

# Procedure file

Basic information		
DEA - Delegated acts procedure	<a href="#">2022/2819(DEA)</a>	Procedure completed - delegated act enters into force
Labelling requirements for unauthorised investigational and unauthorised auxiliary medicinal products for human use		
Supplementing <a href="#">2012/0192(COD)</a>		
Subject		
4.20.02 Medical research		
4.20.02.06 Clinical practice and experiments		
4.20.04 Pharmaceutical products and industry		

Key players			
European Parliament	Committee responsible	Rapporteur	Appointed
	<b>ENVI</b> <a href="#">Environment, Public Health and Food Safety</a>		

Key events			
06/09/2022	Non-legislative basic document published	<a href="#">C(2022)06240</a>	
06/09/2022	Initial period for examining delegated act 2 month(s)		
14/09/2022	Committee referral announced in Parliament		
15/11/2022	Delegated act not objected by Parliament		

Technical information	
Procedure reference	2022/2819(DEA)
Procedure type	DEA - Delegated acts procedure
Procedure subtype	Examination of delegated act
Stage reached in procedure	Procedure completed - delegated act enters into force
Committee dossier	ENVI/9/10019

Documentation gateway				
Non-legislative basic document		<a href="#">C(2022)06240</a>	06/09/2022	EC