The Effectiveness and Safety of Single 6ml Hyaluronic Acid Intra-articular Injection (Suplasyn® 1-shot) in Managing Symptomatic Primary Osteoarthritis of the Knee in Real-life Practice: ESSIK Survey

First published: 27/11/2015 Last updated: 01/04/2024



# Administrative details

#### **EU PAS number**

EUPAS11707

#### **Study ID**

28693

#### DARWIN EU® study

No

#### **Study countries**

⊂Croatia

⊂Czechia



### **Study description**

The current study is designed to assess the effectiveness and safety of one 6ml injection of Suplasyn® 1-shot in a 26-week, international, multicenter, noninterventional observational study of patients recommended with a single intraarticular 6ml injection for the treatment of knee osteoarthritis. The intention is to assess the efficacy and safety of the treatment in real-life practice. Our primary target is to obtain and verify long-term outcomes from a naturalistic primary care experience. Rationale for using viscosupplementation is to restore the protective viscoelasticity of synovial hyaluronic acid, decrease pain and improve mobility. Immediate benefit of viscosupplementation is the relief of pain while long-term benefits results are believed to include the return of joint mobility by the restoration of trans-synovial flow and the metabolic and rheological homeostasis of the joint (31). Short duration of HA within the joint does not fully explain the indisputable long-term clinical efficacy seen in practice (32). In this case, it is appropriate to evaluate, under real-life conditions, the short-term and long-term effectiveness of a single injection of Suplasyn® 1-shot 6ml intra-articular injections and to how the extended treatment regime may impact patient satisfaction or treatment safety.

#### **Study status**

Finalised

# Research institutions and networks

Institutions

## **Krnov Hospital**

Multiple centres: 21 centres are involved in the study

## Contact details

Study institution contact Pavel Martinek clever@recerca.com

Study contact

clever@recerca.com

**Primary lead investigator** Pavel Martinek

Primary lead investigator

# Study timelines

**Date when funding contract was signed** Actual: 22/05/2014

Study start date Actual: 16/06/2014

Date of final study report

Planned: 01/12/2016 Actual: 27/02/2018

# Sources of funding

• Pharmaceutical company and other private sector

## More details on funding

Mylan Institutional International

# Study protocol

ESSIK Protocol\_Adapted\_130614.pdf(266.04 KB)

# Regulatory

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

No

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

Not applicable

## Methodological aspects

Study type

Study type list

### **Study topic:**

Disease /health condition Human medicinal product

### Study type:

Non-interventional study

### Scope of the study:

Assessment of risk minimisation measure implementation or effectiveness Effectiveness study (incl. comparative)

### Data collection methods:

Primary data collection

### Main study objective:

The intention is to assess the efficacy and safety of the treatment in real-life practice. Our primary target is to obtain and verify long-term outcomes from a naturalistic primary care experience.

# Study Design

## Non-interventional study design

Other

Non-interventional study design, other

Prospective Cases Report

# Study drug and medical condition

#### Name of medicine, other

Suplasyn 1-shot

#### Medical condition to be studied

Osteoarthritis

## Population studied

#### Short description of the study population

Patients over 18 years old with Primary Knee Osteoarthritis (Kellgren's grades I to III) without effusion.

#### 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)

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

Other

#### Special population of interest, other

Knee Osteoarthritis patients

#### **Estimated number of subjects**

300

## Study design details

#### Outcomes

To assess changes in clinical outcome of patients with OA recommended with a single injection with Suplasyn® 1-shot viscosupplementation for knee OA during a 26 weeks follow-up. - To evaluate the safety profile and adverse events (AEs) of Suplasyn® 1-shot injection.- To evaluate the concomitant consumption of permitted rescue medications (analgesics, NSAIDS) throughout the study.- To evaluate characteristics of the beneficiary population (Intended to treat).

### Data analysis plan

The primary efficacy endpoint is the change from baseline in the Oxford knee score assessing the function of the OA knee and the change from baseline in the patient-rated knee OA pain assessment at 4 and 26 weeks.Both scores obtained with the Oxford Knee Score and Pain VAS will be registered at 1 and 6 months following viscosupplementation of the knee.Secondary objectives correspond to the evaluation of the safety profile and adverse events (AEs) of Suplasyn® 1-shot 6ml injection, assessment of concomitant consumption of permitted rescue medications (analgesics, NSAIDs) throughout the study and the known the characteristics of the beneficiary population (intend-totreat).Continuous variables will be expressed as mean ± Standard Deviation (SD) while categorical binary variables will be presented as percentages.Both primary and secondary objectives will be analyzed with tests for repeated measures to determinate the evolution of the variables throughout study visits.

## Documents

Study results EUPAS11707-28691.pdf(4.55 MB)

## Data management

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

#### **Check completeness**

Unknown

#### **Check stability**

Unknown

#### **Check logical consistency**

Unknown

## Data characterisation

### **Data characterisation conducted**

No