

# **Quarterly Activities Report and 4C Quarterly Cash Flow Report**

# Highlights:

- GMP production run initiated during the quarter, and successfully completed in April with product characterisation and quality verification on track for completion in Q2 CY2024;
- Positive preclinical data highlights the potential to use deoxymabs to treat vasculitis, and potentially other autoimmune diseases, presented at international workshop;
- Cash and short-term investment balance of \$3 million on 31 March 2024.

**Melbourne, Australia; 30 April 2024:** Patrys Limited (ASX: PAB, "Patrys" or the "Company"), a therapeutic antibody development company, today released its Quarterly Activities Report and Appendix 4C Quarterly Cash Flow report for the quarter ended 31 March 2024.

**Patrys Chief Executive Officer and Managing Director, Dr. James Campbell said:** "The GMP manufacturing run for PAT-DX1 started during the quarter and subsequent to the end of the quarter has recently been successfully completed. The Company expects the additional product characterisation and specification testing will be completed by Q2 CY2024. We were also pleased to announce that after the end of the quarter positive preclinical data concerning deoxymabs in animal models of the autoimmune disease vasculitis was presented during the plenary session at a leading international workshop. This potentially opens up a new therapeutic area for Patrys to develop or partner its deoxymab technology."

## **Operations Update**

Patrys secured a manufacturing slot with its Contract Development and Manufacturing Organization (CDMO) for a GMP (Good Manufacturing Practice) production run that commenced in Q1 CY2024. As subsequent events to the quarter large-scale fermentation was successfully completed in mid-April 2024 and the resultant antibody product was successfully purified in late April 2024. The resulting material will undergo final product characterization and quality verification to ensure it meets product specifications during Q2 CY2024.

As mentioned in the previous quarterly report Patrys has established a Master Cell Bank (MCB) for PAT-DX3 and completed an integration run which combines the upstream fermentation with downstream purification processes. The availability of a GLP manufacturing process for PAT-DX3 is

# patrys

expected to facilitate ongoing partnering discussions for this deoxymab and the Company intends to initiate its own production run of PAT-DX3 at such time that it has access to additional capital.

As a subsequent event, on 9 April 2024 Patrys announced that new data from preclinical studies using PAT-DX1 and PAT-DX3 in animal models of the autoimmune disease anti-neutrophil cytoplasmic antibody (ANCA) vasculitis was presented during the plenary session of the 21<sup>st</sup> International Vasculitis Workshop. Previously Patrys reported results from a range of non-clinical studies which showed that its deoxymabs suppress the formation of neutrophil extracellular traps (NETs). NETs are structures comprised of DNA strands and certain proteins produced by neutrophils (see press release entitled *"New mechanism by which PAT-DX1 may reduce cancer metastasis"*, 14 June 2022). Recent studies in both animal models and patients have indicated that NETs may play an important role in the establishment and maintenance of cancer cells, the spreading of cancer (metastasis) and in regulating inflammation. These data suggest deoxymabs may provide a therapeutic option for autoimmune vasculitis that is able to reduce inflammation without suppressing the immune system. This may open up an additional indication for Patrys to develop or partner its deoxymab technology.

Patrys' maintains an active business development program and is continuing to undertake exploratory work with several global pharmaceutical and biotechnology companies to evaluate development opportunities for Patrys' deoxymab technology for a range of applications from cancer therapies through therapies for inflammatory conditions through to the cellular and nuclear delivery of therapeutic payloads.

### **Corporate Update**

During the quarter ended 31 March 2024, Patrys had net operating cash outflows of A\$1.45 million. At 31 March 2024, Patrys held A\$3.0 million in cash and cash equivalents. During the quarter, Patrys invested A\$0.8 million in R&D activities. Payments to related parties and their associates during the quarter, which are outlined in Section 6 of the accompanying Appendix 4C to this quarterly activity report, were A\$147,000. These payments include non-executive director fees and consulting services as well as salary (including superannuation) for the CEO and Managing Director.

#### -Ends-

This announcement is authorised for release by the Board of Directors of Patrys Limited.

#### For further information, please contact:

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#### **About Patrys Limited**

Based in Melbourne, Australia, Patrys (ASX:PAB) is focused on the development of its deoxymab platform of cell-penetrating antibodies as therapies for a range of different cancers. More information can be found at <u>www.patrys.com</u>.

#### About Patrys' deoxymab 3E10 platform

Patrys' deoxymab platform is based on the deoxymab 3E10 antibody that was first identified as an autoantibody in a mouse model of the human disease systemic lupus erythematosus (SLE). While most antibodies bind to cell surface markers, deoxymab 3E10 penetrates into the cell nuclei and binds directly to DNA where it inhibits DNA repair processes. Cancer cells often have high levels of mutations and underlying deficiencies in the DNA repair mechanisms. For these reasons, the additional inhibition of the DNA repair processes by deoxymab 3E10 can kill cancer cells, but appears to have little impact on normal cells. As a single agent, deoxymab 3E10 has been shown to significantly enhance the efficacy of both chemo- and radiotherapies. Further, deoxymab 3E10 can be conjugated to nanoparticles to target delivery of chemotherapeutics and imaging agents to tumours.

Patrys has developed two humanised forms of deoxymab 3E10, both which have improved activity over the original deoxymab 3E10 antibody. PAT-DX1 is a dimer (two joined subunits) of the short chain from the binding domain of deoxymab 3E10, while PAT-DX3 is a full-sized IgG antibody. In a range of pre-clinical studies, PAT-DX1 has shown significant ability to kill cancer cells in cell models, human tumour explants, xenograft and orthotopic models. PAT-DX1 has been shown to cross the blood brain barrier, reduce tumour size, and increase survival in multiple animal models of brain cancer, other cancers, and cancer metastases. PAT-DX1 is tumour-agnostic, meaning that it can target many different tumour types in the body, regardless of specific tumour antigens. Patrys believes that PAT-DX1 may have application across a wide range of cancers including gliomas, melanomas, prostate, breast, pancreatic and ovarian cancers.

Deoxymabs, such as PAT-DX1 and PAT-DX3, can be used to target nanoparticles carrying a payload of anti-cancer drugs specifically to tumours. This allows specific delivery of cancer drugs to multiple types of cancer while having minimal impact on normal, healthy cells.

Patrys' rights to deoxymab 3E10 are part of a worldwide license to develop and commercialise a portfolio of novel anti-DNA antibodies and antibody fragments, variants and conjugates discovered at Yale University as anti-cancer and diagnostic agents. Six patents covering the unconjugated form of deoxymab 3E10 (and derivatives thereof) have already been granted (Europe, Japan, China, and 3 in the USA), and five patents covering nanoparticle conjugation have been granted (Australia, Canada, China, India and the USA).

# Appendix 4C

# Quarterly cash flow report for entities subject to Listing Rule 4.7B

Name of entity	
PATRYS LIMITED	
ABN	Quarter ended ("current quarter")

Cash flows from operating activities Receipts from customers Payments for a) research and development b) product manufacturing and operating costs	- (794) -	- (2,328) -
Payments for a) research and development b) product manufacturing and operating costs	- (794)	- (2,328) -
<ul> <li>a) research and development</li> <li>b) product manufacturing and operating costs</li> </ul>	(794)	(2,328)
b) product manufacturing and operating costs	(794)	(2,328)
costs	-	-
<ul><li>advertising and marketing</li></ul>	-	-
d) leased assets	-	-
e) staff costs*	(148)	(457)
f) administration and corporate costs	(440)	(874)
Dividends received	-	-
nterest received	29	74
nterest and other costs of finance paid	-	-
ncome taxes paid	-	-
Sovernment grants and tax incentives	-	2,732
Others - IP expenditure	(101)	(199)
let cash from / (used in) operating ctivities	(1,454)	(1,052)
	<ul> <li>d) leased assets</li> <li>e) staff costs*</li> <li>i) administration and corporate costs</li> <li>ividends received</li> <li>interest received</li> <li>interest and other costs of finance paid</li> <li>income taxes paid</li> <li>iovernment grants and tax incentives</li> <li>iothers - IP expenditure</li> <li>iet cash from / (used in) operating</li> <li>ctivities</li> </ul>	d) leased assets-a) staff costs*(148)administration and corporate costs(440)ividends received-interest received29interest and other costs of finance paid-income taxes paid-income taxes paid-interest - IP expenditure(101)iet cash from / (used in) operating(1,454)

2.	Cash flows from investing activities	
2.1	Payments to acquire or for:	
	(g) entities	-
	(h) businesses	-
	(i) property, plant and equipment	(3)
	(j) investments in term deposits	-

Consolidated statement of cash flows		Current quarter \$A'000	Year to date (9 months) \$A'000
	(k) intellectual property	-	-
	(I) other non-current assets	-	-
2.2	Proceeds from disposal of:		
	(a) entities	-	-
	(b) businesses	-	-
	(c) property, plant and equipment	-	-
	(d) investment in term deposits		1,010
	(e) intellectual property	-	-
	(f) other non-current assets	-	-
2.3	Cash flows from loans to other entities	-	-
2.4	Dividends received (see note 3)	-	-
2.5	Other (provide details if material)	-	-
2.6	Net cash from / (used in) investing activities	(3)	1,007

3.	Cash flows from financing activities		
3.1	Proceeds from issues of equity securities (excluding convertible debt securities)	-	-
3.2	Proceeds from issue of convertible debt securities	-	-
3.3	Proceeds from exercise of options	-	5
3.4	Transaction costs related to issues of equity securities or convertible debt securities	-	-
3.5	Proceeds from borrowings	-	-
3.6	Repayment of borrowings	-	-
3.7	Transaction costs related to loans and borrowings	-	-
3.8	Dividends paid	-	-
3.9	Other (provide details if material)	-	-
3.10	Net cash from / (used in) financing activities	-	5

4.	Net increase / (decrease) in cash and cash equivalents for the period		
4.1	Cash and cash equivalents at beginning of period	4,462	3,045
4.2	Net cash from / (used in) operating activities (item 1.9 above)	(1,454)	(1,052)

Con	solidated statement of cash flows	Current quarter \$A'000	Year to date (9 months) \$A'000
4.3	Net cash from / (used in) investing activities (item 2.6 above)	(3)	1,007
4.4	Net cash from / (used in) financing activities (item 3.10 above)	-	5
4.5	Effect of movement in exchange rates on cash held	-	-
4.6	Cash and cash equivalents at end of period	3,005	3,005

5.	Reconciliation of cash and cash equivalents at the end of the quarter (as shown in the consolidated statement of cash flows) to the related items in the accounts	Current quarter \$A'000	Previous quarter \$A'000
5.1	Bank balances	958	2,440
5.2	Call deposits	2,047	2,022
5.3	Bank overdrafts	-	-
5.4	Other (provide details)	-	-
5.5	Cash and cash equivalents at end of quarter (should equal item 4.6 above)	3,005	4,462

6.	Payments to related parties of the entity and their associates	Current quarter \$A'000
6.1	Aggregate amount of payments to related parties and their associates included in item 1	147
6.2	Aggregate amount of payments to related parties and their associates included in item 2	-

7.	<b>Financing facilities</b> Note: the term "facility' includes all forms of financing arrangements available to the entity. Add notes as necessary for an understanding of the sources of finance available to the entity.	Total facility amount at quarter end \$A'000	Amount drawn at quarter end \$A'000	
7.1	Loan facilities	-	-	
7.2	Credit standby arrangements	-	-	
7.3	Other (please specify)	-	-	
7.4	Total financing facilities	-	-	
7.5	Unused financing facilities available at qu	larter end	-	
7.6	Include in the box below a description of each facility above, including the lender, interest rate, maturity date and whether it is secured or unsecured. If any additional financing facilities have been entered into or are proposed to be entered into after quarter end, include a note providing details of those facilities as well.			
	N/A			

8.	Estimated cash available for future operating activities \$A'000			
8.1	Net cash from / (used in) operating activities (item 1.9)	(1,454)		
8.2	Cash and cash equivalents at quarter end (item 4.6)	3,005		
8.3	Unused finance facilities available at quarter end (item 7.5)	-		
8.4	Total available funding (item 8.2 + item 8.3)	3,005		
8.5	Estimated quarters of funding available (item 8.4 divided by item 8.1)	2.07		
	Note: if the entity has reported positive net operating cash flows in item 1.9, answer item 8.5 as "N/A". Other figure for the estimated quarters of funding available must be included in item 8.5			
8.6	If item 8.5 is less than 2 quarters, please provide answers to the following questions:			
	8.6.1 Does the entity expect that it will continue to have the current l cash flows for the time being and, if not, why not?	evel of net operating		
	Answer: N/A			
	8.6.2 Has the entity taken any steps, or does it propose to take any steps, to raise further cash to fund its operations and, if so, what are those steps and how likely does it believe that they will be successful?			
	Answer: N/A			
	8.6.3 Does the entity expect to be able to continue its operations and to meet its business objectives and, if so, on what basis?			
	Answer: N/A			

#### **Compliance statement**

- 1 This statement has been prepared in accordance with accounting standards and policies which comply with Listing Rule 19.11A.
- 2 This statement gives a true and fair view of the matters disclosed.

Date: 30 April 2024

#### Notes

- This quarterly cash flow report and the accompanying activity report provide a basis for informing the market about the entity's activities for the past quarter, how they have been financed and the effect this has had on its cash position. An entity that wishes to disclose additional information over and above the minimum required under the Listing Rules is encouraged to do so.
- 2. If this quarterly cash flow report has been prepared in accordance with Australian Accounting Standards, the definitions in, and provisions of, AASB 107: Statement of Cash Flows apply to this report. If this quarterly cash flow report has been prepared in accordance with other accounting standards agreed by ASX pursuant to Listing Rule 19.11A, the corresponding equivalent standard applies to this report.
- 3. Dividends received may be classified either as cash flows from operating activities or cash flows from investing activities, depending on the accounting policy of the entity.
- 4. If this report has been authorised for release to the market by your board of directors, you can insert here: "By the board". If it has been authorised for release to the market by a committee of your board of directors, you can insert here: "By the [name of board committee eg Audit and Risk Committee]". If it has been authorised for release to the market by a disclosure committee, you can insert here: "By the Disclosure Committee".
- 5. If this report has been authorised for release to the market by your board of directors and you wish to hold yourself out as complying with recommendation 4.2 of the ASX Corporate Governance Council's Corporate Governance Principles and Recommendations, the board should have received a declaration from its CEO and CFO that, in their opinion, the financial records of the entity have been properly maintained, that this report complies with the appropriate accounting standards and gives a true and fair view of the cash flows of the entity, and that their opinion has been formed on the basis of a sound system of risk management and internal control which is operating effectively.