

**Title** 



Sedation vs Axillary Brachial plexus block in

## Consent Form - Adult providing own consent

ritie	Interventional Radiology (SABIR)
Short Title	SABIR
Protocol Number	2021/PID03679
Project Sponsor	None
Coordinating Principal Investigator/ Principal Investigator	Dr Ross Copping
Associate Investigator(s)	Dr Paul Balamon, Dr Shady Osman, Dr Chandra Annabattula, Dr Louise Wei
Location	Liverpool Hospital
Declaration by Participant	
I have read the Participant Information Sheet or someone has read it to me in a language that I understand.	
I understand the purposes, procedures and risks of the research described in the project.	
I give permission for my doctors, other health professionals, hospitals or laboratories outside this hospital to release information to Liverpool Hospital concerning my disease and treatment for the purposes of this project. I understand that such information will remain confidential.	
I have had an opportunity to ask questions and I am satisfied with the answers I have received.	
I freely agree to participate in this research project as described and understand that I am free to withdraw at any time during the study without affecting my future health care.	
I understand that I will be given a signed copy of this document to keep.	
Name of Participant (please print)	
Signature	Date
Name of Witness* to Participant's Signature (please print)	
Signature	Date
* Witness is <u>not</u> to be the investigator, a member of the study team or their delegate. In the event that an interpreter	

<sup>\*</sup> Witness is <u>not</u> to be the investigator, a member of the study team or their delegate. In the event that an interpreter is used, the interpreter may <u>not</u> act as a witness to the consent process. Witness must be 18 years or older.

## Declaration by Study Doctor/Senior Researcher<sup>t</sup>

Name of Study Doctor/
Senior Researcher<sup>†</sup> (please print)

Signature

Date

I have given a verbal explanation of the research project, its procedures and risks and I believe

Note: All parties signing the consent section must date their own signature.

I understand that, if I decide to discontinue the study treatment, I may be asked to attend follow-up visits to allow collection of information regarding my health status. Alternatively, a member of the research team may request my permission to obtain access to my medical records for collection of follow-up information for the purposes of research and analysis.

<sup>&</sup>lt;sup>†</sup> A senior member of the research team must provide the explanation of, and information concerning, the research project.