

# Effectiveness, safety and efficiency of the use of vernakalant in the control of the rhythm of atrial fibrillation in the emergency. Cohort study (VERITA)

**First published:** 15/02/2023

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Study

Finalised

## Administrative details

### EU PAS number

EUPAS103551

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### Study ID

103552

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### DARWIN EU® study

No

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### Study countries

 Spain

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### Study description

Atrial fibrillation (AF) is the most prevalent sustained arrhythmia in clinical practice in hospital emergency departments. Complications of AF include thromboembolism and heart failure. In addition, affected patients may be at increased risk of mortality. Within the acute management of AF, two different strategies should be considered: rate control (usually with a beta-blocker or calcium channel inhibitor) and rhythm control (electrical or pharmacological cardioversion). Rhythm and rate control strategies are associated with similar rates of mortality and morbidity, such as embolic risk. The decision to adopt a therapeutic strategy is usually dictated by the presence of symptoms associated with AF and/or the development of left ventricular systolic dysfunction believed to be secondary to the arrhythmia. According to the ESC guidelines, there are different options for pharmacological cardioversion: flecainide and propafenone (class Ic antiarrhythmics), intravenous amiodarone, and vernakalant. Other alternatives, such as dofetilide and ibutilide, are not marketed in Europe. On June 24, 2010, the Committee for Medicinal Products for Human Use (CHMP) of the European Medicines Agency (EMA) adopted a positive opinion, recommending the marketing authorization of vernakalant, indicated for the rapid conversion to sinus rhythm of AF. of recent onset in adults. For its part, the United States Food and Drug Administration (FDA) denied the marketing application for vernakalant due to safety concerns: severe hypotension, ventricular arrhythmias, conduction abnormalities, and death. Currently, the management in terms of AF rhythm control in emergency services varies depending on the idiosyncrasy of each hospital and its own experience of use. In this context, a retrospective observational study is proposed to evaluate the efficiency and safety of vernakalant in clinical practice.

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## **Study status**

Finalised

## **Research institutions and networks**

# Institutions

## Puerta de Hierro-Majadahonda University Hospital

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Institution

## Contact details

### Study institution contact

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Study contact

[mariabelen.ruiz@salud.madrid.org](mailto:mariabelen.ruiz@salud.madrid.org)

### Primary lead investigator

Ruiz-Antorán Belén

Primary lead investigator

## Study timelines

### Date when funding contract was signed

Planned: 06/06/2022

Actual: 06/06/2022

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### Study start date

Planned: 01/09/2022

Actual: 01/09/2022

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### **Data analysis start date**

Planned: 31/10/2023

Actual: 14/02/2023

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### **Date of interim report, if expected**

Planned: 29/12/2023

Actual: 14/02/2023

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

Planned: 01/06/2024

Actual: 14/02/2023

## Sources of funding

- Other

## More details on funding

The study is not funded

## Study protocol

[PROTOCOLO ESTUDIO VERNAKALANT EN URGENCIAS VERSION 25052022.pdf](#)  
(222.45 KB)

## 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

Disease /health condition

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**Study type:**

Non-interventional study

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**Scope of the study:**

Drug utilisation

Effectiveness study (incl. comparative)

Safety study (incl. comparative)

**Data collection methods:**

Secondary use of data

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**Main study objective:**

To evaluate the effectiveness and safety of vernakalant for the rhythm control of AF in the context of clinical practice, in comparison with its treatment alternatives.

## Study Design

### **Non-interventional study design**

Cohort

Other

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### **Non-interventional study design, other**

Multicenter retrospective, post-authorization drug study

## Study drug and medical condition

### **Anatomical Therapeutic Chemical (ATC) code**

(C01BG11) vernakalant

vernakalant

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### **Medical condition to be studied**

Atrial fibrillation

## Population studied

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

Patients diagnosed with atrial fibrillation received treatment with vernakalant, amiodarone, flecainide, propafenone or electrical cardioversion from 2012 to

2022.

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### **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)
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### **Special population of interest**

Other

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### **Special population of interest, other**

Atrial fibrillation patients

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### **Estimated number of subjects**

352

## Study design details

### **Outcomes**

Percentage of patients with reversion to sinus rhythm in the first hour after treatment, - Percentage of patients with reversion to sinus rhythm in the first 12 hours/24 hours. - Percentage of patients requiring hospitalization. - Hospital stay - Percentage of patients with AF recurrence in the first 5 days/ 30 days/ 6 months

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### **Data analysis plan**

The incidence in exposed (cohort treated with vernakalan and in non-exposed (treated with other cardioversión drug) will be calculated for each of the

effectiveness variables. From these, the measures of association relative risk (RR), absolute risk reduction (RAR) and relative risk reduction (RRR) will be calculated for each of the variables, with their 95% CIs. The RR adjusted for comorbidities and prognostic factors will be estimated.

## 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)

[Electronic healthcare records \(EHR\)](#)

## 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