Head of Paediatric Medicines European Medicines Agency 30 Churchill Place London E14 5EU United Kingdom paediatrics@ema.europa.eu

To:

Email:

Notification of discontinuation of a paediatric development which is covered by an agreed PIP Decision

Actives substances(s): eculizumab Invented name: Soliris Latest Decision number(s): 1) P/0229/2016 2) P/ 3) P/ 4) P/ Corresponding PIP number(s): 1) EMEA-000876-PIP07-15 2) EMEA-3) EMEA-4) EMEA-Please note that development of the medicinal product above in the [condition(s)/indication(s)]: Prevention of delayed graft function after solid organ transplantation \boxtimes has been discontinued has been suspended/put on long-term hold (with possible re-start at a later time) for the following reason(s): (tick all that apply) (possible) lack of efficacy in adults (possible) lack of efficacy in children (possible) unsatisfactory safety profile in adults (possible) unsatisfactory safety profile in children commercial reasons (please specify:) manufacturing / quality problems other regulatory action (please specify:) (e.g. suspension, revocation of M.A.) 🛛 other reason (please specify: internal decision on development portfolio) Please add a brief description (max 2000 characters) of the reason(s) for the discontinuation / suspension: The development of eculizumab for the prevention of delayed graft function after solid organ transplantation has been discontinued. Name and signature of the PIP contact point: Martine Zimmermann Date: Contact for inquiries from interested parties: **Alexion Paediatric Inquiries** Telephone: 0033 1 47 10 06 06

pip.enquiries.eu@alexion.com