

# How many and who are patients with heart failure eligible to SGLT2 inhibitors?

## Responses from the combination of administrative healthcare and primary care databases

**First published:** 15/07/2024

**Last updated:** 15/07/2024

Study

Finalised

### Administrative details

#### EU PAS number

EUPAS1000000259

#### Study ID

1000000259

#### DARWIN EU® study

No

#### Study countries

☐ Italy

## Study description

Background: Recent successful findings (i.e. DAPA-HF trial) in patients with heart failure (HF) with/without diabetes treated with sodium-glucose co-transporter inhibitors (SGLT2-I) have fostered real-world data analyses.

Fondazione Ricerca e Salute's (ReSD) administrative and Health Search's (HSD) primary healthcare databases were combined in the ReS-HS DB Consortium, to identify and characterize HF-patients eligible to SGLT2-I, and assess their costs charged to the Italian National Health Service (INHS).

Methods and results: Eligibility to SGLT2-I was HF diagnosis, age  $\geq 18$  years, reduced ( $\leq 40\%$ ) ejection fraction (HFrEF) and glomerular filtration rate (GFR)  $\geq 30$  ml/min. The HSD, including 13,313 HF-patients (1.5% of the total HSD population) was used to develop and test the algorithms for imputing HFrEF and GFR  $\geq 30$  ml/min, based on a set of covariates, to the ReSD, including 67,369 (1.5% of the total ReSD population). Subjects eligible to SGLT2-I were 2187 in HSD (61.1% of HFrEF); after the imputation, 15,145 in ReSD (58.8% of HFrEF). Prevalence of eligibility to SGLT2-I was higher in males than in females and increased with age; diabetic patients were 44.3% and 33.4% of HSD and ReSD populations eligible to SGLT2-I, respectively. Estimated from ReSD, the mean annual cost charged to the INHS per patient with HF eligible to SGLT2-I was €7122 (68% due to hospitalizations).

Conclusions: Approximately 20% of patients with HF was eligible to SGLT2-I.

Real-world data can identify,

quantify and characterize patients eligible to SGLT2-Is and assess related costs for the health care system, thus

providing useful information to Regulatory Decision makers.

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## Study status

Finalised

## Research institutions and networks

## Institutions

### Fondazione ReS (Ricerca e Salute), CINECA partner

☐ Italy

**First published:** 05/07/2017

**Last updated:** 01/10/2025

**Institution**

Not-for-profit

ENCePP partner

### Health Search, Italian College of General Practicioners

☐ Italy

**First published:** 02/03/2010

**Last updated:** 20/08/2024

**Institution**

Educational Institution

Other

## Contact details

### Study institution contact

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**Study contact**

[piccinni@fondazioneres.it](mailto:piccinni@fondazioneres.it)

### Primary lead investigator

Letizia Dondi

Primary lead investigator

## Study timelines

### **Date when funding contract was signed**

Actual: 12/01/2022

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### **Study start date**

Actual: 10/02/2022

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

Actual: 02/04/2022

## Sources of funding

- Pharmaceutical company and other private sector

## More details on funding

Astra Zeneca SpA

## 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 topic:**

Disease /health condition  
Human medicinal product

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**Study type:**

Non-interventional study

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**Scope of the study:**

Disease epidemiology  
Drug utilisation  
Healthcare resource utilisation

**Data collection methods:**

Combined primary data collection and secondary use of data

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**Study design:**

Retrospective longitudinal cohort study

**Main study objective:**

To identify and characterize HF-patients eligible to SGLT2-I, and  
assess their costs charged to the Italian National Health Service (SSN)

## Study Design

**Non-interventional study design**

Cohort

## Study drug and medical condition

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

(A10BK) Sodium-glucose co-transporter 2 (SGLT2) inhibitors

Sodium-glucose co-transporter 2 (SGLT2) inhibitors

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

Cardiac failure

## Population studied

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

Patients with HF eligible to the SGLT2-Is dapagliflozin, regardless of diabetes (according to the DAPA-HF trial's eligibility criteria) were identified from the Health Search primary care database and from the Fondazione ReS administrative database

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### **Age groups**

- **Adult and elderly population ( $\geq 18$  years)**

## Study design details

### **Setting**

In-hospital and local outpatient setting in public and affiliated with SSN facilities, as regards the Database of Fondazione ReS.

Primary care as regards the Health Search database

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### **Summary results**

The HSD, including 13,313 HF-patients (1.5% of the total HSD population) was used to develop and test the algorithms for imputing HFrEF and  $GFR \geq 30$  ml/min, based on a set of covariates, to the ReSD, including 67,369 (1.5% of the

total ReSD population). Subjects eligible to SGLT2-I were 2187 in HSD (61.1% of HFrEF); after the imputation, 15,145 in ReSD (58.8% of HFrEF). Prevalence of eligibility to SGLT2-I was higher in males than in females and increased with age; diabetic patients were 44.3% and 33.4% of HSD and ReSD populations eligible to SGLT2-I, respectively. Estimated from ReSD, the mean annual cost charged to the INHS per patient with HF eligible to SGLT2-I was €7122 (68% due to hospitalizations).

## Documents

### Study publications

[How many and who are patients with heart failure eligible to SGLT2 inhibitors?...](#)

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## 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)

Database of Fondazione ReS

**Data sources (types)**

Administrative healthcare records (e.g., claims)

Electronic healthcare records (EHR)

## Use of a Common Data Model (CDM)

**CDM mapping**

No

## Data quality specifications

**Check conformance**

Yes

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**Check completeness**

Yes

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**Check stability**

Yes

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**Check logical consistency**

Yes

## Data characterisation

**Data characterisation conducted**

Yes