OSPREY

ONCOSIL PANCREATIC CANCER POST-MARKETING CLINICAL **RE**GISTR**Y**

A post-market, multicentre, observational, prospective registry of data recorded from patients who are treated using the OncoSil[™] device



Purpose:

To collect and assess the performance and safety of the OncoSil[™] device when used within the approved indication of unresectable, locally advanced pancreatic cancer, in combination with gemcitabine-based chemotherapy, within a real-world observational registry.



Location/Participating Centres:

Centres across EU and the UK will participate in the OSPREY Patient Registry.



Duration:

- Recruitment is anticipated to take five years
- Data will be collected over a prescribed 12-month period from enrolment
- Patients will be followed to death or 24 months from date of the last enrolled patient implanted with OncoSil[™]
- Registry therefore expected to run for a seven-year period

Data Collection:

Registry will collect the following real-world observational data and information via medical record review:

- Safety and tolerability of OncoSil[™] within a routine clinical setting
- OncoSil[™] implantation performance
- Overall survival (OS)
- Target (implanted) tumour response (local and distant)
- Progression-free survival (PFS)
- Surgical resection rates and outcome

Registry Treatment:

OncoSil[™] device in combination with gemcitabine-based chemotherapy

Registry Patient Population

Every patient who is intended to undergo implantation of the OncoSil[™] device will be approached by the Treating Physician to consent to participate in the OSPREY Patient Registry.*

The OncoSil[™] device will be used within the approved indication of unresectable, locally advanced pancreatic cancer, in combination with gemcitabine-based chemotherapy.

*Patients who choose not to consent to participate in the OSPREY Patient Registry will not be excluded from being treated with the OncoSil[™] device.

INCLUSION CRITERIA

- Patients eligible for and undergo OncoSil™ implantation at an eligible treatment facility according to the approved Instructions for Use, as part of their clinical care
- Patients who have completed and signed the Patient Informed Consent Form (PICF) for the OSPREY Patient Registry

EXCLUSION CRITERIA

- Patients participating in an interventional clinical study (company or investigator-sponsored)
- Patients using an investigational agent at the time of enrolment



Patient Number:

500 enrolled patients implanted with OncoSil™

Registry Sponsor: OncoSil Medical Ltd. Patient selection will be in accordance with the OncoSil[™] System Instructions for Use.

This is a summary provided for information purposes. Please refer to the OncoSil[™] System Instructions For Use, the OSPREY Registry Protocol (OSPREYO1) and ClinicalTrials.gov (identifier: NCTO4493632) for additional information.

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INTENDED USE/INDICATIONS FOR USE: OncoSil[™] is intended for intratumoural implantation into a pancreatic tumour via injection under endoscopic ultrasound guidance. OncoSil[™] is indicated for the treatment of patients with locally advanced unresectable pancreatic cancer, in combination with gemcitabine-based chemotherapy. The OncoSil[®] System is supplied sterile and is intended for single-patient, single-use.

This information is intended for healthcare professionals only. All medical treatments carry benefits and risks. For safety related information, please refer to the OncoSil[®] System Instructions for Use. OncoSil[®] is a registered trademark of OncoSil Medical Ltd. Level 5, 7 Eden Park Drive, Macquarie Park, NSW 2113 Australia. PRO3-MR-0822 Ver 1.

