Declaration of the End of Trial Form (cf. Section 4.2.1 of the Detailed guidance on the request to the competent authorities for authorisation of a clinical trial on a medicinal product for human use, the notification of substantial amendments and the declaration of the end of the  $trial^{l}$ )

NOTIFICATION OF THE END OF A CLINICAL TRIAL OF A MEDICINE FOR HUMAN US
TO THE COMPETENT AUTHORITY AND THE ETHICS COMMITTEE

For off	icial use					
Date of	receipt:	Competent authority registration num				
		Ethics committee registration number	:			
To be	filled in by the applicant					
v	EMBER STATE IN WHICH THE DEC	TI ADATION IS REING MADE.				
A MII	ENIBER STATE IN WINCH THE DEV	CLARATION IS BEING WADE.				
B TR	IAL IDENTIFICATION					
B.1 Eu	draCT number :	()				
	onsor's protocol code number:	()				
	ll title of the trial :					
C AP	PLICANT IDENTIFICATION (please	tick the appropriate box)				
<b>C.1</b>	DECLARATION FOR THE COMPE	TENT AUTHORITY				
C.1.1	Sponsor					
C.1.2	Legal representative of the sponsor					
C.1.3	Person or organisation authorised by the	sponsor to make the application.				
C.1.4	Complete below:					
	Organisation:					
	Name of person to contact : Address :					
	Telephone number :					
	Fax number:					
	E-mail					
0.11.110						
<b>C.2</b>	DECLARATION FOR THE ETHICS	COMMITTEE				
C.2.1	Sponsor					
C.2.2	Legal representative of the sponsor					
C.2.3	Person or organisation authorised by the					
C.2.4	Investigator in charge of the application	* *	_			
•	Co-ordinating investigator (for multicen					
• G 2 5	Principal investigator (for single centre t	rial):				
C.2.5	Complete below:					
	Organisation: Name:					
	Address:					
	Telephone number :					
	Fax number:					
	E-mail:					
D END OF TRIAL						
D.1	Date of the end of the complete trial in	all countries concerned by the trial?				
D.1.1	(YYYY/MM/DD):					
<b>D.2</b>	Is it an early termination? <sup>3</sup>		yes □ no □			

OJ, C82, 30.3.2010, p. 1; hereinafter referred to as 'detailed guidance CT-1'.

According to national legislation.

<sup>&</sup>lt;sup>3</sup> Cf. Section 4.2. of the detailed guidance CT-1.

- D.2.1 If yes, give date (YYYY/MM/DD):
- D.2.2 Briefly describe in an annex (free text):
- D.2.2.1 The justification for early termination of the trial;
- D.2.2.2 Number of patients still receiving treatment at time of early termination in the MS concerned by the declaration and their proposed management;
- D.2.2.3 The consequences of early termination for the evaluation of the results and for overall risk benefit assessment of the investigational medicinal product.

## E SIGNATURE OF THE APPLICANT IN THE MEMBER STATE

E.1	I hereby	confirm	that/confirm	on be	half of	the spor	sor that	(delete	which is	s not	applicable	e):
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- The above information given on this declaration is correct; and
- That the clinical trial summary report will be submitted within the applicable deadlines in accordance with the applicable guidance by the Commission.<sup>4</sup>

accordance with the applicable guidance by the Commission.					
<b>E.2</b>	<b>APPLICANT TO THE COMPETENT AUTHORITY</b> (as stated in C.1)				
E.2.1	Date:				
E.2.2	Signature:				
E.2.3	Print name:				
E.3	<b>APPLICANT TO THE ETHICS COMMITTEE</b> (as stated in C.2):				
E.3.1	Date:				
E.3.2	Signature:				
E.3.3	Print name:				

Section 4.3. of the detailed guidance CT-1.