

# Surveillance of Adverse Event related to childhood vaccination reported by parents

**First published:** 29/05/2014

**Last updated:** 01/04/2024

Study

Finalised

## Administrative details

### EU PAS number

EUPAS5350

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### Study ID

28000

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### DARWIN EU® study

No

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### Study countries

 Italy

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### Study description

Background: In recent years in Italy the trend of spontaneous reports of suspected adverse reactions from vaccine has been influenced by several factors. In 2009 there was an increase in reporting linked to the pandemic

influenza vaccine, while in 2011-2012 many reports are linked to the active post licensure safety monitoring of HPV vaccine. The new Pharmacovigilance legislation came into force in Europe in July 2012 and one of the indication was to increase the involvement of patients in reporting systems. Objective: the surveillance of adverse event after immunization (AEFI) reported by patients. Methods: Every adverse event should be reported immediately to healthcare professional in the vaccination center. The safety of vaccines is linked to the constant surveillance of adverse events. In this study all vaccinated children until to 15 months age were involved. For each vaccination parents will receive a Vaccination Diary for surveillance of adverse events by the healthcare professional, accompanied by a letter describing this project and reminding the importance of reporting. First of all healthcare professional will complete the Vaccination Diary in the vaccine section and then the parents should complete the Vaccination Diary describing any adverse event occurred after administration of vaccine. This Vaccination Diary may be delivered to next vaccination or sent by fax or e-mail, even if parents don't observe any adverse event. This pilot study lasts three months from 1th October/November 2013 until 31st December/January 2013. All Vaccination Diaries delivered by parents will be collected and sent to the Regional Centre of Pharmacovigilance. These diaries with an adverse events will be included in the Italian Pharmacovigilance database.

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## **Study status**

Finalised

## Research institutions and networks

### Institutions

## Pharmacology Unit - Veneto Pharmacovigilance Centre (Pharmacol UNIVR), University Hospital Verona

 Italy

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**Last updated:** 13/03/2025

**Institution**

Educational Institution

Hospital/Clinic/Other health care facility

ENCePP partner

## Department of Biomedical and Dental Sciences and Morphofunctional Imaging, University of Messina

 Italy

**First published:** 29/11/2021

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**Institution**

Educational Institution

Hospital/Clinic/Other health care facility

One local health unit Veneto Region, One local health unit Liguria Region, One local health unit Emilia -Romagna Region, Three local health units Piemonte Region, Nine local health units Sicily

Region, One local health unit Calabria Region,  
Three local macro-areas Autonomy Trento Region

## Contact details

### Study institution contact

Moretti Ugo [ugo.moretti@univr.it](mailto:ugo.moretti@univr.it)

Study contact

[ugo.moretti@univr.it](mailto:ugo.moretti@univr.it)

### Primary lead investigator

Moretti Ugo

Primary lead investigator

## Study timelines

### Date when funding contract was signed

Planned: 01/01/2011

Actual: 01/01/2011

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### Study start date

Planned: 01/01/2012

Actual: 01/01/2012

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### Data analysis start date

Planned: 01/05/2014

Actual: 01/05/2014

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### **Date of final study report**

Planned: 01/10/2014

Actual: 09/02/2015

## Sources of funding

- Other

## More details on funding

Italian Agency Medicines

## 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:**

Human medicinal product

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**Study type:**

Non-interventional study

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**Scope of the study:**

Assessment of risk minimisation measure implementation or effectiveness

**Data collection methods:**

Primary data collection

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**Main study objective:**

To involve parents in the spontaneous reporting system of Adverse Event following immunization (AEFI). Therefore to propose a routinary procedure in the local health units.

## Study Design

**Non-interventional study design**

Other

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**Non-interventional study design, other**

Intensive monitoring schemes

## Study drug and medical condition

**Anatomical Therapeutic Chemical (ATC) code**

(J07) VACCINES

## Population studied

### **Short description of the study population**

Vaccinated children until to 15 months age.

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### **Age groups**

- Preterm newborn infants (0 - 27 days)
  - Term newborn infants (0 - 27 days)
  - Infants and toddlers (28 days - 23 months)
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### **Estimated number of subjects**

42000

## Study design details

### **Outcomes**

increase of reporting rate of vaccines

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### **Data analysis plan**

All AEFI reports will be introduced in the Italian Pharmacovigilance Database. All data will be evaluated in order to find potential signals. Reports from parents will be compared with reports from health operators

## 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)

[Spontaneous reports of suspected adverse drug reactions](#)

## 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**

Unknown