HORIZONS



Understanding the impact of cancer diagnosis and treatment on everyday life

24 MONTH BREAST CANCER CRF

FOR STAFF USE ONLY

CRF Completion Instructions

- Please complete as much of the CRF as possible
- Please refer to your copies of previous CRFs when completing this 24 month CRF: please complete any additional treatment details that were not captured at 6 or 12 months (for example, additional treatments, or end dates of those ongoing in earlier CRFs)
- 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

| Participant's Study ID |]/[| | /[| | | | | | |
|-----------------------------|-----|---|----|---|---|---|---|---|--|
| Participant's date of birth | d | d | m | m | У | У | У | У | |

Has the participant developed any NEW co-morbidities (which were not recorded in any previous CRFs)? (please tick all that apply in the tables below and overleaf)

| Myocardial infarct | |
|--|--|
| Angina/coronary artery disease | |
| Congestive Heart Failure | |
| Cardiac Arrhythmias | |
| Hypertension | |
| Venous Disease (PE/DVT) | |
| Peripheral Arterial Disease | |
| Restrictive Lung Disease or COPD (chronic bronchitis, emphysema, asthma etc.) | |
| Liver Disease (portal hypertension, chronic/acute hepatitis, cirrhosis etc.) | |
| 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.) | |

| Participant's Study ID | | / | | / | | | |
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| Osteoarthritis | |
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| 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 | | / | | / | | | ı |
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What treatments has the participant received **since those captured in any previous CRFs**, please tick ALL that apply and write details in the spaces provided. Please add end dates for any treatments which were ongoing previously (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 | | |
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| Participant's Study ID | | / | | | | |
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| 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 | ictions, ple | ase enter on line | | |
| | Total radiotherapy dose plo | ease enter | on line | | |
| Chemotherapy | Drug(s), please give details | | // 20 | //20 | |
| | Chemotherapy number of | cycles, plea | ase enter on line | | |
| Ovarian Suppression | Medical, please give details below | | //20 | //20 | |
| | Surgical | | //20 | | |
| | Radiotherapy | | //20 | // 20 | |
| Hormone Therapy | Tamoxifen | | //20 | // 20 | |
| | Anastrazole | | //20 | // 20 | |
| | Letrozole | | // 20 | // 20 | |
| | Exemestane | | //20 | // 20 | |
| | Other, please give details | | // 20 | //20 | |
| | Were bisphosphonates give | | tick)? | I | |

| 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 |
|-------------------------------|---|---------------------------------------|--|---|--|
| 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 | If any additional treatment | | / / 20 | / / 20 | |

Were any of the treatments detailed above given with **palliative intent**? (please tick one box)

| Yes No | Unknown |
|--------|---------|
|--------|---------|

If yes, please indicate which treatments?

has been given please

describe

Treatment

Participant's Study ID

| Participant's Study ID / / |
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| Has the participant had a local recurrence of their breast cancer? (please tick one box) Yes No Unknown |
| If the participant has had a local recurrence, on what date was the recurrence |
| diagnosed? |
| Since the participant's diagnosis of breast cancer, has there been any evidence of distant metastatic disease? (please tick one box) |
| Yes No Unknown |
| If you have answered "yes" to the above question, on what date was the metastation disease diagnosed: |
| d d m m y y y y |
| Please provide details of the site(s) of distant metastatic disease: |
| Is the participant pre or post menopause? (please tick one box) |
| Pre menopause |
| Post menopause |
| 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 question 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 | ne box) | | | | |
| Yes No | Unknown | | | | |
| 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 participant receiving? (please tick ONE box) | | | | | |
| Routine/regular hospital clinic based follow | v-up (medical or nurse led, | | | | |
| face-to-face or by telephone) | | | | | |
| Primary care based follow-up | | | | | |
| Patient initiated follow-up (also known as patient triggered follow-up (PTFU), open access follow-up, or supported self-managed follow-up) | | | | | |
| If the participant is receiving patient-initiated follow-up, on what date were they discharged to this? | | | | | |

| Participant's Study ID / / / / |
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| 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 |
| If available, please give reason for referral (e.g. end of life care, symptom management) |
| 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 / m m / 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 |