# SHP503-803: Drug Utilization Study with Intuniv® in Australia

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# Administrative details

<b>EU PAS number</b> EUPAS40684		
Study ID		
44620		
DARWIN EU® study		
No		
Study countries  Australia		

## **Study description**

This study will evaluate and analyze prescribing behaviors of physicians and determine whether Intuniv was correctly prescribed in Australia.

#### **Study status**

**Finalised** 

## Research institutions and networks

## Institutions

## Shire

First published: 01/02/2024

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Institution

## Contact details

## **Study institution contact**

Study Contact Shire ClinicalTransparency@shire.com

Study contact

ClinicalTransparency@shire.com

## **Primary lead investigator**

Study Contact Shire

**Primary lead investigator** 

# Study timelines

Date when funding contract was signed

Actual: 30/05/2017

#### Study start date

Actual: 11/03/2019

#### Data analysis start date

Actual: 01/05/2019

#### Date of interim report, if expected

Actual: 28/08/2020

#### Date of final study report

Actual: 27/08/2021

# Sources of funding

• Pharmaceutical company and other private sector

# More details on funding

Shire

# Study protocol

SHP503-803- protocol-original-redact.pdf(758.68 KB)

# Regulatory

Was the study required by a regulatory body?

Yes

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

Non-EU RMP only

# Other study registration identification numbers and links

CT.gov: NCT04866030,To obtain more information on the study, click here/on

this link: https://clinicaltrials.takeda.com/study-

detail/609017abf89629001e47b500

# Methodological aspects

# Study type

# Study type list

## **Study topic:**

Disease /health condition

Human medicinal product

## **Study type:**

Non-interventional study

## Scope of the study:

Drug utilisation

#### **Data collection methods:**

Combined primary data collection and secondary use of data

#### Main study objective:

The overall objective is to provide data on an annual basis for 3 years in Australia to evaluate drug utilization and monitor off-label use of Intuniv®

# Study drug and medical condition

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

(C02AC02) guanfacine guanfacine

#### Medical condition to be studied

Attention deficit hyperactivity disorder

# Population studied

#### Short description of the study population

Intuniv® for treatment of ADHD in children and adolescents in Australia in 2018 Patients who have been prescribed Intuniv at least once during the reporting period. For this third (final) study report, the observation period was from 01 February 2020 to 31 January 2021 for the annual and from 01 February 2018 (the launch date in Australia) to 31 January 2021 for the cumulative reporting period.

Subjects and Study Size, Including Dropouts

The following numbers of patients and prescriptions are included in this third (final) report.

- NostraData database:
- o Annual reporting period: 21,028 patients, 152,837 prescriptions
- o Cumulative reporting period: 27,461 patients, 260,460 prescriptions

• Annual physician survey (2021): 26 physicians, 104 patients.

#### Age groups

Children (2 to < 12 years)

Adolescents (12 to < 18 years)

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

Other

#### Special population of interest, other

Patients with attention deficit hyperactivity disorder

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

100

# Study design details

#### **Outcomes**

1.Number of Participants Based on Indication of Use of Intuniv 2.Number of Participants with Presence/Absence of Contraindications, 1.Number of Participants Based on Patterns of Drug Use 2.Number of Participants Stratified by Prescriber Information Based on Physician Survey 3.Frequency of Weight Monitoring of Participants by Physician 4.Frequency of Blood Pressure Monitoring of Participants by Physician 5.Frequency of Heart Rate Monitoring of Participants by Physician

## Data analysis plan

Descriptive analyses will be performed based on the prescriptions contained in the NostraData database from each 12-month reporting period. The use of Intuniv® will be analysed using the prescriptions collected during the 3-year assessment period.

## **Documents**

#### **Study results**

SHP503-803-clinical-study-report-redact.pdf(732.02 KB)

# 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)

Drug dispensing/prescription data
Other

## Data sources (types), other

Prescription event monitoring

# 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