


Basic information	
<b>2018/2108(INI)</b> INI - Own-initiative procedure Implementation of the Cross-border Healthcare Directive See also <a href="#">2008/0142(COD)</a> <b>Subject</b> 2.20 Free movement of persons 2.40 Free movement of services, freedom to provide 4.10.10 Social protection, social security 4.20.06 Health services, medical institutions	Procedure completed

Key players				
European Parliament	<b>Committee responsible</b>		<b>Rapporteur</b>	<b>Appointed</b>
	<b>ENVI</b>	Environment, Climate and Food Safety	BELET Ivo (PPE)	16/04/2018
			Shadow rapporteur WÖLKEN Tiemo (S&D) KRUPA Urszula (ECR) RIES Frédérique (ALDE) PEDICINI Piernicola (EFDD) MÉLIN Joëlle (ENF)	
	<b>Committee for opinion</b>		<b>Rapporteur for opinion</b>	<b>Appointed</b>
	<b>EMPL</b>	Employment and Social Affairs (Associated committee)	The committee decided not to give an opinion.	
	<b>IMCO</b>	Internal Market and Consumer Protection (Associated committee)	GRAPINI Maria (S&D)	23/04/2018
European Commission	<b>Commission DG</b>		<b>Commissioner</b>	
	Health and Food Safety		ANDRIUKAITIS Vytenis Povilas	

Key events			
Date	Event	Reference	Summary
14/06/2018	Committee referral announced in Parliament		
14/06/2018	Referral to associated committees announced in Parliament		

22/01/2019	Vote in committee		
29/01/2019	Committee report tabled for plenary	<a href="#">A8-0046/2019</a>	<a href="#">Summary</a>
12/02/2019	Decision by Parliament	<a href="#">T8-0083/2019</a>	<a href="#">Summary</a>
12/02/2019	Results of vote in Parliament		
12/02/2019	Debate in Parliament	<a href="#">CRE link</a>	
12/02/2019	End of procedure in Parliament		

Technical information	
<b>Procedure reference</b>	2018/2108(INI)
<b>Procedure type</b>	INI - Own-initiative procedure
<b>Procedure subtype</b>	Implementation
	See also <a href="#">2008/0142(COD)</a>
<b>Legal basis</b>	Rules of Procedure EP 55
<b>Other legal basis</b>	Rules of Procedure EP 165
<b>Stage reached in procedure</b>	Procedure completed
<b>Committee dossier</b>	ENVI/8/12701

Documentation gateway				
<b>European Parliament</b>				
Document type	Committee	Reference	Date	Summary
Amendments tabled in committee		<a href="#">PE627.772</a>	18/09/2018	
Committee draft report		<a href="#">PE628.580</a>	30/10/2018	
Committee opinion	<a href="#">IMCO</a>	<a href="#">PE623.966</a>	23/11/2018	
Amendments tabled in committee		<a href="#">PE631.903</a>	05/12/2018	
Committee report tabled for plenary, single reading		<a href="#">A8-0046/2019</a>	29/01/2019	<a href="#">Summary</a>
Text adopted by Parliament, single reading		<a href="#">T8-0083/2019</a>	12/02/2019	<a href="#">Summary</a>
<b>European Commission</b>				
Document type		Reference	Date	Summary
Commission response to text adopted in plenary		<a href="#">SP(2019)327</a>	17/07/2019	

## Implementation of the Cross-border Healthcare Directive

2018/2108(INI) - 29/01/2019 - Committee report tabled for plenary, single reading

The Committee on the Environment, Public Health and Food Safety adopted the report by Ivo BELET (EPP, BE) on the implementation of the Cross-Border Healthcare Directive.

### Implementation

Members noted the benefits of the directive in clarifying the rules on cross-border healthcare and in ensuring access to safe and high-quality cross-border healthcare in the Union, as well as for achieving patient mobility in accordance with the case law of the Court of Justice. They expressed disappointment, however, that a significant number of Member States have not effectively implemented the requirements for guaranteeing patients' rights, urging Member States to transpose the Directive correctly in order to ensure high-quality and accessible cross-border healthcare for patients.

The Commission was invited to proceed with its triennial evaluation reports on the operation of the Directive and to factor patient quality of life and care outcomes into its evaluation of the cost-efficiency of the implementation of the Directive, and establish guidelines on implementation.

### **Funding**

Members recalled that while the financing of cross-border healthcare is the responsibility of Member States, the Commission, through its health programmes, supported the cooperation provided for by the Directive.

Concerned about the proposed reduction in funding for the health programme, they called for the programme to be restored as a robust stand-alone programme with increased funding in the next multiannual financial framework (MFF) (2021-2027).

### **Patient mobility**

The report stated that the reasons for low patient mobility are fourfold: i) some Member States were quite late implementing the Directive; ii) citizens' awareness about their general rights to reimbursement is extremely low, iii) certain barriers limiting cross-border healthcare have been erected by some Member States, and iv) information on patients seeking healthcare in another Member State on the basis of the Directive is missing or incomplete.

The Commission and Member States were asked to work together to assess, realign and simplify reimbursement procedures for patients receiving cross-border care, including by clarifying the reimbursement of follow-up care and procedures, and to set up coordinating one-stop-shop front offices at the relevant healthcare insurers. The report urged the Member States to notify the Commission of any decision to introduce limitations regarding reimbursement of costs, giving their reasons for doing so.

### **Information for patients**

The report encouraged Member States and border regions to deepen cross-border healthcare cooperation, in an efficient and financially sustainable manner, including by providing accessible, sufficient and understandable information, in order to secure the best possible care for patients. It also called on the Commission and the Member States to invest further in the development and promotion of accessible and clearly visible National Contact Points (NCPs) and eHealth platforms for patients, which should provide user-friendly, digitally accessible and barrier-free information for patients and health professionals in multiple languages.

### **Rare diseases, rare cancers and European Reference Networks (ERNs)**

The report stressed the importance of EU-wide cooperation in ensuring the efficient pooling of knowledge, information and resources to tackle rare and chronic diseases, including rare cancers, effectively across the EU. It encouraged the Commission, in that regard, to support the setting up of specialised centres for rare diseases in the EU, which should be fully integrated into the ERNs. It also recommended building on the steps already taken to increase public awareness and understanding of rare diseases and rare cancers and to increase funding for R&D.

### **Mutual recognition of (e-)prescriptions**

The report called on Member States and their respective health authorities to address the legal and practical issues that are hindering the mutual recognition of medical prescriptions across the EU. It also called on the Commission to take steps to ensure that prescriptions issued by ERN-linked centres of expertise are accepted for reimbursement in all Member States.

### **eHealth**

Members acknowledged that eHealth can help to ensure that health systems are sustainable, by reducing certain costs, and can be an important part of the EU's response to current healthcare challenges. They welcomed the creation of the EU-wide eHealth Digital Service Infrastructure, which will foster the cross-border exchange of health data, specifically e-prescriptions and patient summaries. They called on the Commission to address the digital health needs in the Member States as a matter of priority.

### **Brexit**

The Commission was asked to negotiate a solid agreement with post-Brexit UK on health, devoting specific attention to cross-border rights for patients and the functioning of the ERNs.

## **Implementation of the Cross-border Healthcare Directive**

2018/2108(INI) - 12/02/2019 - Text adopted by Parliament, single reading

The European Parliament adopted by 512 votes to 32, with 62 abstentions, a resolution on the implementation of the Cross-Border Healthcare Directive.

Parliament noted the benefits of the Directive in clarifying the rules on cross-border healthcare and in ensuring access to safe and high-quality cross-border healthcare in the Union, as well as for achieving patient mobility in accordance with the case law of the Court of Justice. It expressed disappointment, however, that a significant number of Member States have not effectively implemented the requirements for guaranteeing patients' rights, urging Member States to transpose the Directive correctly in order to ensure high-quality and accessible cross-border healthcare for patients. Parliament recognised that specific improvements could be made with regard to access to prescribed medicines and continuity of treatment. It also called on the Commission to explore the possibility of expanding the scope of the Directive to include vaccination programmes, since the Directive does not cover these.

The Commission was invited to proceed with its triennial evaluation reports on the operation of the Directive and to factor patient quality of life and care outcomes into its evaluation of the cost-efficiency of the implementation of the Directive, and establish guidelines on implementation.

### **Funding**

Members recalled that while the financing of cross-border healthcare is the responsibility of Member States, the Commission, through its health programmes, supported the cooperation provided for by the Directive.

Concerned about the proposed reduction in funding for the health programme, they called for the programme to be restored as a robust stand-alone programme with increased funding in the next multiannual financial framework (MFF) (2021-2027).

### ***Patient mobility***

Parliament considered that the reasons for low patient mobility are fourfold:

- some Member States were quite late implementing the Directive;
- citizens' awareness about their general rights to reimbursement is extremely low;
- certain barriers limiting cross-border healthcare have been erected by some Member States;
- information on patients seeking healthcare in another Member State on the basis of the Directive is missing or incomplete.

The Commission and Member States were asked to work together to assess, realign and simplify reimbursement procedures for patients receiving cross-border care, including by clarifying the reimbursement of follow-up care and procedures, and to set up coordinating one-stop-shop front offices at the relevant healthcare insurers. Parliament urged the Member States to notify the Commission of any decision to introduce limitations regarding reimbursement of costs, giving their reasons for doing so.

### ***Border regions***

The resolution encouraged Member States and border regions to deepen cross-border healthcare cooperation, in an efficient and financially sustainable manner, including by providing accessible, sufficient and understandable information, in order to secure the best possible care for patients.

### ***Information for patients***

Parliament called on the Commission and the Member States to invest further in the development and promotion of accessible and clearly visible National Contact Points (NCPs) and eHealth platforms for patients, which should provide user-friendly, digitally accessible and barrier-free information for patients and health professionals in multiple languages.

### ***Rare diseases, rare cancers and European Reference Networks (ERNs)***

Parliament stressed the importance of EU-wide cooperation in ensuring the efficient pooling of knowledge, information and resources to tackle rare and chronic diseases, including rare cancers, effectively across the EU. It encouraged the Commission, in that regard, to support the setting up of specialised centres for rare diseases in the EU, which should be fully integrated into the ERNs. It also recommended building on the steps already taken to increase public awareness and understanding of rare diseases and rare cancers and to increase funding for R&D.

### ***Mutual recognition of (e-)prescriptions***

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### ***EHealth***

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