PROCEDURAL STEPS TAKEN AFTER GRANTING THE MARKETING AUTHORISATION

For procedures finalised after 1 September 2003 please refer to module 8B.

- On 13 July 2001, the EMEA issued a notification (I/01) for the following type I variation: extension of the shelf-life of the active substance.
- On 13 July 2001, the EMEA issued a notification (I/02) for the following type I variation: extension of the shelf-life of the finished product. The European Commission adopted a Decision on 8 October 2001.

Subsequent post Marketing Authorisation applications agreed upon are summarised in the table below:

Scope	Application number	Type of modification ¹	Notification/ Opinion issued on ²	Commission Decision Issued/amen ded on
Additional indication	II/03	II	28.05.02	22.08.02
Change in the name of a manufacturer of the medicinal product	I/04	Ι	11.04.02	16.05.02
Change in test procedure of active substance	I/05	I	11.04.02	16.05.02
Change in test procedure for starting material/intermediate used in manufacturing of active substance	I/06	Ι	11.04.02	16.05.02
Change in test procedure of active substance	I/07	I	29.05.02	31.05.02
Change in manufacture of active substance	I/08	Ι	27.06.02	30.07.02
Minor change of manufacturing process	I/09	I	27.06.02	10.07.02
Batch size of active substance	I/10	Ι	27.06.02	10.07.02
Change in test procedure of medicinal product	I/11	Ι	21.06.02	03.07.02
Update of Summary of Product Characteristics and Package Leaflet	II/13	II	21.11.02	27.02.03
Extension of Indication	II/15	II	22.05.03	11.08.03
Change in the batch size of finished product	I/16	Ι	19.03.03	02.04.03
Change in or addition of manufacturing site(s) for part or all of the manufacturing process	I/17	Ι	10.04.03	23.04.03
Change in specification of starting material/intermediate used in manuf. of the active substance	I/18	Ι	03.04.03	08.04.03
Update of or change(s) to the pharmaceutical documentation and Change(s) to the test method(s) and/or specifications for the finished product	II/19	II	24.07.03	28.07.03

¹ In accordance with Commission Regulation (EC) No. 542/95 of 10 March 1995, as amended: I refers to a minor variation (Type I variation); II refers to a major variation (Type II variation); I/II refers to a minor variation following the procedure set out in Article 6, 7 and 8 of the Regulation; X refers to an Annex II application.

T refers to a transfer of a Marketing Authorisation in accordance with Commission Regulation (EC) No 2141/96 of 7 November 1996.

N refers to a notification in accordance with Article 10(3) of Council Directive 92/27/EEC of 31 March 1992.

² For Notifications and Type I variations, the date of entry into force of the change is the EMEA Notification date. The Commission Decision will be amended accordingly.