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PI 56 PHARM 68 **COMPET 385** MI 353 **IND 207** IA 89 **CODEC 748**

COVER NOTE

From:	Secretary-General of the European Commission, signed by Ms Martine DEPREZ, Director	
date of receipt:	eceipt: 27 April 2023	
То:	Ms Thérèse BLANCHET, Secretary-General of the Council of the European Union	
No. Cion doc.:	COM(2023) 231 final	
Subject:	ANNEX to the Proposal for a Regulation of the European Parliament and of the Council on the supplementary protection certificate for medicinal products (recast)	

Delegations will find attached document COM(2023) 231 final.

Encl.: COM(2023) 231 final



EUROPEAN COMMISSION

> Brussels, 27.4.2023 COM(2023) 231 final

ANNEX 3

ANNEX

to the

Proposal for a Regulation of the European Parliament and of the Council

on the supplementary protection certificate for medicinal products (recast)

 $\{ SEC(2023) \ 172 \ final \} - \{ SWD(2023) \ 117 \ final \} - \{ SWD(2023) \ 118 \ final \} - \{ SWD(2023) \ 119 \ final \}$

↓ 2019/933 Art. 1 pt. 6

<u>ANNEX III-Ia</u>

Standard form for notification pursuant to Article 5(2), points (b) and (c).

Tick the appropriate box		 New notification Update of an existing notification
(a)	Name and address of the maker	
(b)	Purpose of making	 Export Storing Export and storing
(c)	Member State in which making is to take place and Member State in which first related act (if any) prior to making is to take place	Member State of making
		(Member State of first related act (if any))
(d)	Number of certificate granted in the Member State of making and number of certificate granted in Member State of first related act (if any) prior to making	Certificate of Member State of making
		(Certificate of Member State of first related act (if any))
(e)	For medicinal products to be exported to third countries, reference number of marketing authorisation, or the equivalent of such authorisation, in each third country of export	