

## Patient agreement to investigation or treatment for Cardiac Resynchronisation Therapy Defibrillator (CRT-D)

Designed in compliance with the Department of Health Consent Form 1

Patient details (or pre-printed label)	
Patients NHS Number or Hospital Number	
Patients Surname / Family Name	
Patients First Name(s)	
Date of Birth	
Sex	
Responsible Healthcare Professional	
Job Title	
Special Requirements e.g. other language or other communication method	

Informed consent is obtained in accordance with the requirements of the HT Act 2004, the Human Tissue Regulations 2007 and the HTA's Codes of Practice.

## **Tissue samples**

Tissues may be removed during your procedure for diagnostic examination by a histopathologist (a specialist doctor who looks at tissue from patients). Tissue samples needed for diagnosis are stored by the laboratory for several years. The stored tissue may be anonymously used for laboratory quality control, audit and education. These are essential activities for maintaining high quality diagnostic pathology services. Any remaining excess tissue removed is incinerated.

The specimen may be digitally photographed and the images temporarily stored in the laboratory as part of the diagnostic process. Other completely anonymised images may also be used for quality assurance, audit and education purposes.

Occasionally stored tissues and photographs might be used for research projects. Any such research will have been approved by a research ethics committee (REC). Usually any pathology specimens used for research are made completely anonymous, so that individual patients cannot be identified in any way. If this is not possible, the REC will require the researcher to contact you and ask permission to use your stored tissue or photographs. You would then be free to decide whether or not to allow the use of the material. Your decision would not in any way affect your medical care.

Notes Copy		Pati	ent identifier/label
Name of Proposed Procedure (include if medical term not clear)	a brief explanatio	n Anaesthetic	
Insertion of CRT-D		Local Sedation	
Statement of health professional (To be fille procedure, as specified in the consent policy). I have explained the procedure to the patient.  The intended benefits: To treat life-threaten	In particular, I	have explained:	of the proposed
Significant, unavoidable or frequently occ	urring risks initial		initial
Death < 1 in 1000		LV lead displacement 5 - 10 in 100	
Lead displacement 1 in 100		Haematoma	
Infection < 1 in 200			
Pneumothorax < 1 in 100			
Document any consent variations here			
Any extra procedures which may become need blood transfusion  other procedure (please specify)  The following leaflet / tape has been given:  I have discussed what the procedure is like treatments (including no treatment) and are has no outstanding queries and all their queries information sheet(s) as detailed above	The consent  The consent  in t	the benefits and risks of any available oncerns of this patient. I have checked	d that the patient
Signed:		Date:	
Name (PRINT)		Job Title:	
Contact Details (if patient wishes to discuss o  Statement of interpreter (where appropositive and in a way I believe s/he can use	oriate). I have in nderstand.	terpreted the information above to the p	
Signature of Interpreter	Name (print)	Date	

Patient's copy	Patient identifier/label			
Name of Proposed Procedure (include a if medical term not clear)	a brief explanation	Anaesthetic		
Insertion of CRT-D		Local Sedation		
tatement of health professional (To be fille rocedure, as specified in the consent policy).	d in by health pr	ofessional with appropriate knowledg	e of the proposed	
have explained the procedure to the patient.	In particular, I ha	ave explained:		
he intended benefits: To treat life-threateni	ing heart rhythm	disturbances and improve prognosis		
Significant, unavoidable or frequently occu	<u> </u>			
Death < 1 in 1000	initial	V lead displacement 5 - 10 in 100	initial	
Lead displacement 1 in 100	<del>                                     </del>			
		aematoma	<u> </u>	
Infection < 1 in 200			<u> </u>	
Pneumothorax < 1 in 100				
Document any consent variations here				
blood transfusion  other procedure (please specify)  the following leaflet / tape has been given:  have discussed what the procedure is like reatments (including no treatment) and an as no outstanding queries and all their queries.	The consent f  to involve, the particular con	orm  e benefits and risks of any availabl cerns of this patient. I have check	ed that the patient	
ne information sheet(s) as detailed above.  Signed:		Date:	- Thave given them	
		Job Title:		
Name (PRINT)  Contact Details (if patient wishes to discuss operations of interpreter (where approping my ability and in a way I believe s/he can ur	riate). I have inte		patient to the best	
ignature of Interpreter	Name (print)	Date		

Patient identifier/label						
Statement of patient						
Please read this form carefully. If your own copy of page 2, which describes the offered a copy now. If you have any functioning your mind at any time, including	ne benefits and ri	sks of the proposed treatm lo ask - we are here to help	ent. If not, you will be			
agree to the procedure or course of treatment described on this form.						
understand that you cannot give me person will, however, have appropriate	-	a particular person will per	form the procedure. The			
understand that I will have the oppor the procedure, unless the urgency of n regional anaesthesia).	•					
understand that any procedure in ad- necessary to save my life or prevent se have been told about additional procedures which I do not we	erious harm to my cedures which ma	health. y become necessary during	my treatment. I have listed			
Patient's signature	Name (PRINT)		Date:			
A witness should sign below if the pat beople/children may also like a paren			or her consent. Young			
Signature	Name (PRINT)		Date:			
Confirmation of consent (to be completed by a health professional when the patient is admitted for the procedure, if the patient has signed the form in advance).  On behalf of the team treating the patient, I have confirmed with the patient that s/he has no further questions and wishes the procedure to go ahead.  Signed:  Date						
Name (DDINIT)		tala Titla				
Name (PRINT)		Job Title				
mportant notes: (tick if applicable)  See also advance decision to ref  Patient has withdrawn consent (  Patient agrees to the use of surp	(ask patient to sig	n/date here)				