## F10 FORMAL REQUEST AND PRESCRIPTION FOR HPC, MARROW; HPC, APHERESIS AND/OR MNC, APHERESIS

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PATIENT DATA							
Patient name:							
Patient registry:							
Diagnosis:							
Patient ID:			Patient ID:				
(assigned by patient registry)			(assigned by donor registry)				
Transplant centre:							
Date of birth: (YYYY-MM-DD)	Gender:	Weight: (kg)		CMV:	Blood	d group/RhD:	
DONOR DATA	-						
Donor registry:						ION:	
Donor ID:						I	
GRID:							
Date of hirth:	Candar	Moight. "	<u></u>		Dloor	d aroun /DbD.	
(YYYY-MM-DD)	Gender:	Weight: (kg	į)	CMV:	B1000	d group/RhD:	
Product ships	Product shipping address: Invoice(s) to be sent to:			:			
Institution:			Institution:				
Address:			Address:				
ZIP code:			ZIP code:				
City:			City:				
Country:			Country:				
Attention:			Attention:				
Phone:			Phone:				
Fax:			Fax:				
E-mail: E-mail:							
PRODUCT REQUEST							
HPC, Marrow ONLY			○ HP	C. Marrow	second onti	on: HPC, Apheresis	
HPC, Apheresis ONLY			_		•	otion: HPC, Marrow	
MNC, Apheresis, please specify number of DLI (e.g. 1st, 2nd):							
Reason for product preference			<u></u>				
DONOR PREFERENCE (in case of HPC, Marrow and/or HPC, Apheresis)							
Are any other donors under consideration for donation of behalf of this patient?   Yes   No							
Are any other donors in process of physical examination on behalf of this patient?							
If you have answered yes to either of these questions above, is this donor requested for stem cell collection on this form the preferred donor?							
If no, please explain:							



## F10 FORMAL REQUEST AND PRESCRIPTION FOR HPC, MARROW; HPC, APHERESIS AND/OR MNC, APHERESIS

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PATIENT DATA	
Patient name:	
Patient registry:	
Patient ID: Patient ID:	
(assigned by patient registry) (assigned by donor registry)	
DONOR DATA	
Donor registry:	ION:
Donor ID:	
GRID:	
PROTOCOL DATA please enclose a brief protocol flow chart if applicable	
Products that are included in the protocol and therefore may later be requested:	
Additional HPC, Marrow Additional HPC, Apheresis MNC, Apheresis, please spec	ify number of DLL
Other, please specify:	ny nambor or ben
Total days of conditioning regimen the patient will receive prior to infusion:	
This includes chemotherapy for days, and radiation for days	
	_
TRANSPLANT HISTORY	
Has this patient received any previous stem cell transplants? OYes No	
If yes, please include WMDA Form F20 and answer following transplant histo	ry questions.
List types and dates of previous (allogenic) transplants:	
Specify source of stem cells :	
Reason for subsequent transplant:	
In case the current request is for a MNC apheresis answer the following transplant	history questions:
Did the donor being requested above previously donate stem cells on behalf of this patient?	Yes ○No
Was any of the original stem cell product cryopreserved for later infusion?	○Yes ○No
If yes, was that product infused?	⊝Yes ⊝No
DDEFEDDED DATES (in order of professions)	
PREFERRED DATES (in order of preference)  (First) collection date: (YYYY-MM-DD)  Corresponding infusion date: (YYYY-MM-DD)	ANY MAA DD)
, , , , , , , , , , , , , , , , , , ,	(YY-MIM-DD)
1 1 2	
3	
Minimum number of days prior to collection that donor clearance must be received:	
PICK UP PREFERENCE	
Pick up preference, if one apheresis is sufficient:	
Pick up at the end of the first collection day	
○No pick up preference	
Comments:	
PRE-COLLECTION SAMPLES	
Are pre-collection samples required?	
Sample type: ml heparin ml EDTA ml ACD	
ml no anticoagulant ml other:	



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PATIENT DATA					
Patient name:					
Patient registry:					
Patient ID:		Patient ID:			
(assigned by patient registry)		(assigned by donor registry	<u>()                                    </u>		
DONOR DATA					
Donor registry:				ION:	
Donor ID:				ION.	
GRID:					
GKID:					
PRE-COLLECTION SAMPLES TO BE	SHIPPED TO:				
Institution:					
Attention:					
Address:					
ZIP code:					
City:		Country:			
Phone:		Fax:			
Email:		<u>'</u>			
STEM CELL AND/OR LYMPHOCYT	E COLLECTION				
Product type:					
Cell type:					
Required cells/kg					
x Patient weight (kg)					
= Total number of cells					
+ Cells for quality assurance tes	sting				
= Total number of cells					
Please provide explanation for high	gh number of cells:	Please provide exp	lanation for hi	gh number of cells:	
IRB/Ethics board approval (or equivalent):		IRB/Ethics board approval (or equivalent):			
Date:		Date:			
(YYYY-MM-DD)			(YYYY-MM-DD)		
ADDITIONAL SAMPLES TO ACCOM	/IPANY STEM CELL OR	LYMPHOCYTE PRODUC	Γ		
Peripheral blood samples:					
ml heparin	ml ACD	ml EDTA		ml no anticoagulant	
ml product tube, type:		ml other	r:		
Samples to be taken on collection	day:				
Additional					
comments:					



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Pag	je 4 0ī 4			
PATIENT DATA				
Patient name:				
Patient registry:				
Patient ID:	Patient ID:			
(assigned by patient registry)	(assigned by donor registry)			
DONOR DATA				
Donor registry:		ION:		
Donor ID:		ION.		
GRID:				
OND.				
TRANSPORT DATA				
Product type:	Product type:			
Required anticoagulant:	Required anticoagulant:			
Heparin TEDTA	Heparin EDTA			
ACD	ACD			
Other:	Other:			
Donor plasma required?	Donor plasma required? OYes	s		
If yes, please indicate the desired final concentration:	If yes, please indicate the desired fina	al concentration:		
Transport temperature:	Transport temperature:	,		
Preferred method of overnight	Preferred method of overnight			
storage of product(s) (if needed):	storage of product(s) (if needed):			
Additional instructions:	Additional instructions:			
Additional matractions.	Additional matractions.			
REQUIRED DOCUMENTATION TO ACCOMPANY THIS REQ	NIEST			
In case of HPC, Marrow and/or HPC, Apheresis:	10L31			
WMDA Form F30 Final Compatibility Test Results, or equivalents.	ent			
In case of MNC, Apheresis:				
1. Summary of transplant protocol to be used with the most rec	sent protocol review date			
WMDA Form F20 Transplant History, or equivalent	sent protocorreview date			
DISCLAIMER:				
The cell products collected from this donor are intended solely for the purpos	se of immediate therapeutic treatment for the above m	entioned patient Any		
planned cryopreservation of the cell products prior to initial infusion to the pa	atient may only occur with the advance written approva	from the donor centre .		
<ul> <li>Excess cells may be stored for future therapeutic treatment for this patient. N treatment of the above mentioned patient must be disposed of properly and</li> </ul>		d for the therapeutic		
The donor centre must be provided detailed information concerning the use a		epting these cells, the		
transplant physician also accepts these terms and conditions. Deviations from				
<ul><li>centre.</li><li>Any serious product events and/or adverse reactions must be reported both t</li></ul>	to the donor's registry and transplant center. Correspon	ding S(P)EAR reports must be		
completed by the registry providing the product submitted to the WMDA office		•		

Date: (YYYY-MM-DD)



Person completing form:

Signature: