

Bell Potter Healthcare Conference
Positive interim data from phase 2 cancer study



developing breakthrough treatments for fibrosis and inflammation

Investor Presentation | November 2022
Gary Phillips CEO

Forward looking statement

This document contains forward-looking statements, including statements concerning Pharmaxis' future financial position, plans, and the potential of its products and product candidates, which are based on information and assumptions available to Pharmaxis as of the date of this document. Actual results, performance or achievements could be significantly different from those expressed in, or implied by, these forward-looking statements. All statements, other than statements of historical facts, are forward-looking statements.

These forward-looking statements are not guarantees or predictions of future results, levels of performance, and involve known and unknown risks, uncertainties and other factors, many of which are beyond our control, and which may cause actual results to differ materially from those expressed in the statements contained in this document. For example, despite our efforts there is no certainty that we will be successful in developing or partnering any of the products in our pipeline on commercially acceptable terms, in a timely fashion or at all. Except as required by law we undertake no obligation to update these forward-looking statements as a result of new information, future events or otherwise.

Shareholders & cash



Financial Information	8 Nov 22
ASX Code	PXS
Share price	\$0.066
Liquidity (turnover last 12 months)	92m shares
Market Cap	A\$42m
Pro forma ¹ cash balance (30 September 2022)	A\$26m
Enterprise value	A\$16m
Clinical development program supported by:	
• R&D tax credits • Strategy of partnering deals with pipeline assets	
1. Proforma cash includes cash of \$12m, estimated 2022 R&D tax credit of \$5m (expected receipt H2 CY22), and capital raising of \$10m less offer costs	

Institutional Ownership	27 October 22
BVF Partners LP	17%
Karst Peak Capital Limited	11%
D&A Income Limited	7%
Regal Funds Management Pty Ltd	6%
Platinum Investment Management Limited	5%
Total Institutional Ownership	~46%

Share Price



Pharmaxis Overview

Executive Summary

- Pharmaxis is a clinical stage drug development company targeting inflammation, fibrosis and selected cancer indications with first in class or best in class small molecule drugs in markets of high value
- Global leader in fibrosis driven by lysyl oxidase enzymes having invested in a multi year research program leveraged with extensive external scientific collaborations
- 5 studies recruiting in 2022 and 2023 that will provide near term value opportunities
- Pro forma cash position at 30 September 2022 of A\$26m¹, funding the company's clinical programs into early 2024

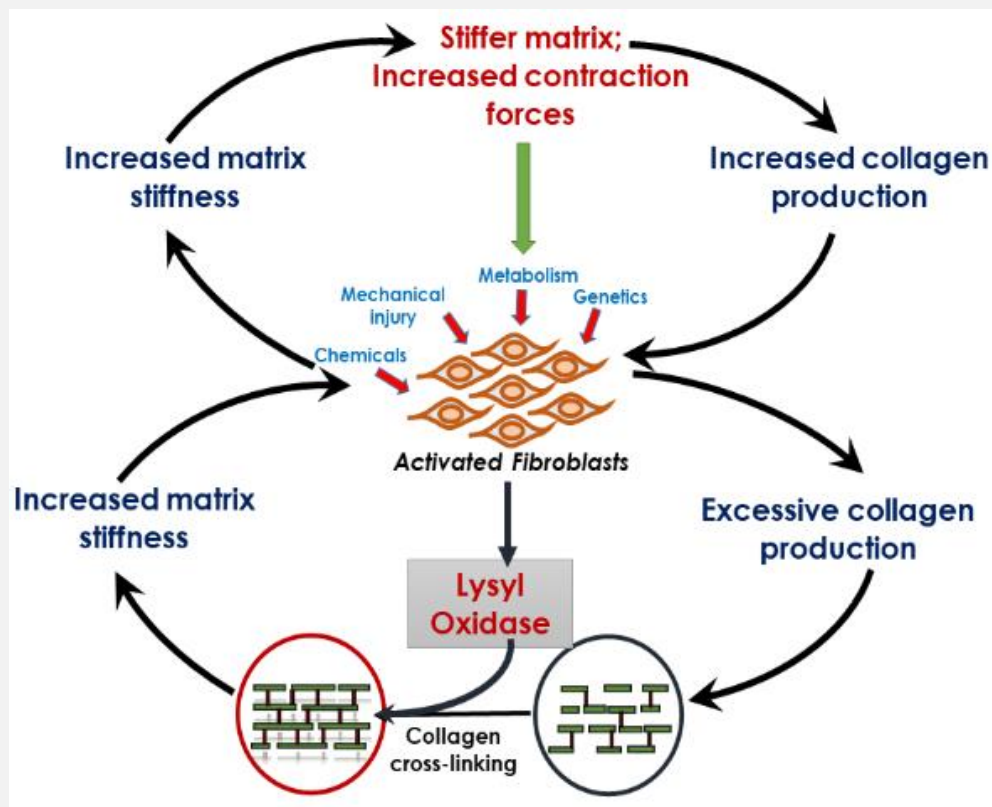
Pipeline creates multiple opportunities in high value markets

1. Lead asset PXS-5505 is in a multinational phase 2 trial – a breakthrough clinical program with disease modifying potential in Myelofibrosis. 15 out of 24 targeted patients recruited
2. US investigator led phase 1/2 trial in liver cancer with PXS-5505 as first line treatment added to existing chemotherapy to commence Q4 2022
3. Topical drug PXS-6302 trial in patients with potential to improve function and appearance of established scars. >80% recruited
4. Additional PXS-6302 trial in scar prevention to commence recruitment in 1H 2023
5. Neuro inflammation drug PXS-4728 in phase 2 trial of patients with severe sleep disorder that can lead to neurodegenerative diseases e.g. Parkinson's

Pharmaxis is the global leader in lysyl oxidase chemistry and biology

Multi year research program leveraged with extensive scientific collaborations worldwide has delivered 2 drugs in the clinic

Lysyl oxidases are the final stage in fibrosis



Tissue stiffening due to increases in collagen and number of cross-links which is a hallmark of fibrosis, is preventable through lysyl oxidase inhibition; at the heart of a true anti-fibrotic therapy

■ PXS-5505

- Oral dosage form – four capsules twice a day
- Patent filed - priority date 2018
- Strong pre clinical evidence in models of fibrosis and cancer
- INDs approved for myelofibrosis and hepatocellular carcinoma
- Potential in multiple cancer indications
- Phase 1 data demonstrates a safe, well tolerated drug that gives >90% inhibition of LOX enzymes

■ PXS-6302

- Topical dosage form
- Patent filed - priority date 2019
- Strong pre clinical evidence in models of skin fibrosis and scarring
- Potential in prevention of scar formation and modification of existing scars
- Phase 1a (healthy volunteer) data demonstrates a safe, well tolerated drug that gives full inhibition of LOX enzymes in the skin with minimal systemic exposure

Program Update



Myelofibrosis

A rare type of bone marrow cancer that disrupts the body's normal production of blood cells

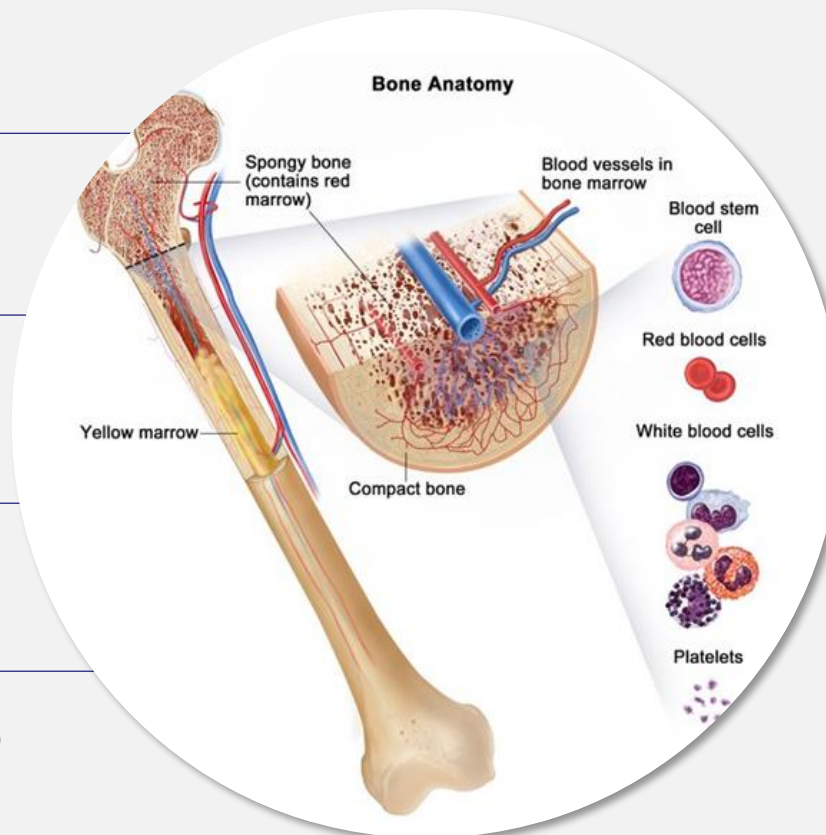
KEY FACTS

- Affects ~15 in 1m people worldwide

- 5 Years Median survival

- Age of onset typically from age 50

- 11% transformation to leukemia



Primary Myelofibrosis is caused by a build up of scar tissue (fibrosis) in bone marrow reducing the production of blood cells:

- Reduced red blood cells can cause extreme tiredness (fatigue) or shortness of breath
- Reduced white blood cells can lead to an increased number of infections
- Reduced platelets can promote bleeding and/or bruising
- Spleen increases blood cell production and becomes enlarged
- Other common symptoms include fever, night sweats, and bone pain

Current Standard of Care; JAK inhibition

- Symptomatic relief plus some limited survival improvement. 75% discontinuation at 5 years
- Median overall survival is 14 – 16 months after discontinuation

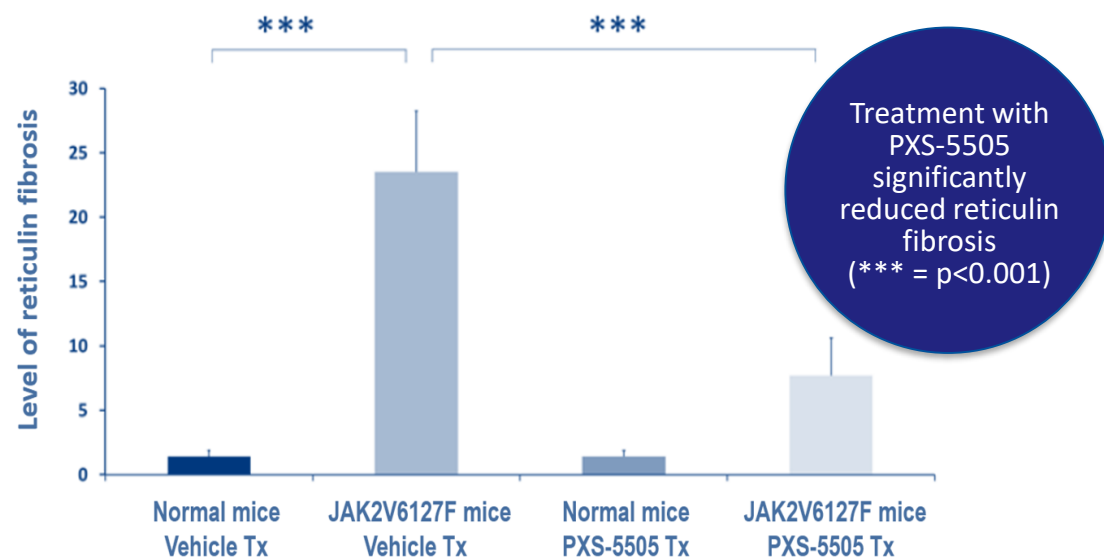
Commercial Opportunity

- Current standard of care ; revenue ~US\$1b per annum

Myelofibrosis - PXS-5505; an effective and safe inhibitor of LOX in myelofibrosis patients

Pre clinical and clinical studies strongly support entry into long term phase 2 patient studies

PXS-5505 attenuates hallmarks of primary myelofibrosis in mice³

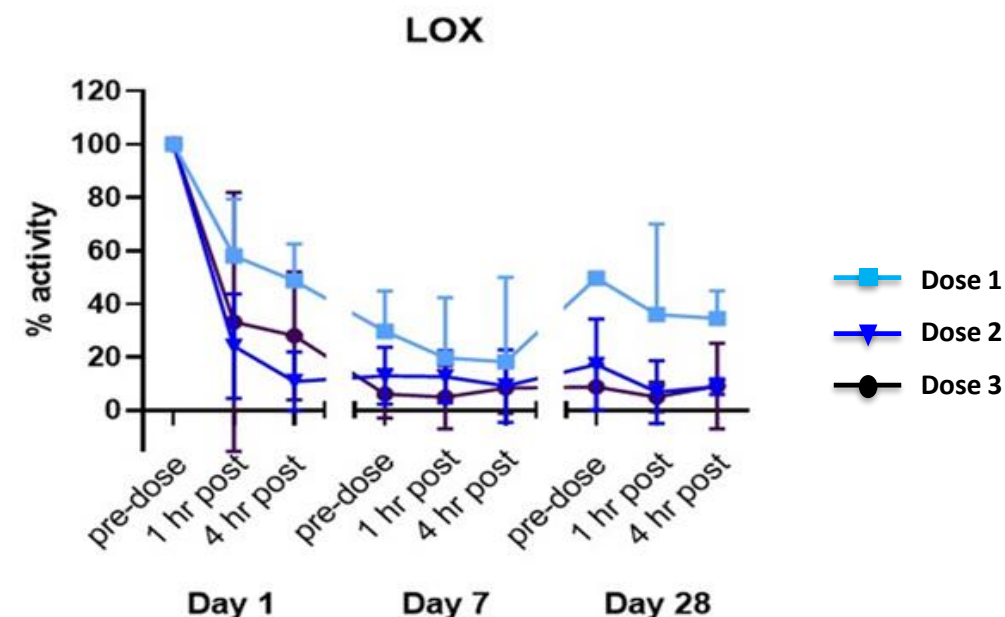


"None of the drugs approved to date consistently or meaningfully alter the fibrosis that defines this disease. PXS-5505 has a novel mechanism of action by fully inhibiting all LOX enzymes.

Preliminary data thus far, demonstrate that PXS-5505 leads to a dramatic, >90% inhibition of LOX and LOXL2 at one week and 28 days. This confirms what's been shown in healthy controls as well as mouse models, that this drug can inhibit the LOX enzymes in patients. Inhibiting these enzymes is a novel approach to the treatment of myelofibrosis by preventing the deposition of fibrosis and ultimately reversing the fibrosis that characterizes this disease"

Dr Gabriela Hobbs¹

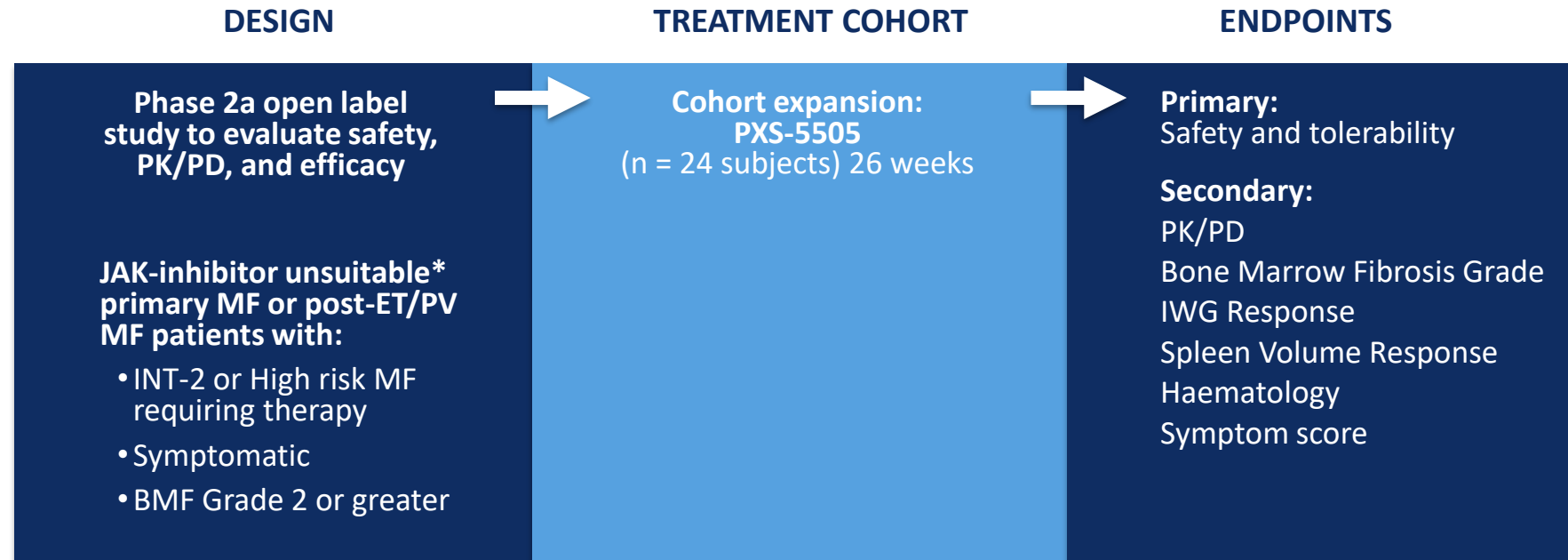
PXS-5505 – Phase 1c dose escalation in MF patients



- Open label dose escalation in JAK-inhibitor unsuitable² primary MF or post-ET/PV MF patients
- Maximum of 3 patients on each dose for 28 days
- Good safety profile with no adverse events at highest dose
- >90% inhibition of LOX and LOXL2 at trough on highest dose at day 7 and 28

Myelofibrosis – PXS-5505 Phase 1/2a Trial

6 month monotherapy study with meaningful safety and efficacy endpoints



FDA granted orphan drug designation July 2020 and IND approved August 2020

20 sites across 4 countries
(Australia, South Korea, Taiwan, USA)

Study budget to spend
~A\$6.2m

Study recruitment commenced Q4 2021, study targeted to report mid 2023

Myelofibrosis – PXS-5505 Phase 2a Trial (*INTERIM DATA*)

Very well tolerated with encouraging signs of clinical efficacy in JAK inhibitor unsuitable patients

DESIGN	TREATMENT COHORT	ENDPOINTS
Phase 2a open label study to evaluate safety, PK/PD, and efficacy JAK-inhibitor unsuitable* primary MF or post-ET/PV MF patients with: • INT-2 or High risk MF requiring therapy • Symptomatic • BMF Grade 2 or greater • Median survival after JAK-inhibitor discontinuation; approximately 1 year	Cohort expansion: PXS-5505 (n = 24 subjects) 26 weeks • A total of 15 patients have been enrolled • 6 patients having completed 24 weeks of treatment. • 4 patients have dropped out of the study due to a lack of clinical response.	Primary: PXS5505 has been well tolerated with no serious treatment related adverse events reported. Secondary: • 2/6 patients show clinically important improvement in symptoms. • 5/6 patients show either stable or improved bone marrow fibrosis scores of ≥ 1 grade. • 5/6 have stable or improved platelet and/or haemoglobin scores • No reductions were seen in spleen volume



“PXS-5505 continues to be very well tolerated in the clinic with no serious treatment related adverse events reported.

Though still early in the dose expansion phase of the study, PXS5505 appears to be stabilising and in some cases, improving the hemoglobin and platelet counts, which has also been associated with symptom improvements in those patients that were treated to 24 weeks.

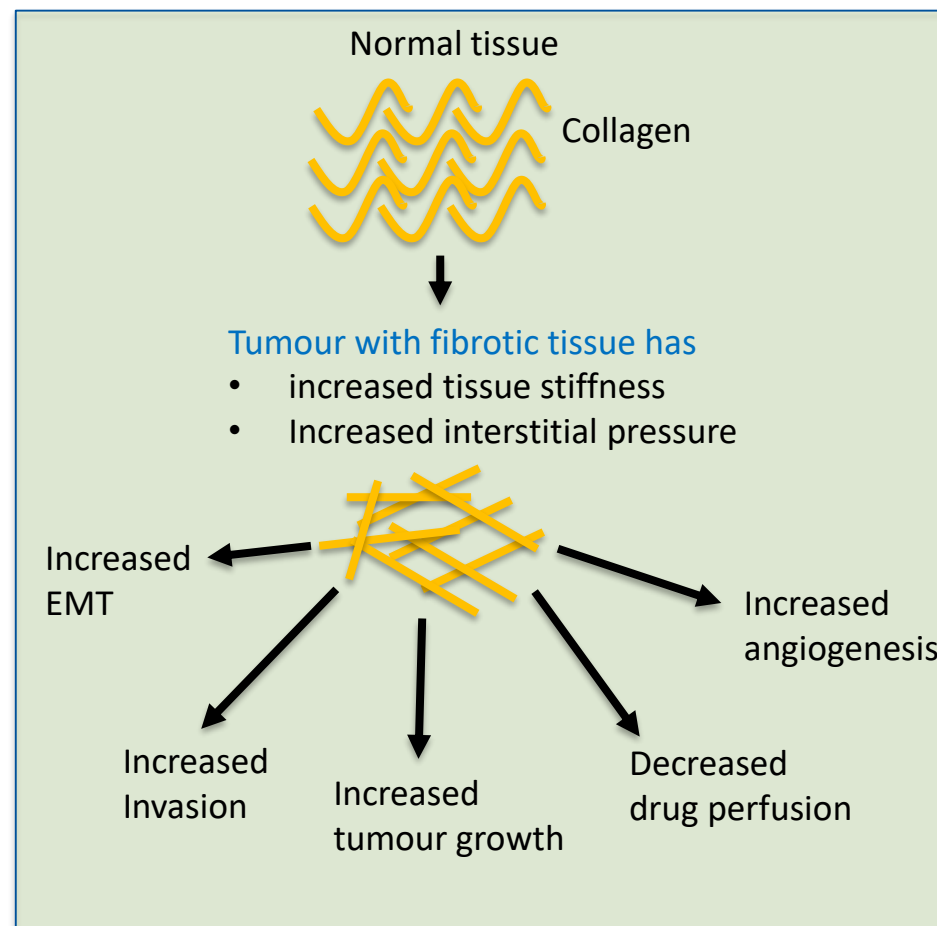
This is encouraging given the poor prognosis seen after ruxolitinib discontinuation with a median overall survival of only 11-14 months typical of this study population. These results support further clinical investigation of PXS5505 in myelofibrosis.”

Dr Gabriela Hobbs MD,
Assistant Professor, Medicine, Harvard
Medical School & Clinical Director,
Leukemia Service, Massachusetts General
Hospital

Hepatocellular Carcinoma (HCC)

4th leading cause of cancer-related mortality worldwide with a 19.6% 5-year relative survival

- Primary liver malignancies have doubled in incidence over the last two decades.
- 4th leading cause of cancer-related mortality worldwide with a 19.6% 5-year relative survival
- HCC is a stromal (fibrotic) tumour
 - Accumulation of collagen cross-links increases stromal stiffening and interstitial fluid pressure reducing delivery of chemotherapy and immunotherapy
- Current standard of care
 - 20-30% are resectable at presentation with many patients relying on systemic therapy:
 - Tyrosine kinase inhibitors
 - PD-L1 inhibitors + anti-VEGF



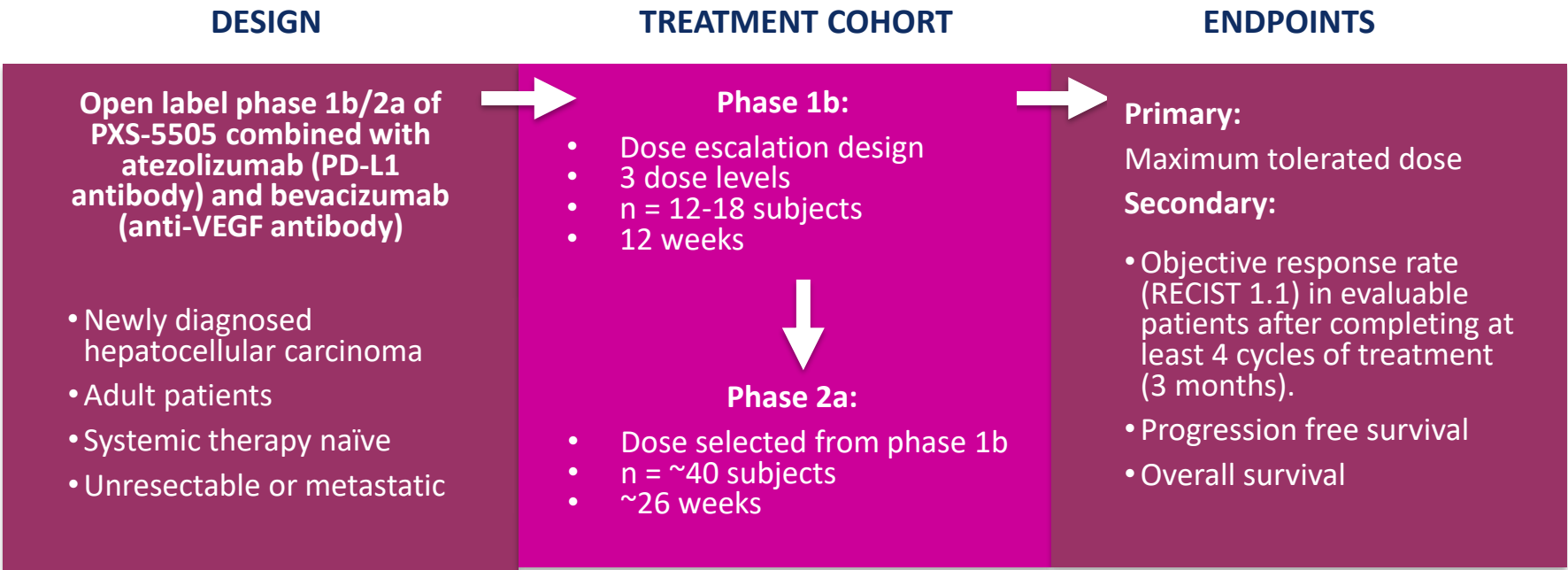
- Pre-clinical data (Rochester Uni; Aug 2021)
 - Tumour tissue specimens show LOX enzymes are significantly elevated in human liver cancer and correlate with poor prognosis.
 - PXS-5505 with or without chemotherapy treatment in a pre-clinical model significantly **improves survival, delays tumour growth, and reduces intratumoral pressure.**

Commercial Opportunity

Drugs market currently worth ~US\$2bn with rising incidence forecasted to drive growth to ~US\$7bn by 2027

Hepatocellular carcinoma – PXS-5505 Phase 1b/2a Trial

First line combination treatment with standard of care to assess safety and efficacy endpoints



IND (University of Rochester) reviewed by FDA November 2021

<https://clinicaltrials.gov/ct2/show/NCT05109052>

Phase 1b trial being conducted by Rochester Medical Centre, New York State, USA (Study sponsor).
Opened for enrollment 23 Sept

Phase 1b study budget to spend ~A\$1.7m

Study recruitment to commence Q4 2022, study targeted to conclude H1 2024

Hypertrophic and keloid scarring

Cutaneous scarring following skin trauma or a wound is a major cause of morbidity and disfigurement

KEY FACTS

100m patients develop scars in the developed world alone each year as a result of elective operations and operations after trauma

Hypertrophic scars and keloids are fibroproliferative disorders that may arise after any deep cutaneous injury caused by trauma, burns, surgery, etc.

Hypertrophic scars and keloids are cosmetically and functionally problematic significantly affecting patients' quality of life



"In (preclinical) models of scarring we found that topical application of PXS-6302 reduces collagen deposition and cross-linking and improves scar appearance without reducing tissue strength. This is a unique way of modulating a critical stage in scar formation and maintenance and holds out great promise for the treatment of scars."
- Dr Mark Fear,
UWA

■ Mechanisms underlying scar formation are not well established; prophylactic and treatment strategies remain unsatisfactory

■ Current standard of care includes:

- Corticosteroids
- Surgical revision
- Cryotherapy
- Laser therapy
- 5-fluorouracil



■ Pre clinical evidence

- Treatment with PXS-6302 monotherapy demonstrates cosmetic and functional improvements to scarring in pre clinical models¹

■ Clinical evidence

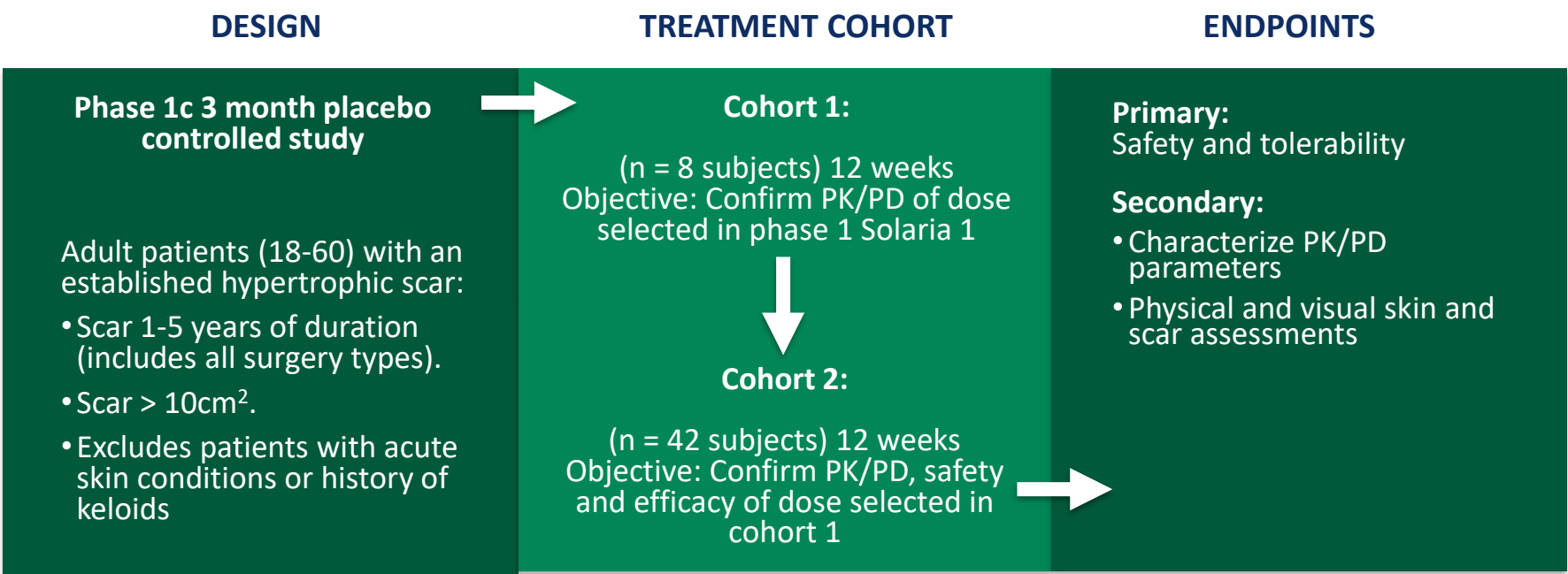
- 1 month phase 1a in healthy volunteers demonstrates good tolerability and full inhibition of LOX in skin.

■ Commercial Opportunity

- Total scar treatment market in 2019 exceeded US\$19b. Keloid and hypertrophic scar segment ~US\$3.5b

Established Hypertrophic Scarring – PXS-6302 Phase 1c Trial (Solaria 2)

3 month monotherapy study to assess dosage, tolerability and efficacy endpoints



Investigator initiated study (sponsor UWA) - long term collaboration with UWA to research and develop PXS-6302 supported by Australian NHMRC grants

Single site study in Perth Australia

Study budget to spend; A\$0.3m

Study recruitment commenced Q1 2022, study targeted to report H1 2023

Established Hypertrophic Scarring – PXS-6302 Phase 1c Trial (Solaria 2)
3 month monotherapy study to assess dosage, tolerability and efficacy endpoints

DESIGN	TREATMENT COHORT	ENDPOINTS
Phase 1c 3-month placebo controlled study Adult patients (18-60) with an established hypertrophic scar: • Scar 1-5 years of duration (includes all surgery types). • Scar > 10cm². • Excludes patients with acute skin conditions or history of keloids	Cohort 1: Fully recruited Cohort 2: • A total of 24 out of 42 patients have been enrolled • Dosage regimen modified to reduce drug exposure but still maintain the overall high level of enzyme inhibition.	Cohort 1: • Skin biopsies show skin penetration and high inhibition of LOX • Reduction in biomarkers of the scarring process suggests a disease modifying effect. • Clinician notes positive changes in appearance and pliability • Four patients withdrew after experiencing redness & itchiness at the site of application that resolved on treatment cessation and informed the decision to reduce dosage frequency for Cohort 2

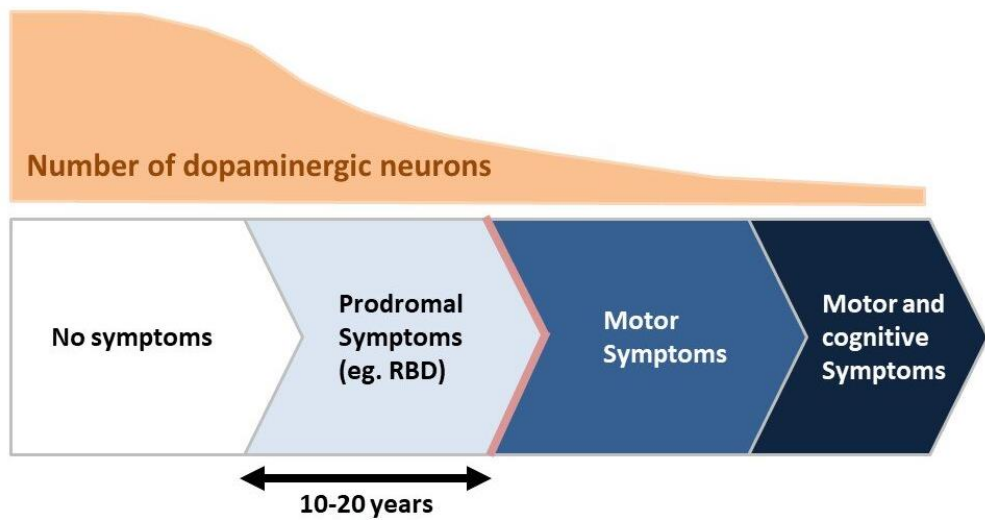
“We have noted positive changes in appearance and pliability of scars in those patients on active drug that now need to be confirmed by the results from the placebo controlled phase of this trial later this year.

We are learning a lot as we move from the promising pre-clinical work done at UWA and into the clinic where we have many patients who are in great need of a treatment that can improve both the cosmetic appearance of their scars and improve the functionality of their scarred skin; factors that have a huge impact on patient’s wellbeing.”

Professor Fiona Wood
Burns Service of Western Australia
Director of the Burn Injury Research Unit
University of Western Australia

iRBD & Neuro Inflammation - Using a sleep disorder to target Parkinson's Disease

SSAO inhibition proven effective mechanism against neuro inflammation and is neuro protective in pre clinical models

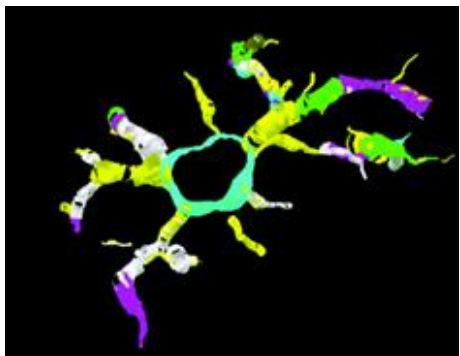


Parkinson's Disease and RBD

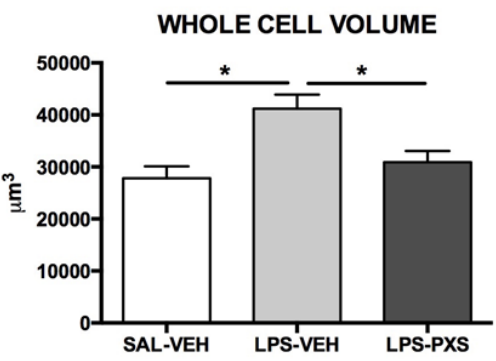
- More than 50% of dopaminergic neurons in the substantia nigra are lost at the onset of motor symptoms in Parkinson's Disease.
- Prodromal symptoms, such as isolating REM sleep behavior disorder (iRBD), proceed the onset of motor cognitive dysfunction by 10-20 years.
- 70% of iRBD patients transition to a neurodegenerative disease such as Parkinson's disease and Dementia with Lewy Bodies

PXS-4728 and neuro inflammation

- PXS-4728 has already undergone extensive development by Boehringer Ingelheim
- PXS-4728 inhibits SSAO and MAOB in the brain – both of which play a role in neurodegenerative diseases such as Parkinson's.
- Dual SSAO & MAO-B inhibition protects against neuronal degradation in pre clinical models²
- MAO-B inhibition alone (selegiline) does not offer any protection in the same model²



Activated microglia - reconstruction



Change in Microglia whole cell volume in the Substantia Nigra (SN) after LPS¹

iRBD and Neuro Inflammation – Parkinson's UK Funding

PXS-4728 to proceed to phase 2 trial

Short and longer term commercial opportunities

- Current standard of care for iRBD is melatonin. There remains a high unmet need.
- >8% of 70 – 89 year olds have iRBD
- >70 % of iRBD patients develop Parkinson's disease and the related α -synuclein deposition disorders, dementia with Lewy bodies (DLB) and multiple system atrophy (MSA).
- Parkinson's market ~3.5bn in 2019

Clinical Trial

- 40 patients
- Randomised, double blind, placebo controlled clinical trial with iRBD
- 12 weeks of treatment with oral PXS-4728
- Two sites - University of Sydney and the University of Oxford
- Expected to commence dosing in H1 2023
- Efficacy endpoints for iRBD and neuroinflammation

Parkinson's UK Funding Agreement

Clinical trial in precursor to Parkinson's Disease

- The funding agreement with Parkinson's UK entails up to £2.9m (~A\$4.9m) to be paid to Pharmaxis to run the phase 2 trial with advance payments received as the trial progresses.
- Pharmaxis is providing the study drug and the compound that will be used to measure inflammation in the brain scan of trial participants. The total is expected to cost approximately A\$5.8 million.
- Parkinson's UK will receive a return of up to 4 times their funding from royalties on future revenue Pharmaxis receives from commercialising PXS-4728 in neurological diseases and up to 2 times in other indications.

Parkinson's
Virtual Biotech

Upcoming News Flow

Five trials to deliver near term value

Pipeline creates multiple opportunities in high value markets

	Indication	Addressable market (US\$)	Trial design	# patients	Status	Data
PXS-5505	Myelofibrosis (MF)	\$1 billion	Phase 2 open label 6 month study in JAK intolerant / ineligible myelofibrosis patients	24	Recruiting	Interim data released Full data mid 2023
	Hepatocellular Carcinoma (HCC)	\$7 billion	Phase 1c open label dose escalation study in newly diagnosed patients with unresectable HCC on top of standard of care (PD-L1 inhibitor + anti VEGF)	18	First Patient Q4 2022	H1 2024
PXS-6302	Modification of established scars	\$3.5 billion	Phase 1c 3 month placebo controlled study in patients with established scars (>1 year old)	50	Recruiting	H1 2023
	Scar prevention post surgery	\$3.5 billion	Phase 1c 3 month placebo controlled study in patients with scarring subsequent to a burns injury	50	First patient H1 2023	H1 2024
PXS-4728	Isolated REM sleep behaviours disorder (iRDB) and neuro inflammation	\$3.5 billion	Phase 2 double blind, placebo controlled study in patients with iRBD	40	First patient H1 2023	H1 2025

News flow

Q4 2022 and H1 2023 anticipated news flow

Strong and growing pipeline with advancement in studies expected to provide value inflection points in FY23



Q4 2022

- PXS-5505 phase 2a myelofibrosis study – interim data
- PXS-5505 phase 1c liver cancer (HCC) study – starts recruitment
- PXS-5505 phase 2a myelofibrosis study – fully recruited
- PXS-5505 publications by KOL's in other cancers
- Two presentations at ASH (American Society of Haematology) conference in November
 - PXS-5505 phase 1c/2 study in myelofibrosis
 - PXS-5505 pre-clinical data in Myeloid Neoplasms (e.g. myelodysplastic syndrome)



Q1 2023

- LOX topical drug PXS-6302 commences independent investigator patient studies – scar prevention
- LOX topical drug PXS-6302 top line data from established scars study
- PXS-5505 publications by KOL's in other cancers



Q2 2023

- PXS-5505 phase 2a myelofibrosis study completed and reports safety and efficacy data
- PXS-4728 iRBD / neuro inflammation study commences recruitment



Appendix

Board

Significant international pharmaceutical experience



Malcolm McComas – Chair

- Former investment banker and commercial lawyer
- Former MD Citi Group
- Has worked with many high growth companies across various industry sectors and has experience in equity and debt finance, acquisitions and divestments and privatisations
- Joined Pharmaxis Board in 2003
- Chair since 2012



Gary Phillips – Chief Executive Officer

- 30+ years' of operational management experience in the pharmaceutical and healthcare industry in Europe, Asia and Australia
- Joined Pharmaxis in 2003 and was appointed Chief Executive Officer in March 2013 at which time he was Chief Operating Officer
- Previously held country and regional management roles at Novartis – Hungary, Asia Pacific and Australia



Dr Kathleen Metters – Non-Executive Director

- Former Senior Vice President and Head of Worldwide Basic Research for Merck & Co. with oversight of all the company's global research projects
- In a subsequent role at Merck & Co she led work on External Discovery and Preclinical Sciences
- Former CEO of biopharmaceutical company Lycera Corp



Dr Neil Graham – Non-Executive Director

- Former VP of immunology and inflammation responsible for strategic program direction overseeing pipeline development and clinical programs at Regeneron (REGN:US)
- Former SVP program and portfolio management at Vertex Pharmaceuticals
- Former Chief Medical Officer at Trimeris Inc and Tibotec Pharmaceuticals

Experienced senior management team

Significant global experience in drug development, commercialisation and partnering



Gary Phillips – CEO and Managing Director

- 30+ years' of operational management experience in the pharmaceutical and healthcare industry in Europe, Asia and Australia
- joined Pharmaxis in 2003 and was appointed Chief Executive Officer in March 2013 at which time he was Chief Operating Officer
- Previously held country and regional management roles at Novartis – Hungary, Asia Pacific and Australia



Jana Baskar – Chief Medical Officer

- 20+ years' experience both in clinical medicine and the biopharmaceutical industry
- Broad therapeutic knowledge and significant clinical research expertise having worked in several different specialties
- Former Medical Director at Novartis Oncology in Australia; former Medical Director for IQVIA in Australia and New Zealand



Wolfgang Jarolimek – Drug Discovery

- 20+ years' experience in pharmaceutical drug discovery and published more than 30 peer reviewed articles
- Previously Director of Assay Development and Compound Profiling at the GlaxoSmithKline Centre of Excellence in Drug Discovery in Verona, Italy
- Spent 8 years as post-doc at the Max-Planck Institute in Munich, Germany; Baylor College of Medicine, Houston, Texas; Rammelkamp Centre, Cleveland Ohio; and University of Heidelberg, Germany



David McGarvey – CFO

- more than 30 years' experience building Australian based companies from inception to globally successful enterprises
- joined Pharmaxis as Chief Financial Officer and Company Secretary in December 2002
- previously Chief Financial Officer of the Filtration and Separations Division of US Filter (1998-2002), and Memtec Limited (1985-1998)
- commenced career at PricewaterhouseCoopers



Kristen Morgan – Alliance Management

- more than 20 years' experience in the pharmaceutical industry having previously held a senior role in medical affairs at Sanofi-Aventis, and a commercial sales role at GlaxoSmithKline
- responsibility for alliance management and medical and regulatory affairs



Dieter Hamprecht – Head of Chemistry

- 20+ years' experience with small molecule and peptide drug discovery, contributed to greater than 10 drug candidates brought to development and co-inventor of 50 patent families, co-author of 30+ scientific publications
- Previously Managing Director – Boehringer Ingelheim's research group in Milan
- Senior medicinal chemistry positions at GSK

Mannitol respiratory business (Bronchitol® and Aridol®)

Sales growth expected from Bronchitol sales in US and Russia

Sales

- Bronchitol > 75% of sales
- Strong short term growth from Russia
- Sales growth expected in approved markets as patients access hospitals again post COVID-19 restrictions
- Strong longer term growth contribution expected from US

Expenses

- Relatively fixed production cost base

Segment EBITDA

- Negative EBITDA for FY 2022 \$1.3m
- Forecast positive EBITDA as CF clinics reopen post COVID
- US volumes contribute to mannitol segment generating profit



Bronchitol in US

- US CF market >65% of global market in value
 - US market doubles global cystic fibrosis patient opportunity with attractive pricing
- US sales commenced in Q1 CY 2021 – delay in patient initiation due to COVID
- High teens % of Chiesi sales + supply contract - ~20% of Chiesi US Bronchitol net sales flow directly to the Pharmaxis bottom line

Financials

Income statement highlights

Periods ended (A\$'000)	Three months		Twelve months	
	Sep-22	Sep-21	Jun-22	Jun-21
Segment Financials				
New drug development				
Oral LOX (external costs)	(1,009)	(1,467)	(5,431)	(2,521)
Other program external costs (net of grants)	(300)	(303)	(1,712)	(1,850)
Employee costs	(891)	(715)	(2,943)	(3,270)
Overhead	(161)	(102)	(374)	(396)
R&D tax credit and other income	-	-	5,600	148
EBITDA	(2,361)	(2,587)	(4,859)	(7,889)
Mannitol respiratory business				
Sales	760	3,272	7,427	6,680
Other revenue and income	7,192	2,342	2,342	15,986
	7,952	5,614	9,769	22,666
Expenses – employee costs	(1,119)	(1,197)	(4,760)	(5,558)
Expenses – manufacturing purchases	(648)	(1,205)	(2,729)	(1,168)
Expenses – other	(806)	(1,142)	(3,584)	(4,483)
EBITDA	5,379	2,070	(1,304)	11,457
Corporate – EBITDA	(333)	(678)	(4,080)	(3,793)
Total Adjusted EBITDA	2,685	(1,195)	(10,243)	(225)
Net profit (loss)	943	(3,029)	(1,934)	(2,970)

Financials

Cash

Periods ended (A\$'000)	Three months		Twelve months	
	Sep-22	Sep-21	Jun-22	Jun-21
Cash				
Cash at period end	11,597	18,712	11,597	18,712
Cash received/receivable post period end				
2022 R&D tax credit - expected H2 CF 2022	4,900			
Placement (\$5.1m subject to shareholder approval)	9,235			
Proforma cash at period end	25,732			

Cash Flow Statement Highlights

Operations

Receipts from customers	404	1,498	8,313	7,242
R&D tax incentive	-	-	-	5,048
Milestone payments	-	-	-	13,844
Sale of Orbital/distribution rights	7,192	2,340	2,902	1,365
Other	17	17	1,005	236
Payments to suppliers, employees etc (net)	(4,951)	(5,757)	(28,322)	(24,663)
Total operations	2,662	(1,902)	(16,102)	3,072
Investing (capex & patents)	(26)	(40)	(306)	(644)
Finance lease payments ¹	(545)	(596)	(2,379)	(2,305)
Financing agreement payments ²	(14)	(43)	(62)	(240)
Share issue - net	-	-	9,074	4,065
Net increase (decrease) in cash	2,660	(2,581)	(9,775)	3,948

1. Lease over 20 Rodborough Rd (to May 2024) – total liability at 30 June 2022: \$4.4 million
2. NovaQuest financing – not repayable other than as % of US Bronchitol revenue through to March 2028



developing breakthrough treatments for fibrosis and inflammation

Pharmaxis Ltd ABN 75 082 811 630

www.pharmaxis.com.au



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