CLINUVEL

From Orphan to Large

CAPITAL MARKETS BRIEFING

Strategic Update VIII | Sydney, 01 May 2024

ASX: CUV | Börse Frankfurt: UR9 | ADR Level 1: CLVLY

Forward-looking statement CLINUVEL GROUP

This release contains forward-looking statements, which reflect the current beliefs and expectations of CLINUVEL's management. Statements may involve a number of known and unknown risks that could cause our future results, performance, or achievements to differ significantly from those expressed or implied by such forward-looking statements. Important factors that could cause or contribute to such differences include risks relating to: our ability to develop and commercialise pharmaceutical products; the COVID-19 pandemic and/or other world, regional or national events affecting the supply chain for a protracted period of time, including our ability to develop, manufacture, market and sell biopharmaceutical products; competition for our products, especially SCENESSE® (afamelanotide 16mg), CYACÊLLE, PRÉNUMBRA® or NEURACTHEL®; our ability to achieve expected safety and efficacy results in a timely manner through our innovative R&D efforts; the effectiveness of our patents and other protections for innovative products, particularly in view of national and regional variations in patent laws; our potential exposure to product liability claims to the extent not covered by insurance; increased government scrutiny in either Australia, the U.S., Europe, Israel, China and Japan of our agreements with third parties and suppliers; our exposure to currency fluctuations and restrictions as well as credit risks; the effects of reforms in healthcare regulation and pharmaceutical pricing and reimbursement; that the Company may incur unexpected delays in the outsourced manufacturing of SCENESSE[®], CYACÊLLE, PRÉNUMBRA[®] or NEURACTHEL[®] which may lead to it being unable to supply its commercial markets and/or clinical trial programs; any failures to comply with any government payment system (i.e. Medicare) reporting and payment obligations; uncertainties surrounding the legislative and regulatory pathways for the registration and approval of biotechnology and consumer based products; decisions by regulatory authorities regarding approval of our products as well as their decisions regarding label claims; our ability to retain or attract key personnel and managerial talent; the impact of broader change within the pharmaceutical industry and related industries; potential changes to tax liabilities or legislation; environmental risks; and other factors that have been discussed in our 2023 Annual Report. Forward-looking statements speak only as of the date on which they are made, and the Company undertakes no obligation, outside of those required under applicable laws or relevant listing rules of the Australian Securities Exchange, to update or revise any forward-looking statement, whether as a result of new information, future events or otherwise. More information on preliminary and uncertain forecasts and estimates is available on request, whereby it is stated that past performance is not an indicator of future performance.

Introduction

Malcolm Bull

From Orphan to Large

- 09:00 Introduction
- 09:05 Chairman
- 09:10 **Commercial Update**
- 09:35 **R&D Pipeline**
- 10:00 *Break*
- 10:20 Focus on Vitiligo, Stroke & DNA Repair
- **11:10 Diversification Strategy and Decision-Making**
- 12:00 **Q&A**
- 12:45 Buffet Lunch
- 14:00 *Close*

Chairman

Prof Jeffrey Rosenfeld

Longevity

Board (5) median tenure 4.4 yrs	Executive Management (9) median tenure 14.9 yrs		anagement (10) lian tenure 4.5 yrs
Accountancy, b	Integrated skill set ookkeeping, legal, investor relations	12	
Medicine, pharma. sciences, chemistry, engineering Gen mgmt., comms, back office		52 21	
Creative, branding		13 98	

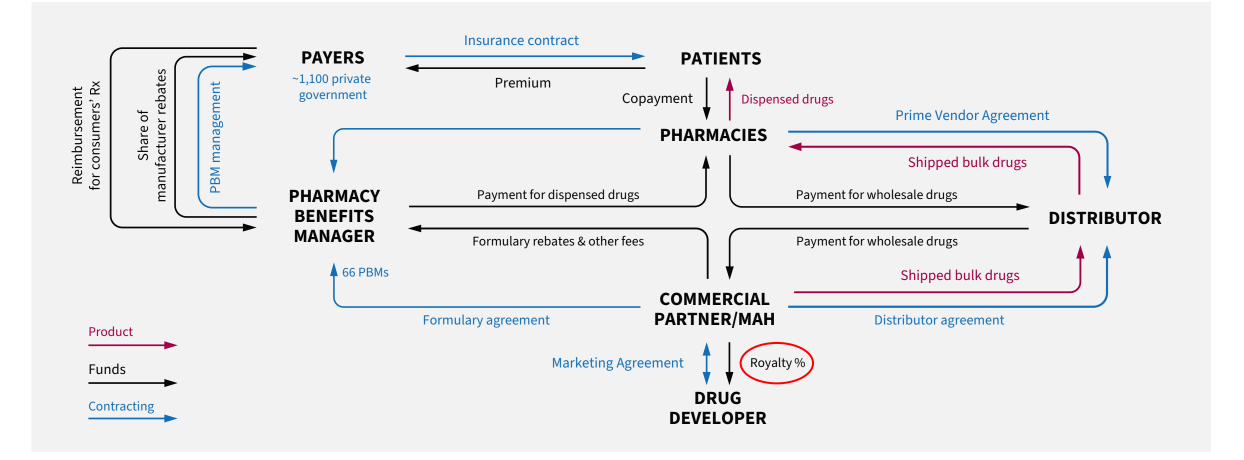
Commercial Update

Lachlan Hay

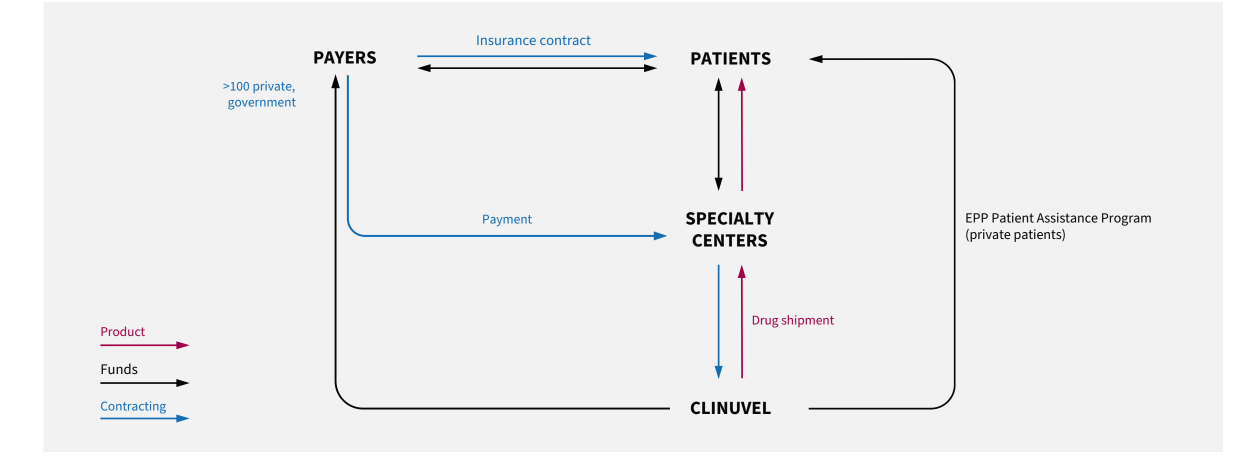
SCENESSE® (afamelanotide) for EPP

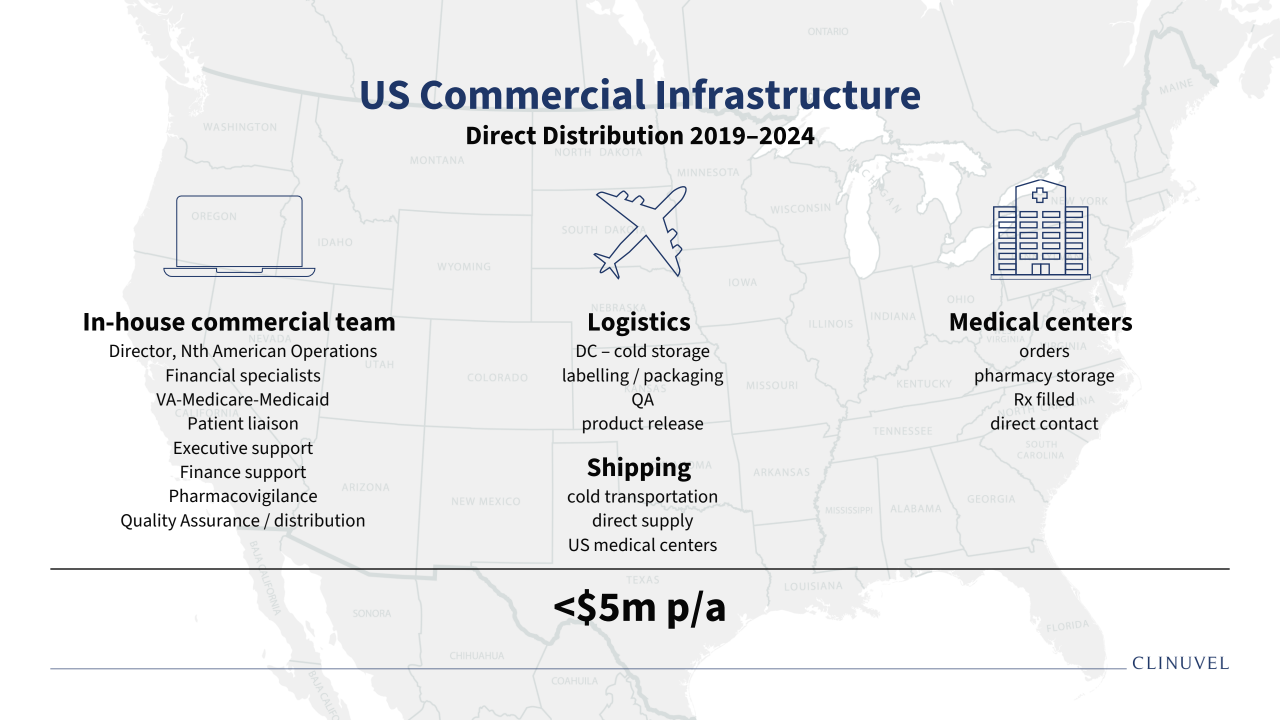
2006 2007 2009 2010 2011 2012 2013 2014 2015	Ph II CUV010 commences Ph III CUV017 commences CUV010 published <i>NEJM</i> Italian 648/96 – first reimbursement Ph III CUV029 completed Swiss Special Access (reimbursed) launched Ph III CUV039 completed EMA approval CUV029/039 published <i>NEJM</i> Long-term observational study <i>BJD</i>	R&D '06-'15	SCENESSE [®] first-in-class (EMA-FDA) 18 years of clinical experience • >30 publications • >14,500 doses
2016 2017 2019 2020 2021 2022 2023	EU launch GKV pricing agreement (DE) FDA approval US launch TGA approval Long-term post-authorisation data JAMA Israeli National Health Basket First adolescent patients treated First Canadian patients treated First US Medicare-Medicaid-VA treatments	COMMERCIAL '16-'24	• >300 EPP patients, ≥15 doses longitudinal use is foundation for expansion

Traditional US commercial model

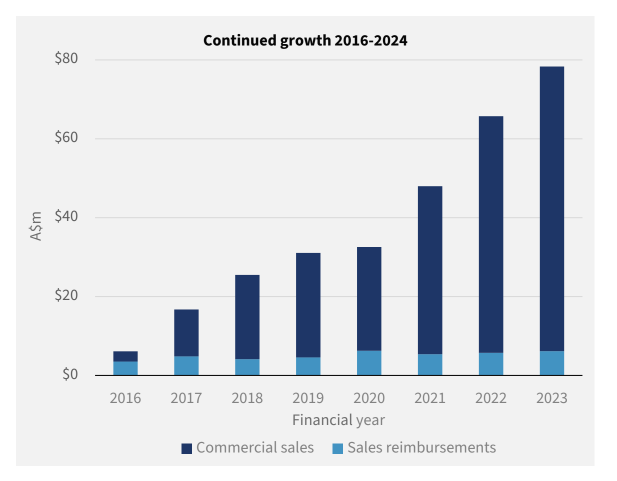


CLINUVEL's US commercial model





SCENESSE® (Rx) for EPP



CLINUVEL self-distributing EU-USA-ISL

- 7% increase filled prescriptions (CY23 vs CY22)
- 4% volume growth (1H FY24 vs 1H FY23)

Distribution agreement in new region negotiated

Revenues >A\$330m since launch (June 2016)

North American distribution 2020-2024

Patient centricity



Objective: 3–4hrs access (cluster distribution)

95% access for patients (pursued PA process)
80 Specialty Centers (trained and accredited)
30 States
100+ private insurers
CMS, VA
CPT[®] code (11981) & J-Code (J7352) established
AMA CPT[®] 12 months
CMS J-Code 18-24 months

Canada special access '24/'25 Health Canada filing

Target 120 Specialty Centers

European distribution 2016-2024

- CY23 highest number of patients treated95% treatment continuation (2022-23)*
- FY245 new European EPP Expert CentresEMA submission label expansion

Future claims

Hepatoprotective effect published**, research ongoing

In our clinical experience afamelanotide treatment is **much more effective** in clinical practice than demonstrated in clinical trials and **should be made available for all EPP patients** meeting inclusion-criteria. Wensink et al, 2021 – Rotterdam (NL)

SCENESSE® for adolescent EPP



EU+US regulatory pathway 7 adolescents (15-17 yrs) off-label treatment

- ≥4 implants
- fully reimbursed

Estimated 60 patients (15-17 yrs) FR-DE-NL-IT

CUV052 pharmacokinetic study underway

- (n=28), 9 patients treated
- first results in 2024

Attempts to date

Experimental therapies	Target, endpoint focus	Status
Beta carotene	Anti-oxidant, no RCT	Use largely discontinued, deployed in absence of therapy access (i.e. paediatric)
Cimetidine, H ₂ -receptor antagonist	Inhibit ALAS, reduce PPIX	Lack of clinical, academic support for use, lack of evidence of safety-efficacy
Dersimelagon, MC1R agonist	Activate melanin, time to prodrome	Phase III (2 doses) failed to meet primary endpoint, compassionate use discontinued, new Phase III recruiting treatment naïve patients
Bitopertin, GlyT1 inhibitor	Limit glycine, reduce PPIX	Phase II failed to meet time in sunlight endpoint, program "pending regulatory feedback"

R&D Pipeline

Dennis Wright

Melanocortin peptides for skin and brain

Afamelanotide (α-MSH analogue) SCENESSE[®] & PRÉNUMBRA[®]



Photoprotective Assist DNA repair Anti-oxidative Anti-oncotic Melanogenesis (bronzing, repigmentation)

Adrenocorticotropic hormone (ACTH) NEURACTHEL[®]



Anti-inflammatory Immune modulation

Pharmaceutical Pipeline

		Preclinical	Phase I	Phase II	Phase II	I Commercial
	SCENESSE [®] (afamelanotide	e 16 mg) in adult EPP (EEA, U	K, CH, USA, ISL, CAN, AUS)			
	SCENESSE [®] (afamelanotide	e 16 mg) in adolescent EPP				
SKIN	SCENESSE [®] (afamelanotide	e 16 mg) in adolescent and ad	dult vitiligo			
SK	SCENESSE [®] (afamelanotide	e 16 mg) in adolescent and ad	dult XP			
	SCENESSE [®] (afamelanotide	e 16 mg) in variegate porphyr	ia			
	CUV9900 transdermal					
	PRÉNUMBRA [®] in arterial iso	chaemic stroke				
BRAIN	PRÉNUMBRA [®] to be disclos	ed				
BR	NEURACTHEL [®] instant – IS,	MS				
	NEURACTHEL® modified re	lease – CNS				

NEURACTHEL[®]

Unmet needs with adrenocorticotropic hormone (ACTH)



Manufacturing agreements (2, exclusive) Analytical methods advanced DMF in development Generic (Instant), Branded (Modified-release) in development sNDA submission 2026

Clinical program planned for

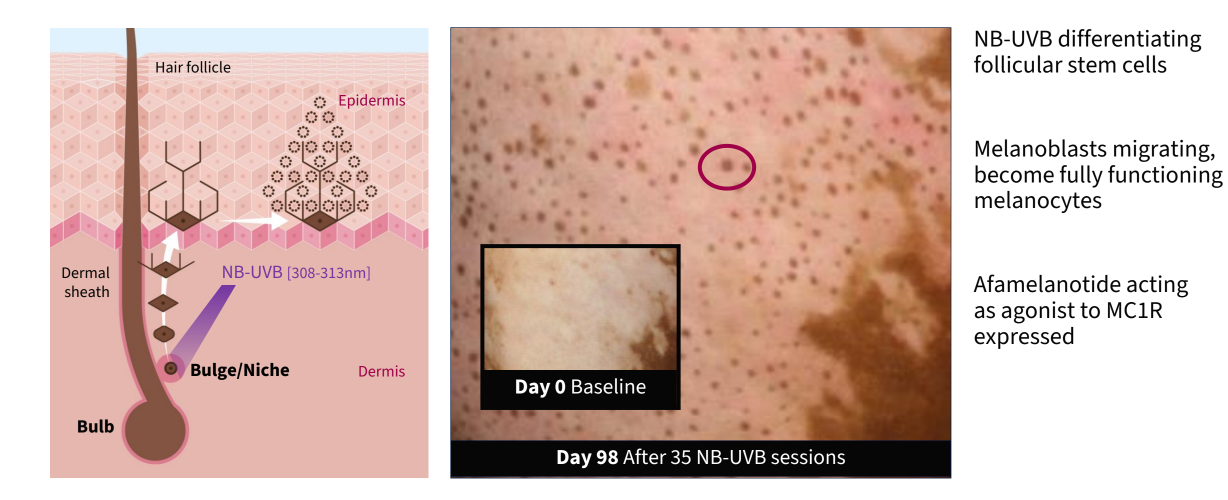
- 1. West Syndrome
- 2. Relapsing MS



Dennis Wright

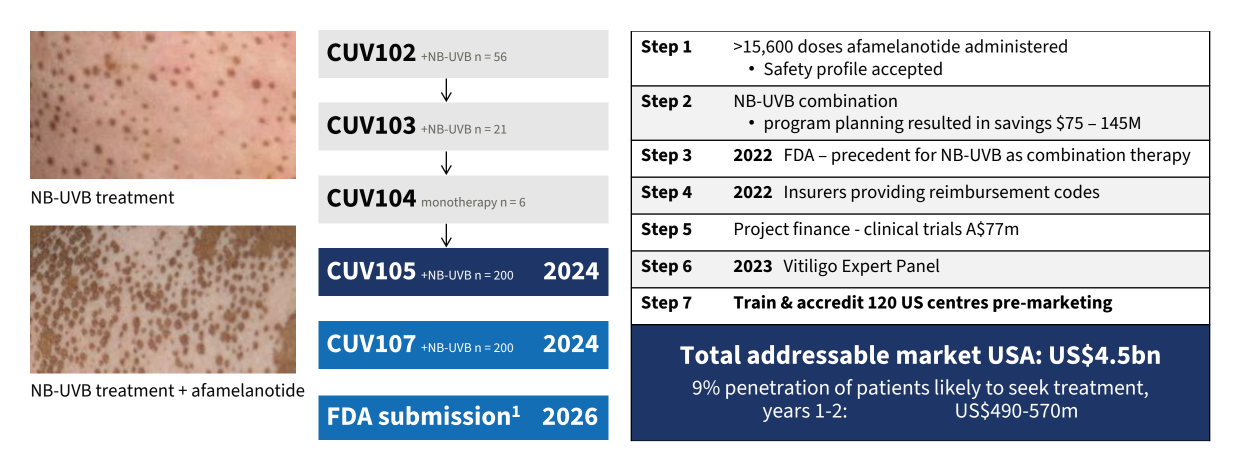


NB-UVB – follicular repigmentation



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Vitiligo Path to market



Vitiligo Global Phase III study (CUV105)

	CLINUVEL CUV105 Phase III	Pfizer pivotal Phase III oral JAK inhibitor*
Study population	N=200, adults and adolescents (≥12 years) highest unmet need: darker skin (Fitzpatrick IV-VI)	
Inclusion	≥0.3% body surface area with facial vitiligo: T-VASI ≥0.3 & F-VASI ≥0.3	
Primary endpoint	repigmentation of total body surface (T-VASI50)	proportion of participants achieving F-VASI75
Secondary endpoint/s	evaluate repigmentation of the face, maintenance of repigmentation	proportion of participants achieving T-VASI50
Randomisation	1:1 to SCENESSE [®] + NB-UVB vs NB-UVB monotherapy	
Treatment duration	20-week treatment phase, six-month follow up	
Sites	expert treatment centres globally	
Status	12-month recruitment (to October 2024)	

"Once on the market, SCENESSE" will clinically become the pigment booster for every dermatologist in North America"

Vitiligo Expert Panel member

SCENESSE® (afamelanotide) for vitiligo

Case study presented to 2024 American Academy of Dermatology (FST IV)





Day 0 baseline

Day 134 7 implants, 39 NB-UVB sessions

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Images have been amended/cropped and pixelated to protect the patient's privacy but are otherwise unaltered. Images courtesy of the investigator.

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Philippe Wolgen

Arterial Ischaemic Stroke

Targeted product position

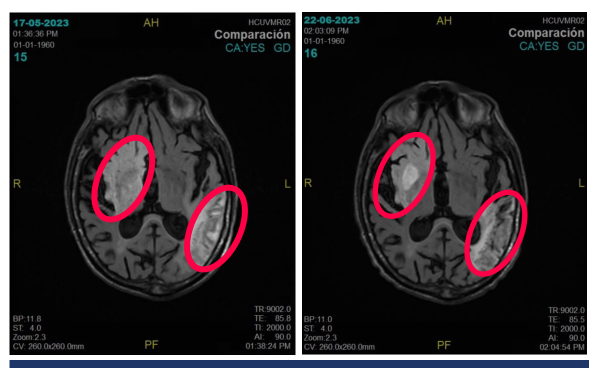
A hormonal treatment to assist hypoxic brain

Study CUV801 (n = 6) proof of concept – afamelanotide

- open-label, up to 4 doses: days 0, 1, 7, 8; evaluation at day 42
- occlusion higher regions: > M1
- functional recovery in 5 patients; NIHSS \geq 4 (4/6)
- cerebral perfusion improved per MRI-FLAIR (CBF, Tmax)

Study CUV803 (n = 12) 9 patients on treatment

- moderate & severe patients
- occlusion higher regions: > M2/A2/P2
- higher, more frequent dosing, PRÉNUMBRA®
- safety
- neurological functionality (NIHSS)
- perfusion of penumbra, oligemic zone



Total addressable market: US\$31B Penetration US\$1.98-2.23B - USA + EU + AU

DNA Repair

Philippe Wolgen

The Photomedicine Foundation



Evaluating XP – DNA Repair – objectives



Reducing DNA damage

Decreasing photodamage

Reducing the risk of skin cancer

29 April 2024

Afamelanotide granted ODD in Europe

- first clinical data on efficacy
- lack of therapy,
- high unmet need,
- medical plausibility of treatment

• potential to "extend" label

- reduction in fees
- 10 years market exclusivity

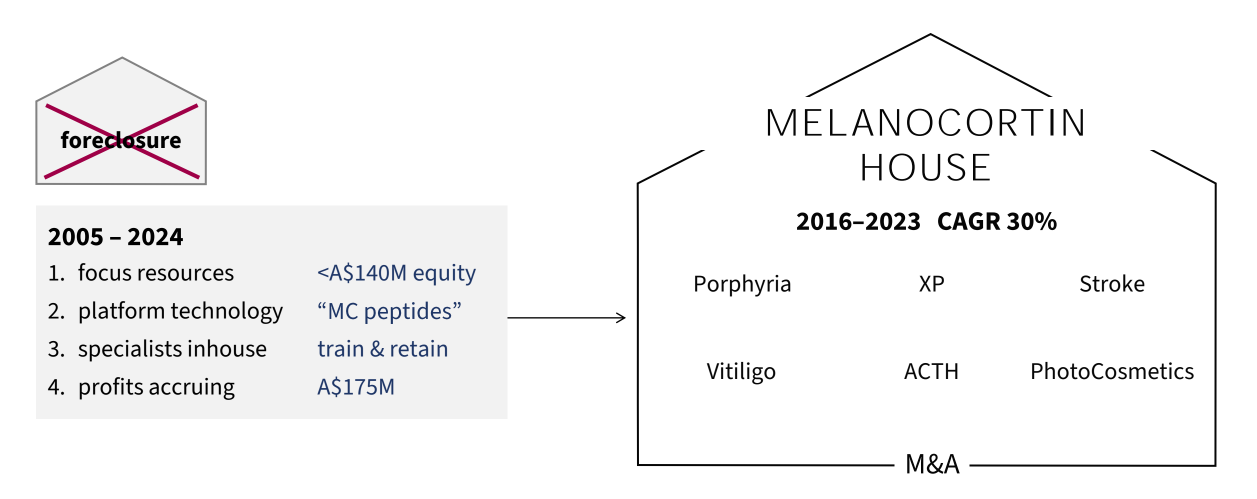
Evaluating XP – DNA Repair – clinical endpoints

		CUV156	CUV151	
DNA repair markers	 'photoproducts' (CPDs) γ-H2AX P53 Ki67 	- reduction - increase - variable, increase	 reduction at 15 mins (p=0.0039) reduction at 24 hours (p=0.0078) no change no change 	
Quality of life	validated questionnaires	pending		
Safety	treatment-emergent adverse events clinical & laboratory evaluations	safety profile maintained	no SADRs	
Erythema response Severity	minimal Erythemal Dose (MED) patient & physician assessments	- variable - pending	- reduction (p=0.018)	
Melanin density	spectrophotometry	- increase	- increase (p<0.05)	
Total addressable market: <us\$100m< b=""> Europe USA Africa Middle East S&C America ~1,300 XP patients worldwide</us\$100m<>				

Decision Model & Diversification Strategy

Philippe Wolgen

CUV Construct



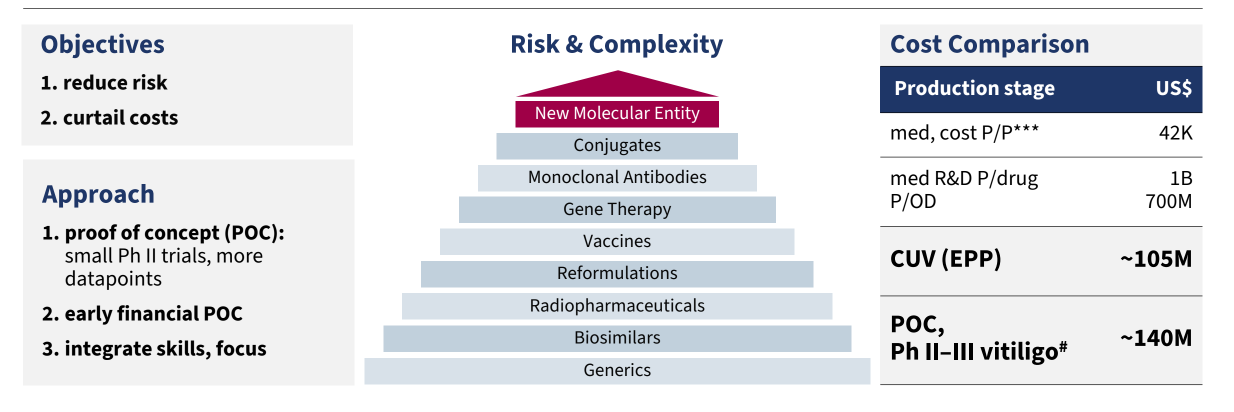
CUV Differentiated Decision Model

no alternativ		ROPEPTIDES eatening or severe academic/clin	ical belief reimbursable
TECHNOLOGY	PROOF OF PRINCIPI	LE —_\$\$ → Ph III	- COMMERCIALISATION
1. SEVERE/LIFE-TH	HREATENING – Brain		
2. EXTREME COND	ITIONS – Skin	TRANSLATION	PHOTOCOSMETICS
I EPP	– absolute intolerance to UV + HEV λ		CYACÊLLE (UV–HEV)
II XP-DNA repair	 photodamage from UV 		DNA assisted repair, DECREASE photodamage
III Vitiligo	 loss of pigmentation 		Risk-free-sunless bronzing
	Pharmaceuticals – Core	Cosmetics – Complemen	tary
			CLINUVEL

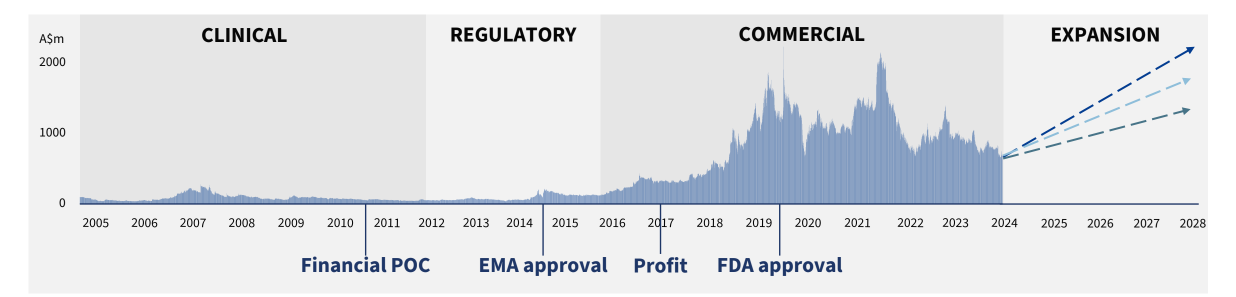
Risk Management

Global Biopharmaceutical Market US\$1.7 trillion*

New compounds 1:5,000–10,000 | 9,000 products in clinical development | 21,000 compounds** (406,000 centres) **13.8% success rate:** Meeting objectives (endpoints)



Longitudinal Valuation CUV

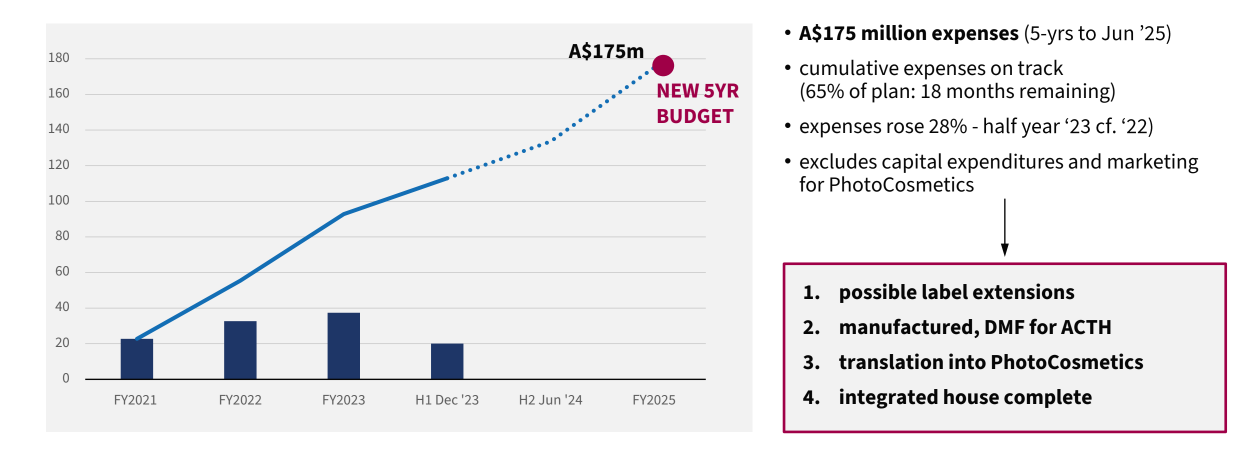


FINANCIAL STRENGTH

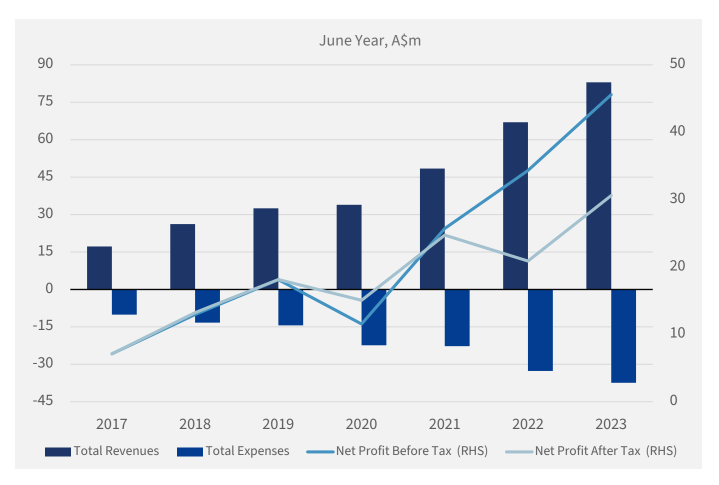
- 1. Phase III vitiligo commercial
- 2. DNA repair
- 3. Neurodegenerative Disease
- 4. PhotoCosmetics
- 5. Manufacturing

Controlled Expenses

Tracking 5-year expense projections 2021–2025



Earnings Growth



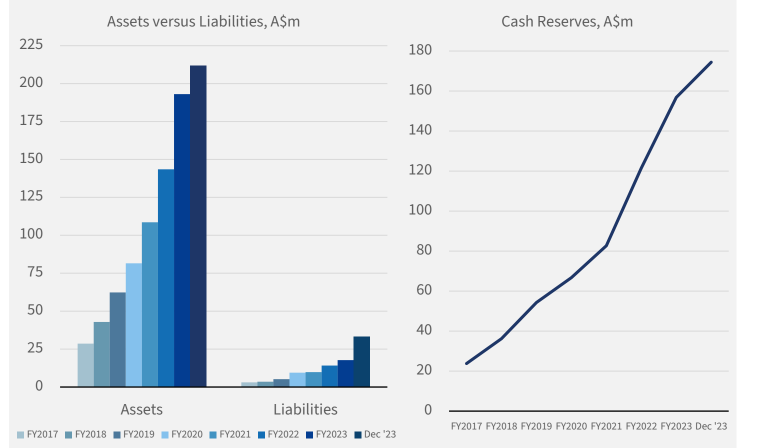
30 June 2023

- 7 yrs annual growth revenues, profits, net cashflow
- 6 yrs annual dividends (< 5% of NPAT)
- cash reserves financing expansion
- earnings per share A\$0.62
- return on equity 19%

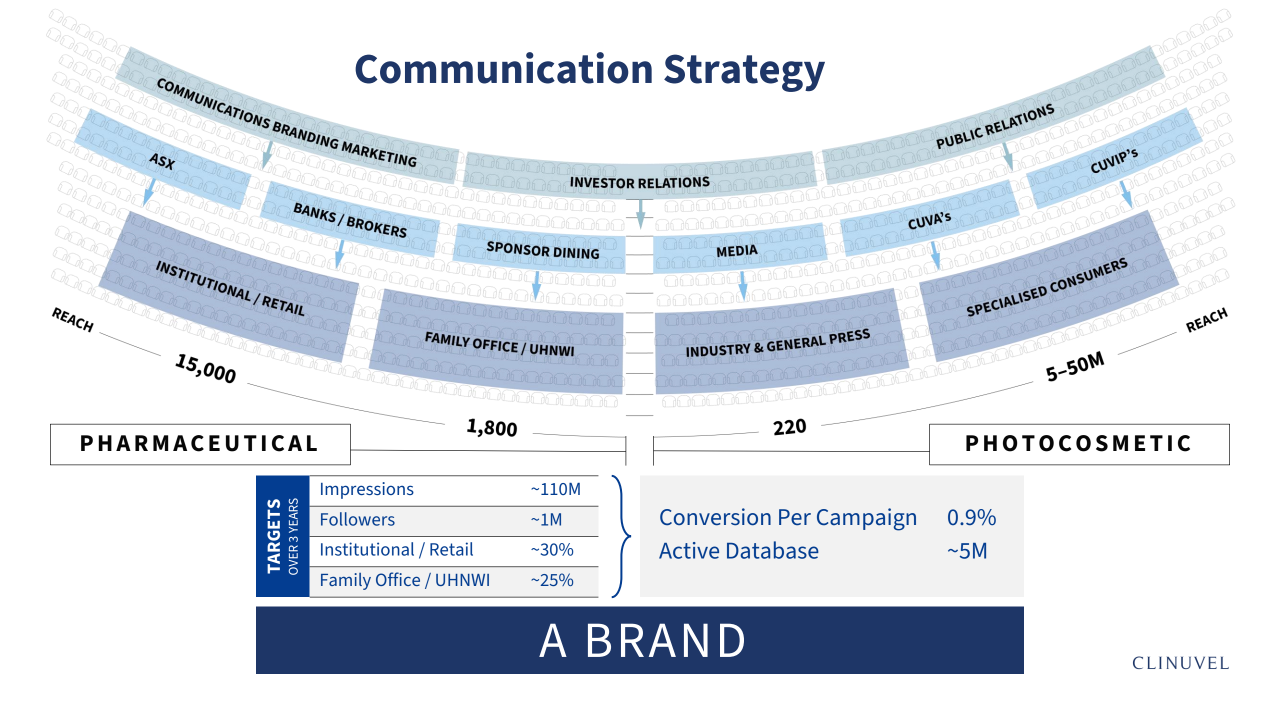
Half Year to December 2023

- revenues up 10% (cf. half yr Dec '22)
- expenses up 28% supporting growth
- NPBT up 1% to A\$14.8 M
- NPAT down 4% to A\$10.9 M [due to increased tax expenses]
- earnings per share A\$0.22

Strong Balance Sheet



AS OF	30 Jun '23	31 Dec '23
Total Assets	\$193.7m	\$211.7m
Total Liabilitiesno debt	\$29.1m	\$33.0m
Net Assets	\$164.6m	\$178.7m
Cash Reserves	\$156.8m	\$174.5m
 cyclical buffer 		
 financing organic growth 		
 share buy-back (12 months 28/03/25) 		\$20m
 acquisition (opportunistic) 		





The Future

MELANOCORTIN HOUSE

TAM>US\$44b | Penetration >US\$2b

TAM US\$300m Porphyria FY23 US\$53m

TAM US\$4.5b Vitiligo US\$490-570m TAM US\$100m **XP** ~US\$50m

TAM US\$1.29b **ACTH**

US\$150m

M&A

TAM US\$31b Stroke US\$1.98-2.23b

TAM US\$6.2b PhotoCosmetics

US\$60m

Objectives CY 2024

1	SCENESSE [®] adolescent outcome EMA	9
2	SCENESSE [®] Canada Health submission	1
3	Vitiligo CUV105 completion recruitment	1
4	Vitiligo CUV107 start recruitment	1
5	XP-DNA Repair CUV151 read out complete (selected markers)	1
6	CUV156 read out complete (selected markers)	1
7	CUV154 start	1
8√	Paediatric PK study CUV052 start – March 2024	1

9√	VP CUV040 complete results – March 2024
10	CNS CUV803 completed with final results
11	CNS and/or New Indication
12	NEURACTHEL [®] manufacturing progress
13√	Website launch
14	PhotoCosmetics E-shop launched
15	CYACÊLLE global launch
16 🗸	Financial growth earnings: half year (Feb 2024), final year end

Q&A with the audience

CLINUVEL Chair and Executives

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Thank you for your attendance

Authorised for ASX release by the Board of Directors of CLINUVEL PHARMACEUTICALS LTD Head of Investor Relations: Mr Malcolm Bull, CLINUVEL PHARMACEUTICALS LTD Investor Enquiries: <u>https://www.clinuvel.com/investors/contact-us</u> Level 22, 535 Bourke Street, Melbourne – Victoria, Australia, 3000 | T+61 3 9660 4900 | F+61 3 9660 4909

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