My infor	mation						
Name:							
Date of birth		DD / MM / YYYY					
Phone number	r:						
Emergency co	ntact (name):						
Emergency co (phone numbe							
NHS number:							
My treat			ır doc	tor to (	do it.		
Start date:	D/MM/YYY	Y En	d dat	e: D	D <b>/</b> MN	//	
It is important to may interfere will least 6 months at	th the indirect an iter the last infusi	tiglobulin tes on.	st (indir	rect Cod	ombs tes	st) for	at
Blood type:							
А	В	AB	0		Rh+		Rh-
The result of m	Positive fo			ect Co	ombs t	est) v	was:
Other							
Contact detail where blood t		med:					
My haer In case of en contact my h Haematolog	nergency, or inaematologis	if you find	d this	card,	pleas	9	
Haematolog	ist's phone n	umber:					
Name of hos	pital:						
MAT-GB-200000 Date of prep: Mo			N	IHRA ap	oproval:	Marc	h 2025



This material fulfills the conditions of the marketing authorisation and has been approved by the Medicines and Healthcare products Regulatory Agency (MHRA).

## SARCLISA

(isatuximab)

## Patient card

## For patients receiving isatuximab

- Provide this card to healthcare providers before blood transfusion
- Keep this card with you at all times and for at least 6 months after the last dose of isatuximab
- If you notice any side effects, talk to your doctor or pharmacist
- Please report suspected side effects to the MHRA through the Yellow Card Scheme, via the Yellow Card website www.mhra.gov.uk/yellowcard or the free Yellow Card app available in Apple App Store or <u>Google Play Store</u>. Alternatively, you can call 0800 731 6789 for free, Monday to Friday between 9am and 5pm.
  - By reporting side effects, you can help provide more information on the safety of this medicine.
  - Side effects can also be reported to Sanofi: Tel: 0800 090 2314. email: uk-drugsafety@sanofi.com
- For further information about isatuximab, you can consult the Patient Information Leaflet (PIL)

## Warning message for healthcare professionals treating the patient at any time, including in emergency situations

- Please note this patient is receiving treatment with isatuximab for the treatment of multiple myeloma
- This patient card contains important safety information that you need to be aware of before, during, and after treatment with isatuximab
- Isatuximab can bind to CD38 on red blood cells (RBCs) and is associated with a risk of interference with blood typing (positive indirect Coombs test), which may persist for at least 6 months after the last isatuximab infusion
- To avoid potential problems with RBC transfusion, you should perform blood type and screen tests prior to the first infusion of isatuximab. Phenotyping may be considered as per local practice
- If treatment with isatuximab has already started and in the event of a planned transfusion, you should notify the blood bank that the patient is receiving isatuximab and alert them to the risk of interference with the indirect antiglobulin test (indirect Coombs test)
- For additional information about isatuximab, please refer to the Summary of Product Characteristics (SmPC).