



31 March 2020

Sydney, Australia

Noxopharm Virtual Roadshow Corporate Presentation

Sydney, 31 March 2020: Noxopharm (ASX: NOX) is pleased to provide shareholders and the market the attached Noxopharm "Corporate Presentation".

This document is being used by Noxopharm for presentation during a non-deal virtual roadshow being held by the company from 31st March to 2nd April 2020.

The presentation can be found at www.noxopharm.com

About Noxopharm

Noxopharm is a clinical-stage Australian drug development company with offices in Sydney and New York. The Company has a primary focus on the development of Veyonda® and is the major shareholder in Nyrada Inc. (ASX:NYR).

www.noxopharm.com

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Graham Kelly, CEO and Chairman of Noxopharm has approved the release of this document to the market.

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March/April 2020



Noxopharm Limited

Veyonda[®]

Non-Deal Roadshow Presentation




NOXOPHARM

ASX: NOX



DISCOVER



DEVELOP



DELIVER

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Contents

Introduction

Investment Highlights

Clinical Strategy

Commercial Strategy

Market Opportunities



Veyonda[®]

Introduction

Technology platform

- ❑ isoflavonoid molecular structure offering new generation of therapeutics built around polypharma actions (tyrosine and serine-threonine kinase inhibition) and restriction to prion-like (abnormal functioning) kinase targets

Proprietary IP

- ❑ structure/functional relationship; delivery technology

Veyonda®

- ❑ dual cytotoxic/immuno-oncology drug for late-stage prostate cancer

Drug pipeline

- ❑ oral cytotoxic for pancreatic/gall-bladder cancers
- ❑ first-in-class glutamate G-protein receptor inhibitor for GBM

We are a **drug discovery and drug development company**. We will seek strategic partnerships for pipeline drugs when they attain **key valuation points**

Investment Case

Veyonda® being positioned for the largest sector in the oncology market

- **end-stage cancer where treatment limited to palliative care**
- **little competition**
- **multi-billion \$ market opportunity**

Veyonda® immediate goal is late-stage prostate cancer

- **estimated 300,000 p.a. deaths globally**
- **estimated 33,000 in the U.S. in 2020**
- **U.S. market alone estimated at US\$1 billion +**

Veyonda® considerably de-risked

- **safety confirmed**
- **evidence of meaningful clinical efficacy in Phase I/II trials**
- **multiple programs (DARRT, LuPIN, CEP, IONIC)**

Commercial outreach commenced

- **GenesisCare relationship**
- **territorial carve-outs being explored**

Lean operation. Virtual company

** American Cancer Society Cancer Statistics Centre 2020*

Clinical Strategy for Prostate Cancer



➤ Main objective:

To provide a clinical data package that is attractive for future commercial partners

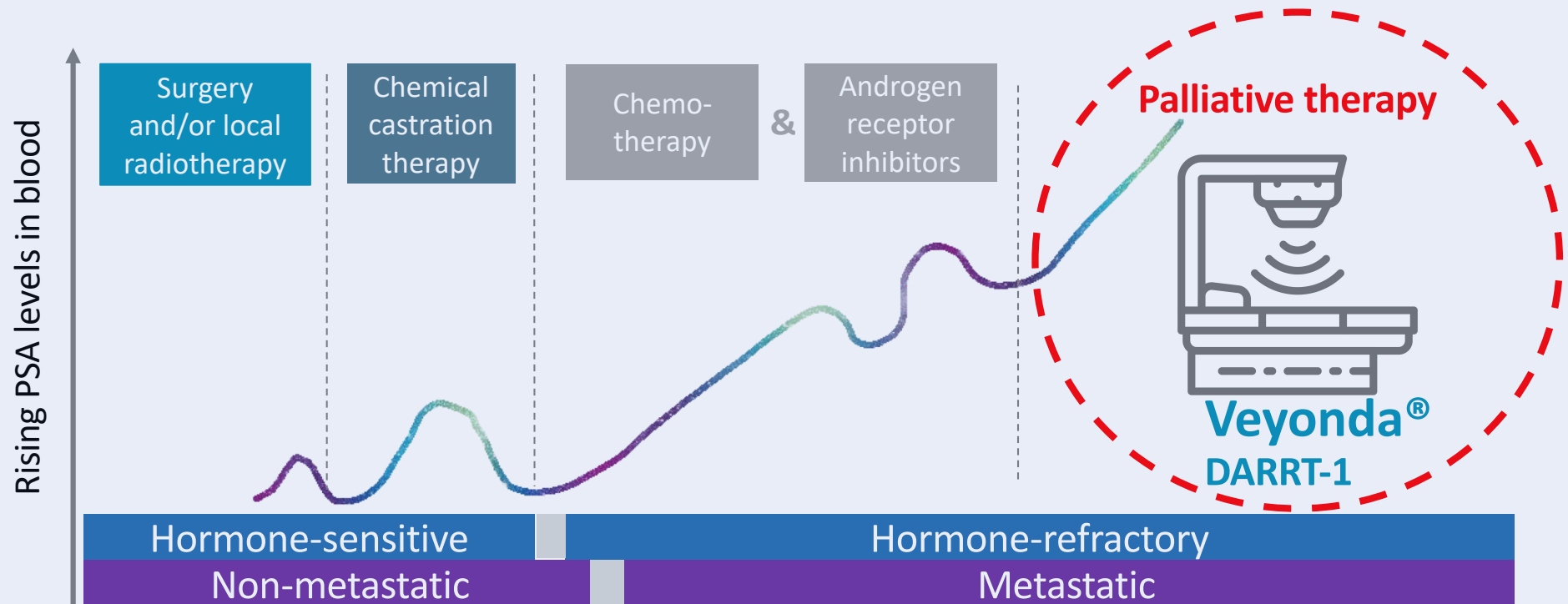
➤ Implementation Steps:

- First Prostate Cancer clinical trial (DARRT-1) has finished and was successful
- Second Prostate Cancer clinical trial (LuPIN-1) is ongoing and showing encouraging interim results
- Next clinical trial (DARRT-2) will build on these results and will include more patients
- News flow and a growing data portfolio will be ensured

Which Market Segment does Veyonda occupy?



Course of Disease and Treatment Journey for Stage I - IV Prostate Cancer

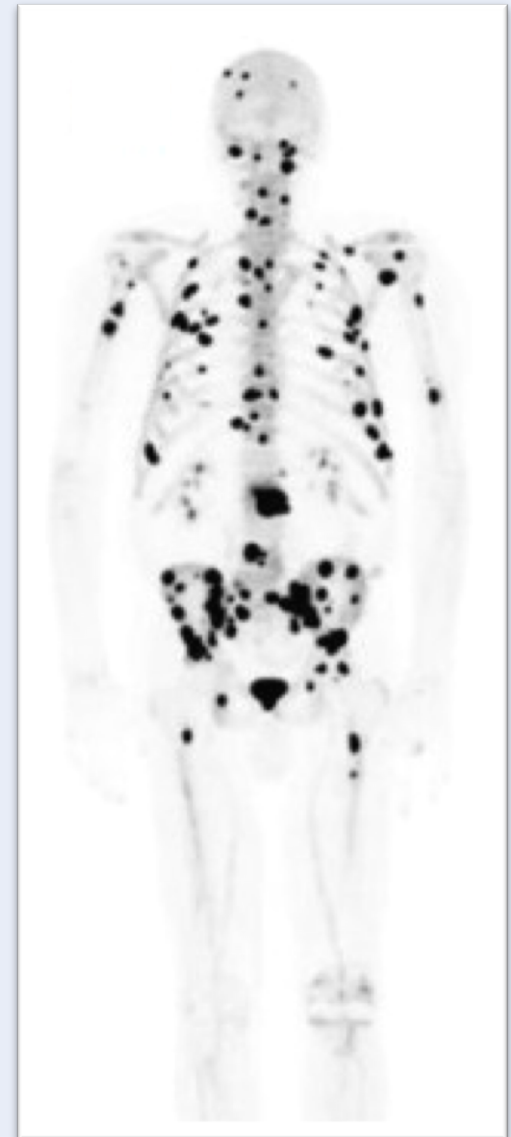


Veyonda[®] – What does the Clinical Data show?



- 26 men enrolled with late-stage **prostate cancer**
- Metastatic castration-resistant prostate cancer (mCRPC)
- Progressive disease
- No remaining standard treatment options
- Eligible for palliative RT for symptomatic relief

- Treatment with low-dose RT (20Gy in 5 fractions) and 14 days of NOX66 (400, 800, 1200 mg)



Bone scan with metastatic disease

DARRT = Direct and Abscopal Response to Radiation Therapy; RT = Radiation Therapy



DARRT-1: Does Veyonda work?

- In the 15* patients who were evaluable at 6 months¹

The Tumours stopped growing or reduced in size in 10 patients

(1 patient achieved a partial response and 9 achieved stable disease at 6 months)

6-months follow up	First part 400mg, 800mg & 1200mg (Reported on 12 November 2019)	Expansion part 1200mg	Overall All doses (Reported on 2 December 2019)
Overall (RECIST1.1)	N=10	N=5	N=15
Complete response	0	0	0
Partial response	1 (10%)	0	1 (7%)
Stable disease	7 (70%)	2 (40%)	9 (60%)
Progressive disease	2 (20%)	3 (60%)	5 (33%)

* 10 patients lost to follow-up, were not measurable, withdrew from study or died (unrelated to treatment)

1. Noxopharm. Data on file.

What is the Aim of the next Trial?



➤ Objectives:

- The DARRT-2 trial is designed to provide the data that Commercial Partners are looking for
- It also aims to satisfy the Regulatory Authorities
- Building on the experience and data of DARRT-1
- Phase 2 trial; multinational
- Medical Advisory Boards established
- Anticipated regulatory submissions late-2020
- Study expected to commence in early-2021

✓ **We are developing the most efficient and impactful study possible**

Additional Opportunity in Prostate Cancer



External Radiation

- Standard-of-Care
- Widely used



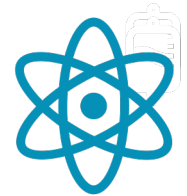
DARRT 

Internal Radiation

- Experimental
- Billion-dollar Acquisition by Novartis



LuPIN 



LuPIN Trial: Key Interim Results

❖ Veyonda[®] + ¹⁷⁷Lu-PSMA-617

- Overall Survival (OS) is a measure of the time from the start of treatment until death.



- ✓ Median OS was **17.1 months** – a remarkable result at this late stage of the disease
- ✓ The combination therapy was well tolerated, pointing to Veyonda[®] being safe to use in combination with intravenous radiotherapy

In summary, combination therapy of Veyonda[®] and ¹⁷⁷Lu-PSMA-617 showed major benefits to patients and underscores the Company's confidence in Veyonda[®] eventually becoming a standard drug in the management of prostate cancer

Commercial Strategy

Noxopharm commercial priorities

1. Attract potential partners by continuing to undertake clinical trials to develop Veyonda[®] as standard of care in treatment of prostate cancer
2. Develop alliances to strengthen the commerciality of Veyonda[®]
3. Investigate commercial agreements to build revenue prior to commercialisation of Veyonda[®]



DARRT-2 Phase 2 clinical trial planning underway



GenesisCare use of Veyonda[®] for compassionate treatment



Pursuing regional licencing agreements for Veyonda[®]

Market Opportunity

Prostate cancer market opportunity

- 33,330 prostate cancer deaths are forecast in the US in 2020*
- Potentially all of these patients could benefit from treatment with Veyonda®
- The potential global demand of late stage cancer patients for multiple cycles of Veyonda® indicates potential for multi-million dollar revenue from Veyonda®
 → positioning Veyonda® for acquisition by big pharmaceutical companies

Recent acquisitions in the prostate cancer space

Recent acquisitions	Buyer	Seller	Price range
XTANDI® mCRPC (2016)			US\$14 billion
¹⁷⁷ Lu-PSMA-617 mCRPC (2018)			US\$2.1 billion
¹⁷⁷ Lu-PSMA-617 & others mCRPC (2018)			US\$3.9 billion



* American Cancer Society Cancer Statistics Centre 2020



**For further information please visit
www.noxopharm.com**

Veyonda[®]

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