Donor last name	Donor first name	Donor ID	an_gridformatte
lastname	Firstname		

# **CONSENT FORM FOR BLOOD STEM CELL DONATION (UK)**

The original consent form should be retained by the Collection Centre. One copy should then be retained by the donor and a copy forwarded to Anthony Nolan

## A. STATEMENT BY HEALTHCARE PROFESSIONAL (Please tick the boxes)

I confirm that the donor for whom consent is being taken has identified themselves by confirming their name, date of birth and home address information supplied to me by Anthony Nolan.

I have explained the proposed procedure of peripheral blood stem cell mobilisation and collection to the volunteer donor and briefly discussed the intended benefits to the patient. In particular, I have explained to the donor:

- 1. the use of Granulocyte-Colony Stimulating Factor (G-CSF) to mobilise the donor's stem cells and any serious or potential side effects from this drug
- 2. the need for microbiology testing and in particular the need to test the donor's blood for markers of infection including Syphilis, HIV, HTLV, Hepatitis B, C & E
- **3.** the use of a blood cell separator to collect the donor's stem cells and any serious or potential occurring side effects involved in the procedure
- 4. the potential need to insert a femoral, internal jugular or sub-clavian central venous line if peripheral access is not adequate, as well as any serious or frequently occurring risks associated with such a procedure. I have also explained that separate donor consent for this procedure would be required
- 5. the possible short and long-term risks associated with donating peripheral blood stem cells including:
  - hypocalcaemia (sudden drop of calcium in the bloods) due to the citrate (ACD-A) used in the
    apheresis procedure, which can cause transient paraesthesia (pins and needles, numbness),
    muscle spasms, cramps, and in severe untreated cases risk of seizures (extremely rare). This
    may require calcium tablets or occasionally IV calcium replacement
  - risks associated with G-CSF such as bone ache, myalgia, headache, fatigue, fever, chest pain and thrombocytopenia (low platelets). I have explained these will usually require analgesia(paracetamol)
  - that in extremely rare cases the following G-CSF side effects may occur; vascular event, splenic rupture, sore eyes and anaphylaxis (allergic reaction)
  - bruising and bleeding at the site of venepuncture or central line site
  - the possibility of infection of the venepuncture site
- **6.** To reduce risk of possible exposure to transmissible infections ahead of donation, including unprotected sex with a new or high-risk sexual partner or intravenous drug use, and if such activity occurs to inform Anthony Nolan to facilitate further testing
- 7. the requirement to store confidential information in accordance with applicable data protection and related laws and guidance (see section G below)
- **8.** the possible storage of cells, the need for discard of stored material as well as the possible use of cells for research purposes by the transplant centre (which depending on the circumstances, may be outside of the UK and the EEA) ("the Transplant Centre").
- 9. that a copy of all test results and findings will be sent to the volunteer donor's GP and to Anthony Nolan
- 10. the potential need for cryopreservation should the transplant centre request this for patient safety

Donor last name lastname	Donor first name Firstname	е	Donor ID	an_gridforma	atte
Please tick this box to confirm you Please tick this box to confirm you	•				
provided and can freely give conse		underste	inds the inio	imation	
<ul> <li>I confirm that I have read and understood:         <ul> <li>The current versions of the HTA's Codes of Practice on the Donation of Allogeneic Bone Marrow and Peripheral Blood Stem Cells for Transplantation, and on Consent</li> <li>The current version of the HTA's Guidance for Transplant Teams and Accredited Assessors and have applied the principles and procedures accordingly.</li> </ul> </li> </ul>					
Signed by Healthcare Professional	I	Date of	assessment		
First name		Last nar	ne		
Job title		Collection	on centre		

	r last name name	Donor first name Firstname	Donor ID	an_gridformatte	
B. ST	ATEMENT BY DONOR PRO	OCEDURE INFORMATION	<b>V</b> (Please tick	the boxes)	
l've b to co consi mobi	een told I'm a match for a pa nfirm compatibility, and I've k deration I've voluntarily chos lised peripheral blood stem c umber of stem cells my body	tient in need of a stem cell been asked to donate haen en to donate my cells thro ell collection (PBSC), whicl	transplant. I p natopoietic (bl ugh the proced n involves takil	rovided blood sampl ood) stem cells. Afte dure known as a ng a drug to increase	r
The h	ealthcare professional named	d in section A has clearly ex	xplained to me	2:	
•	the donation procedure, inc and the administration of th				П
•	the possible short and long	-term risks related to the c	collection		
•	that if sexually active to take contracting an infection that			on to reduce the risk	of
•	if I have any new sexual pa via my coordinator	rtners between now and th	e donation, to	inform Anthony Nola	an
oppor	received and understood the tunity to ask questions. Any conjure sufficient information to	questions have been answe	red to my sati	sfaction. I believe I ha	ave
1.	undergo blood tests to asce evidence of important infect & E viruses. I understand the also understand that further Nolan as necessary	tions including those cause at if the results of any of th	d by the Syph ese tests are a	ilis, HIV, HTLV and He abnormal, I will be inf	epatitis B, C ormed. I
2.	receive G-CSF in order to pr	oduce sufficient stem cells	in my circulat	ing blood	
3.	donate stem cells to a patier	nt, collected by the use of	the apheresis r	machine	
Pleas	e tick this box to confirm you	ır agreement with points 1	to <b>3</b> above		
I unde	rstand that:				
4.	There is a possibility I may be to be approached in the future decline a request for a further	ure to discuss and consider			
5.	I may withdraw my consents staff at the donor collection fully understand the life-thre commenced pre-transplant	centre. The basic risks to t eatening implications for th	he patient hav	e been explained to	me and I
6.	Following my cells being information recovery. These tests may in In rare cases these tests may and I may be contacted by A	nclude genetic screening, a y result in findings which m	s well as scree ay be relevant	ning for other blood	disorders.
Pleas	e tick this box to confirm you	ır agreement with points <b>4</b>	to <b>6</b> above		

	r last name name	Donor first name Firstname	Donor ID	an_gridformatte
n add	ition, I understand that:			
7.	I cannot be given a guarante procedure, although the hea			
8.	my recovery will be monitor post-donation, as well as an donation			icipate in routine follow-ups n be at eight and 10 years after
9.	my stem cells will be given t and who may remain anony		nity will be mai	ntained for at least two years,
10.	the patient who receives my world	cells may be of any age, I	race or religion	and be living in any part of the
11.	the primary responsibility fo medical and other professio			G-CSF therapy rests with the
12.	this consent is automatically blood cell separator machin		ot to be fit to d	onate blood stem cells using a
13.	Transplant is carried out in t be cured and may not surviv		ne patient. Sadl	y however, the patient may not

Please tick this box to confirm your agreement with points 7 to 13 above

	or last name name	Donor first name Firstname	Donor ID	an_gridformatte
C.STA	ATEMENT BY DONOR: STO	PRAGE, USE AND DISCA	RD OF CELL	S AT TRANSPLANT CENTRE
I unde	rstand that:			
1.	some of my blood, cells or D at the time of, donation) ma appropriately treat the patie	y be stored for the purpos	es of undertak	cells provided by me prior to, or ing tests to monitor and
2.	a small part of my donation patient after the transplant i		of therapeutic	c cells to be administered to the
3.				ne purposes of quality control future testing relevant to the
4.	my cells will be disposed of, for research, if I have provid disposal of biohazardous ma	led consent), in a manner w		ve unsuitable for clinical use (or oplicable regulations for the
Pleas	e tick this box to confirm you	ur agreement with points 1	to <b>4</b> above	
D. ST.	ATEMENT OF DONOR : CF	RYOPRESERVATION OF	PBSC DONA	TION
	casion, a transplant centre ma patient on a later date. This r			lonated stem cells, to be infused or logistics issues.
In add	ition to consenting to the dor	nation procedure in the ter	ms set above i	in section B:
1.				, and understand that the stem ved for infusion at a later date
2.	If my cells are cryopreserved required or prove unsuitable Anthony Nolan			
3.	If discarded, I understand th for the disposal of biohazard		ropriately acco	ording to applicable regulations
	Please tick this box to confin	rm your agreement with po	oints <b>1</b> to <b>3</b> abo	ove
	OR			

I do not consent to my cells being cryopreserved

Donor last name lastname	Donor first name Firstname	Donor ID	an_gridformatte

#### E. STATEMENT BY DONOR: USE OF CELLS FOR RESEARCH

On occasion, there may be cells remaining in the product bag post-transplant and Anthony Nolan or transplant centres may request to use these remaining cells for research purposes. This may also be the case with the full donation if, for any reason, the transplant cannot take place. In these cases, requests are assessed and approved by a properly constituted research ethics committee and undertaken in accordance with appropriate ethical, legal and professional standards.

I understand that:

- 1. Some or all of my blood, cells or DNA from this collection could be used in a non-identifiable way for future medical research projects. I will not benefit financially from any research undertaken and I waive all rights to any registered patents
- 2. My participation in the storage of my blood, cells or DNA for research is voluntary. Refusal to participate will not affect my status on the Anthony Nolan register as a stem cell donor or result in the loss of any benefits such as follow-up care following my donation
- **3.** My pseudonymised data may be used to support such research and will be used in accordance with the Anthony Nolan Privacy Policy
- 4. I have the right to withdraw consent for the use of my blood, cells or DNA for research without it affecting my status on the Anthony Nolan register as a stem cell donor or resulting in the loss of any benefits, such as follow-up care post-donation. I understand that once my cells have been used for a research study, they will not be able to be withdrawn from that study.

research study, they will not be usic to be withdrawn from that study.	
Please tick this box to confirm your agreement with points 1 to 4 above	
OR	
Please tick this box to confirm that you do not want your blood, cells or DNA to be used for future research	

Donor last name lastname	Donor first name Firstname	Donor ID	an_gridformatte

## F. STATEMENT BY DONOR: ANTHONY NOLAN PATIENT DONOR PROJECT

Anthony Nolan is undertaking a research study that we would like you to consider joining. This study is investigating the importance of HLA matching (tissue typing) and other genetic factors that have been shown to influence the outcome of unrelated stem cell transplants.

Although this research will not directly alter results in this specific transplant, it is hoped that in future it will enable us to advise which donor should be chosen in the event that no fully matched donor is available, but where there is a choice of partially matched donors.

We are asking UK donors and all patients who receive stem cells from a UK donor to join this research project. The DNA extracted from this sample will only be used for matching studies in our laboratory (i.e. only looking for factors to do with outcome in haematopoietic stem cell transplants). It will be stored within the Research Institute, with a unique coding number for the duration of the study (i.e. only the researchers will be able to link the sample to the person who provided it).

After the study is completed, we would like to store the donor/patient sample pairs in a anonymised form (i.e. the details cannot be traced back to an individual person). The purpose of this is to enable us to test these samples for any genetic factors related to stem cell transplantation that may be discovered in years to come. These samples will be owned by Anthony Nolan. All that will be required from you will be a blood and/or a buccal swab sample (mouth swab). If you choose not to join this study, it will not affect your treatment/donation in any way.

I understand the following:

- 1. I have read and fully understood the above information regarding participating in an Anthony Nolan research study.
- 2. I have had the opportunity to ask questions and have received satisfactory answers.
- **3.** my participation is voluntary and if I choose not to provide a blood and/or buccal cell sample (mouth swab), my treatment/donation will not be affected in any way.
- 4. I agree to take part in the study by providing a blood and/or buccal cell sample (mouth swab).
- 5. I agree that my blood and/or buccal cell sample (mouth swab) can be retained after the study completes (in a anonymised form).
- **6.** Anthony Nolan will use and store my personal data in accordance with the Anthony Nolan Privacy Policy and that I may withdraw my consent to the use of my personal data, at any time, in accordance with the terms of this policy.

Please tick this box to confirm your agreement with points 1 to 6 above	
OR	
I do not want to be part of this study	Ш

Donor last name lastname	Donor first name Firstname	Donor ID	an_gridformatte

# G. STATEMENT BY DONOR: PRIVACY

I give my consent to Anthony Nolan processing and storing the following data as per the Anthony N privacy policy (available at <b>anthonynolan.org/privacy</b> ), specifically:	lolan
The data I have provided in this form	
Any analysis of the blood sample I donate, which I understand will be tested for markers of infection including syphilis, HIV, HTLV, and Hepatitis B, C & E	
The results of such blood tests which I specifically consent to Anthony Nolan sharing with my GP	
Any analysis of the stem cells I donate, which I understand may be stored by the transplant centre and/or Anthony Nolan for patient transplant and, if I have agreed, for research purposes	
All health and medical information I provide, which I understand may be stored by the Transplant Centre and Anthony Nolan in order to establish I am medically fit to donate to a patient	
My pseudonymised personal data that may be shared with third party organisations including but not limited to the European Group for Blood and Marrow Transplant registry, to analyse factors that contribute to the outcome of transplants, in accordance with applicable data protection and related laws and guidance	
I consent to Anthony Nolan's transfer of my data (in pseudonymised form) to countries without the same data protection laws as the UK/EU for the purposes stated in the Anthony Nolan privacy policy. Anthony Nolan agrees to protect my data as described in its Privacy Policy and provide adequate protection for transfers to countries outside the UK	
I understand that I have the right to access my medical information in accordance with applicable data protection and related laws and guidance	
Additional statement only relevant to participants in the Anthony Nolan Patient Donor Project:	
Additionally, and only where I have agreed to participate in the research detailed in Section F, I give my consent to Anthony Nolan to use the data provided in this form and a sample of my DNA for the purposes of the research outline at section F above.	

lastname	Firstname	e Donor ID an_gri	arormatte			
H. DONOR AND HEALTH	CARE PROFESSIONAL	DECLARATION				
<b>DONOR</b> I confirm that I have	read and completed part	ts B, C, D, E and F of this form.				
Signed by Donor		Date				
Donor first name		Donor last name				
<b>HEALTHCARE PROFESSION</b> and F of this form.	IAL I confirm that I have v	vitnessed the above donor com	pleting parts B, C, D, E			
Signed by Healthcare Profin section A)	iessional (usually same individual	Date				
Healthcare Professional fir	Healthcare Professional first name  Healthcare Professional last name					
Healthcare Professional tit	le (and email if not the Healthc	are Professional mentioned in section A	)			
I.CONFIRMATION OF CONSENT						
TO BE COMPLETED BY THE DONOR IS ADMITTED FOR T		RE PROFESSIONAL WHEN THE	Ē			
DONOR please tick the relev	ant box					
	her questions and that I w	vish to proceed with stem cell	П			
donation. I confirm that I have not be to this donation.	en coerced, paid, or recei	ved any inducement in relation				
OR						
I withdraw my consent a	nd will not be proceedi	ng				
Signed by Donor		Date				
Donor first name		Donor last name				

HEALTHCARE PROFESSIONAL

Donor last name	Donor first name	Donor ID	an_gridformatte
lastname	Firstname		

Signed by Healthcare Professional	Date
Healthcare Professional first name	Healthcare Professional last name
Job title	Collection centre