Post-marketing Surveillance of Spesolimab I.V. Infusion in improvement of Generalized Pustular Psoriasis (GPP) with acute symptoms in Japan (PMS for GPP with acute symptoms)

First published: 25/11/2022 Last updated: 02/09/2024





Administrative details

EU PAS number
EUPAS49953
Study ID
49954
DARWIN ELLO study
DARWIN EU® study
No
Charden accombala a
Study countries
Japan

Study description

The primary outcome is the incidence of investigator/BI-defined drug reactions (including safety concerns: Serious infections, Serious hypersensitivity).

Study status

Planned

Research institutions and networks

Institutions

Boehringer Ingelheim

First published: 01/02/2024

Last updated: 01/02/2024

Institution

Contact details

Study institution contact

Yukako Ogi zzCDMJP_PV_PMS@boehringer-ingelheim.com

Study contact

 $zz CDMJP_PV_PMS@boehringer-ingelheim.com$

Primary lead investigator

Yukako Ogi

Primary lead investigator

Study timelines

Date when funding contract was signed

Planned: 22/09/2022 Actual: 22/09/2022

Study start date

Planned: 01/03/2023

Date of final study report

Planned: 31/12/2026

Sources of funding

Pharmaceutical company and other private sector

More details on funding

Nippon Boehringer Ingelheim Co., Ltd.

Regulatory

Was the study required by a regulatory body?

Yes

Is the study required by a Risk Management Plan (RMP)?

Non-EU RMP only

Methodological aspects

Study type

Study type list

Study type:

Non-interventional study

Scope of the study:

Assessment of risk minimisation measure implementation or effectiveness

Disease epidemiology

Drug utilisation

Main study objective:

The primary outcome is the incidence of investigator/BI-defined drug reactions (including safety concerns: Serious infections, Serious hypersensitivity).

Study Design

Non-interventional study design

Cohort

Study drug and medical condition

Medicinal product name

SPEVIGO

Medical condition to be studied

Pustular psoriasis

Additional medical condition(s)

acute symptoms of generalized pustular psoriasis

Population studied

Age groups

- 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

40

Study design details

Outcomes

The incidence of adverse drug reactions (ADRs)

Data analysis plan

analyses are descriptive in nature, including confidence intervals.

Data management

ENCePP Seal

The use of the ENCePP Seal has been discontinued since February 2025.

The ENCePP Seal fields are retained in the display mode for transparency but are no longer maintained.

Data sources

Other	(types)				
Data sources	(types), othe	r			
Prospective pa	ient-based dat	a collectio	n		
Use of a (Common	Data N	Model (CDM)	
CDM mapping					
No					
Data qua	ity spacit	fication	2.5		
Data qua	ity specii	icatioi	15		
Check confor		icatioi	15		
•		icatioi	15		
Check confor	nance	icatioi	15		
Check confor	nance	icatioi	15		
Check conford Unknown Check comple	nance teness	icatioi	15		

Data characterisation

Data characterisation conducted

No