



EUROPEAN MEDICINES AGENCY
SCIENCE MEDICINES HEALTH

16 October 2017
EMA/PRAC/38618/2017 Rev. 3

Timetable for the procedure

Referral under Article 31 of Directive 2001/83/EC resulting from
pharmacovigilance data

Quinolone and fluoroquinolone containing medicinal products

Procedure no: EMEA/H/A-31/1452

Quinsair EMEA/H/A-31/1452/C/002789/0010

Procedural step:	Date
Notification:	01 February 2017
Start of the procedure (PRAC):	February 2017 PRAC
List of questions:	09 February 2017
Submission of responses:	07 August 2017
Re-start of the procedure:	31 August 2017
Rapporteur/co-rapporteur assessment reports circulated to PRAC and to CHMP ¹ :	08 September 2017
Comments:	15 September 2017
Updated Rapporteur/co-rapporteur assessment reports circulated to PRAC and to CHMP:	21 September 2017
PRAC list of outstanding issues:	06 October 2017
Submission of responses:	28 December 2017

¹ Committee for Medicinal Products for Human Use



Procedural step:	Date
Re-start of the procedure:	11 January 2018
Rapporteur/co-rapporteur joint assessment report(s) circulated to PRAC and to CHMP:	24 January 2018
Comments:	29 January 2018
Updated Rapporteur/co-rapporteur joint assessment report(s) circulated to PRAC and to CHMP	01 February 2018
PRAC recommendation:	February 2018 PRAC