A Non-Interventional Post-Authorization Safety Study (NI-PASS) as an effectiveness check of a Patient Card for Padcev™

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Administrative details

EU PAS number	
EUPAS104456	
Study ID	
104457	
DARWIN EU® study	
No	
Study countries	
France	
Germany	
☐ Italy	
Poland	

Spain		
Sweden		
Switzerland		

Study description

Padcev is a treatment for cancer in the bladder lining (urothelial cancer). Padcev is injected through a vein (known as an infusion). Padcev is now available in some countries in Europe.

People in this study will be adults with locally advanced (the cancer has spread to nearby tissue) or metastatic urothelial cancer. Metastatic means the cancer has spread to other parts of the body.

During their care, the person's doctor will have prescribed Padcev treatment and other medicines to treat their cancer.

People in the study will be treated according to their clinic's standard practice. This study is about collecting information only.

This study will survey people who know they are receiving Padcev treatment and have been given an information leaflet called a Patient Card.

The aim of the study is to check how well people understand the information on the Patient Card, including the risk of skin reactions that can be caused by Padcev treatment. Once a doctor has prescribed Padcev treatment, the patient (or their caregiver) will be asked if they would like to complete a survey. The survey will be completed on a computer and should take about 15 minutes to complete.

Study status

Ongoing

Contact details

Study institution contact

Clinical Trial Registration Department clinicaltrialregistration@astellas.com

Study contact

clinicaltrialregistration@astellas.com

Primary lead investigator

Samantha Kimball

Primary lead investigator

Study timelines

Date when funding contract was signed

Planned: 01/01/2023

Actual: 21/09/2022

Study start date

Planned: 02/01/2024

Actual: 27/03/2024

Data analysis start date

Planned: 07/05/2025

Date of final study report

Planned: 29/07/2025

Sources of funding

• Pharmaceutical company and other private sector

More details on funding

Astellas

Regulatory

Was the study required by a regulatory body?

Yes

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

EU RMP category 3 (required)

Other study registration identification numbers and links

7465-PV-0002

Methodological aspects

Study type

Study type list

Study type:

Non-interventional study

Main study objective:

The objectives of this NI-PASS are to assess patients' (or caregivers') knowledge of the risk of skin reactions associated with the use of Padcev[™], as well as awareness of the Padcev[™] Patient Card (PC), knowledge of the content of the PC, and reported behaviors to minimise the risk of skin reaction.

Study Design

Non-interventional study design

Cross-sectional

Population studied

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)

Estimated number of subjects

62

Study design details

Outcomes

- Knowledge levels of patients, or their caregivers, of the risk of skin reactions associated with the use of Padcev $^{\text{\tiny TM}}$,
- Knowledge of the symptoms and recommended care for skin reactions associated with the use of Padcev $^{\text{\tiny TM}}$
- Knowledge levels of patients, or their caregivers, of the symptoms and recommended care for skin reactions associated with the use of Padcev $^{\text{\tiny TM}}$
- Reported behaviors of patients, or their caregivers, to minimise the risk associated with the use of Padcev™.

Data analysis plan

A statistical analysis plan will be developed to describe all planned analyses in detail, along with shells for variable lists, tables and figures.

All analyses will be performed using SAS® V9.0 or later.

A final report summarizing the results of the survey will be developed.

The study population included in the data analysis will include patients/caregivers who completed the question in the survey associated with the primary endpoint.

A threshold of success for the primary endpoints is to have 80% or more patients/caregivers providing a correct response to this question. Descriptive data analyses will be conducted. Levels of awareness and knowledge, self-reported behaviors as well as patient characteristics will be calculated with 95% two-sided Confidence Intervals. Descriptive results will be reported for the overall population of respondents and by country and, if the data permit, for sub-groups such as patients versus caregivers or length of time receiving Padcev™ treatment.

Data management

Data sources

Other
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