

30 January 2020 EMA/135954/2020 Human Medicines Evaluation Division

Statement indicating compliance with the agreed completed paediatric investigation plan

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|-----------------------------------|---------------------------------|
| Medicinal product | |
| MabThera | |
| rituximab | |
| Pharmaceutical form(s): | See Annex A of the CHMP Opinion |
| Strength(s): | See Annex A |
| Route(s) of administration: | See Annex A |
| Packaging and package | See Annex A |
| size(s): | |
| Number(s)in the Community | See Annex A |
| Register of Medicinal Products: | |

Marketing Authorisation Holder (MAH):

Name and address of the MAH: Roche Registration GmbH

Emil-Barell-Strasse 1 79639 Grenzach-Wyhlen

GERMANY

Procedure

Procedure number: EMEA/H/C/000165/II/0162

Further to the compliance check performed under Article 23 of Regulation (EC) No 1901/2006, and in accordance with Article 28(3) and Article 45(3) of the said Regulation it is concluded that:

- -the development of this product has complied with all measures in the agreed paediatric investigation plan P/0060/2016. All studies in the agreed paediatric investigation plan P/0060/2016 were conducted after the entry into force of that Regulation,
- -the Summary of Product Characteristics reflects the results of studies conducted in compliance with this agreed paediatric investigation plan.

In accordance with Article 23a of Regulation (EC) No 1234/2008, this statement indicating



| compliance with the agreed completed paediatric investigation plan P/0060/2016 is included in the technical dossier. |
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