

28 November 2017

Market Announcements Office Australian Securities Exchange Limited 20 Bridge Street SYDNEY NSW 2000

Dear Sir,

Results of Annual General Meeting CLINUVEL PHARMACEUTICALS LIMITED

In accordance with Listing Rule 3.13.2 and section 251AA of the Corporations Act, we advise details of the resolutions and the proxies received in respect of each resolution are set out in the attached proxy summary.

Yours faithfully,

Darren Keamy

Company Secretary

CLINUVEL PHARMACEUTICALS LIMITED

Annual General Meeting Tuesday, 28 November 2017 Voting Results

The following information is provided in accordance with section 251AA(2) of the Corporations Act 2001 (Cth).

Resolution details		
Resolution	Resolution Type	
1 Adoption of the Remuneration Report	Ordinary	
2 Re-election of Mr Willem Blijdorp	Ordinary	
3 Re-approval of Performance Rights Plan	Ordinary	

Instructions given to validly appointed proxies (as at proxy close)			
For	Against	Proxy's Discretion	Abstain
13,576,915	218,926	275,243	50,926
96.48%	1.56%	1.96%	
14,349,990	12,023	2,866,067	3,392
83.29%	0.07%	16.64%	
12,079,072	1,565,125	275,243	202,570
86.78%	11.24%	1.98%	

ľ	lumber of votes cast on the (where applicable)	e poll
For	Against	Abstain*
14,265,300 92.87%	1,095,772 7.13%	136,206
17,860,738 96.13%	718,671 3.87%	195,273
12,696,389 83.01%	2,598,319 16.99%	202,570

Resolution Result
Carried / Not Carried
Carried
Carried
Carried

^{*} Votes cast by a person who abstains on an item are not counted in calculating the required majority on a poll.

About CLINUVEL PHARMACEUTICALS LIMITED

CLINUVEL PHARMACEUTICALS LTD (ASX: CUV; NASDAQ INTERNATIONAL DESIGNATION ADR: CLVLY; XETRA-DAX: UR9) is a global biopharmaceutical company focused on developing and delivering treatments for patients with a range of severe genetic and skin disorders. As pioneers in understanding the interaction of light and human biology, CLINUVEL's research and development has led to innovative treatments for patient populations with a clinical need for photoprotection and repigmentation. These patient groups range in size from 5,000 to 45 million worldwide. CLINUVEL's lead product, SCENESSE® (afamelanotide 16mg), was approved by the European Commission in 2014 for the prevention of phototoxicity (anaphylactoid reactions and burns) in adult patients with erythropoietic protoporphyria (EPP). More information on EPP can be found at http://www.epp.care.

Headquartered in Melbourne, Australia, CLINUVEL has operations in Europe, Switzerland, the US and Singapore.

For more information go to http://www.clinuvel.com.

SCENESSE® is a registered trademark of CLINUVEL PHARMACEUTICALS LTD.

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Forward-Looking Statements

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, including our ability to develop, manufacture, market and sell biopharmaceutical products; competition for our products, especially SCENESSE® (afamelanotide 16mg); our ability to achieve expected safety and efficacy results 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 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; 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 based products; decisions by regulatory authorities regarding approval of our products as well as their decisions regarding label claims; any failure 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 2017 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 the forecasts and estimates is available on request. Past performance is not an indicator of future performance.

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