui(003.80 KB)
Regulatory ——	
Was the study red	quired by a regulatory body?
No	
Is the study requi	red by a Risk Management Plan (RMP)?
is the study requi	red by a Kisk Management Fran (Kivir):
Non-EU R	MP only
Other study racio	stration identification numbers and links —
20230026	stration identification numbers and miks
Methodological asp	pects
Study type ——	
Study type list	
Study topic: Human medici	nal product
Study type:	nui product
Non-interventi	onal study
Scope of the st	udy:
	tudy (incl. comparative)
Main study obj	
	aracteristics associated with avacopan prescription and initiation, and outcomes
	ag avacopan initiators. Among avacopan initiators: (1) investigate factors
resource utiliza	a glucocorticoid discontinuation and relapse risk; and (2) evaluate healthcare
resource utiliza	
Study Design —	
Non-intervention	al study design
Cohort	

Name of medicine

TAVNEOS

Study drug International non-proprietary name (INN) or common name

AVACOPAN

Anatomical Therapeutic Chemical (ATC) code

(L04AJ05) avacopan avacopan

Medical condition to be studied

Granulomatosis with polyangiitis Microscopic polyangiitis

Population studied

Age groups

Adults (18 to < 65 years)

Adults (18 to < 46 years)

Adults (46 to < 65 years)

Adults (65 to < 75 years)

Adults (75 to < 85 years)

Adults (85 years and over)

Estimated number of subjects

305

Study design details

Outcomes

Primary Outcomes:

- Number or participants achieving discontinuation of glucocorticoids for treatment of GPA or MPA 1 month before Month 6 and disease remission within 6 months of avacopan initiation.
- Number of participants achieving sustained remission at Month 12, defined as remission at Month 6 and remission at Month 12, with no glucocorticoid treatment for GPA/MPA 1 month before Month 12, and no relapse between Months 6 and 12.

Data analysis plan

No

There are three primary analyses. Analysis 1 will describe the baseline characteristics of participants by avacopan prescription status (yes/no), as well as the baseline characteristics of participants prescribed avacopan by initiation status (yes/no). Analyses 2 and 3 will examine the 6- and 12-month outcomes among initiators of avacopan, respectively. Analysis 1 will be performed at least six months after the study start date when the final cohort of avacopan initiators is identified and baseline data are collected. Analyses 2 and 3 will begin once at least 6 months after the last avacopan participant is identified during the inclusion period and at least 6 months of follow-up data are collected. The primary outcomes will be analysed using a descriptive approach. Remission will be based on the Birmingham Vasculitis Activity Score version 3 with/out glucocorticoids for the treatment of GPA/MPA in the 4 weeks prior to the month 6 or 12 visits. Summary statistics for continuous variables, including the number of patients, mean, standard deviation (SD), median, Q1, Q3, minimum and maximum, will be calculated after excluding missing/unknown values.

Data management —	
— Data sources —	
Data source(s), other	
Data source(s), other	
Mass General Brigham (MGB) Electronic Data Warehouse (EDW)	
Research patient data registry (RPDR)	
Cleveland Clinic Foundation's Center for Vasculitis Care and Research	
Social Security Master Death File	
NIH United States Renal Data Set	
National Death Index	
Data sources (types)	
Zum sources (cypes)	
Electronic healthcare records (EHR)	
Other	
Use of a Common Data Model (CDM)	
(02 1/2)	
CDM mapping	
No	
110	
— Data quality specifications ————————————————————————————————————	
Check conformance	
Check Comformance	

Check completeness
No
Check stability
No
Check logical consistency
No
Data characterisation
Data characterisation conducted
No