

Procedure file

Basic information		
CNS - Consultation procedure Decision	1998/0830(CNS)	Procedure lapsed or withdrawn
EEA Agreement: amend. protocol 37 and annex II (technical regulations, standards, testing, certification)		
Subject 2.10.03 Standardisation, EC/EU standards and trade mark, certification, compliance 6.40.01 Relations with EEA/EFTA countries		

Key players			
European Parliament	Committee responsible	Rapporteur	Appointed
	RELA External Economic Relations		
Council of the European Union			

Key events			
25/03/1998	Legislative proposal published	SEC(1998)0434	Summary
10/11/1998	Vote in committee		
16/11/1998	Committee referral announced in Parliament		
18/11/1998	Decision by Parliament	T4-0659/1998	Summary

Technical information	
Procedure reference	1998/0830(CNS)
Procedure type	CNS - Consultation procedure
Procedure subtype	International agreement
Legislative instrument	Decision
Legal basis	EC before Amsterdam E 098; EC before Amsterdam E 000; Rules of Procedure EP 52-p1
Stage reached in procedure	Procedure lapsed or withdrawn
Committee dossier	RELA/4/10545

Documentation gateway					
Legislative proposal		SEC(1998)0434	26/03/1998	EC	Summary
Text adopted by Parliament, 1st reading/single		T4-0659/1998	18/11/1998	EP	Summary

EEA Agreement: amend. protocol 37 and annex II (technical regulations, standards, testing, certification)

OBJECTIVE: the draft decision by the EEA Joint Committee concerns the amendment of Protocol 37 and Annex II to the EEA Agreement relating to technical regulations, standards, testing and certification, with the aim of integrating the Community acquis recently adopted in this field. SUBSTANCE: in order to ensure the requisite legal security and homogeneity, the EEA Joint Committee is to integrate all the relevant Community legislation into the EEA Agreement as soon as possible after its adoption. The draft decision is intended to amend Protocol 37 and Annex II to the EEA Agreement to take account of the Community acquis with regard to authorisation and supervision of medicines for human and veterinary use and the setting-up of a European Agency for the Evaluation of Medicinal Products (the EEA/EFTA States are specifically invited to participate in the Agency). No transition period is provided for.?

EEA Agreement: amend. protocol 37 and annex II (technical regulations, standards, testing, certification)

The European Parliament adopted the simplified procedure amending Protocol 37 and Annex II to the EEA Agreement (procedure without report).?