

## Kiadis Pharma announces results of Annual General Meeting

**Amsterdam, The Netherlands, June 5, 2018 - Kiadis Pharma N.V. ("Kiadis Pharma" or the "Company") (Euronext Amsterdam and Brussels: KDS)**, a clinical stage biopharmaceutical company developing a T-cell immunotherapy product candidate designed to reduce Graft versus Host Disease (GVHD) and relapse after hematopoietic stem cell transplantations (HSCT), today announces that at its Annual General Meeting held yesterday at 10:00 CEST at the Amsterdam Stock Exchange (Euronext), Beursplein 5, 1012 JW Amsterdam, The Netherlands, all resolutions were duly passed by the shareholders.

Dr. Otto Schwarz, previously Chief Operating Officer at Actelion, was appointed as a member of the Supervisory Board and will replace Mr. Stuart Chapman, who has resigned as a member of the Supervisory Board following the Meeting. The Board would like to thank Mr. Chapman for his contribution to the Company over several years. Mr. Subhanu Saxena, previously Chief Executive Officer of Cipla and member of the Executive Committee at Novartis, was also appointed as a member of the Supervisory Board.

**For more information, please contact:**

**Kiadis Pharma:**

Karl Hård, Head of IR & Communications  
Tel. +31 611 096 298  
[k.hard@kiadis.com](mailto:k.hard@kiadis.com)

**Optimum Strategic Communications:**

Mary Clark, Supriya Mathur, Hollie Vile  
Tel: +44 203 714 1787  
[kiadis@optimumcomms.com](mailto:kiadis@optimumcomms.com)

### **About**

### **Kiadis**

### **Pharma**

Kiadis Pharma's allodepleted T-cell immunotherapy product candidate, given after a haploidentical hematopoietic stem cell transplantation (HSCT), is designed to reduce Graft versus Host Disease (GVHD) and relapse. Single dose Phase 2 data with lead product candidate ATIR101 has demonstrated substantial and clinically relevant improvements over historical observational cohort data for a similar HSCT without ATIR101, and also shows an improvement over the Post-Transplant Cyclophosphamide (PTCy), or Baltimore protocol, data reported in scientific literature. Based on the positive results from the Phase 2 trial, the Company submitted a Marketing Authorization Application (MAA) to the European Medicines Agency (EMA) in April 2017, for approval of ATIR101 across the EU as an adjunctive treatment in HSCT for adult malignant disease. Kiadis Pharma submitted responses to the Day 120 List of Questions in March 2018 and is on track to obtain a CHMP opinion for ATIR101 in Q4 2018 and, if positive, (conditional) approval from the European Commission in Q1 2019, which would allow for a European launch in H2 2019. Kiadis Pharma is conducting a Phase 3 trial with ATIR101 across Europe and North America (head to head against the PTCy/Baltimore protocol). The first patient was enrolled in December 2017.

In September 2017 the U.S. Food and Drug Administration (FDA) granted ATIR101 the Regenerative Medicine Advanced Therapy (RMAT) designation. ATIR101 has been granted Orphan Drug Designations both in the U.S. and Europe.

The Company's shares are listed on Euronext Amsterdam and Brussels under the ticker KDS.

### **Forward**

### **Looking**

### **Statements**

*Certain statements, beliefs and opinions in this press release are forward-looking, which reflect Kiadis Pharma's or, as appropriate, Kiadis Pharma's directors' current expectations and projections about future events. By their nature, forward-looking statements involve a number of risks, uncertainties and assumptions that could cause actual results or events to differ materially from those expressed or implied by the forward-looking statements. These risks, uncertainties and assumptions could adversely affect the outcome and financial effects of the plans and events described herein. A multitude of factors including, but not limited to, changes in demand, competition and technology, can cause actual events, performance or results to differ significantly from any anticipated development. Forward looking statements contained in*

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