HORIZONS



Understanding the impact of cancer diagnosis and treatment on everyday life

12 MONTH BREAST CANCER CRF

FOR STAFF USE ONLY

CRF Completion Instructions

- Please complete as much of the CRF as possible
- Please refer to your copy of the 6 month CRF when completing this 12 month CRF: please complete any additional treatment details that were not captured at 6 months (for example, additional treatments, or end dates of those ongoing at 6 months)
- If you have any queries, please contact the HORIZONS
 Co-ordinating Centre, email address HORIZONS@soton.ac.uk
- Please tick boxes when appropriate
- When you have completed the CRF, please keep a copy for your own records and return a copy to us, by post, fax or email along with the completed return cover sheet

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Participa	nt's Study ID / /			
Participa	nt's date of birth	уууу	У	
Has the រ	participant been tested for BRCA1 or BR	CA2 (please	tick one box)	
	Yes (already recorded in previous CRF)			
	Yes (but not recorded in previous CRF)			
	No			
	Unknown			
Negativ	e for a mutation in BRCA1 or BRCA2 e for a mutation in BRCA1 or BRCA2 ous or uncertain			
Unknow	vn			
Awaitin	g result			
Has the box)	participant had any other genetic tests f	or inherited	cancers? (please	tick one
	Yes (already recorded in previous CRF)			
	Yes (but not recorded in previous CRF)			
	No			
	Unknown			

If you answered "Yes (but not recorded in previous CRF)" to the above question, please provide some information about the participant's other genetic test(s) by completing the table(s) overleaf

Participant's Study ID /	
Name of genetic test for cancer (1)	Result of genetic test
	Positive
	Negative
	Ambiguous/uncertain
	Awaiting result
	Unknown
Name of constint at favorages (2)	Decult of constintent
Name of genetic test for cancer (2)	Result of genetic test Positive
	Negative
	Ambiguous/uncertain
	Awaiting result
	Unknown
Has the participant developed any NEW of the baseline or 6 month CRFs)? (please tick overleaf)	o-morbidities (which were not recorded in ck all that apply in the tables below and
Myocardial infarct	
Angina/coronary artery disease	
Congestive Heart Failure	
Cardiac Arrythmias	
Hypertension	
Venous Disease (PE/DVT)	
Peripheral Arterial Disease	
Restrictive Lung Disease or COPD (chronic bron	chitis, emphysema, asthma etc.)

Liver Disease (portal hypertension, chronic/acute hepatitis, cirrhosis etc.)

Participant's Study ID	
Stomach Ulcers or Inflammatory Bowel Disease	
Acute or Chronic Pancreatitis	
End-stage Renal Disease (chronic renal insufficiency, dialysis etc.)	
Thyroid problems (hyperthyroidism, hypothyroidism etc.)	
Diabetes Mellitus Type 1	
Diabetes Mellitus Type 2	
Stroke/TIA	
Dementia	
Paralysis (paraplegia or hemiplegia)	
Neuromuscular Condition (multiple sclerosis, Parkinson's, myasthenia gravis, other chronic neuromuscular disorder)	
Clinical diagnosis of anxiety	
Clinical diagnosis of depression	
Other Psychiatric Diagnosis (eg. schizophrenia, bipolar disorder etc.)	
Osteoarthritis	
Rheumatoid Arthritis	
Other Rheumatological Disease (systemic lupus, mixed connective tissue disorder,	
polymyositis, rheumatic polymyositis, scleroderma etc.)	
HIV/AIDS	
Alcohol Abuse (or history of, must be accompanied by social, behavioural or medical complications)	
Drug/Substance Abuse (or history of, must be accompanied by social, behavioural or medical complications)	
Morbid Obesity	
Other (please give details)	
Other (please give details)	
Other (please give details)	

Participant's Study ID	
at treatments has the	participant received since those captured at 6 month

What treatments has the participant received **since those captured at 6 months**, please tick ALL that apply and write details in the spaces provided. Please add end dates for any treatments which were ongoing at 6 months (table continued overleaf).

Treatment type	Specific treatment details	Tick if patient has received	Date of surgery or start date of other treatment (dd/mm/yyyy)	End date of treatment (if finished) (dd/mm/yyyy)	If course of treatment was not completed as planned, please give a reason why
Surgery	Wide local excision (breast conserving surgery)		// 20		
	Mastectomy		//20		
	Sentinel node biopsy (SNBx)		// 20		
	Axillary node clearance (ANC)		// 20		
	Other axillary treatment please describe on line below)		// 20		
Breast Reconstruction	Immediate reconstruction		// 20		
	Delayed reconstruction		//20		
	Delayed reconstruction is planned but has not yet taken place				
Reconstruction Type	Implant		//20		
	Latissimus dorsi (LAD)		// 20		
	Deep inferior epigastric perforator artery (DIEP)		//20		
	Tissue reconstruction with abdominal tissue (TRAM)		//20		
	Nipple reconstruction		// 20		
	Nipple/Areola Tattoo		// 20		
	Other (please describe on line below)		// 20		

Participant's Study ID		/		/			

Treatment type	Specific treatment details	Tick if patient has re- ceived	Date of surgery or start date of other treatment (dd/mm/yyyy)	End date of treat- ment (if finished) (dd/mm/yyyy)	If course of treatment was not completed as planned, please give a reason why
Radiotherapy	Breast		//20	// 20	
	Chest wall		//20	//20	
	Supraclavicular fossa (SCF)		//20	//20	
	Axilla		//20	// 20	
	Number of radiotherapy fra	ctions, ple	ease enter on line		
	Total radiotherapy dose pl	ease enter	r on line		
Chemotherapy	Drug(s), please give details below		//20	//20	
	Chemotherapy number of	cycles, ple	ease enter on line	1	
Ovarian Suppression	Medical, please give details below		// 20	// 20	
	Surgical		//20		
	Radiotherapy		//20	// 20	

Participant's Study ID		/		/			
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Treatment type	Specific treatment details	Tick if patient has received	Date of surgery or start date of other treatment (dd/mm/yyyy)	End date of treatment (if finished) (dd/mm/yyyy)	If course of treatment was not completed as planned, please give a reason why
Hormone	Tamoxifen		//20	// 20	
Therapy	Anastrazole		// 20	// 20	
	Letrozole		// 20	// 20	
	Exemestane		//20	// 20	
	Other, please give details below		//20	// 20	
	Were bisphosphonates give Yes No Unknow		ick)?		
Symmeterisation Operations	Contralateral risk reducing mastectomy		// 20	// 20	
	Other symmeterisation operation (please give details)		//20	// 20	
	Other risk reducing surgery (please give details)		// 20	// 20	
Immunotherapy	Trastuzumab (Herceptin)		//20	// 20	
	Pertuzumab (Perjeta)		//20	// 20	
	Other immunotherapy (please give details		//20	// 20	
Additional Treatment	If any additional treatment has been given please describe		// 20	// 20	

Par	ticipant's S	Study ID			/						
	as the part ox)	icipant I	nad a loc	al recui		of the	eir breast ca Unknowi		plea	se tick one	3
If	the partici	pant has	s had a lo	cal rec	urrenc	e, on	what date v	vas the	recu	rrence	
di	agnosed?	d d	m m	У	уу	У					
	e the parti ant metast	-	_				has there bo	een any	evic	lence of	
		Yes		No			Unknown				
-	ou have an		"yes" to	the abo	ove que	stion	, on what da	ate was	the	metastatio	C
		d d	m m	у	УУ	У					
Ple	ase provid	e details	of the s	ite(s) o	f distan	it met	tastatic dise	ase:			
ls t	he particip	oant pre	or post	menop	ause? (pleas	e tick one b	ox)			
	Pre meno	pause									
	Post mend	pause									
	Unknown										

Participant's Study ID / /	
Is the participant taking part in a clinical tri	al? (please tick one box)
Yes No	Unknown
If you answered "yes" to the above questic clinical trial the participant is taking part in	
Name of clinical trial	
Since the participant's diagnosis of breast ca another new primary cancer? (please tick on Yes	,
If you answered "yes" to the above questic about the participant's new cancer diagnos	
Type of cancer	
Date of diagnosis	// 20
Treatment received	
Date treatment ended (if finished)	// 20
What type of follow-up care is the participa	nt receiving? (please tick ONE box)
Routine/regular hospital clinic based follow face-to-face or by telephone)	v-up (medical or nurse led,
Primary care based follow-up	
Patient initiated follow-up (also known as patient), open access follow-up, or supported	
If the participant is receiving patient-initiat they discharged to this?	ed follow-up, on what date were

Participant's Study ID / / / /
Has the participant been referred to any of the following services and/or had a
Holistic Needs Assessment? (please tick all that apply)
Participant has been referred to palliative care services
Participant has been referred to psychological services
If ticked, please provide route to referral (e.g. GP, Improving Access to Psychological Therapies)
Participant has been referred to community services
Participant has been referred for treatment related problems (e.g. pain clinic) If ticked, please provide more details below:
Participant has had an HNA (holistic needs assessment)
If the participant has died please give the date and cause of death:
Participant's date of death d d / m m / y y y y
Cause of participant's death
1) a)
1) b)
1) c)
2)
Cause of death not known
Please add your name and signature and the date that you completed this CRF
Name Signature
Date CRF completed dd / m m / y y y y