Breast (BRCA)

<u>Instructions:</u> The Enrollment Form should be completed for each TCGA qualified case, upon qualification notice from the BCR. All information provided on this form should include activity from the date of initial diagnosis to the most recent date of contact with the patient ("Date of Initial Pathologic Diagnosis" and "Date of Last Contact" on this form).

Questions regarding this form should be directed to the Tissue Source Site's primary Clinical Outreach Contact at the BCR.

Please note the following definitions for the "Unknown" and "Not Evaluated" answer options on this form.

Unknown: This answer option should only be selected if the TSS does not know this information after all efforts to obtain the data have been exhausted. If this answer option is selected for a question that is part of the TCGA required data set, the TSS must complete a discrepancy note providing a reason why the answer is unknown.

Tissue Source Site (TSS): ______TSS Identifier: _____TSS Unique Patient Identifier: ____

Not Evaluated: This answer option should only be selected by the TSS if it is known that the information being requested cannot be obtained. This could be because the test in question was never performed on the patient or the TSS knows that the information requested was never disclosed.

Completed By (Interviewer Name in OpenClinica):			Completed Date:
Gene	ral Information		
#	Data Element	Entry Alternatives	Working Instructions
1	Has this TSS received permission from the NCI to provide time intervals as a substitute for requested dates on this form?	☐ Yes ☐ No	If the answer to this question is yes, time intervals must be provided instead of dates, as indicated throughout this form. Provided time intervals must begin with the date of initial pathologic diagnosis (e.g. biopsy). Only provide interval data if you have received permission from the NCI to provide time intervals as a substitute for requested dates on this form.
2	Is this a prospective tissue collection?	☐ Yes ☐ No	Indicate whether the TSS providing tissue is contracted for prospective tissue collection. If the submitted tissue was collected for the specific purpose of TCGA, the tissue has been collected prospectively. 3088492
3	Is this a retrospective tissue collection?	☐ Yes ☐ No	Indicate whether the TSS providing tissue is contracted for retrospective tissue collection. If the submitted tissue was collected prior to the date the TCGA contract was executed, the tissue has been collected retrospectively. 3088528

Patient Information

#	Data Element	Entr	y Alternatives		Working Instructions
4	Month of Birth	□ 01 □ 04 □ 02 □ 05 □ 03 □ 06	□ 07 □ 08 □ 09	□ 10 □ 11 □ 12	Provide the month the patient was born. 2896950
5	Day of Birth	□ 01 □ 08 □ 02 □ 09 □ 03 □ 10 □ 04 □ 11 □ 05 □ 12 □ 06 □ 13 □ 07	□ 14 □ 20 □ 15 □ 21 □ 16 □ 22 □ 17 □ 23 □ 18 □ 24 □ 19 □ 25	□ 26 □ 27 □ 28 □ 29 □ 30 □ 31	Provide the day the patient was born. 2896952
6	Year of Birth			_	Provide the year the patient was born. 2896954

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#	Data Element	Entry Alternatives	Working Instructions
7	Number of Days from Date of Initial Pathologic Diagnosis to Date of Birth		Provide the number of days from the date the patient was initially diagnosed pathologically with the disease to the patient's date of birth. 3008233 Only provide Interval data if you have received permission from the NCI to provide time intervals as a substitute for requested dates on this form.
8	Gender	☐ Female ☐ Male	Provide the patient's gender using the defined categories. 2200604
9	Menopause Status (at time of diagnosis)	□ Premenopausal <6 months since LMP AND no prior bilateral oophorectomy AND not on estrogen replacement □ Perimenopausal 6-12 months since last menstrual period □ Postmenopausal Prior bilateral oophorectomy OR >12 months since LMP with no prior oophorectomry □ Indeterminate or Unknown □ Not Evaluated	Using the patient's medical records, indicate menopause status at the time the patient was diagnosed with the malignancy submitted for TCGA. 2957270
10	Race	 □ American Indian or Alaska Native A person having origins in any of the original peoples of North and South America (including Central America), and who maintains tribal affiliation or community attachment. □ Asian A person having origins in any of the original peoples of the far East, Southeast Asia, or in the Indian subcontinent including, for example, Cambodia, China, India, Japan, Korea, Malaysia, Pakistan, the Philippine Islands, Thailand, and Vietnam. □ White	Provide the patient's race using the defined categories. 2192199
11	Ethnicity	 Not Hispanic or Latino: A person not meeting the definition of Hispanic or Latino. Hispanic or Latino: A person of Mexican, Puerto Rican, Cuban, Central or South American or other Spanish culture or origin, regardless of race. Not Evaluated Not provided or available. Unknown	Provide the patient's ethnicity using the defined categories. 2192217
12	History of Other Malignancy	☐ Yes ☐ No	Indicate whether the patient was, at any time in their life, diagnosed with a malignancy prior to the diagnosis of the specimen submitted for TCGA. If the patient has had a prior malignancy, an additional form (the "Other Malignancy Form") must be completed for each prior malignancy. If the OMF was completed and submitted with the Initial Case Quality Control Form, the OMF does not need to be submitted a second time. 3382736 If this question cannot be answered because the answer is unknown, the case will be excluded from TCGA. If the patient has a history of multiple diagnoses of basal or squamous cell skin cancer, complete an OMF for the first diagnosis for each of these types.

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#	Data Element	Ent	ry Alternatives		Working Instructions
13	History of Neo-adjuvant Treatment for Sample Submitted for TCGA	☐ Yes ☐ No			Indicate whether the patient received neo-adjuvant treatment (radiation, pharmaceutical, or both) prior to the collection of the sample submitted for TCGA. 3382737 Systemic therapy and certain localized therapies (those administered to the same site as the TCGA submitted sample) given prior to the collection of the sample submitted for TCGA is
14	Tumor Status (at time of last contact or death)	☐ Tumor free☐ With tumor☐ Unknown			exclusionary. Indicate whether the patient was tumor/disease free at the date of last contact or death. 2759550
15	Vital Status (at date of last contact)	☐ Living ☐ Deceased			Indicate whether the patient was living or deceased at the date of last contact. 2939553
16	Month of Last Contact	□ 01 □ 04 □ 02 □ 05 □ 03 □ 06	□ 07 □ 08 □ 09	□ 10 □ 11 □ 12	If the patient is living, provide the month of last contact with the patient (as reported by the patient, medical provider, family member, or caregiver). 2897020
17	Day of Last Contact	□ 01 □ 08 □ 02 □ 09 □ 03 □ 10 □ 04 □ 11 □ 05 □ 12 □ 06 □ 13 □ 07	□ 14 □ 20 □ 15 □ 21 □ 16 □ 22 □ 17 □ 23 □ 18 □ 24 □ 19 □ 25	□ 26 □ 27 □ 28 □ 29 □ 30 □ 31	Do not answer if patient is deceased. If the patient is living, provide the day of last contact with the patient (as reported by the patient, medical provider, family member, or caregiver). 2897022 Do not answer if patient is deceased.
18	Year of Last Contact			-	If the patient is living, provide the year of last contact with the patient (as reported by the patient, medical provider, family member, or caregiver). 2897024 Do not answer if patient is deceased.
19	Number of Days from Date of Initial Pathologic Diagnosis to Date of Last Contact			-	Provide the number of days from the date the patient was initially diagnosed pathologically with the disease described on this form to the date of last contact. 3008273 Only provide Interval data if you have received permission from the NCI to provide time intervals as a substitute for requested dates on this form.
20	Month of Death	□ 01 □ 04 □ 02 □ 05 □ 03 □ 06	□ 07 □ 08 □ 09	□ 10 □ 11 □ 12	If the patient is deceased, provide the month of death. 2897026
21	Day of Death	□ 01 □ 08 □ 02 □ 09 □ 03 □ 10 □ 04 □ 11 □ 05 □ 12 □ 06 □ 13 □ 07	□ 14 □ 20 □ 15 □ 21 □ 16 □ 22 □ 17 □ 23 □ 18 □ 24 □ 19 □ 25	□ 26 □ 27 □ 28 □ 29 □ 30 □ 31	If the patient is deceased, provide the day of death. 2897028
22	Year of Death			-	If the patient is deceased, provide the year of death. 2897030
23	Number of Days from Date of Initial Pathologic Diagnosis to Date of Death			-	Provide the number of days from the date the patient was initially diagnosed pathologically with the disease described on this form to the date of death. 3165475 Only provide Interval data if you have received permission from the NCI to provide time intervals as a substitute for requested dates on this form.

#	Data Element	Entry Alternatives	Working Instructions
24	Adjuvant (Post- Operative) Radiation Therapy	☐ Yes ☐ No ☐ Unknown	Indicate whether the patient had adjuvant/ post- operative radiation therapy <i>for the tumor submitted for TCGA</i> 2005312 If the patient did have adjuvant radiation, the Radiation Supplemental Form should be completed.
25	Adjuvant (Post- Operative) Pharmaceutical Therapy	☐ Yes ☐ No ☐ Unknown	Indicate whether the patient had adjuvant/ post- operative pharmaceutical therapy <u>for the tumor</u> <u>submitted for TCGA</u> . 3397567 If the patient did have adjuvant pharmaceutical therapy, the Pharmaceutical Supplemental Form should be completed.

Patno	athologic/Prognostic Information							
#	Data Element	Entry Alternatives	Working Instructions					
26	Primary Site of Disease	□ Breast	Using the patient's pathology/laboratory report, select the anatomic site of disease of the tumor submitted for TCGA. 2735776 The tumor submitted for TCGA must be located in the endometrium; indicate other involvement, as initially diagnosed.					
27	Anatomic Organ Sub- Division	☐ Right Breast ☐ UIQ ☐ UOQ ☐ LIQ ☐ LOQ ☐ Left Breast ☐ UIQ ☐ UOQ ☐ LIQ ☐ LOQ	Using the patient's pathology/laboratory report, select the anatomic organ sub-division of the tumor used for TCGA. Include all areas of tumor invasion. 2008006					
28	Histological Subtype	☐ Infiltrating Ductal Carcinoma ☐ Infiltrating Lobular Carcinoma ☐ Infiltrating Carcinoma, NOS ☐ Mucinous Carcinoma ☐ Medullary Carcinoma ☐ Mixed Histology, specify ☐ Other, specify	Using the patient's pathology/laboratory report, select the histology and/or subtype of the tumor submitted for TCGA. <i>Mixed Histology</i> : The specimen is mixed with ductal and lobular carcinomas only. <i>Other</i> : Any other histology mixed with ductal and/or lobular OR rare/special histological types. 2549638					
29	Other Histological Subtype or Mixed Diagnosis		If the histological subtype on the pathology/laboratory report does not fall under the provided histological types, describe the histology and/or subtype here. 3124492					
30	Month of Initial Pathologic Diagnosis	$\begin{array}{c ccccc} & \Box & 01 & & \Box & 04 & & \Box & 07 & & \Box & 10 \\ \hline \Box & 02 & & \Box & 05 & & \Box & 08 & & \Box & 11 \\ \hline \Box & 03 & & \Box & 06 & & \Box & 09 & & \Box & 12 \\ \end{array}$	Provide the month the patient was initially diagnosed with the malignancy submitted for TCGA. 2896956					
31	Day of Initial Pathologic Diagnosis	□ 01 □ 08 □ 14 □ 20 □ 26 □ 02 □ 09 □ 15 □ 21 □ 27 □ 03 □ 10 □ 16 □ 22 □ 28 □ 04 □ 11 □ 17 □ 23 □ 29 □ 05 □ 12 □ 18 □ 24 □ 30 □ 06 □ 13 □ 19 □ 25 □ 31 □ 07	Provide the day the patient was initially diagnosed with the malignancy submitted for TCGA. 2896958					
32	Year of Initial Pathologic Diagnosis		Provide the year the patient was initially diagnosed with the malignancy submitted for TCGA. 2896960					
33	Age at Initial Diagnosis		Provide the age of the patient in years, at the time the patient was initially pathologically diagnosed. 2006657 Only complete this question if you have received permission from the NCI to provide time intervals as a substitute for requested dates on this form.					
34	Method of Initial Pathologic Diagnosis	☐ Cytology ☐ Fine needle aspiration biopsy ☐ Core needle biopsy ☐ Incision biopsy ☐ Excisional biopsy ☐ Tumor resection ☐ Other, specify	Provide the procedure used to initially diagnose the patient. 2757941					

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#	Data Element	Entry Alternatives	Working Instructions	
35	Other Method of Initial Pathologic Diagnosis		If the procedure used to initially diagnose the patient was not included in the list provided, please describe the method used. 2757948	
36	First Surgical Procedure	☐ Lumpectomy ☐ Simple mastectomy ☐ Modified radical mastectomy ☐ Other, specify	Provide the first procedure used after the initial diagnosis. 2739580	
37	Other First Surgical Procedure		If the first procedure used after the initial diagnose was not included in the list provided, please describe the method used. 3020338	
38	Margin Status after First Surgical Procedure	☐ Positive (+) ☐ Negative (-) ☐ Close	Provide the margin status after the patient's first surgical procedure. 3114007	
39	If margins were positive after first surgical resection, what was the surgical procedure performed to achieve negative margins?	□ Surgery not performed □ Lumpectomy □ Mastectomy □ Modified radical mastectomy □ Other, specify	If margins were positive after the first surgical resection, provide the additional surgery performed to ensure negative margins. 1218	
40	Other Surgical Method Performed to Achieve Negative Margins		If the additional procedure used after the first surgery resulted in positive margins was not included in the list provided, please describe the method used. 3124493	
41	Margin Status after second surgical resection	☐ Positive (+) ☐ Negative (-) ☐ Close	Provide the margin status after the additional procedure used after the first surgery resulted in positive margins. 2241252	
42	Axillary Staging Method	 □ No axillary staging □ Sentinel lymph node biopsy alone □ Sentinel lymph node biopsy plus axillary dissection □ Axillary lymph node dissection alone □ Other, specify 	Using the pathology/laboratory report, provide the axillary staging method used to detect nodal involvement. 2516112	
43	Other method of Axillary Staging		If the axillary staging method used was not included in the list provided, please describe the method used. 3124496	
44	Was IHC Staining used to Detect Micro metastasis?	☐ Yes ☐ No ☐ Unknown	Indicate whether immunohistochemistry (IHC) staining was performed to detect micro metastasis. 3086152	
Lyn	iph Node Status			
45	Were Lymph Nodes Examined at the Time of Primary Resection?	☐ Yes ☐ No	Indicate whether any lymph nodes were examined at the time of the primary resection. 2200396	
46	Number of Lymph Nodes Examined		Provide the number of lymph nodes examined, if one or more lymph nodes were removed. 3	
47	Number of Lymph Nodes Positive by H&E light microscopy		Provide the number of lymph nodes positive through hematoxylin and eosin (H&E) staining and light microscopy. 3086388	
48	Number of Lymph Nodes Positive by IHC Keratin Staining only		Provide the number of lymph nodes positive through keratin immunohistochemistry (IHC) staining. 3086383	
AJC	C Staging			
49	AJCC Cancer Staging Edition	☐ 1st Edition (1978-1983) ☐ 2nd Edition (1984-1988) ☐ 3rd Edition (1989-1992) ☐ 4th Edition (1993-1997) ☐ 5th Edition (1998-2002) ☐ 6th Edition (2003-2009) ☐ 7th Edition (2010-present)	Based on the date the patient was staged select the AJCC edition used to stage the patient. 2722309	

#	Data Element	Entry Alternatives	Working Instructions
50	Pathologic T Stage	□ TX □ T1a □ T3a □ T0 □ T1b □ T3b □ Tis □ T1c □ T4 □ Tis (DCIS) □ T2 □ T4a □ Tis (LCIS) □ T2a □ T4b □ Tis □ T2b □ T4c (Paget's) □ T3 □ T4d □ T1mic □ T1	Using the patient's pathology/laboratory report, select the code for the pathologic T (primary tumor) defined by the American Joint Committee on Cancer (AJCC). 3045435
51	Pathologic N Stage	□ NX □ N1a □ N2 □ N0 □ N1b □ N2a □ N0 (i-) □ N1bi □ N2b □ N0 (i+) □ N1bii □ N3 □ N0 (mol-) □ N1biii □ N3a □ N0 (mol+) □ N1biv □ N3b □ N1 □ N1c □ N3c □ N1mi □ N1mi	Using the patient's pathology/laboratory report, select the code for the pathologic N (nodal) defined by the American Joint Committee on Cancer (AJCC). 3203106
52	Pathologic M Stage	□ MX □ cM0 (i+) □ M1	Using the patient's pathology/laboratory report, select the code for the pathologic M (metastasis) defined by the American Joint Committee on Cancer (AJCC). 3045439
53	Stage	□ Stage X □ Stage IIA □ Stage Tis □ Stage IIB □ Stage 0 □ Stage III □ Stage I □ Stage IIIA □ Stage IA □ Stage IIIB □ Stage IB □ Stage IIIC □ Stage II □ Stage IV	Using the patient's pathology/laboratory report, select the stage defined by the American Joint Committee on Cancer (AJCC). 3203222
54	Site of First Non-Nodal Metastatic Tumor If metastasis were found at multiple sites simultaneously, check all that apply	□ Lung □ Bone □ Liver □ Brain □ Other, specify	If the patient had a non-nodal metastasis associated with the diagnosis of the tumor submitted for TCGA, provide the site of the first non-nodal metastasis. Only select more than one site if there were synchronous metastasis where the first non-nodal met was found at multiple sites. 3124499
55	Other Site of First Non- Nodal Metastatic Tumor		If the site of the first non-nodal metastasis was not included in the list provided, please provide the site. 3124503
Prir	nary Tumor Molecular M	arkers Used for Tumor Prognosis	
56	Estrogen Receptor (ER) Status by IHC for this patient	□ Positive (1%-100%) □ Negative (0%) □ Indeterminate □ Performed but not available □ Not performed (skip to next molecular marker)	If IHC estrogen receptor testing was performed, provide the result of the test. If this test was not performed, selected "not performed," and continue to the progesterone receptor questions. 2957359
57	IHC ER Percent Positive for this patient	□ <10% (1-9%) □ 10-19% □ 20-29% □ 30-39% □ 40-49% □ 90-100%	If IHC estrogen receptor testing was performed, provide the percent of estrogen receptor positive by IHC. 3128341
58	IHC Intensity: Scale Used to determine ER Positivity for this Patient	☐ 4 Point Scale ☐ 3 Point Scale	Using the pathology/laboratory report, indicate the intensity scale used for the estrogen receptor positivity score. 3203081

#	Data Element	Entry Alternatives	Working Instructions
59	IHC Intensity: ER Positivity Score for this patient	$\begin{array}{c c} \square \ 0 & \square + 3 \\ \square + 1 & \square + 4 \\ \square + 2 & \square \ \text{Other, specify in next question} \end{array}$	Provide the positivity score using the IHC intensity scale, found on the pathology/laboratory report. 2230166
60	IHC Intensity: Other Method Used to Determine ER Positivity For this Patient		If another scale was used to measure the estrogen receptor positivity, please describe the scale used. 3086851
61	Define Method of Calculation for ER Positivity if Other than IHC		If a special method was used to calculate estrogen receptor status (e.g. dextran coated charcoal), describe the method used. 69
62	Progesterone Receptor (PR) Status by IHC for this patient	 □ Positive (1%-100%) □ Negative (0%) □ Indeterminate □ Performed but not available □ Not performed (skip to next molecular marker) 	If IHC progesterone receptor testing was performed, provide the result of the test. If this test was not performed, select "not performed," and continue to the HER2/ERBB2 IHC questions. 2957357
63	IHC PR Percent Positive for this patient	□ <10% (0-9%) □ 50-59% □ 10-19% □ 60-69% □ 70-79% □ 30-39% □ 80-89% □ 40-49% □ 90-100%	If IHC progesterone receptor testing was performed, provide the percent of progesterone receptor positive nuclei by IHC. 3128342
64	IHC Intensity: Scale Used for PR Positivity	☐ 4 Point Scale ☐ 3 Point Scale	Using the pathology/laboratory report, indicate the intensity scale used for the progesterone receptor positivity score. 3203083
65	IHC Intensity: PR Positivity Score for this Patient	□ 0 □ +3 □ +1 □ +4 □ +2	Provide the positivity score using the IHC intensity scale, found on the pathology/laboratory report. Only answer this question if PR status is considered positive; if the PR status was negative, continue to the HER2/ERBB2 IHC questions. 3133874
66	IHC Intensity: Other Method Used to Determine PR Positivity		If another scale was used to measure the progesterone receptor positivity, please describe the scale used. 3086857
67	Define Method of Calculation for Positivity if Other Than IHC		If a special method was used (other than IHC) to calculate progesterone receptor status (e.g. dextran coated charcoal), describe the method used. 785
68	HER2/ERBB2 Status by IHC for this Patient	□ Positive □ Negative □ Equivocal □ Indeterminate □ Performed but not available □ Not performed(skip to next molecular marker)	If IHC HER2/ERBB2 testing was performed, provide the result of the test. If this test was not performed, select "not performed," and continue to the HER2/ERBB2 FISH questions. 2957563
69	IHC HER2/ERBB2 Percent Positive for this patient	□ <10% □ 50-59% □ 10-19% □ 60-69% □ 20-29% □ 70-79% □ 30-39% □ 80-89% □ 40-49% □ 90-100%	If IHC HER2/ERBB2 testing was performed, provide the percent of HER2/ERBB2 positive by IHC. If HER2/ERBB2 was negative, continue to the HER2/ERBB2 FISH questions. 3086980
70	IHC Intensity: HER2/ERBB2 Positivity Score for this Patient	□ 0 □ +2 □ +3	Provide the positivity score using the IHC intensity scale, found on the pathology/laboratory report. 2178402
71	Other Scale Used to Measure HER2/ERBB2 Positivity		If an additional scale was used to measure HER2/ERBB2 positivity, please describe the scale used. 3087479
72	Define method of calculation for HER2/ERBB2 Positivity		If a special method was used to calculate HER2/ERBB2 status, describe the method used. 3087487

#	Data Element	Entry Alternatives	Working Instructions
73	HER2/ERBB2 Status by FISH for this Patient	 □ Positive □ Negative □ Equivocal □ Indeterminate □ Performed but not available □ Not performed (skip to next molecular marker) 	If HER2/ERBB2 FISH testing was performed, provide the result of the test. If this test was not performed, select "not performed." 2854089
74	Number of HER2 FISH Signals for this Patient		If HER2 copy number testing was performed by FISH, provide the average number of HER2 FISH signals for this patient. If this test was not performed, leave this question blank and move to the next question. 3133738
75	Number of Centromere 17 Signals for this Patient		If Centromere 17 copy number testing was performed by FISH, provide the average number of Centromere 17 signals for this patient. If this test was not performed, leave this question blank and move to the next question. 3104295
76	Number of Cells Counted for HER2 & Centromere 17 by FISH for this Patient		Indicate the total number of cells counted by FISH for HER2 & Centromere 17 copy numbers. If these tests were not performed, leave this question blank and move to the next question. 3087902
77	HER2/Centromere 17 Ratio for this Patient		If HER2 copy number and Centromere 17 copy number testing was performed by FISH, provide the ratio of the outcomes of these tests. (For example, if both the HER2 copy number and the Centromere 17 copy number equal 2, the ratio would be 2÷2 or 1.0.) 2497552
78	Other Scale Used to Measure HER2 & Centromere 17 Positivity (Please Include Score)		If an additional scale was used to measure HER2 & Centromere 17 positivity, please describe the scale used. 3087923
79	Define Method of Calculation for HER2/ERBB2 Positivity if other than IHC or FISH		If a special method was used to calculate HER2 & Centromere 17 positivity, describe the method used. 3087929

New Tumor Event Information Complete this section if the patient had a new tumor event. If the patient did not have a new tumor event (or if the TSS does not know) indicate this in the question below, and the remainder of this section can be skipped.

#	Data Element	Entry Alternatives	Working Instructions
80	New Tumor Event After Initial Treatment?	☐ Yes ☐ No ☐ Unknown	Indicate whether the patient had a new tumor event (e.g. metastatic, recurrent, or new primary tumor) after initial treatment. 3121376 If the patient did not have a new tumor event or if this is unknown, the remaining questions can be skipped.
81	Type of New Tumor Event	☐ Locoregional Recurrence ☐ Distant Metastasis ☐ New Primary Tumor	Indicate whether the patient's new tumor event was a locoregional recurrence, a distant metastasis or a new primary tumor. A new primary tumor is a tumor with a different histology as the tumor submitted to TCGA. 3119721
82	Anatomic Site of New Tumor Event	☐ Lung ☐ Brain ☐ Other, specify ☐ Liver	Indicate the site of this new tumor event. 3108271
<u>83</u>	Other Site of New Tumor Event		If the site of the new tumor event is not included in the provided list, describe the site of this new tumor event. 3128033
<u>84</u>	Month of New Tumor Event	$\begin{array}{c ccccc} \square \ 01 & \square \ 04 & \square \ 07 & \square \ 10 \\ \square \ 02 & \square \ 05 & \square \ 08 & \square \ 11 \\ \square \ 03 & \square \ 06 & \square \ 09 & \square \ 12 \\ \end{array}$	If the patient had a new tumor event, provide the month of diagnosis for this new tumor event. 3104044

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#	Data Element	Entry Alternatives					Working Instructions
<u>85</u>	Day of New Tumor Event	□ 01 □ 02 □ 03 □ 04 □ 05 □ 06 □ 07	□ 08 □ 09 □ 10 □ 11 □ 12 □ 13	☐ 14 ☐ 15 ☐ 16 ☐ 17 ☐ 18 ☐ 19	□ 20 □ 21 □ 22 □ 23 □ 24 □ 25	□ 26 □ 27 □ 28 □ 29 □ 30 □ 31	If the patient had a new tumor event, provide the day of diagnosis for this new tumor event. 3104042
<u>86</u>	Year of New Tumor Event						If the patient had a new tumor event, provide the year of diagnosis for this new tumor event. 3104046
<u>87</u>	Number of Days from Date of Initial Pathologic Diagnosis to Date of New Tumor Event After Initial Treatment	_					Provide the number of days from the date the patient was initially diagnosed pathologically with the disease to the date of new tumor event after initial treatment. 3392464 Only provide Interval data if you have received permission from the NCI to provide time intervals as a substitute for requested dates on this form.
88	Additional treatment for New Tumor Event: Surgery	☐ Yes ☐ No ☐ Unkno	wn				Using the patient's medical records, indicate whether the patient had surgery for the new tumor event in question. 3427611
<u>89</u>	Month of Additional Surgery for New Tumor Event	□ 01 □ 02 □ 03	□ 04 □ 05 □ 06	I	□ 07 □ 08 □ 09	□ 10 □ 11 □ 12	If the patient had surgery for the new tumor event, provide the month this surgery was performed. 3427612
90	Day of Additional Surgery for New Tumor Event	□ 01 □ 02 □ 03 □ 04 □ 05 □ 06 □ 07	□ 08 □ 09 □ 10 □ 11 □ 12 □ 13	☐ 14 ☐ 15 ☐ 16 ☐ 17 ☐ 18 ☐ 19	☐ 20 ☐ 21 ☐ 22 ☐ 23 ☐ 24 ☐ 25	□ 26 □ 27 □ 28 □ 29 □ 30 □ 31	If the patient had surgery for the new tumor event, provide the day this surgery was performed. 3427613
91	Year of Additional Surgery for New Tumor Event					-	If the patient had surgery for the new tumor event, provide the year this surgery was performed. $\underline{3427614}$
<u>92</u>	Number of Days from Date of Initial Pathologic Diagnosis to Date of Additional Surgery for New Tumor Event					-	Provide the number of days from the date the patient was initially diagnosed pathologically with the disease described on this form to the date of additional surgery for new tumor event (loco-regional). 3008335 Only provide Interval data if you have received permission from the NCI to provide time intervals as a substitute for requested dates on this form.
<u>93</u>	Additional treatment for New Tumor Event: Radiation Therapy	☐ Yes ☐ No ☐ Unkno	wn				Indicate whether the patient received radiation treatment for this new tumor event. 3427615
94	Additional treatment for New Tumor Event: Pharmaceutical Therapy	☐ Yes ☐ No ☐ Unkno	wn				Indicate whether the patient received pharmaceutical treatment for this new tumor event. 3427616
Nev	v Tumor Event: Molecula	r Markers	Used for	Tume	or Progno	sis	
95	Estrogen Receptor (ER) Status by IHC for this patient	☐ Positiv ☐ Negati ☐ Indete ☐ Perfor	☐ Positive (1%-100%) ☐ Negative (0%) ☐ Indeterminate ☐ Performed but not available ☐ Not performed (skip to next molecular marker)				If IHC estrogen receptor testing was performed, provide the result of the test. If this test was not performed, selected "not performed," and continue to the progesterone receptor questions. 3131865
96	IHC ER Percent Positive for this patient	□ <10% □ 10-199 □ 20-299 □ 30-399 □ 40-499	(1-9%) % % %		□ 50-59% □ 60-69% □ 70-79% □ 80-89% □ 90-1009		If IHC estrogen receptor testing was performed, provide the percent of estrogen receptor positive by IHC. 3131869

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#	Data Element	Entry Alternatives		Working Instructions
97	IHC Intensity: Scale Used to determine ER Positivity for this Patient	☐ 4 Point Scale		Using the pathology/laboratory report, indicate the intensity scale used for the estrogen receptor positivity score. 3203082
98	IHC Intensity: ER Positivity Score for this patient	□ 0 □ +1 □ +2	□ +3 □ +4	Provide the positivity score using the IHC intensity scale, found on the pathology/laboratory report. 3131873
99	IHC Intensity: Other Method Used to Determine ER Positivity For this Patient			If another scale was used to measure the estrogen receptor positivity, please describe the scale used. 3131877
100	Define Method of Calculation for ER Positivity if Other than IHC			If a special method was used to calculate estrogen receptor status (e.g. dextran coated charcoal), describe the method used. 3131881
101	Progesterone Receptor (PR) Status by IHC for this patient	☐ Positive (1%-100%) ☐ Negative (0%) ☐ Indeterminate ☐ Performed but not av ☐ Not performed (skip t		If IHC progesterone receptor testing was performed, provide the result of the test. If this test was not performed, select "not performed," and continue to the HER2/ERBB2 IHC questions. 3131884
102	IHC PR Percent Positive for this patient	□ <10% (0-9%) □ 10-19% □ 20-29% □ 30-39% □ 40-49%	□ 50-59% □ 60-69% □ 70-79% □ 80-89% □ 90-100%	If IHC progesterone receptor testing was performed, provide the percent of progesterone receptor positive nuclei by IHC. 3131891
103	IHC Intensity: Scale Used for PR Positivity	☐ 4 Point Scale ☐ 3 Point Scale		Using the pathology/laboratory report, indicate the intensity scale used for the progesterone receptor positivity score. 3203085
104	IHC Intensity: PR Positivity Score for this Patient	□ 0 □ +1 □ +2	□ +3 □ +4	Provide the positivity score using the IHC intensity scale, found on the pathology/laboratory report. Only answer this question if PR status is considered positive; if the PR status was negative, continue to the HER2/ERBB2 IHC questions. 3131988
105	IHC Intensity: Other Method Used to Determine PR Positivity			If another scale was used to measure the progesterone receptor positivity, please describe the scale used. 3131992
106	Define Method of Calculation for Positivity if Other Than IHC			If a special method was used (other than IHC) to calculate progesterone receptor status (e.g. dextran coated charcoal), describe the method used. 3131993
107	HER2/ERBB2 Status by IHC for this Patient	☐ Positive ☐ Negative ☐ Equivocal ☐ Indeterminate ☐ Performed but not available ☐ Not performed(skip to next molecular marker)		If IHC HER2/ERBB2 testing was performed, provide the result of the test. If this test was not performed, select "not performed," and continue to the HER2/ERBB2 FISH questions. 3131997
108	IHC HER2/ERBB2 Percent Positive for this patient	□ <10% □ 10-19% □ 20-29% □ 30-39% □ 40-49%	□ 50-59% □ 60-69% □ 70-79% □ 80-89% □ 90-100%	If IHC HER2/ERBB2 testing was performed, provide the percent of HER2/ERBB2 positive by IHC. If HER2/ERBB2 was negative, continue to the HER2/ERBB2 FISH questions. 3132322
109	IHC Intensity: HER2/ERBB2 Positivity Score for this Patient	□ 0 □ +1	□ +2 □ +3	Provide the positivity score using the IHC intensity scale, found on the pathology/laboratory report. 3132444
110	Other Scale Used to Measure HER2/ERBB2 Positivity			If an additional scale was used to measure HER2/ERBB2 positivity, please describe the scale used. 3132448

#	Data Element	Entry Alternatives	Working Instructions
111	Define method of calculation for HER2/ERBB2 Positivity		If a special method was used to calculate HER2/ERBB2 status, describe the method used. 3132452
112	HER2/ERBB2 Status by FISH for this Patient	☐ Positive ☐ Negative ☐ Equivocal ☐ Indeterminate ☐ Performed but not available ☐ Not performed (skip to next molecular marker)	If HER2/ERBB2 FISH testing was performed, provide the result of the test. If this test was not performed, select "not performed." 3132455
113	Number of HER2 FISH Signals for this Patient		If HER2 copy number testing was performed by FISH, provide the average number of HER2 FISH signals for this patient. If this test was not performed, leave this question blank and move to the next question. 3133734
114	Number of Centromere 17 Signals for this Patient		If Centromere 17 copy number testing was performed by FISH, provide the average number of Centromere 17 signals for this patient. If this test was not performed, leave this question blank and move to the next question. 3132887
115	Number of Cells Counted for HER2 & Centromere 17 by FISH for this Patient		Indicate the total number of cells counted by FISH for HER2 & Centromere 17 copy numbers. If these tests were not performed, leave this question blank and move to the next question. 3132899
116	HER2/Centromere 17 Ratio for this Patient		If HER2 copy number and Centromere 17 copy number testing was performed by FISH, provide the ratio of the outcomes of these tests. (For example, if both the HER2 copy number and the Centromere 17 copy number equal 2, the ratio would be 2÷2 or 1.0.) 3132903
117	Other Scale Used to Measure HER2 & Centromere 17 Positivity (Please Include Score)		If an additional scale was used to measure HER2 & Centromere 17 positivity, please describe the scale used. 3132907
118	Define Method of Calculation for		If a special method was used to calculate HER2 & Centromere 17 positivity, describe the method used. 3132910

Print Name

Date

Principal Investigator or Designee Signature