

Cleansing Notice - Notice under ASIC Corporations (Share and Interest Purchase Plans) Instrument 2019/547

MELBOURNE Australia, 26 August 2026: Prescient Therapeutics Limited (ASX: PTX) a clinical stage oncology company developing personalised therapies for cancer, is conducting a share purchase plan (**SPP**) as part of its capital raising initiative announced by the Company on 25 August 2026. The Company is targeting a raise of up to \$7 million. Under the SPP, eligible shareholders may apply to purchase up to \$30,000 worth of fully paid ordinary shares (**Shares**) at an offer price of \$0.065 (6.5 cents) per Share, potentially issuing approximately 107.7 million Shares.

Under ASIC Corporations (Share and Interest Purchase Plans) Instrument 2019/547 (**ASIC Instrument 2019/547**), the Company gives notice that:

(a) the Company will make offers to issue Shares under the SPP without disclosure to investors under Part 6D.2 of the *Corporations Act 2001 (Cth)* (**Corporations Act**);

(c) the Company is providing this notice under paragraph 7(f)(ii) of ASIC Instrument 2019/547, which requires the Company to give this notice to ASX within 24 hours before the SPP offer is made.

(d) as at the date of this notice the Company has complied with:

- i. the provisions of Chapter 2M of the Corporations Act as they apply to the Company;
- ii. and sections 674 and 674A of the Corporations Act; and

(e) as at the date of this notice, there is no excluded information of the type referred to in Sections 708A(7) and 708A(8) of the Corporations Act which is required to be disclosed pursuant to section 708A(6)(e).

The Board of Prescient Therapeutics Limited has approved the release of this announcement.

For more information please contact:

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About Prescient Therapeutics Limited (Prescient)

Prescient Therapeutics is a clinical-stage oncology company developing personalised cancer treatments through targeted therapies and advanced cell therapy enhancement platforms.

Targeted Therapy

PTX-100 is a first-in-class compound with the ability to block an enzyme that can contribute to cancer growth known as geranylgeranyl transferase-1 (GGTase-1). Blocking this target disrupts oncogenic Ras pathways by inhibiting the activation of Rho, Rac and Ral signaling circuits in cancer cells, leading to their apoptosis (death). Mutations in Ras pathways are believed to be involved in up to 22% of all cancers.

PTX-100 demonstrated safety and early clinical activity in a previous Phase 1 study and in a recent PK/PD basket study of hematological and solid malignancies. PTX-100 has recently completed a Phase 1b expansion cohort study in T-cell lymphomas, where it showed encouraging efficacy and safety. The US FDA has granted PTX-100 Orphan Drug Designation for all T-Cell Lymphomas and Fast Track Designation for the treatment of adults with relapsed or refractory (r/r) mycosis fungoides, the most common subtype of CTCL. A Phase 2 study in Cutaneous T-cell lymphoma (CTCL) is recruiting globally and expects to enrol up to 40 patients in the Phase 2a part of the trial.

Cell Therapy Platforms

OmniCAR: is a universal immune receptor platform enabling controllable T-cell activity and multi-antigen targeting with a single cell product. OmniCAR's modular CAR system decouples antigen recognition from the T-cell signalling domain. It is the first universal immune receptor allowing post-translational covalent loading of binders to T-cells. OmniCAR is based on technology licensed from Penn; the SpyTag/SpyCatcher binding system licensed from Oxford University; and other assets. OmniCAR is in pre-clinical development.

The targeting ligand can be administered separately to CAR-T cells, creating on-demand T-cell activity post infusion and enables the CAR-T to be directed to an array of different tumour antigens. OmniCAR provides a method for single-vector, single cell product targeting of multiple antigens simultaneous or sequentially, whilst allowing continual re-arming to generate, regulate and diversify a sustained T-cell response over time.

CellPryme-A: CellPryme-A is an adjuvant therapy designed to be administered to patients alongside cellular immunotherapy to help them overcome a suppressive tumour microenvironment. CellPryme-A significantly decreases suppressive regulatory T cells; increases expansion of CAR-T cells in vivo; increases tumour penetration of CAR-T cells. CellPryme-A improves tumour killing and host survival of CAR-T cell therapies, and these benefits are even greater when used in conjunction with CellPryme-M pre-treated CAR-T cells.

CellPryme-M: Prescient's novel, ready-for-the-clinic, CellPryme-M technology enhances adoptive cell therapy performance by shifting T cells towards a central memory phenotype, improving persistence, and increasing the ability to find and penetrate tumours. CellPryme-M is a 24-hour, non-disruptive process during cell manufacturing. Cell therapies that could benefit from additional productivity in manufacturing or increased potency and durability in-vivo, would be good candidates for CellPryme-M.

Find out more at www.ptxtherapeutics.com or connect with us via [LinkedIn](#).

Forward-Looking Statements

This announcement contains forward-looking statements based on current expectations, estimates, and assumptions. These statements are subject to risks and uncertainties that may cause actual results to differ materially. Readers are cautioned not to place undue reliance on forward-looking statements, which speak only as of the date of this announcement. Prescient undertakes no obligation to revise forward-looking statements except as required by law. Find out more at www.ptxtherapeutics.com or connect with us via [LinkedIn](#).

Disclaimer and Safe Harbor Statement

Certain statements made in this document are forward-looking statements within the meaning of the safe harbor provisions of the United States Private Securities Litigation Reform Act of 1995. These forward-looking statements are not historical facts but rather are based on the current expectations of Prescient Therapeutics Limited ("Prescient" or the "Company"), their estimates, assumptions, and projections about the industry in which Prescient operates. Material referred to in this document that use the words 'estimate', 'project', 'intend', 'expect', 'plan', 'believe', 'guidance', and similar expressions are intended to identify forward-looking statements and should be considered an at-risk statement. These forward-looking statements are not a guarantee of future performance and involve known and unknown risks and uncertainties, some of which are beyond the control of Prescient or which are difficult to predict, which could cause the actual results, performance, or achievements of Prescient to be materially different from those which may be expressed or implied by these statements. These statements are based on our management's current expectations and are subject to a number of uncertainties and risks that could change the results described in the forward-looking statements. Risks and uncertainties include, but are not limited to, general industry conditions and competition, general economic factors, global pandemics and related disruptions, the impact of pharmaceutical industry development and health care legislation in the United States and internationally, and challenges inherent in new product development. In particular, there are substantial risks in drug development including risks that studies fail to achieve an acceptable level of safety and/or efficacy. Investors should be aware that there are no assurances that results will not differ from those projected and Prescient cautions shareholders and prospective shareholders not to place undue reliance on these forward-looking statements, which reflect the view of Prescient only as of the date of this announcement. Prescient is not under a duty to update any forward-looking statement as a result of new information, future events or otherwise, except as required by law or by any appropriate regulatory authority.

Certain statements contained in this document, including, without limitation, statements containing the words "believes," "plans," "expects," "anticipates," and words of similar import, constitute "forward-looking statements." Such forward-looking statements involve known and unknown risks, uncertainties and other factors that may cause the actual results, performance or achievements of Prescient to be materially different from any future results, performance or achievements expressed or implied by such forward-looking statements. Such factors include, among others, the following: the risk that our clinical trials will be delayed and not completed on a timely basis; the risk that the results from the clinical trials are not as favourable as we anticipate; the risk that our clinical trials will be more costly than anticipated; and the risk that applicable regulatory authorities may ask for additional data, information or studies to be completed or provided prior to their approval of our products. Given these uncertainties, undue reliance should not be placed on such forward-looking statements. The Company disclaims any obligation to update any such factors or to publicly announce the results of any revisions to any of the forward-looking statements contained herein to reflect future events or developments except as required by law.

This document may not contain all the details and information necessary for you to make a decision or evaluation. Neither this document nor any of its contents may be used for any other purpose without the prior written consent of the Company.