PATIENT CONSENT FORM - PART I

Patient Trial Number



PATIENT CONSENT FORM - PART I Version 4.0 Date 01.08.2007

Study title: A Phase II Single Arm Study of the use of CODOX-M/IVAC with Rituximab (R-CODOX-M/IVAC) in the treatment of patients with Diffuse Large B-Cell Lymphoma (DLBCL) of International Prognostic Index (IPI) High or High-Intermediate Risk (MREC reference 05/Q0201/81)

PATIENT CONSENT FORM PART I* (please read carefully)

Name of Researcher: _____

Please initial

I confirm that I have read and understand the information dated	
01.08.2007 (version 4.0) for the above study and have had an	
opportunity to ask questions.	
I understand that my participation in this study is voluntary and that I	
am free to withdraw at any time without giving a reason, without my	
medical care or legal rights being affected.	
I understand that sections of my medical notes may be looked at by	
responsible individuals involved in this research or from regulatory	
authorities where it is relevant to my taking part in research. I give	
permission for these individuals to have access to my records.	
I understand that samples of tissue taken will be reviewed by a	
pathologist in another hospital and that additional tests for biological	
risk factors will be performed	
I agree to take part in the above study.	

OPTIONAL

I agree for my GP to be informed	
I agree for information to be obtained from the Office of National Statistics	

Name of Patient

Date

Signature

Name of person taking consent Date (if different from researcher)

Signature

^{*} Three copies required: one each for the patient, researcher and hospital case notes

Name of Researcher Date		Signature	
Patient Trial Number			
PATIENT CONSENT FOR	M - PART I	I	
Study title: A Phase II Sin M/IVAC with Rituximab (R patients with Diffuse La International Prognostic Ind (MREC reference 05/Q0201/	R-CODOX-M arge B-Cel lex (IPI) Hig	/IVAC) in the tre l Lymphoma (I	eatment of DLBCL) of
PATIENT CONSENT FORM PART	II* (please re	ead carefully)	
Name of Researcher:		, 1Per	
VOLUNTARY DONATION OF	TISSUF S Á M		Please initial
be linked to the trial data throug other personal data will be held. approval will be sought before the studies.	I understand	separate ethics	
			1]
Name of Patient	Date	Signature	
Name of Patient Name of person taking consent (if different from researcher)	Date Date	Signature Signature	

^{*} Three copies required: one each for the patient, researcher and hospital case notes

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