

Transforming Patient Outcomes with Superior Vision Gains

Corporate Overview | October 2024 NASDAQ (OPT); ASX (OPT.AX)

Disclaimer and Forward-looking Statements

This presentation includes general background information about the activities of Opthea Limited (ABN 32 006 340 567) ("Opthea" or "Company") and its affiliates and subsidiaries (together, the "Opthea Group"). The information contained in this presentation is in summary form and does not purport to be complete or to contain all material information about the Opthea Group which a prospective investor or purchaser may require in evaluating a possible investment in Opthea or acquisition of securities in Opthea. The information in this presentation remains subject to change without notice. No member of the Opthea Group nor any director, officer, employee, adviser, agent or representative of any member of the Opthea Group (each an Opthea Party and together, the Opthea Parties) has any obligation to update or correct this presentation.

This presentation contains forward-looking statements within the meaning of the U.S. federal securities laws that involve substantial risks and uncertainties. All statements, other than statements of historical facts, contained in this presentation, including statements regarding the therapeutic and commercial potential and size of the estimated market opportunity of the Company's product in development, the viability of future opportunities, future and the expected timing of top-line data, our expectations about topline data based on masked pooled data, the future cash runway, the financial condition, results of operations and business of Opthea, certain plans, objectives and strategies of management of Opthea, including with respect to the current and planned clinical trials of its product candidate and the future performance of Opthea, are forward-looking statements, and you should not place undue reliance on the forward-looking statements contain these identifying words. Opthea may not actually achieve the plans, intentions or expectations disclosed in the forward-looking statements contained in this presentation reflect Opthea's current views with respect to future events, and we assume no obligation to update any forward-looking statements events could differ materially from the plans, intentions on dexpectations disclosed in the forward-looking statements events as required by applicable law. Please refer to information, including risk factors, set forth in Opthea's Annual Report on Form 20-F filed with the U.S. Securities and Exchange Commission on August 30, 2024, and other future capital requirements and ability to continue as a going concern, the development, testing, production, marketing and sale of drug treatments, regulatory risk and potential loss of regulatory approvals, ongoing clinical studies to demonstrate sozinibercept safety, tolerability and therapeutic efficacy, additional analysis of data from Opthea's Phase 3 clinical trials, including once masked pooled data is unmasked, clinical res

This presentation includes statistical and other industry and market data that we obtained from industry publications and research, surveys and studies conducted by third parties as well as our own estimates of potential market opportunities. All of the market data used in this presentation involves a number of assumptions and limitations, and you are cautioned not to give undue weight to such data. Industry publications and third-party research, surveys and studies generally indicate that their information has been obtained from sources believed to be reliable, although they do not guarantee the accuracy or completeness of such information. Our estimates of the potential market opportunities for our product candidates include several key assumptions based on our industry knowledge, industry publications, third-party research and other surveys, which may be based on a small sample size and may fail to accurately reflect market opportunities. While we believe that our internal assumptions are reasonable, no independent source has verified such assumptions.

The information contained in this presentation does not constitute investment or financial product advice (nor taxation or legal advice) and is not intended to be used as the basis for making an investment decision. The presentation is for informational purposes only and is not a prospectus or other disclosure document under Australian law or the law of any other jurisdiction and does not contain all the information which would be required to be disclosed in a prospectus or other disclosure document. The information presented in this presentation may differ materially from that presented in any disclosure document prepared in connection with any offer of securities. It does not take into account the investment objectives, financial situation, taxation position or needs of any particular investor, which should be considered when deciding if an investment is appropriate. You must consider your own investment objectives, financial situation and needs and conduct your own independent investigations and enquiries, including obtaining taxation, legal, financial or other professional advice in relation to the information contained in this presentation as appropriate to your jurisdiction. This presentation should not be relied upon by the Recipient in considering the merits of any particular transaction.

This presentation does not constitute an offer to sell, or the solicitation of an offer to buy, any securities in the United States or any other jurisdictions in which such an offer would be unlawful prior to registration or qualification under the U.S. Securities Act of 1933, as amended, or the securities laws of any state or other jurisdiction of the United States.

This presentation may contain trademarks and trade names of third parties, which are the property of their respective owners. Third party trademarks and trade names used in this presentation belong to the relevant owners and use is not intended to represent sponsorship, approval or association by or with any of the Opthea Group.

Sozinibercept Has the Potential to Be the First Product in 20 Years to Improve Visual Outcomes

Addressing High Unmet Need	 Despite wide use of anti-VEGF-A therapy, wet AMD patients still experience loss in vision long term¹ Every letter of vision counts to improve quality of life and reduce mortality
Proprietary Technology	 First-in-class VEGF-C/D 'trap' inhibitor intended for combination with standard of care anti-VEGF-A therapies Composition of Matter and Methods of Use Patents through 2034; opportunities to extend beyond 2034*
Superior Lead Asset	 Phase 2b demonstrated superiority in combination with SOC therapy, with well tolerated safety profile Sozinibercept has the potential to improve vision for millions of patients with wet AMD
Topline Data from Pivotal Trials in 2025	 Topline data anticipated for COAST (n=998) in early 2Q CY2025 and ShORe (n=986) in mid-CY2025 Current cash expected to fund operations into 3Q CY2025²
Substantial Market Opportunity	 Multibillion dollar commercial opportunity in a growing market with an established clinical practice Sozinibercept developed for use in combination with any anti-VEGF-A; will not compete directly with SOC therapies

 $\mathsf{AMD}-\mathsf{age}\text{-related}\ \mathsf{macular}\ \mathsf{degeneration};\ \mathsf{MOA}-\mathsf{Mechanism}\ \mathsf{of}\ \mathsf{Action};\ \mathsf{SOC}-\mathsf{Standard}\ \mathsf{of}\ \mathsf{care}$

¹CATT Research Group; Maguire MG et al. Ophthalmology. 2016 Aug.

²Additional funding will be required to reach commercialization of sozinibercept and to meet obligations under the Development Funding Agreement ("DFA"). As a result of obligations under the DFA and applicable law regarding liquidity, the Company expects to raise or obtain additional capital in one or more transactions, earlier than 3Q CY 2025 or anticipated topline data readout dates.

*Potential for Patent Term Extensions & Data and Market Exclusivity (12 Years for Biologic)

Sozinibercept Designed to Improve Visual Outcomes in Combo with VEGF-A Inhibitors; Potential to Create New Multi-Billion Dollar Class



Experienced Leadership Team

Expertise and Track Record to Make a Positive Impact on the Retinal Community

Management Team Chief Medical Advisor Arshad M. Khanani, MD, MA, FASRS ^ograybug Managing Partner, Director of Clinical Research Fred Guerard, PharmD, MS **U**NOVARTIS and Director of Fellowship at Sierra Eye Chief Executive Officer Alcon Associates, and Clinical Professor at the University of Nevada, Reno School of Medicine AMARIN CCARA **Clinical Advisory Board Tom Reilly** 🔅 Allergan **Chief Financial Officer U**NOVARTIS Charles C. Wykoff, MD, PhD Director of Research, Retina Consultants of Texas, Chairman of Research and Clinical Trials Complement graybug Parisa Zamiri, MD, PhD Committee, Retina Consultants of America **U**NOVARTIS **Chief Medical Officer** Tim Jackson, PhD, MB, ChB, FRCophth National Health Service, Consultant at Kings Megan Baldwin, PhD, MAICD Genentech Hospital College Hospital, London Founder, Chief Innovation Officer VIATRIS" OYSTER Jason Slakter, MD Clinical Profession at New York University School Mike Campbell **U**NOVARTIS **Shire** of Medicine and partner at Vitreous Retina Macula **Chief Commercial Officer** Genentech Consultants of New York

Despite Treatment with Standard of Care Anti-VEGF-A Therapies, the Majority of Patients Achieve Suboptimal Vision Outcomes

Despite treatment with anti-VEGF-A therapy*

>45% do not achieve significant vision gains

>60% will have persisting macular fluid

25% will have **further vision loss at 12+ months**



The majority of patients fail to achieve 20/40 vision¹



Suboptimal vision is associated with decrease in Instrumental Activities of Daily Living (IADL) skills²

*Based on randomised, controlled clinical trial data; >45% fail to achieve ≥ 2 lines improvement in Best Corrected Visual Acuity (BCVA); Persisting fluid: SD-OCT CST ≥ 300 µM or Time-Domain OCT CST ≥ 250 µM
 IADL: Instrumental activities of daily living (complex activities related to the ability to live independently)
 ¹Mettu PS, et al. Prog Retin Eye Res. 2021
 ²Hochberg C, et al. Invest Ophthalmol Vis Sci. 2012 May 31.

Every Letter Counts When Loss of Vision Potentially Leads to Increased Mortality Risk



Decrease of 1 ETDRS letter per year increases mortality risk by 16%² associated exclusively with IADL levels

IADL – Instrumental activities of daily living; ETDRS – Early Treatment Diabetic Retinopathy Study chart

¹Ehrlich JR et al. "Association between vision impairment and mortality: a systematic review and meta-analysis." Lancet Glob Health. 2021.

²Christ SL, et al. "Longitudinal relationships among visual acuity, daily functional status, and mortality: the Salisbury Eye Evaluation Study." JAMA Ophthalmol. 2014.

Sozinibercept Has Demonstrated Improvement in Vision Gains and Reduction in Vision Loss

Opportunity in Wet AMD Market for an *Overall Shift* **Towards Superior Visual Outcomes**



Jackson, Timothy L., et al. "A randomized controlled trial of OPT-302, a VEGF-C/D inhibitor for neovascular age-related macular degeneration." Ophthalmology. June 2023.

Sozinibercept, a Proprietary VEGF-C/D "Trap" Inhibitor, Has the Potential to Address the Limitations of Anti-VEGF-A Therapies



The Problem

Wet AMD is a **multi-factorial disease**. Treatment with VEGF-A inhibitors **upregulates VEGF-C/D**, driving angiogenesis and vascular permeability.





When used in combination with any VEGF-A inhibitor, **sozinibercept completely blocks** VEGFR-2 and VEGFR-3 signaling.

^a Bevacizumab is used 'off-label' for the treatment of neovascular (wet) AMD

Published Evidence Supports Broader VEGF Pathway Inhibition with Sozinibercept



Sozinibercept Is Designed to Integrate into Current Anti-VEGF-A Clinical Practice



Concentrated prescriptions in U.S. enables potential self-commercialization opportunity with lean and targeted organization

Physicians Willing to Administer Second Injection to up to 41% of Their Patients for Additional BCVA Improvement



What percentage of your Wet AMD patients would you use a second injection (anti-VEGF C/D) immediately after an anti-VEGF-A injection at various levels of BCVA improvement of the combination over SoC? (Among Total Respondents, Avg. % of Patients*, n=125)

Estimate 1% Share of Wet AMD TAM Equals ~\$100M+ in Sales Per Annum

Long-term Value Opportunities for Sozinibercept

Main Patent Family Extends through 2034, with Expansion Opportunities Beyond 2034*

	DEVELOPMENT PHASE				ANTICIPATED
PROGRAM	RESEARCH / PRECLINICAL	PHASE 1	PHASE 2	PHASE 3	MILESTONES
Wet Age-Related Macul	ar Degeneration (W	(et AMD)			
Sozinibercept For use in combination with anti-VEGF-A therapies					Topline data: COAST (in early 2Q CY2025) ShORe (in mid-CY2025)
Diabetic Macular Edem	a (DME)				
Sozinibercept For use in combination with anti-VEGF-A therapies					Phase 3 ready
Co-formulation (Sozinil	bercept + VEGF-A Ir	nhibitor)			
Sozinibercept Co-formulation with VEGF-A Inhibitor					Feasibility underway

Robust Phase 2b Trial in Wet AMD Demonstrated Superiority in Visual Outcome



CNV – choroidal neovascularisation; IVT – intravitreal; Q4W – once very 4 weeks; ITT – Intent to Treat Population, all participants who were randomised into the study irrespective of whether study medication was administered or not; Safety Population - all participants in the ITT but excluding those who did not receive at least one dose of study medication; mITT – Modified ITT Population, all participants in the Safety Population but excludes any participant without a Baseline VA score and/or any participant who did not return for at least one post-baseline visit

Phase 2b Primary and Secondary Endpoints

Primary Endpoint

Mean change from baseline in BCVA at week 24

Key Secondary Endpoints

Proportion of patients gaining ≥15 letters from baseline at week 24

Change in central subfield thickness (CST) from baseline at week 24

Change in intra-retinal and sub-retinal fluid from baseline to week 24

Safety and tolerability

Select Pre-specified Subgroups

Predominantly classic, minimally classic, & occult lesions (Stratification Factor)

> Retinal Angiomatous Proliferation (RAP) detected/not detected at baseline

Polypoidal Choroidal Vasculopathy (PCV) detected/not detected at baseline

Phase 2b Trial Demographics and Baseline Characteristics

Demographic/Ba	seline Disease Characteristic	Sham + ranibizumab n=121	0.5 mg sozinibercept + ranibizumab n=122	2 mg sozinibercept + ranibizumab n=123
Mean Age – years ± SD		76.1 ± 9.48	78.8 ± 8.16	77.8 ± 8.82
Male		48 (39.7%)	49 (40.2%)	45 (36.6%)
Sex – n (%)	Female	73 (60.3%)	73 (59.8%)	78 (63.4%)
Caucasian Race – n	(%)	117 (99.2%)	119 (99.2%)	117 (97.5%)
Mean Visual Acuity (BCVA) – letters ± SD		50.7 ± 10.21	51.1 ± 8.96	49.5 ± 10.26
Mean Total Lesion Area - mm ² ± SD		6.08 ± 3.21	6.48 ± 3.30	6.62 ± 3.39
	Predominantly classic - n (%)	15 (12.4%)	15 (12.3%)	16 (13.0%)
	Minimally classic – n (%)	53 (43.8%)	51 (41.8%)	53 (43.1%)
Lesion Type	Occult - n (%)	53 (43.8%)	56 (45.9%)	54 (43.9%)
	PCV detected ¹ -n (%)	20 (16.5%)	24 (19.7%)	22 (17.9%)
	RAP detected ² -n (%)	15 (12.7%)	22 (18.5%)	14 (11.8%)
Mean central subfield thickness (CST) - mm ±SD		412.10 ± 110.62	425.18 ± 120.45	414.12 ± 123.25
Sub-retinal fluid (SRF) present – % participants		89.3%	84.4%	87.8%
Intra-retinal cysts present –% participants		57.9%	63.9%	56.1%

Intent-to-Treat (ITT) population; SD – standard deviation; BCVA – Best Corrected Visual Acuity ¹PCV - polypoidal choroidal vasculopathy, detected by SD-OCT, FA and fundus photography. ²RAP - retinal angiomatous proliferation, detected by SD-OCT, FA and fundus photography.

Sozinibercept 2 mg Combination Therapy Demonstrated Superiority in Visual Acuity over Ranibizumab Monotherapy



To Maximize Probability of Success, Best Responding Phase 2b Patients Represents Primary Analysis Population in the Pivotal Phase 3 Trials



Phase 2b demonstrated **superior efficacy** of +5.7 letter gain over standard of care, based on a pre-specified analysis

This patient population (minimally classic & occult) represents ~75% of wet AMD patients

Control Arm in Phase 2b Overperformed MARINA Trial at Week 24 in in Similar Lesion Type Patient Population

Mean Change in BCVA from Baseline at Week 24 – Sozinibercept Phase 2b vs. MARINA Trial Occult and Minimally Classic Lesions



MARINA was a Phase 3 registrational trial. Baseline BCVA values across trials vary. Number of patients randomised to treatment group (n, bottom table). Mean change in Best Corrected Visual Acuity (BCVA) from baseline shown in ETDRS letters (top of bars).

In a Disease Where Every Letter Counts, a Greater Proportion of Sozinibercept Patients Gained Substantial Vision and Fewer Experienced Vision Loss



Legend (ETDRS Letter Change from Baseline to Week 24)

Greater Proportion of Sozinibercept Patients Achieved Minimum Driving-level of Vision (≥20/40)

Percentage of Participants with 20/40 Vision or Greater at Week 24



Sozinibercept Reduced the Proportion of Patients Experiencing Vision Loss by 82%

Percentage of Participants with any Vision Loss ≥1 ETDRS Letter at Week 24

Occult and Minimally Classic Lesions (RAP Absent) 82% relative decrease compared to ranibizumab control



Decrease of 1 ETDRS letter per year increases mortality risk by 16%² associated exclusively with IADL levels

Modified Intent-to-Treat (mITT) population; as observed.

IADL – Instrumental activities of daily living; ETDRS – Early Treatment Diabetic Retinopathy Study chart

¹Christ SL, et al. "Longitudinal relationships among visual acuity, daily functional status, and mortality: the Salisbury Eye Evaluation Study." JAMA Ophthalmol. 2014.

Sozinibercept Reduced Retinal Thickness and Dried the Retina Better With Combination Therapy in Occult & Minimally Classic (RAP Absent) Patients



Sozinibercept Demonstrated Greater CNV and Lesion Regression With Combination Therapy in Occult & Minimally Classic (RAP Absent) Patients



mITT; as observed; top of bar - statistic, bottom of bar - n. CNV: Choroidal Neovascular.

Sozinibercept Demonstrated Superior Vision Gains in a Pre-Specified Subgroup of Hard-To-Treat PCV Lesion Patients



Polypoidal Choroidal Vasculopathy (**PCV**) is a difficult-to-treat wet AMD subtype; it is often described as the **most prevalent form of wet AMD worldwide**

PCV is **highly prevalent in Asian populations** (up to ~60%), while ~8-13% prevalence in Caucasians

Phase 3 ShORe and COAST trials enrolled patients with PCV¹

*Unadjusted p-value

¹ Evaluated by color fundus photography (FP), fluorescein angiography (FA), and spectral domain optical coherence tomography (SD-OCT)

Pooled Safety for Completed Sozinibercept Trials

Combination Therapy Well Tolerated and Comparable to Standard of Care Monotherapy

N Participants (%)	Sozinibercept Any dose* N=399 (N=1,842 injections)	Sozinibercept 2 mg N=263 (N=1,121 injections)	Sham + anti-VEGF-A control N=170 (N=854 injections)
Ocular TEAEs - Study Eye – related to study product(s)	41 (10.2%)	22 (8.4%)	20 (11.8%)
Ocular TEAEs - Study Eye – Severe	4 (1.0%)	2 (0.8%)	2 (1.2%)
Intraocular inflammation – Study Eye	71,2,3 (1.8%)	3 ¹ (1.1%)	31 (1.8%)
Participants with AEs leading to treatment discontinuation	42,4-6 (1.0%)	14 (0.4%)	27,8 (1.2%)
Any APTC event	4 ^{4,5,9,10} (1.0%)	35,9,10(1.1%)	211,12 (1.2%)
Deaths	210,13 (0.5%)	210,13 (0.8%)	214,15 (1.2%)

- Pooled safety analysis of 399 patients for completed sozinibercept trials
- Data Monitoring Committee ("DMC") regularly reviews data from ongoing Phase 3 COAST and ShORe studies
- Safety data from our completed sozinibercept trials show sozinibercept combination therapy has a safety and tolerability profile comparable to standard of care anti-VEGF-A monotherapy.
- Masked data from patients that have completed the week 52 visit in the ongoing Phase 3 clinical trials show greater mean BCVA increases from baseline than results with standard of care anti-VEGF-A monotherapy from Opthea's Phase 2b study**

¹Transient anterior chamber cell (trace 1-4 cells); ² SAE of endophthalmitis, with AE's of hypopyon and anterior chamber cell (n=1; 0.5 mg); ³ SAE of vitritis (n=1; 0.5 mg); ⁴Non-fatal myocardial infarction; ⁵Cerebrovascular accident; ⁶Enteritis; ⁷Abdominal pain; ⁸Increased IOP; ⁹ Non-fatal angina pectoris; ¹⁰Fatal congestive heart failure/myocardial infarction; ¹¹Non-fatal arterial embolism; ¹²Embolic stroke; ¹³Metatstaic ovarian cancer; ¹⁴ Pneumonia; ¹⁵ infective endocarditis. *Any dose (sozinibercept 0.3 mg, 0.5 mg, 1 mg or 2 mg)

**Masked data represent pooled data from both sozinibercept combination and standard of care monotherapy treatment arms. The Phase 3 clinical trial masked data are incomplete and subject to additional analysis once unmasked. There is no assurance that standard of care monotherapy in our Phase 3 clinical trials will yield similar results to our prior clinical trials or previously published clinical trials with anti-VEGF-A monotherapies. As a result, there can be no assurance that topline results for sozinibercept from the Phase 3 clinical trial, if completed, will be consistent with results from masked data available to date.

Intraocular Inflammation Observed in Combination Therapy Across Completed Sozinibercept Trials Similar to Standard of Care

N Participants (%)	Sozinibercept Any dose* N=399 (N=1,842 injections)	Sozinibercept 2 mg N=263 (N=1,121 injections)	Sham + anti-VEGF-A control N=170 (N=854 injections)
Intraocular Inflammation ¹	7 (1.8%)	3 (1.1%)	3 (1.8%)
OPT-302-1001 (Phase 1/2a wet AMD)	2	0	0
Uveitis with anterior chamber cell 1+	1	0	0
Uveitis with anterior chamber cell 2+	1	0	0
OPT-302-1002 (Phase 2b wet AMD)	3	1	2 ^a
Endophthalmitis with anterior chamber 1+ and hypopyon	1	0	0
Vitritis	1	0	0
Anterior chamber cell, trace	1	1	2 ^a
OPT-302-1003 (Phase 1b/2a DME)	2 ^b	2 ^b	1
Iritis with keratic precipitates and anterior chamber cell 2+	1	1	0
Iritis with anterior chamber cell 2+	0	0	1
Anterior chamber cell 4+, associated with cataract extraction/ intraocular lens implant and hyphema	1 ^b	1 ^b	0

Safety population

¹AEs observations considered to be indicative of intraocular inflammation, defined prior to database lock

^aObserved during ophthalmic examination, but not reported as TEAEs

^bConsidered associated with lens extraction and not reported as TEAEs

Phase 3 Clinical Program Is Informed by Phase 2b Results and Optimized for Success



Hierarchical primary analysis first conducted in the high-responding occult and minimally classic population (RAP absent), followed by total patient population



Two robust pivotal trials studying sozinibercept in combination with Eylea® and Lucentis® in treatment naïve patients with wet AMD



Phase 3 designed to support broad label for use in combination with any VEGF-A inhibitor for all wet AMD patients (treatment naïve and prior treated)

Phase 3 Wet AMD Trials COAST and ShORe Are Well Advanced Topline Data Anticipated for COAST in Early 2Q CY 2025 and ShORe in Mid-CY2025

Design	 Multi-center, double-masked, randomized (1:1:1), sham control Treatment naïve wet AMD patients
Sample Size	 COAST n=998; ShORe n=986
Comparators	 2 mg Eylea[®] q8w (COAST) & 0.5 mg Lucentis[®] q4w (ShORe)
Regulatory Quality	 ~90% power, 5% type I error rate

Phase 3 Primary and Secondary Endpoints Primary Efficacy Endpoint at Week 52 to Support BLA Submission

Primary Endpoint

Mean change from baseline in BCVA at week 52

Key Secondary Endpoints (Baseline to Week 52)

Proportion of participants gaining \geq 15 letters

Proportion of participants gaining \geq 10 letters

Change in choroidal neovascularization area

Proportion of participants with absence of both sub-retinal fluid and intra-retinal cysts

Phase 3 Trial Design Supports Potential Broad Label for Use With Any Anti-VEGF-A Therapy



Standard of care administered according to approved dosing schedule: aflibercept (2 mg IVT q8w after 3 loading doses) and ranibizumab (0.5 mg IVT q4w after 3 loading doses). Sozinibercept dosed at 2 mg. Note that Sham administered at visits when sozinibercept is not administered. Maintenance dosing continued through end of the safety follow-up.

Enrollment Complete

Advancing Therapeutic Innovations to Transform Patient Outcomes with Superior Vision Gains

We are dedicated to advancing sozinibercept to **improve patients' visual outcomes**

iteps	Clinical Milestones	 Phase 3 program enrolled 1,984 patients across COAST and ShORe Topline data anticipated for COAST in early 2Q CY2025 and ShORe in mid-CY2025
	Manufacturing Scale-up	 DS PPQ campaign completed Sep-2024; update on DP PPQ in early CY2025 PPQ validation batches supportive of BLA filing and launch
Next §	Regulatory Preparations	 FDA Fast Track designation allows rolling submission of completed BLA modules
	Commercial Readiness	 Strengthen medical expert engagement and develop market access strategy Complete development of product launch plan

Recent Financings Anticipated to Provide Cash Runway Through Both Pivotal Topline Data Readouts

Financial Overview	
Ticker	OPT (ASX/NASDAQ)
Shares Outstanding ¹	Ordinary Shares: 1,231.1M ADS equivalents: 153.9M
Cash/Cash Equivalents ²	US\$207.3M
Offices	Melbourne, Australia Princeton, NJ

Development Funding Agreement (DFA)

- Total funding drawn under DFA: US\$170M
- Provides non-dilutive funding for development of sozinibercept
- If sozinibercept is approved, repayment is capped at 4x investment and split between fixed payments and variable payments at 7% of revenues
- No amounts owed if the clinical trials do not meet the primary endpoint or if regulatory approval is not received³

¹As of June 30, 2024, pro-forma for the 2024 Retail Entitlement Offer which closed in July 2024.

²Includes \$172.5M as of June 30, 2024 and \$34.8M net proceeds from the 2024 Retail Entitlement Offer which closed in July 2024.

³In certain circumstances, upon or following the termination of the DFA, the Company may owe the DFA investors a multiple of amounts paid to the Company under the DFA. Please refer to the description of the DFA included in the Company's Form 6-K filed with the SEC on August 15, 2022 and the DFA filed as Exhibit 4.14 to the Company's Annual Report on Form 20-F filed with the SEC on September 29, 2022 for more information. Note: Additional funding will be required to reach commercialization of sozinibercept and to meet obligations under the Development Funding Agreement ("DFA"). As a result of obligations under the DFA and applicable law regarding liquidity, the Company expects to raise or obtain additional capital in one or more transactions, earlier than 3Q CY 2025 or anticipated topline data readout dates.

Sozinibercept Will Not Compete Head-to-Head with Anti-VEGF-A Differentiated Combination Approach Targeting Better Visual Outcomes Drives Commercial Value



Addressing unmet medical need of improved efficacy in large wet AMD patient population in a potential ~\$15B market



First and only therapy to have demonstrated superior visual outcomes over anti-VEGF-A therapy with a novel and highly differentiated MOA



Only asset in near or long-term pipeline with potential to disrupt treatment paradigm on basis of efficacy in wet AMD



Concentrated prescriptions in U.S. enables potential selfcommercialization opportunity with lean and targeted organization