## On headed paper



## POSNOC - A randomised trial of armpit (axilla) treatment for women with early stage breast cancer

**REC ref:** 13/EM/0459

Name of Researcher: Local details to be added

**Patient Name:** 

Please initial boxes (To be completed by the patient)		
		YES
1.	I confirm that I have read, understood and received a copy of the information sheet dated 17-Dec-2019 (Final version 4.1a) for the above study and have been given a copy to keep. I have had the opportunity to consider the information and to ask questions and have had these questions answered satisfactorily.	
2.	I understand that my participation is voluntary, and that I am free to withdraw my consent at any time, without giving any reason, without my medical care or legal rights being affected.	
3.	I understand that information from my medical notes and the data collected during the study, may be looked at by individuals from University Hospitals of Derby and Burton NHS Foundation Trust (sponsor) and other third parties appointed by the sponsor where it is relevant to my taking part in this research. I give permission for these individuals to have access to these records and for a copy of this signed consent form to be sent to the Nottingham Clinical Trials Unit. I understand that my personal details will be kept confidential.	
4.	I give permission to my GP being contacted.	
5.	I understand that information held by the NHS and records maintained by NHS Digital may be used to check my health status during the study and for future research. I give my permission to register my identifiable details with NHS Digital.	
6.	I understand that my personal details will be held by SHORE-C, (University of Sussex) and NCTU (University of Nottingham) and will be used to contact me by post and telephone. I give permission for this information to be kept and for these individuals to contact me. This is optional and you can still take part in the study if you answer 'No".	YES NO
7.	I agree to donate tissue routinely obtained to make the diagnosis or treat cancer for possible use in future ethically approved research. I understand that these samples will be stored and may be transferred to a licensed biobank. This is optional and you can still take part in the study if you answer 'No".	YES NO
8.	I would like to receive a summary of the results at the end of the study. This is optional and you can still take part in the study if you answer 'No".	YES NO
9.	I agree to take part in the above study	
Your signature confirms that you have had an opportunity to ask questions and all questions have been answered. You will be given a signed and dated copy of this consent form to take away with you.		
Name o	of patient (Print) Signed Date	
Name o	of Person taking consent Signed Date	

When completed, 1 Copy for Patient, 1 Copy for Researcher, 1 Copy for Hospital Notes & 1 copy to the NCTU

The views expressed are those of the author(s) and not necessarily those of the NHS, the NIHR or the Department of Health.

