








Procedure file

Basic information	
COD - Ordinary legislative procedure (ex-codecision procedure) Regulation	2022/0216(COD) Awaiting committee decision
Standards of quality and safety for substances of human origin intended for human application	
Subject 4.20.01 Medicine, diseases 4.20.02 Medical research 4.20.02.06 Clinical practice and experiments 4.20.04.02 Safety of blood and transfusion 4.20.05 Health legislation and policy	

Key players			
European Parliament	Committee responsible	Rapporteur	Appointed
	 Environment, Public Health and Food Safety	 COLIN-OESTERLÉ Nathalie	20/10/2022
		Shadow rapporteur	
		 GONZÁLEZ CASARES Nicolás	
		 GLÜCK Andreas	
		 METZ Tilly	
		 KOPCIŃSKA Joanna	
		 KONEČNÁ Kateřina	
Council of the European Union			
European Commission	Commission DG Health and Food Safety	Commissioner KYRIAKIDES Stella	
European Economic and Social Committee			
European Committee of the Regions			

Key events			
14/07/2022	Legislative proposal published	COM(2022)0338	Summary
12/09/2022	Committee referral announced in Parliament, 1st reading		

Technical information	
Procedure reference	2022/0216(COD)

Procedure type	COD - Ordinary legislative procedure (ex-codicedecision procedure)
Procedure subtype	Legislation
Legislative instrument	Regulation
Legal basis	Treaty on the Functioning of the EU TFEU 168-p4
Mandatory consultation of other institutions	European Economic and Social Committee European Committee of the Regions
Stage reached in procedure	Awaiting committee decision
Committee dossier	ENVI/9/09602

Documentation gateway

Legislative proposal	COM(2022)0338	14/07/2022	EC	Summary
Document attached to the procedure	SEC(2022)0304	14/07/2022	EC	
Document attached to the procedure	SWD(2022)0189	14/07/2022	EC	
Document attached to the procedure	SWD(2022)0190	14/07/2022	EC	
Document attached to the procedure	SWD(2022)0191	14/07/2022	EC	
Document attached to the procedure	N9-0084/2022 OJ C 450 28.11.2022, p. 0007	07/09/2022	EDPS	
Economic and Social Committee: opinion, report	CES4815/2022	26/10/2022	ESC	
Committee draft report	PE738.661	18/01/2023	EP	

Standards of quality and safety for substances of human origin intended for human application

PURPOSE: to ensure a high level of health protection for EU citizens and ensure access to safe and effective substances of human origin (blood, tissue and cells) (SoHOs).

PROPOSED ACT: Regulation of the European Parliament and of the Council.

ROLE OF THE EUROPEAN PARLIAMENT: the European Parliament decides in accordance with the ordinary legislative procedure and on an equal footing with the Council.

BACKGROUND: every year, EU patients are treated with 25 million blood transfusions (during emergency surgery, cancer or other care), a million cycles of medically assisted reproduction, over 35 000 transplants of stem cells (mainly for blood cancers) and hundreds of thousands of replacement tissues (e.g., for orthopaedic, skin, cardiac or eye problems).

The EU framework for safety and quality of substances of human origin (SoHOs) has currently three main Directives, respectively for Blood, Tissues and Cells, and Organs, together with implementing legislation.

Although these Directives have harmonised to a certain degree the rules of Member States in the area of safety and quality of blood, tissues and cells, they include a significant number of options and possibilities for Member States to implement the rules they laid down. This results in divergences between national rules, which can create obstacles to cross-border sharing of these substances.

A fundamental revision of those Directives is needed for a robust, transparent, up-to-date and sustainable regulatory framework for these substances, which achieves safety and quality for all parties involved, enhances legal certainty and supports continuous supply, whilst facilitating innovation for the benefit of public health. In addition, the proposal aims to tackle concerns regarding the sufficiency of supply highlighted by the COVID-19 pandemic.

This initiative is part of the EU's ambition to build a stronger European Health Union.

CONTENT: the proposed Regulation aims to establish measures setting high standards of quality and safety for all substances of human origin intended for human applications and for activities related to these substances, in order to ensure a high level of human health protection, in particular for donors of substances of human origin, recipients of substances of human origin and offspring of medically assisted reproduction.

All substances of human origin would be covered, with the exception of solid organs. Human breast milk is one of the new substances covered by the proposal.

More specifically, the proposal provides for measures to:

- ensure safety and quality for patients receiving SoHO therapies and fully protect them from avoidable risks related to substances of human origin;

- ensure safety and quality for donors of substances of human origin and for children born from donated eggs, sperm or embryos;
- empower the EU and its Member States to better prevent and combat future pandemics (surveillance, data analysis, risk assessment, early warning and rapid response);
- facilitate the development of innovative, safe and effective SoHO therapies;
- improve the resilience of European health systems (sufficient supply of human-derived substances) by mitigating the risk of shortages.

Specifically, the proposal:

- improves the protection of patients treated with SoHO (recipients) and offspring from medically assisted reproduction, with standards, and how to implement these standards concerning recipient and offspring protection;
- contains provisions on the competent authorities for substances of human origin, which are responsible for the supervisory activities;
- covers all activities that competent authorities undertake in relation to SoHO entities or registration procedures, with the obligation to maintain a register of SoHO entities and to establish a procedure for their registration;
- describes all general obligations on SoHO entities, namely their registration, the nomination of a Responsible Person if they release substances of human origin for clinical use, as well as obligations regarding the export of SoHOs;
- requires blood and tissue establishments to meet safety and quality standards by following guidelines developed and updated by designated expert bodies such as the European Centre for Disease Prevention and Control (ECDC) and the European Directorate for the Quality of Medicines and Health Care (Council of Europe);
- lays down provisions to ensure the continuity of supply of SoHOs. It covers the obligation for Member States to have national SoHO emergency plans (for SoHOs that are critically important for patients) and the responsibilities of competent authorities and entities regarding supply alerts for critical SoHOs.

A SoHO Coordination Board (SCB) would be established with and for the Member States to support a common implementation of the new Regulation.

The creation of the EU SoHO Platform, to gather all required information, streamline reporting and increase visibility to citizens will give a new impetus to digitalisation.

BUDGETARY IMPLICATIONS: the financial impact of the proposal is estimated at EUR 55.411 million in commitment appropriations for the period 2024-2027. The appropriations will be reallocated within the financial envelope of the EU Health Programme in the Multiannual Financial Framework (MFF) 2021-2027.

Transparency				
GONZÁLEZ CASARES Nicolás	Shadow rapporteur	ENVI	29/09/2022	European Blood Alliance
GONZÁLEZ CASARES Nicolás	Shadow rapporteur	ENVI	11/10/2022	Grifols, S.A.
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	12/10/2022	EUCOPE
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	12/10/2022	Grifols, S.A. Plasma Protein Therapeutics Association Europe, international Association without lucrative purpose
GONZÁLEZ CASARES Nicolás	Shadow rapporteur	ENVI	12/10/2022	EUCOPE
GONZÁLEZ CASARES Nicolás	Shadow rapporteur	ENVI	12/10/2022	PPTA
GONZÁLEZ CASARES Nicolás	Shadow rapporteur	ENVI	12/10/2022	DG SANTE
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	19/10/2022	Commissionner Kyriakides
GONZÁLEZ CASARES Nicolás	Shadow rapporteur	ENVI	19/10/2022	Spanish Permanent Representation
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	24/10/2022	Blood and Beyond
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	25/10/2022	European Blood Alliance

COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	25/10/2022	DG Santé
GONZÁLEZ CASARES Nicolás	Shadow rapporteur	ENVI	25/10/2022	French Blood Establishment
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	26/10/2022	Macopharma
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	26/10/2022	Représentation permanente de la France - chargée Santé
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	27/10/2022	ECDC
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	04/11/2022	IRIS/IPOPI
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	04/11/2022	Pr Mahlaoui - Hôpital Necker
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	04/11/2022	Etablissement français du sang
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	07/11/2022	Représentation permanente de l'Allemagne
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	07/11/2022	Blood Transfusion Association
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	08/11/2022	ESHRE
GONZÁLEZ CASARES Nicolás	Shadow rapporteur	ENVI	08/11/2022	Blood and Beyond
GONZÁLEZ CASARES Nicolás	Shadow rapporteur	ENVI	08/11/2022	Cerus Europe B.V.
GONZÁLEZ CASARES Nicolás	Shadow rapporteur	ENVI	10/11/2022	Alliance of Regenerative Medicine
GONZÁLEZ CASARES Nicolás	Shadow rapporteur	ENVI	10/11/2022	Terumo Blood and Cell Technologies
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	14/11/2022	DG Santé
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	14/11/2022	EPODIN/AFNP
GONZÁLEZ CASARES Nicolás	Shadow rapporteur	ENVI	14/11/2022	DG SANTE
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	15/11/2022	Centre de Transfusion des Armées (CTSA)
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	15/11/2022	LFB
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	15/11/2022	EMA
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	16/11/2022	HERA
GONZÁLEZ CASARES Nicolás	Shadow rapporteur	ENVI	17/11/2022	Dr. Esteve Trias Dr. Silvia Cufi
COLIN-OESTERLÉ	Rapporteur	ENVI	18/11/2022	Agence française de la

Nathalie				Biomédecine
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	23/11/2022	ANSM
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	24/11/2022	Alliance for Regenerative Medicine
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	24/11/2022	International Plasma and Fractionation Association
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	24/11/2022	Présidence tchèque du Conseil
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	28/11/2022	European Sperm Bank
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	30/11/2022	MedTech Europe
GONZÁLEZ CASARES Nicolás	Shadow rapporteur	ENVI	19/12/2022	European Sperm Bank
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	04/01/2023	European Economic and Social Committee
GLÜCK Andreas	Shadow rapporteur	ENVI	10/01/2023	Takeda
GLÜCK Andreas	Shadow rapporteur	ENVI	11/01/2023	IPFA
COLIN-OESTERLÉ Nathalie	Rapporteur	ENVI	12/01/2023	EDQM
GLÜCK Andreas	Shadow rapporteur	ENVI	18/01/2023	Bundesärztekammer
SOLÍS PÉREZ Susana	Member	12/10/2022	Grifols, S.A.	
KYMPUROPOULOS Stelios	Member	12/01/2023	CSL Behring	