

# An Immuno-Dermatological disease registry to understand the burden of Atopic dermatitis (AD), Alopecia areata (AA), and Vitiligo in Indian Patients

**First published:** 17/04/2023

**Last updated:** 19/01/2026

Study

Ongoing

## Administrative details

### EU PAS number

EUPAS48566

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### Study ID

106116

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### DARWIN EU® study

No

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### Study countries

India

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### Study description

A prospective observational multi-center study in India to evaluate the epidemiological burden of mild, moderate and severe atopic dermatitis, vitiligo and alopecia areata.

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### **Study status**

Ongoing

## Research institutions and networks

### Institutions

[Pfizer](#)

**First published:** 01/02/2024

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**Institution**

- [DY Patil Hospital](#)
- [Father Muller Medical College Hospital](#)
- [Wizderm Specialty Skin and Hair Clinic](#)
- [Vibrance Skin Clinic](#)
- [Indushree Skin Clinic](#)
- [Lokmanya Tilak Municipal General Hospital and Lokmanya Tilak Municipal Medical College](#)

- Bharati Vidyapeeth (deemed to be University)  
Medical College Hospital & Research Centre
- KPC Medical College and Hospital
- Ayana Clinic
- B. J. Medical College & Civil Hospital
- Venkateshwar Hospital
- Dermaworld Skin Clinic
- Dr. D.Y.Patil Medical College, Hospital and  
Research
- Maharishi Markandeshwar Institute of Medical  
sciences and Research
- Indus International Hospital
- Anisha Private Clinic

## Networks

Insignia

## Contact details

**Study institution contact**

Sreenath Hariharan Sreenath.Hariharan@pfizer.com

Study contact

[Sreenath.Hariharan@pfizer.com](mailto:Sreenath.Hariharan@pfizer.com)

**Primary lead investigator**

Lavinia Popa

Primary lead investigator

## Study timelines

**Date when funding contract was signed**

Planned: 02/05/2022

Actual: 28/06/2022

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**Study start date**

Planned: 29/09/2023

Actual: 28/08/2023

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**Date of interim report, if expected**

Planned: 31/01/2026

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**Date of final study report**

Planned: 30/12/2026

## Sources of funding

- Pharmaceutical company and other private sector

## More details on funding

Pfizer

## Study protocol

[B7451103 Non-Interventional Study Protocol Version 1.0\\_ 30 September 2022\\_Redacted.pdf](#) (3.71 MB)

[B7451103\\_NIS Protocol v3.0\\_Amendment 2 Clean Copy\\_23DEC2024\\_Redacted.pdf](#) (4.07 MB)

## Regulatory

**Was the study required by a regulatory body?**

No

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

Not applicable

## Methodological aspects

Study type

Study type list

**Study topic:**

Disease /health condition

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**Study type:**

Non-interventional study

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**Scope of the study:**

Disease epidemiology

**Data collection methods:**

Primary data collection

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**Main study objective:**

The objective of this study is to evaluate the epidemiological burden of mild, moderate and severe atopic dermatitis, vitiligo, and alopecia areata across enrolled dermatology centers.

## Study Design

**Non-interventional study design**

Cohort

Cross-sectional

Other

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**Non-interventional study design, other**

Prospective longitudinal study

## Study drug and medical condition

## **Medical condition to be studied**

Dermatitis atopic

Alopecia areata

Vitiligo

## Population studied

### **Age groups**

- Children (2 to < 12 years)
  - Adolescents (12 to < 18 years)
  - Adults (18 to < 46 years)
  - Adults (46 to < 65 years)
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### **Estimated number of subjects**

3000

## Study design details

### **Outcomes**

To evaluate the epidemiological burden of mild, moderate and severe atopic dermatitis, vitiligo and alopecia areata in patients across dermatology centers in India.

Explain the current diagnostic criteria and grading modalities for AD, vitiligo and AA, to understand the burden of disease with a demographic overview Better understand the treatment for the diseases: topical, advanced therapies, surgical interventions and cosmetic procedures, the Unmet needs in diagnosis and management of dermatological disorders. Focus on patients' perspectives on benefits.

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## Data analysis plan

Descriptive statistics will be presented to describe patient characteristics. Categorical covariates will be described by frequency distribution while continuous covariates expressed in terms of their mean and standard deviation or median and interquartile range (IQR) as appropriate.

The bivariate analysis will be conducted to determine if there is any association between the outcome and the exposure (the covariates).

Unadjusted comparisons between groups of covariates and outcomes will be evaluated using appropriate tests: Pearson's chi-square test will be used for dichotomous categorical variables, t-test will be used for continuous variables, and the Mann-Whitney U test for non-parametric variables, p-values will be generated and 5% level of significance will be considered.

In case of multiple testing or stepdown procedure either Tukey HSD or Bonferroni adjustment to p-value will be applied.

## Documents

### Study, other information

[B7451103 Non Interventional Study Protocol Abstract V2.0\\_19 April 2023\\_Redacted.pdf](#) (719.94 KB)

[B7451103 Protocol Abstract 30 September 2022\\_Redacted.pdf](#) (1.89 MB)

[B7451103\\_NIS Protocol Abstract V3.0\\_Clean Copy\\_23DEC2024\\_Redacted.pdf](#) (732.41 KB)

## 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

### Data sources (types)

Other

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### Data sources (types), other

Prospective patient-based data collection

## Use of a Common Data Model (CDM)

### CDM mapping

No

## Data quality specifications

### Check conformance

Unknown

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### Check completeness

Unknown

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### Check stability

Unknown

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**Check logical consistency**

Unknown

**Data characterisation**

**Data characterisation conducted**

No