

# UK NEQAS Histocompatibility & Immunogenetics

# UK NEQAS H&I

Educational Scheme (iED) Scenario 2: HSCT Scenario Feedback





### **Our iED Schemes**





#### Report deadline: 29th September 2020

Please consider the potential cardiothoracic transplant case detailed below and complete your answers to potential cardiotriviavio transplant asset detailed bases and samp questions 1-5 using a <u>maximum of 40 words for each answer</u>.

A potential cardio-thoracic donor is offered to your centre on 07/01/2020.

The donor is Female, 64 years old and ABO blood group O.

The donor HLA type is: HLA-A2, A11; -827; -Cw1; -DR15, DR103; DR51; -DQ6, DQ7; -DP81\*03-01, DP81\*10:01

The Transplant Co-ordinator asks you to assess the following recipients. All patients have the same clinics

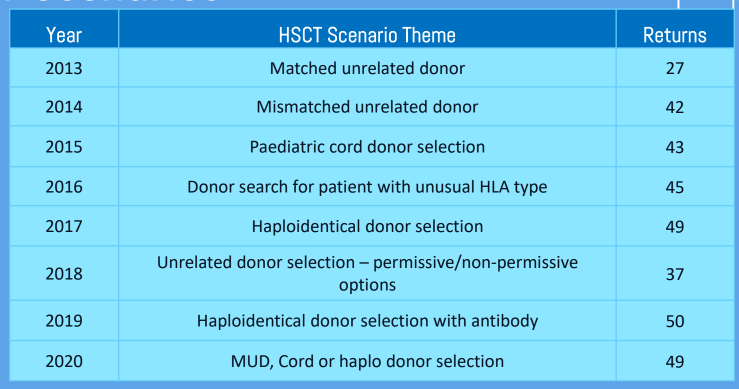
(FI) Date of Last	Control (Deak MFI)				urgency:
	Donor Directed (Peak MFI)	Antibody	Organ Req'd	-	
26/11/2019	Yes (DR15 - 12500)	Positive	Organ Red a	ABO	Recipient
03/01/2020	Yes (DR15 - 12300)	Yes			
27/11/2019	No	Yes	Heart	A	A
	Yes {Cw1 - 1989}		Heart	0	R
2 2 4 2 4 4 2 4 4 4	No		Double Lung	0	C
20/21/20	Yes {B27 - 13716, A2 -		Heart	Α	0
02/01/2020	3095, A11 - 1662)	Yes	Single Lung	0	5
03/01/2020		_		1	E
31/10/2019	2450)		Heart	0	_
	165 (67.5	Yes	Heart	1	F
14/10/2019	105	Yes No Yes Yes Yes	Double Lung Heart Single Lung Heart	0 A 0	B C D E

ank the 3 most suitable recipients based on the information provided and give reasons for the choices made.

1.	Rank the	3 most suitable recip	plents based on the international
	Rank	Recipient G	Reason Click or tap here to enter text.
	2 <sup>nd</sup>	Choose an item	Click or tap here to enter text.
	3 <sup>1d</sup>	Choose an item.	Click or tap here to enter text.

- 3 clinical scenarios a year
  - Solid organ, HSCT, platelet/transfusion
- Based on patient cases
  - Provide relevant clinical details and test results
  - o Questions on interpretation of results and clinical advice
- Not assessed
- Provided free of charge

#### **HSCT Scenarios**





47 Responses

18 from UK and Ireland (UK&I)

29 from the Rest of the World (RoW)



### Case History

A 35 year old male patient suffering from Acute Myeloid Leukaemia (AML) received a haplo-identical transplant from their brother in 2018

Patient's HLA type:

HLA-A\*02:01, A\*03:01; B\*47:01, B\*51:01; C\*06:02, C\*15:13; DRB1\*04:02, DRB1\*14:54;

DRB3\*02:02, DRB4\*01:03; DQB1\*03:02, DQB1\*05:03; DPB1\*04:01, DPB1\*20:01



HLA-A\*02:01, A\*-; B\*51:01, B\*-; C\*14:02, C\*15:13; DRB1\*04:02, DRB1\*04:03;

DRB4\*01:03, DRB4\*-; DQB1\*03:02, DQB1\*-; DPB1\*04:01, DPB1\*-



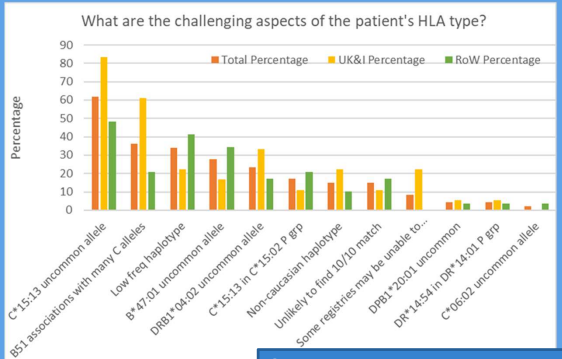




• What aspects of the patient's HLA type make this a challenging unrelated donor search?

Reason	Т	otal	U	K&I	R	oW
	Number	%	Number	%	Number	%
C*15:13 uncommon allele	29	62	15	83	14	48
B51 associations with many C alleles	17	36	11	61	6	21
Low frequency haplotype	16	34	4	22	12	41
B*47:01 uncommon allele	13	28	3	17	10	34
DRB1*04:02 uncommon allele	11	23	6	33	5	17
C*15:13 in C*15:02 P grp	8	17	2	11	6	21
Non-Caucasian haplotype	7	15	4	22	3	10
Unlikely to find 10/10 match	7	15	2	11	5	17
Some registries may be unable to distinguish DR*14:01/DR*14:54	4	9	4	22	0	0
DPB1*20:01 uncommon	2	4	1	6	1	3
DR*14:54 in DR*14:01 P grp	2	4	1	6	1	3
C*06:02 uncommon allele	1	2	0	0	1	3

## Q1: Challenging HLA Type?





#### **Comments:**

B47 is generally low frequency in our local population. The highest known frequencies are found in Central and Western Africa. This could mean it is challenging to find a 10/10 donor for this patient.





#### **Further information**

In November 2020, a post-transplant peripheral blood sample was received in the laboratory for HLA typing with suspected relapse.

HLA typing of the patient was performed post-transplant on a peripheral blood sample using next generation sequencing:

HLA-A\*02:01, A\*-; B\*51:01, B\*-; C\*15:13, C\*-;

DRB1\*04:02, DRB1\*-; DRB4\*01:03, DRB4\*-; DQB1\*03:02, DQB1\*-





### **Further information**

#### Patient's original HLA type:

HLA-A\*02:01, A\*03:01; B\*47:01, B\*51:01; C\*06:02, C\*15:13; DRB1\*04:02, DRB1\*14:54;

DRB3\*02:02, DRB4\*01:03; DQB1\*03:02, DQB1\*05:03

#### Latest HLA type:

HLA-A\*02:01, A\*-; B\*51:01, B\*-; C\*15:13, C\*-;

DRB1\*04:02, DRB1\*-; DRB4\*01:03, DRB4\*-; DQB1\*03:02, DQB1\*-



## Q2: What might these results indicate?



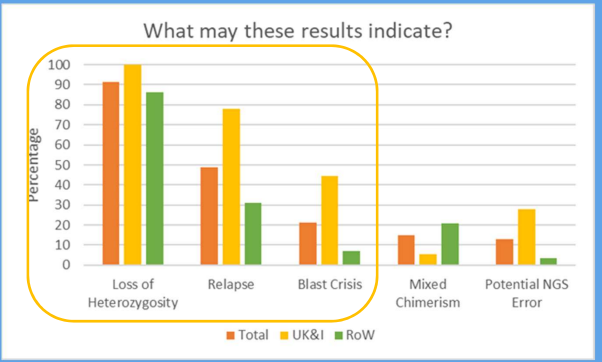
Reason	Tota	Total		d	RoW		
	Number	%	Number	%	Number	%	
Loss of Heterozygosity	43	91	18	100	25	86	
Relapse	23	49	14	78	9	31	
Blast Crisis	10	21	8	44	2	7	
Mixed Chimerism	7	15	1	6	6	21	
Potential NGS Error	6	13	5	28	1	3	



#### Q2: What might these results indicate?

#### **Comments:**

The results indicate a possible loss of heterozygosity (LoH) post-transplant with the loss of mismatched haplotype as part of GVL immune evasion by leukemic stem cells. It would be good practice to request a buccal swab sample from the patient to confirm LoH. If this is the case you would expect the HLA type from the buccal swab to correspond to pre-transplant HLA type. Also, information on proportion of blast cells can be used to confirm LoH.

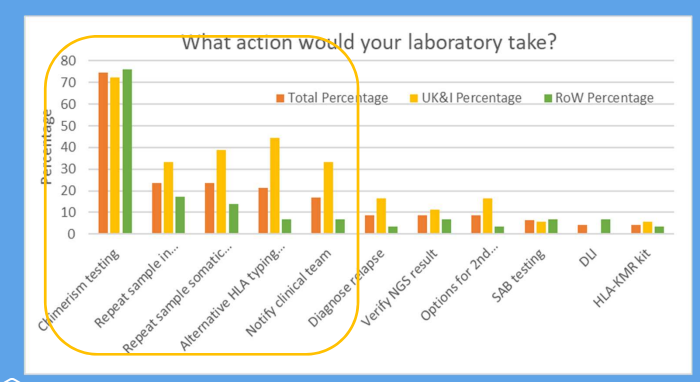


## Q2: What action would your lab take?

Reason	Total		UK8	d	Ro	W
	Number	%	Number	%	Number	%
Chimerism testing	35	74	13	72	22	76
Repeat sample in remission	11	23	6	33	5	17
Repeat sample somatic cells	11	23	7	39	4	14
Use alternative HLA typing method	10	21	8	44	2	7
Notify clinical team	8	17	6	33	2	7
Diagnose relapse	4	9	3	17	1	3
Verify NGS result	4	9	2	11	2	7
Investigate options for 2nd	4	9	3	17	1	3
transplant						
Perform single antigen bead testing	3	6	1	6	2	7
Donor lymphocyte infusion	2	4	0	0	2	7
HLA-KMR kit	2	4	1	6	1	3



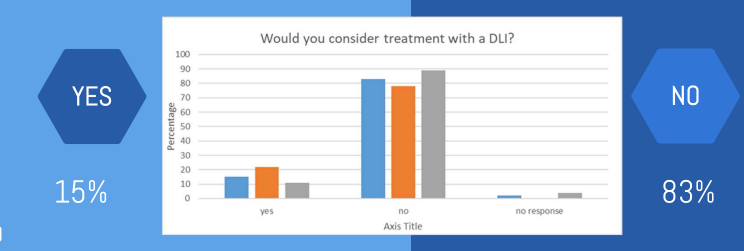
#### Q2: What action would your lab take?



## Q3: Treat patient with DLI?



Post-transplant chimerism results for the patient showed 20% donor cells in the whole blood samples and 95% donor cells in the T cell sample.



## Q3: Treat patient with DLI?

Decision	Reasons	Total		Total UK&I		RoW				
		Number	%	Number	%	Number	%			
Yes	To increase GvL effect	7	15	4	22	3	11			
	Aim to achieve complete chimerism									
	To avoid relapse						J			
	More useful for graft failure than active relapse									
	May be less effective as no mm for donor cells to target									
	Considered for all relapse patients unless have GvHD									
No	Increased risk of GvHD	39	83	14	78	25	89			
	No benefit - no mismatch for donor cells to target									
	A second allograft from another donor with mismatched haplotype should be considered over DLI (EBMT Guidelines)									
	Relapse is extensive so unlikely to respond to DLI									
	95% of T cells are donor derived so no benefit of DLI									
	DLI only indicated if the lymphocyte fraction low									
	Relapse in myeloid line									
	Patient could benefit from NK cell cellular immunotherapy									
	Decision made by clinical team									
No response	We do not give advice to clinicians regarding post-transplant treatment	1	2	0	0	1	4			



#### **Comments:**

An attempt to induce remission by the infusion of donor T-lymphocytes (DLI) would be expected to be ineffective against the leukemic cells due to the LoH and therefore mismatched HLA target antigens. A DLI may also potentially be harmful to the patient due to the risk of inducing GvHD.

### **Further information**

The patient is treated with chemotherapy and a second haplo-identical transplant is planned.

HLA typing reports from four further siblings are received from the family's country of origin:

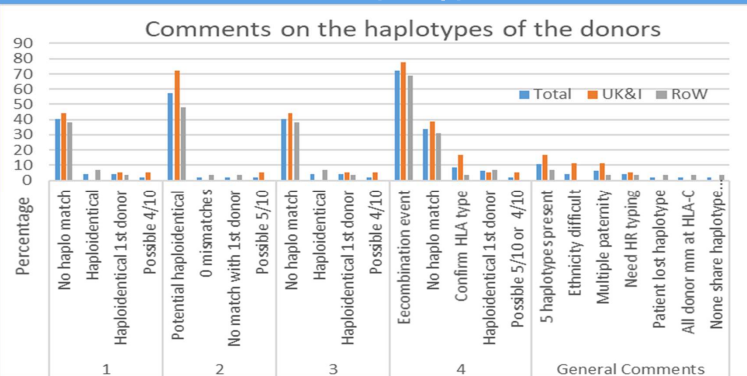
ID	Gender	Age	HLA Type
Patient	Male	35	A*02:01, B*51:01, C*15:13, DRB1*04:02, DQB1*03:02
			A*03:01, B*47:01, C*06:02, DRB1*14:54, DQB1*05:03
1 <sup>st</sup> Donor (Brother)	Male	50	A*02:01, B*51:01, C*15:13, DRB1*04:02, DQB1*03:02 A*02:01, B*51:01, C*14:02, DRB1*04:03, DQB1*03:02
Donor 1	Male	50	A*02, B*51, C*14, DRB1*04, DQB1*03 A*24, B*38, C*12, DRB1*13, DQB1*06
Donor 2	Female	46	A*03, B*47, C*06, DRB1*14, DQB1*05:03
			A*24, B*38, C*12, DRB1*13, DQB1*06
Donor 3	Female	47	A*02, B*51, C*14, DRB1*04, DQB1*03 A*24, B*38, C*12, DRB1*13, DQB1*06
Donor 4	Female	49	A*02, B*38, C*12, DRB1*13; DQB1*06 A*02, B*51, C*14, DRB1*04, DQB1*03

## Q4: Comment on the haplotypes of the donors?

Donor	Comments	Total		UK&	d	RoW	1
		Number	%	Number	%	Number	%
1	Does not share a haplotype with patient	19	40	8	44	11	38
	Haploidentical	2	4	0	0	2	7
	Shares haplotype with 1st donor	2	4	1	6	1	3
	Possible 4/10	1	2	1	6	0	0
2	Potentially shares a haplotype with patient	27	57	13	72	14	48
	0 mismatches	1	2	0	0	1	3
	No match with 1st donor	1	2	0	0	1	3
	Possible 5/10	1	2	1	6	0	0
3	Does not share a haplotype with patient	19	40	8	44	11	38
	Haploidentical	2	4	0	0	2	7
	Shares haplotype with 1st donor	2	4	1	6	1	3
	Possible 4/10	1	2	1	6	0	0
4	Potential recombination event HLA-A	34	72	14	78	20	69
	Does not share a haplotype with patient	16	34	7	39	9	31
	Confirm HLA type	4	9	3	17	1	3
	Haploidentical with 1st donor	3	6	1	6	2	7
	Possible 5/10 HvG or 4/10 GvH direction	1	2	1	6	0	0
General	5 haplotypes present	5	11	3	17	2	7
Comments	Ethnicity difficult to define	2	4	2	11	0	0
	Potential multiple paternity	3	6	2	11	1	3
	Need high resolution typing	2	4	1	6	1	3
	Patient lost haplotype	1	2	0	0	1	3
	All donor mm at HLA-C with patient post-tx	1	2	0	0	1	3
	None share haplotype between patient and 1st donor	1	2	0	0	1	3



#### Q4: Comment on the haplotypes of the donors?





Donor 2 is the only haplo-identical donor.

Donor 4 has a HLA-A\*02 which could be attributed to a recombination event based on segregation of haplotypes.

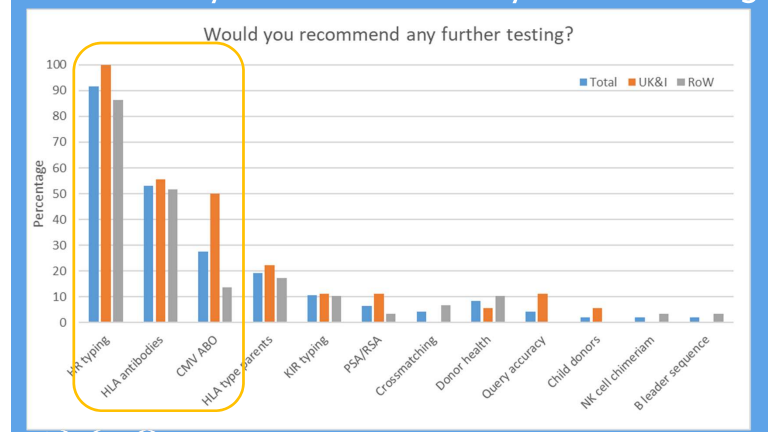


## Q4: Would you recommend any further testing?



	Further Testing	Tota		UK&	l	RoW	1
		Number	%	Number	%	Number	%
<b>\</b>	High resolution typing of donors	43	91	18	100	25	86
<b>\</b>	Test patient for HLA antibodies	25	53	10	56	15	52
	CMV and/or Blood Group	13	28	9	50	4	14
	HLA type parents	9	19	4	22	5	17
	KIR typing	5	11	2	11	3	10
	Test donors for patient specific HLA antibodies	3	6	2	11	1	3
	Crossmatching	2	4	0	0	2	7
	Donor health / pregnancy info	4	9	1	6	3	10
	Query accuracy of donor HLA typing	2	4	2	11	0	0
	Enquire if patient has children able to donate	1	2	1	6	0	0
	NK cell chimerism	1	2	0	0	1	3
	B leader sequence of donor and patient	1	2	0	0	1	3

#### Q4: Would you recommend any further testing?



#### **Comments:**

Further tests could include verification of the donor HLA type including DP genotyping.

## Q4: Rank the donors

Choice	Donor	Reason	Tota	l	UK&	l	RoW	1		
			Number	%	Number	%	Number	%		
First	2	Potential shared haplotype	38	81	15	83	23	79		
		Mismatch haplotype different to lost haplotype								
		Possible enhanced GvL effect								
		Youngest donor								
	1	Male donor	5	5	5 <b>11</b>	.1 1	6	4	4	14
		Haploidentical								
		Likely less sensitising events								
	4	Closest match to first donor	2	4	0	0	2	7		
		Least mismatches of all donors								
	None	Wait until NGS typing performed	1	2	1	6	0	0		

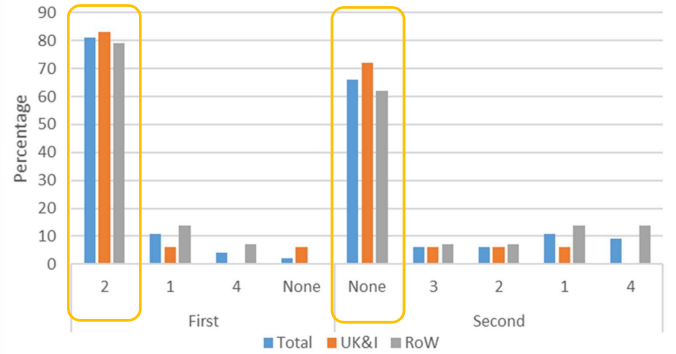
## Q4: Rank the donors



Second	None	No other haploidentical donors	31	66	13	72	18	62
		All other donors HLA-C mismatched						
		Re-graft original donor						
		Use matched unrelated donor						
		Wait until NGS typing performed						
		Don't use original donor due to LoH						
	3	Haploidentical	3	3 <b>6</b>	1	6	2	7
		Younger than Donor 4						
		Avoid donor-recipient mismatch linked with GvHD	-					
	2	Best HLA match - one mismatch at HLA-C	3	3 <b>6</b>	1	6	2	7
		Youngest donor						
	1	Male donor	5	11	1	6	4	14
		Antigen match						
		May get NK alloreactivity						
	4	Possibility of NK cell reactivity	4	9	0	0	4	14
		Almost haploidentical						

## Q4: Rank the donors





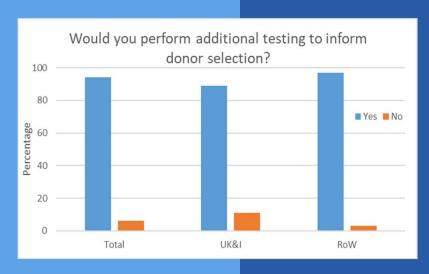


#### **Comments:**

We suggest that Donor 2 would be the preferred option as this donor is haplo-identical. The mismatched haplotype differs to the haplotype lost by the leukaemic cells signifying a potential graft versus leukaemic (GvL) effect. This donor is also the youngest sibling.

# Q5: Additional testing required?





YES

94%

NO

6%

# Q5: Additional testing required?



Decision	Reason	Tota	I	UK&	.I	RoW	/
		Number	%	Number	%	Number	%
Yes	High resolution HLA genotype / family pedigree	44	94	16	89	28	97
	KIR genotyping						
	ABO and/or CMV testing						
	Screen patient for HLA antibodies						
	Sensitisation history including pregnancies						
	Medical fitness incl. IDM, weight						
	Crossmatching if DSA present						
	Chimerism testing						
	Perform search for an unrelated donor						
	Test donor for patient specific HLA antibodies						
	Check B leader sequences						
	Consider NK cell immunotherapy						
No	Donor 2 the only acceptable related donor	3	6	2	11	1	3
	Due to limits donor options additional testing of limited benefit						

#### **Comments:**

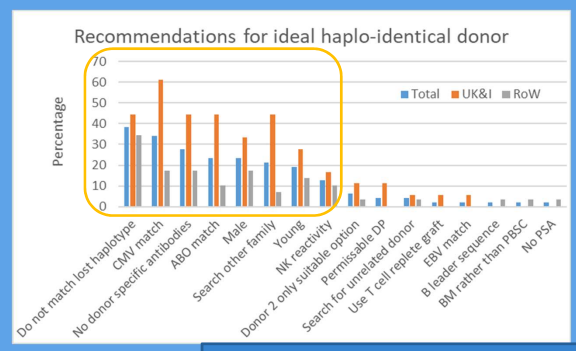
Further tested is recommended. This could include testing for HLA antibodies, high resolution HLA typing including HLA-DPB1, verification typing and KIR typing.

# Q6: Recommendations for ideal haplo-identical donor?

Reason	Total		UK&I		RoW	
	Number	%	Number	%	Number	%
New haploidentical donor should not match the lost haplotype	18	38	8	44	10	34
CMV match	16	34	11	61	5	17
No donor specific antibodies	13	28	8	44	5	17
ABO match	11	23	8	44	3	10
Male	11	23	6	33	5	17
Search other family members (NIMA or NIPA)	10	21	8	44	2	7
Young	9	19	5	28	4	14
NK reactivity	6	13	3	17	3	10
Donor 2 only suitable option	3	6	2	11	1	3
Permissable DP	2	4	2	11	0	0
Search for unrelated donor	2	4	1	6	1	3
Use T cell replete graft for GvL effect	1	2	1	6	0	0
EBV match	1	2	1	6	0	0
B leader sequence compatible	1	2	0	0	1	3
BM rather than PBSC	1	2	0	0	1	3
No patient specific HLA antibodies	1	2	0	0	1	3

# Q6: Recommendations for ideal haplo-identical donor?





#### **Comments:**

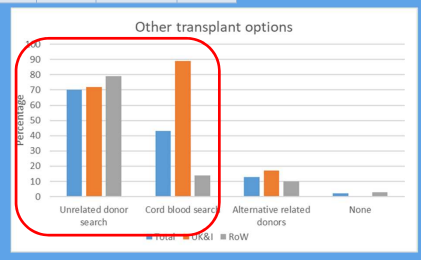
We would recommend a second transplantation from a different HLA-haploidentical donor, selected for being mismatched against the HLA haplotype retained by leukaemic blasts.

# Q7: Any other transplant options?

|--|

Option	Total		UK8	ķΙ	RoW		
	Number	%	Number	%	Number	%	
Unrelated donor search	33	70	13	72	23	79	
Cord blood search	20	43	16	89	4	14	
Alternative related donors	6	13	3	17	3	10	
None	1	2	0	0	1	3	



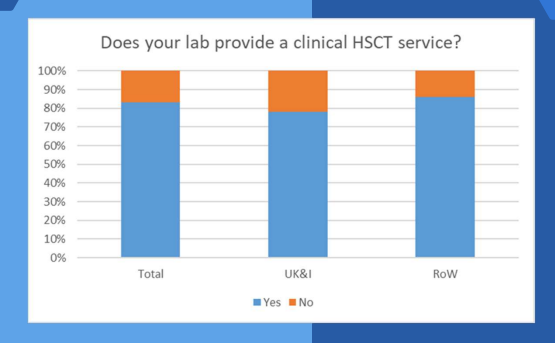


# Q8: Does your lab provide a clinical HSCT service?



YES

83%



NO

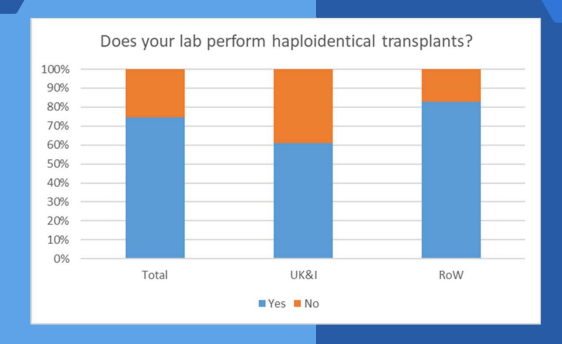
17%

# Q9: Does your lab routinely perform haplo-identical transplants?



YES

74%



NO

26%

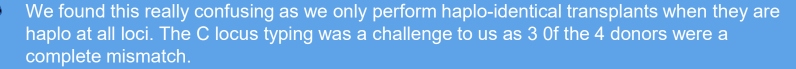
#### **Further Comments**

- Are they all full siblings? There is a large age gap >10 years between patient and siblings, who
  are all of a similar age. If parents available would request samples to confirm family haplotypes.
- We only assist for HLA typing, Ab detection, KIR genotyping and time to time for donor selection (depending on the responsible physician). The HSCT performed within the Haematology clinics.
- While we don't perform transplants ourselves, or make decisions regarding donor selections, we
  provide HLA typing, antibody screening and clinical data for transplantation. We don't perform all
  post-transplant monitoring, but we do perform post-transplant HLA antibody monitoring as well as
  platelet transfusion support. We do a lot of testing for haplo-identical HSCT transplants, but only
  for a few hospitals.
- We do not have experience of allogenic bone marrow transplant. We only perform familial HLA typing study.
- Our laboratory does not perform chimerism and does not have any physician or contact with patients. Both are done by the hospital.
- We do not perform chimerism in our lab and do not give advice to clinicians regarding posttransplant treatment.

#### **Further Comments**



- Aware of the potential use of NK cells in haplo-transplantation, and their use in triggering graft vs leukaemia effect, in AML. However, this would not be undertaken in our routine setting.
- Haplo-identical transplants are performed at this centre. Haplo-identical transplant would be the option selected when the patient does not have a suitable matched sibling or 10/10 unrelated donor.
- In recent years, haplo-identical transplants are being used more frequently, but they are not the preferred transplant option if a suitable HLA matched related, HLA matched unrelated or 9/10 unrelated donor is available.





## Patient Follow Up



WMDA search revealed no 10/10 matched unrelated donors available.

HLA antibody screening of the patient revealed he was negative for the present of donor specific antibodies against the potential sibling donors.

Following chemotherapy to reduce disease burden, he was transplanted with Donor 2 (female, 46 years old) who was chosen to harness the potential GVL effect against HLA retained by the leukaemic cells which experienced LoH to prevent another relapse.

The patient had full donor chimerism 100 days post-transplant.





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