Completed By:	lissue Sou	lissue Source Site (ISS) Name: ISS Identifier: ISS Unique Patient #:					
upsations continued cont	Completed By: Completion Date (MM/DD/YYYY):						
Please note that time intervals must be recorded in place of feets where designed throughout this form if wou have selected "yes" in the box to the left. Note 1: Provided time intervals as a substitute for requested dates on this form? Yes No No Note 2: Only provide interval date if you have received permission from the NCI to provide time intervals as a substitute for requested dates on this form? No Note 2: Only provide interval date if you have received permission from the NCI to provide time intervals as a substitute for requested dates on this form. 3233305 Indicate the reason for submission of this follow-up form. It additional new Tumor Event Scheduled (Routine) Follow-up Submission Indicate the reason for submission of this follow-up form. It additional new tumor. 6.1333 Indicate whether the patient is lost to follow-up as defined by the ACOS Commission on cancer. This only includes cases where updated information has not been collected within the last 15 months. If the patient is deceased and a TCGA Follow-up Form has not yet been completed, the answer to this questions should be "No" of and the remaining updicable questions should be "No" of and the remaining updicable questions should be "No" of and the remaining updicable questions should be "No" of and the remaining updicable questions should be "No" of and the remaining updicable questions should be "No" of and the remaining updicable questions should be "No" of and the remaining updicable questions should be "No" of an experience of the patient is deceased and a TCGA Follow-up Form has not yet been completed. Yes	Form Notes: A Follow-up Form is to be completed for any of the following reasons: 1) For each additional new tumor event identified at the time of enrollment or follow-up submission; or 2) 12 months after a case is shipped to the Biospecimen Core Resource (BCR) for cases that have qualified. All information provided on this form includes activity from the "Date of Last Contact" provided on the TCGA Enrollment Form to the most recent date of contact with the patient. This form should only be completed by the Tissue Source Site if updated information can be provided to TCGA. Questions regarding this form should be directed to the Tissue Source Site's (TSS) primary Clinical Outreach Contact at the BCR. The following definitions for the use of "Unknown" and "Not Evaluated" on this form are as follows: Unknown: This answer option should only be selected if the TSS cannot answer the question because the answer is not known at the TSS. 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 the reason why the answer is unknown. Not evaluated: This answer option should be selected by the TSS if it is known that the information being requested cannot be obtained due to the test not being						
Please note that time intervals must be recorded in place of deswhere designated throughout this place where designated throughout this form from the NCI to provide time intervals as a substitute for requested dates on this form? Yes	Ouestion#	Data Element Label	Data Entry Alternatives	CDE ID With Working Instructions			
Reason For Follow-up Form Submission Scheduled (Routine) Follow-up Submission Indicate the reason for submission of this follow-up form. If scheduled follow-up, complete entire form. If additional new tumor event, complete only questions pertaining to new tumor. Additional New Tumor Event Reading the patient is lost to follow-up as defined by the ACOS Commission on Cancer. This only includes cases where updated information has not been collected within the last 15 months. If the patient is lost to follow-up, the remaining questions may be left unanswered. No		Has this TSS received permission from the NCI to provide time intervals as a substitute for	Yes	Please note that time intervals must be recorded in place of dates where designated throughout this form if you have selected "yes" in the box to the left. Note 1: Provided time intervals must begin with the date of initial pathologic diagnosis. (i.e., biopsy or resection) Note 2: 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.			
Indicate whether the patient is lost to follow-up as defined by the ACOS Commission on Cancer. This only includes cases where updated information has not been collected within the last 15 months. If the patient is lost to follow-up, the remaining questions may be left unanswered. Note: If the patient is deceased and a TCGA Follow-up Form has not been collected within the last 15 months. If the patient is lost to follow-up, the remaining questions may be left unanswered. Note: If the patient is deceased and a TCGA Follow-up Form has not been collected within the last 15 months. If the patient is lost to follow-up, the remaining questions may be left unanswered. Note: If the patient is deceased and a TCGA Follow-up Form has not been collected within the last 15 months. If the patient is lost to follow-up is not been collected within the last 15 months. If the patient is lost to follow-up is not been collected within the last 15 months. If the patient is lost to follow-up is not been collected within the last 15 months. If the patient is lost to follow-up and be to be patient is lost to follow-up and be to make the cases where updated information has not been collected within the last 15 months. If the patient is lost to follow-up and phermital file file patient is lost to follow-up and be completed. Primary Treatment Yes	2	•		Indicate the reason for submission of this follow-up form. If scheduled follow-up, complete entire form. If additional new tumor event, complete only			
Primary Treatment Adjuvant Post-operative Radiation Therapy Adjuvant Post-operative Radiation No Operative radiation therapy. Note: If the patient did have adjuvant radiation, the Radiation Supplemental Form should be completed. Yes Operative Radiation Supplemental Form should be completed. Yes Operative Pharmaceutical Therapy No Operative Pharmaceutical Therapy Note: If the patient had adjuvant/ post-operative pharmaceutical therapy. Note: If the patient did have adjuvant pharmaceutical therapy, Note: If the patient did have adjuvant pharmaceutical therapy, the Pharmaceutical Supplemental Form should be completed. Measure of Success of Outcome at the Completion of Initial First Course Treatment (surgery and adjuvant therapies) Stable Disease Not Applicable Course Treatment (surgery and/or adjuvant therapies). Patient Status Deceased Deceased Supplemental Form should be completed Stable Disease Onte of Not Applicable Course treatment (surgery and/or adjuvant therapies).	3	Is This Patient Lost to Follow-up?		Indicate whether the patient is lost to follow-up as defined by the ACoS Commission on Cancer. This only includes cases where updated information has not been collected within the last 15 months. If the patient is lost to follow-up, the remaining questions may be left unanswered. Note: If the patient is deceased and a TCGA Follow-up Form has not yet been completed, the answer to this question should be "No" and the remaining			
Adjuvant Post-operative Radiation Therapy No	Primary Trea	atment					
Adjuvant Post-operative Pharmaceutical Therapy Measure of Success of Outcome at the Completion of Initial First Course Treatment (surgery and adjuvant therapies) Patient Status Ves	4		□ No	Indicate whether the patient had adjuvant/ post- operative radiation therapy. Note: If the patient did have adjuvant radiation, the			
the Completion of Initial First Course Treatment (surgery and adjuvant therapies) Stable Disease Deceased Not Applicable Unknown Provide the patient's response to their initial first course treatment (surgery and/or adjuvant therapies). Patient Status Provide the patient's response to their initial first course treatment (surgery and/or adjuvant therapies). Patient Status Deceased Provide the patient's response to their initial first course treatment (surgery and/or adjuvant therapies). Patient Status Deceased Deceased Indicate whether the patient was living or deceased	5	- · · · · · · · · · · · · · · · · · · ·	□ No	Indicate whether the patient had adjuvant/ post- operative pharmaceutical therapy. Note: If the patient did have adjuvant pharmaceutical therapy, the Pharmaceutical			
7 Vital Status Deceased 2939553 Indicate whether the patient was living or deceased		the Completion of Initial First Course Treatment (surgery and adjuvant therapies)	Stable Disease Not Applicable	Provide the patient's response to their initial first course treatment (surgery and/or adjuvant			
7 Vital Status	Patient Statu	IS		2030553			
	7	Vital Status	Living Deceased	Indicate whether the patient was living or deceased			

Tissue Source Site (TSS) Name: _____ TSS Identifier: ____ TSS Unique Patient #: ____

Question#	Data Element Label	Data Entry Alternatives	CDE ID With Working Instructions		
-	Contact(or date of death, if deceased)	Data Entry Atternatives	CDE 1D WITH WORKING HISTIACTIONS		
8	Month Of Last Contact	□□ (MM)	2897020 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). Note: Do not answer this question if the patient is deceased.		
9	Day Of Last Contact	□□ (DD)	2897022 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). Note: Do not answer this question if the patient is deceased.		
10	Year Of Last Contact		2897024 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). Note: Do not answer this question if the patient is deceased.		
11	Number of Days from Date of Initial Pathologic Diagnosis to Date of Last Contact		3008273 Provide the number of days from the date the patient was initially diagnosed pathologically with the disease to the date of Last Contact. Note 1: Do not answer this question if the patient is deceased. Note 2: 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.		
Date of Deat	th	Not Applicable (Patient is Alive)			
			2897026		
12	Month of Death	□□ (MM)	If the patient is deceased, provide the month of death.		
13	Day of Death	□□ (DD)	2897028 If the patient is deceased, provide the day of death.		
14	Year of Death		2897030 If the patient is deceased, provide the year of death.		
15	Number of Days from Date of Initial Pathologic Diagnosis to Date of Death		3165475 Provide the number of days from the date the patient was initially diagnosed pathologically with the disease to the date of Death. Note: 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.		
16	Tumor Status	Tumor Free Unknown Tumor Status	2759550 Indicate whether the patient was tumor/disease free from the tumor submitted for TCGA at the date of last contact or death.		
New Tumor	New Tumor Event: Please verify that new tumor event information has not previously been reported on the Enrollment Form or on a prior Follow-up Form				
17	New Tumor Event After Initial Treatment?	☐ Yes ☐ No ☐ Unknown	3121376 Indicate whether the patient had a new tumor event (e.g. metastatic, recurrent, or new primary tumor) after their initial treatment for the tumor submitted to TCGA. Note: If the patient had multiple new tumor events, a follow-up form should be completed for each new tumor event.		

Tissue Source Site (TSS) Name: ______ TSS Identifier: _____ TSS Unique Patient #: _____

Question#	Data Element Label	Data Entry Alternatives		CDE ID With Working Instructions		
-	Tumor Event After Initial Treatment	•		<u> </u>		
18	Month of New Tumor Event After Initial Treatment	(MM)		3104044 If the patient had a new tumor event, provide the month of diagnosis for this new tumor event.		
19	Day of New Tumor Event After Initial Treatment	□□ (DD)		3104042 If the patient had a new tumor event, provide the day of diagnosis for this new tumor event.		
20	Year of New Tumor Event After Initial Treatment	(YYYY)		3104046 If the patient had a new tumor event, provide the year of diagnosis for this new tumor event.		
21	Number of Days from Date of Initial Pathologic Diagnosis to Date of New Tumor Event After Initial Treatment			3392464 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. Note: 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.		
22	Additional Surgery for New Tumor Event Loco-Regional	☐ Yes ☐ No	Unknown	3008755 Using the patient's medical records, indicate whether the patient had surgery for the new locoregional tumor event in question.		
Date Additional Surgery for New Tumor Event – Loco–Regional Not Applicable						
23	Month of Additional Surgery for New Tumor Event Loco–Regional	ПП (ММ)		2897032 If the patient had surgery for the new loco-regional tumor event, provide the month of surgery for this new loco-regional tumor event.		
24	Day of Additional Surgery for New Tumor Event Loco–Regional	[DD]		2897034 If the patient had surgery for the new loco-regional tumor event, provide the day of surgery for this new loco-regional tumor event.		
25	Year of Additional Surgery for New Tumor Event Loco–Regional			2897036 If the patient had surgery for the new loco-regional tumor event, provide the year of surgery for this new loco-regional tumor event.		
26	Number of Days from Date of Initial Pathologic Diagnosis to Date of Additional Surgery for New Tumor Event Loco–Regional		-	3408572 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 (locoregional). Note: 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.		
27	Residual Tumor after Surgery for New Tumor Event Loco–Regional	RX RO	□ R1 □ R2	33104061 If the patient had surgery for the new loco-regional tumor event, provide the status of any residual tumor after this surgery.		
28	Additional Surgery for New Tumor Event Metastasis	Yes No	Unknown	3008757 Using the patient's medical records, indicate whether the patient had surgery for the new metastatic tumor event in question.		
29	Site of Additional Surgery for New Tumor Event Metastasis	Liver Lung	Lymph Nodes Other	1611 Indicate the location of additional surgery for the new metastatic tumor event which has spread from original tumor located in the large intestine or rectum.		

Tissue Source Site (TSS) Name:		TSS Identifier: TS	S Unique Patient #:	
Question#	Data Element Label	Data Entry Alternatives	CDE ID With Working Instructions	
Date of Add	itional Surgery for New Tumor Event N	Metastasis Not Applicable (No Surgical Procedure for	Metastatic Tumor Event)	
30	Month of Additional Surgery for New Tumor Event Metastasis	(MM)	2897038 If the patient had surgery for the new metastatic tumor event, provide the month of surgery for this new metastatic tumor event.	
31	Day of Additional Surgery for New Tumor Event Metastasis	□□ (DD)	2897040 If the patient had surgery for the new metastatic tumor event, provide the day of surgery for this new metastatic tumor event.	
32	Year of Additional Surgery for New Tumor Event Metastasis		2897042 If the patient had surgery for the new metastatic tumor event, provide the year of surgery for this new metastatic tumor event.	
33	Number of Days from Date of Initial Pathologic Diagnosis to Date of Additional Surgery for New Tumor Event Metastasis		3408682 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 (metastasis) Note: 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.	
34	Residual Tumor after surgery for New Tumor Event Metastasis	□ RX □ R1 □ R0 □ R2	3104081 If the patient had surgery for the new metastatic tumor event, provide the status of any residual tumor after this surgery.	
35	Additional Treatment of New Tumor Event Radiation Therapy	Yes No Unknown	3008761 Indicate whether the patient received radiation treatment for this new tumor event.	
36	Additional Treatment of New Tumor Event Pharmaceutical Therapy	☐ Yes ☐ No ☐ Unknown	2650646 Indicate whether the patient received pharmaceutical treatment for this new tumor event.	
37	Measure of Success of Outcome at the Completion of this Follow-up Submission	Progressive Disease Complete Response Partial Response Unknown	3104050 Provide the patient's outcome of treatment up to the point of the current follow-up data submission	
Comments:				
Principal Investigator Name: Principal Investigator Signature: Date Signed (MM/DD/YYYY):				