

Basic information	
<p><b>2008/0126(COD)</b></p> <p>COD - Ordinary legislative procedure (ex-codecision procedure) Regulation</p>	Procedure completed
<p>Supplementary protection certificate for medicinal products. Codification</p> <p>Amended by <a href="#">2018/0161(COD)</a></p> <p><b>Subject</b></p> <p>3.50.01.05 Research specific areas 3.50.16 Industrial property, European patent, Community patent, design and pattern 4.20.04 Pharmaceutical products and industry</p>	

Key players			
European Parliament	<b>Committee responsible</b>	<b>Rapporteur</b>	<b>Appointed</b>
	<span style="border: 1px solid red; padding: 2px;">JURI</span> Legal Affairs	WALLIS Diana (ALDE)	25/06/2008
Council of the European Union	<b>Council configuration</b>	<b>Meetings</b>	<b>Date</b>
	Justice and Home Affairs (JHA)	2936	2009-04-06
European Commission	<b>Commission DG</b>	<b>Commissioner</b>	
	Financial Stability, Financial Services and Capital Markets Union	MCCREEVY Charlie	

Key events			
Date	Event	Reference	Summary
17/06/2008	Legislative proposal published	COM(2008)0369 	Summary
19/06/2008	Committee referral announced in Parliament, 1st reading		
07/10/2008	Vote in committee, 1st reading		Summary
09/10/2008	Committee report tabled for plenary, 1st reading	<a href="#">A6-0385/2008</a>	
21/10/2008	Decision by Parliament, 1st reading	<a href="#">T6-0482/2008</a>	Summary
21/10/2008	Results of vote in Parliament		
06/04/2009	Act adopted by Council after Parliament's 1st reading		
06/05/2009	Final act signed		
06/05/2009	End of procedure in Parliament		

16/06/2009	Final act published in Official Journal		
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Technical information	
Procedure reference	2008/0126(COD)
Procedure type	COD - Ordinary legislative procedure (ex-codecision procedure)
Procedure subtype	Codification
Legislative instrument	Regulation
Amendments and repeals	Amended by <a href="#">2018/0161(COD)</a>
Legal basis	EC Treaty (after Amsterdam) EC 095
Stage reached in procedure	Procedure completed
Committee dossier	JURI/6/64375

Documentation gateway				
<b>European Parliament</b>				
Document type	Committee	Reference	Date	Summary
Committee report tabled for plenary, 1st reading/single reading		<a href="#">A6-0385/2008</a>	09/10/2008	
Text adopted by Parliament, 1st reading/single reading		<a href="#">T6-0482/2008</a>	21/10/2008	<a href="#">Summary</a>
<b>Council of the EU</b>				
Document type	Reference	Date	Summary	
Draft final act	<a href="#">03602/2009/LEX</a>	06/05/2009		
<b>European Commission</b>				
Document type	Reference	Date	Summary	
Legislative proposal	<a href="#">COM(2008)0369</a> 	17/06/2008	<a href="#">Summary</a>	
<b>Other institutions and bodies</b>				
Institution/body	Document type	Reference	Date	Summary
EESC	Economic and Social Committee: opinion, report	<a href="#">CES1510/2008</a>	17/09/2008	

Additional information		
Source	Document	Date
National parliaments	<a href="#">IPEX</a>	

**Final act**

[Regulation 2009/0469](#)  
[OJ L 152 16.06.2009, p. 0001](#)

[Summary](#)

## Supplementary protection certificate for medicinal products. Codification

2008/0126(COD) - 21/10/2008 - Text adopted by Parliament, 1st reading/single reading

The European Parliament adopted, by 654 votes to 14 with 8 abstentions, under 1st reading of the codecision procedure, a legislative resolution approving the proposal for a regulation of the European Parliament and of the Council concerning the supplementary protection certificate for medicinal products (codified version). The report had been tabled for consideration in plenary by Diana **WALLIS** (ALDE, UK) on behalf of the Legal Affairs Committee. The Commission proposal was approved as adapted to the recommendations of the Consultative Working Party of the Legal Services of the European Parliament, the Council and the Commission.

## Supplementary protection certificate for medicinal products. Codification

2008/0126(COD) - 17/06/2008 - Legislative proposal

**PURPOSE:** to codify Council Regulation (EEC) No 1768/92 concerning the creation of a supplementary protection certificate for medicinal products.

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

**CONTENT:** the purpose of this proposal is to undertake a codification of Council Regulation (EEC) No 1768/92 concerning the creation of a supplementary protection certificate for medicinal products. The new Regulation will supersede the various acts incorporated into it. This proposal fully preserves the content of the acts being codified and hence does no more than bring them together with only such formal amendments as are required by the codification exercise itself.

The codification proposal was drawn up on the basis of a preliminary consolidation, in all official languages, of Regulation (EEC) No 1768/92 and the instruments amending it, carried out by the Office for Official Publications of the European Communities, by means of a data-processing system. Where the Articles have been given new numbers, the correlation between the old and the new numbers is shown in a table set out in Annex II to the codified Regulation.

## Supplementary protection certificate for medicinal products. Codification

2008/0126(COD) - 06/05/2009 - Final act

**PURPOSE:** to codify Council Regulation (EEC) No 1768/92 concerning the creation of a supplementary protection certificate for medicinal products.

**LEGISLATIVE ACT:** Regulation (EC) No 469/2009 of the European Parliament and of the Council concerning the supplementary protection certificate for medicinal products

**CONTENT:** the aim of this Regulation is to codify Council Regulation (EEC) No 1768/92 of 18 June 1992 concerning the creation of a supplementary protection certificate for medicinal products. The new Regulation will supersede the various acts incorporated into it. It fully preserves the content of the acts being codified and hence does no more than bring them together with only such formal amendments as are required by the codification exercise itself.

Regulation (EEC) No 1768/92, as amended by the acts listed in Annex I, is repealed.

**ENTRY INTO FORCE:** 6 July 2009.