

# Long-term follow-up of pediatric patients exposed to nivolumab + relatlimab fixed-dose combination (FDC) enrolled in the Dutch Melanoma Treatment Registry (DMTR) (CA224-122)

**First published:** 13/12/2023

**Last updated:** 20/02/2024

Study

Planned

## Administrative details

### EU PAS number

EUPAS107906

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### Study ID

107907

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### DARWIN EU® study

No

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### Study countries

Netherlands

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## Study description

This post-authorization safety study (PASS) will collect long-term follow-up data in pediatric patients exposed to nivolumab + relatlimab fixed-dose combination, and is part of EMA approved European Union (EU) Risk Management Plan. Data will be collected through the Dutch Melanoma Treatment Registry

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## Study status

Planned

## Research institutions and networks

### Institutions

#### Bristol-Myers Squibb (BMS)

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Institution

### Networks

#### Dutch Melanoma Treatment Registry (DMTR)

## Contact details

### Study institution contact

Elise Roy [ctt.group@bms.com](mailto:ctt.group@bms.com)

Study contact

[ctt.group@bms.com](mailto:ctt.group@bms.com)

**Primary lead investigator**

Elise Roy

Primary lead investigator

## Study timelines

**Date when funding contract was signed**

Planned: 29/02/2024

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**Study start date**

Planned: 30/06/2025

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**Data analysis start date**

Planned: 30/06/2025

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**Date of interim report, if expected**

Planned: 31/12/2026

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**Date of final study report**

Planned: 31/12/2038

## Sources of funding

- Pharmaceutical company and other private sector

## More details on funding

Bristol-Myers Squibb

## Regulatory

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

Yes

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

EU RMP category 3 (required)

## Methodological aspects

### Study type

### Study type list

#### **Study type:**

Non-interventional study

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#### **Main study objective:**

The main objective is to evaluate Grade 3 to 4 adverse drug reactions (including immune-related adverse reactions) experienced by pediatric patients treated with nivolumab + relatlimab FDC, and their management.

## Study Design

## Non-interventional study design

Cohort

# Study drug and medical condition

## Medicinal product name

OPDUALAG

# Population studied

## Age groups

- Adolescents (12 to < 18 years)

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## Estimated number of subjects

20

# Study design details

## Outcomes

Number of participants with Frequency of Grade 3 to 4 adverse drug reactions (ADRs) as assessed by the Common Terminology Criteria for Adverse Events (Version 6) criteria ADR management, Participant baseline demographic, comorbidities, disease characteristics, treatment history Dose levels and frequency of nivolumab+relatlimab (nivo+rela) FDC treatment Number of nivo+rela FDC treatment infusions Number of participants with nivo+rela FDC treatment dose interruptions or discontinuations, subsequent therapies, or growth/development disorders Overall survival Time to progression

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## Data analysis plan

A detailed Statistical Analysis Plan will be developed for this study. General descriptive statistics will include mean, median, minimum, maximum, and standard deviation for continuous variables, count and percentages will be used to examine categorical variables. The time to event endpoint will be analyzed according to the Kaplan-Meier method, as data allow. Data from all patients who receive at least 1 dose of treatment will be analyzed. The treated set is defined as all patients enrolled in the registry and meeting the study eligibility criteria and receiving at least 1 dose of treatment. The data collected at baseline will be used to characterize the population. Descriptive statistics will be provided to assess demographic information, disease characteristics and other clinical characteristics, and treatment history.

## 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 source(s), other

DMTR Netherlands

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### Data sources (types)

Other

## 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