## Consent to Procedure or Surgical Treatment

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Surgical Treatment	DOB		
Part A - To be completed by Treating Medical Practitioner			
Interpreter used: ☐ Not required ☐ Yes If Yes, w	hich language was the information translated to:		
Interpreter service used:			
Has anyone been appointed as a person responsibl	e¹ for this patient's care? ☐ No ☐ Yes		
If Yes, please specify who:			
Does the patient have any written requests / requirements / instructions relating to their care²? ☐ No ☐ Yes			
If Yes, please specify:			
Where applicable, all written requests / requiremen	nts / instructions relating to the patient's care must be sighted		
A person responsible may include an appointed medical treatmen power to make medical treatment decisions appointed under the	t decision maker under the Medical Treatment Planning and Decisions Act 2016 (Vic) or a guardian with Guardianship and Administration Act 2019 (Vic)		
2. Advanced Care Directives including any Instructional directives or	r Values directives under the Medical Treatment Planning and Decisions Act 2016 (Vic) or similar document		
Description of the procedure or surgery, noting of	correct side / correct site. List all anticipated procedures or treatments		

The risks of this procedure / treatment have been discussed with the patient and these include:

(including the expected transfusion of blood or blood products and any possible secondary procedures).

- Infection
- Bruising or bleeding
- Pain / swelling / scars
- Risks associated with anaesthesia / sedation (if applicable you will have the opportunity to discuss these in more detail prior to your procedure)
- Other (please specify including any risks specific to the patient):

Signature of MEDICAL PRACTITIONER

## To be completed by Treating Medical Practitioners

I have explained the nature and purpose of the procedure / treatment detailed above and what it entails for the patient, the known benefits and risks of the procedure / treatment, the risks of not having the procedure / treatment, and the alternatives to having the procedure / treatment.

Full Name:	Name of <b>MEDICAL PRACTITIONER</b>	
Signature:		Date of signature: DD/MM/YYY

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## Consent to Procedure or **Surgical Treatment**

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## Part B - To be completed by Patient or Person Responsible

I consent to the procedure and associated treatments as detailed in **Part A** of this form (overleaf).

In providing my consent to treatment, I acknowledge that the treatment has been explained to me by my treating medical practitioner and that I understand and have had the opportunity to ask questions about:

- The need for the procedure(s) or treatment(s);
- The known expected benefits and possible risks of the procedure(s) or treatment(s), including any risks specific to me;
- Alternative treatment options available and the expected benefits and possible risks of not having this procedure or treatment.

I further understand and agree to the following:

- My health information will be collected and used in accordance with Cabrini's Privacy Policy and applicable privacy laws.
- I may need additional procedure(s) / treatment(s) as is necessary in the reasonable opinion of my treating medical practitioner to preserve my health or life. This may include the transfusion of blood or blood products. \*If you refuse the emergency / life-saving transfusion of blood or blood products, please notify your treating doctor so that your preferences can be documented.
- If a staff member is exposed to my blood, a sample of my blood may be collected and tested for infectious diseases and that I will be informed of the test and results.
- Clinical information, including clinical photography / videography, blood or tissue specimens, may be collected during my procedure or treatment for diagnostic and treatment purposes. Additionally, this clinical information may be de-identified and

' '	oose of education and / or ethically approved research. *If you do r ate your preferences below and notify your treating doctor:	not want your clinical info	rmation used in this
☐ I DO NOT conser	nt to my de-identified clinical information being used for the purpo	ose of education.	
☐ I DO NOT conser	nt to my de-identified clinical information being used for the purpo	ose of ethically approved r	esearch.
Full Name: Signature:	Name of <b>PATIENT</b> or parent / guardian / person responsible  Signature of <b>PATIENT</b> or parent / guardian / person responsible	Date of signature:	DD/MM/YYYY
Part C - Consent to the transfusion of Blood and Blood Products			

(To be completed by Patient or Person Responsible, where indicated as appropriate by the treating Medical Practitioner. Strike out Part C if not applicable.)

	_ have discussed	l the following with m	ıy doctor:
Name of <b>PATIENT</b> or parent / guardian / person responsible	_	J	•

- The likelihood that I require / may require a transfusion of blood or blood products in association with this treatment
- The reason(s) why I require / may require a transfusion of blood or blood products and the type of blood components and / or products required
- The general risks and benefits of receiving / not receiving blood or blood products
- The alternative treatments to having a blood transfusion and alternative blood management strategies

I have also been provided with written information about blood transfusions and have had the opportunity to ask my doctor any questions.

I understand the information provided to me and consent to the transfusion of blood or blood products as required, in association with the procedure / treatment outlined in Part A of this form (overleaf)

with the pr	occadicy treatment outlined in the A of this form (overlear).		
Signature:		Date of signature:	DD/MM/YYYY
	Signature of <b>PATIENT</b> or parent / guardian / person responsible		

 $^st$  If you REFUSE to consent to the transfusion of blood or blood products (or specific products) please discuss this with your treating doctor so that your preferences can be documented

