Tissue Sour	rce Site (TSS) Name:	TSS Identifier:	TSS Unique Patient #:
Completed By:		Completion Date (MM/DD/YYYY):	
Form Notes: Tissue Source Site (TSS) acknowledges that the Biospecimen Core Resource (BCR) may confirm that the diagnosis of the frozen biospecimen is consistent with the primary diagnosis reported by the TSS through histopathology examination in the BCR laboratory. If the BCR identifies a possible discrepancy, the TSS authorizes the BCR to report these patient results to the TSS by means of a formal report in confidential email format for the quality assurance program of the TSS to address.			
Question #	Data Element Label	Data Entry Alternatives	CDE ID With 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	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.
2	Tumor Identifier		3288096 Provide the TSS unique tumor ID. If multiple pieces of tumor are submitted, each tumor needs a unique ID. Note: If submitting multiple pieces of the same primary tumor for this case, complete the tumor information for each piece of tumor sent to the BCR.
3	Lung Squamous: Histologic Subtype	Papillary Squamous Cell Carcinoma Clear Cell Squamous Cell Carcinoma Small Cell Squamous Cell Carcinoma Basaloid Squamous Cell Carcinoma Squamous Cell Carcinoma, Not Otherwise Specified (NOS)	3081934 Indicate the histologic subtype for the lung squamous cell tumor sample being submitted to TCGA. Note 1: The listed histologies are the only squamous cell histologies being accepted for the TCGA Project. Note 2: Squamous Cell Carcinoma tumors are allowed a minor component of < or = 5% Adenocarcinoma.
4	Lung Adeno: Histologic Subtype	Adenocarcinoma, Mixed Subtype Acinar Adenocarcinoma Papillary Adenocarcinoma Bronchioloalveolar Carcinoma, Mucinous Bronchioloalveolar Carcinoma, Non-Mucinous Solid Pattern Predominant Adenocarcinoma Micropapillary Adenocarcinoma Fetal Adenocarcinoma Mucinous Cystadenocarcinoma Mucinous (Colloid) Adenocarcinoma Signet Ring Adenocarcinoma Clear Cell Adenocarcinoma Adenocarcinoma, Not Otherwise Specified (NOS)	3081934 Indicate the histologic subtype for the lung adenocarcinoma tumor sample being submitted to TCGA. Note: The listed histologies are the only adenocarcinoma histologies being accepted for the TCGA Project.
5	Tumor Type	Primary	3288124 Confirm that the tumor being submitted to TCGA is a primary untreated malignant biospecimen.
6	Tumor Site (Anatomic Site of Frozen Biospecimen)	Right Upper Lobe Lung Right Middle Lobe Lung Right Lower Lobe Lung Left Upper Lobe Lung Left Lower Lobe Lung Bronchus Other (please specify below)	2008006 Indicate the tumor site (anatomic site of the frozen tumor) submitted for TCGA.

Question #	Data Element Label	Data Entry Alternatives	CDE ID With Working Instructions
7	Data Element Laber		3320289
	Other Anatomic Site of Frozen		If the anatomic site of the frozen biospecimen is not included in the
	Biospecimen		provided list, indicate the other anatomic site of the frozen tumor submitted to TCGA.
Date of Cance	er Sample Procurement		Submitted to TCGA.
Date of cance			3008197
8	Month of Cancer Sample	□ □ (MM)	Provide the month of the procedure performed to obtain the
	Procurement		malignant tissue submitted for TCGA.
_	Day of Cancer Sample		3008195
9	Procurement	□□ (DD)	Provide the day of the procedure performed to obtain the malignant
			tissue submitted for TCGA. 3008199
10	Year of Cancer Sample		Provide the year of the procedure performed to obtain the
	Procurement		malignant tissue submitted for TCGA.
			3288495
			Provide the number of days from the date the patient was initially
	Number of Days from Date of		diagnosed pathologically with the disease described on this form to
11	Initial Pathologic Diagnosis to Date of Cancer Sample		the date of the procedure that produced the malignant sample submitted for TCGA.
	Procurement		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.
		☐ Cytology	
		Fine Needle Aspiration Biopsy	
12	Method of Cancer Sample	Incisional Biopsy	3103514
12	Procurement	☐ Excisional Biopsy	Indicate the procedure performed to obtain the malignant tissue submitted for TCGA.
			Submitted for 1887.
		Tumor Resection	
		Other Method (please specify below)	
	Other Method of Cancer Sample		2006730
13	Procurement		If the procedure performed to obtain the malignant tissue is not
			included in the provided list, specify the procedure.
14	Country of Cancer Sample		3203072 Provide the country where the tissue submitted for TCGA was
1.	Procurement		procured.
		☐ American Indian or Alaska Native	2192199
		A person having origins in any of the	Provide the patient's race using the defined categories.
		original peoples of North and South	
		America (including Central America), and who maintains tribal affiliation or	
	Race	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.	
15		☐ White	
15		A person having origins in any of the	
		original peoples of the far Europe, the Middle East, or North Africa.	
		☐ Black or African American	
		A person having origins in any of any of	
		the black racial groups of Africa. Terms	
		such as "Haitian" or "Negro" can be	
		used in addition to "Black or African	
		American." ☐ Native Hawaiian or other Pacific Islander:	
		A person having origins in any of the	
		original peoples of Hawaii, Guam,	
		Samoa, or other Pacific Islands.	
		□ Not Reported: Not provided or available.	
		Unknown: Could not be determined.	

Question #	Data Element Label	Data Entry Alternatives	CDE ID With Working Instructions
16	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 Could not be determined or unsure.	2192217 Provide the patient's ethnicity using the defined categories.
17	Vessel Used	☐ Cryovial ☐ Cryomold ☐ Cassette ☐ Biospecimen Storage Bag ☐ Other vessel (please specify below)	3081940 Indicate the type of vessel used to ship the tissue to the Biospecimen Core Resource (BCR) for TCGA.
18	Other Vessel Used		3288137 If the vessel used to ship tissue to the Biospecimen Core Resource (BCR) is not included in the provided list, specify the other type of vessel used.
19	Weight of Frozen Tumor		3081946 Provide the weight of the tumor sample submitted for TCGA. Note: (0.2cm³ (0.6cm * 0.6cm * 0.6cm) = ~200mg
20	Is Tumor Sample being Submitted for Laser Cryo Enrichment (LCE) Processing?	Yes No	3288488 Indicate if the tumor sample being submitted is to be processed using Laser Cryo Enrichment (LCE).
21	Tumor Nuclei %		2841225 Provide the percent of tumor nuclei for the sample submitted for TCGA. Note: Check with the BCR to confirm the current acceptable TCGA metrics.
22	Tumor Necrosis %		2841237 Provide the percent of necrosis for the sample submitted for TCGA. Check Note: Check with the BCR to confirm the current acceptable TCGA metrics.
23	Was sample prescreened at site?	Yes No	3081942 Indicate whether the sample submitted to the BCR was prescreened at the TSS.
24	Will Top Slide be submitted to the BCR?	Yes No	3081944 Indicate whether a physical top slide for the sample submitted to the BCR will be shipped with the tissue sample. Note: Top slide definition: Slide cut directly from frozen biospecimen = mirror image of inked surface.
25	Will Digital Slide Image be submitted to the BCR?	Yes No	3081948 Indicate whether a digital slide image for the sample submitted to the BCR will be shipped with the tissue sample. Note: Physical top slides are preferred.
26	Top Slide / Digital Slide Image ID #		2321277 Provide the slide ID for the physical top slide OR the digital slide image being sent to the BCR.
Normal Information: A normal control must be present to qualify			
27	Normal Identifier		3288138 Provide the TSS unique normal ID. If multiple normal control samples are submitted, each normal control needs a unique ID.
28	Type of Normal Control	☐ Whole Blood ☐ Lymphocytes (Buffy Coat) ☐ Extracted DNA from Blood ☐ Normal Tissue	3081936 Indicate the type of normal control submitted for this case. Note: Whole blood is preferred. Normal tissue is only allowable with NCI approval.

Question #	Data Element Label	Data Entry Alternatives	CDE ID With Working Instructions
29	Anatomic Site of Normal Tissue	Right Upper Lobe Lung Right Middle Lobe Lung Right Lower Lobe Lung Left Upper Lobe Lung Left Lower Lobe Lung Bronchus Other (please specify below)	3081938 If the normal control type is normal tissue, indicate the anatomic site of the non-neoplastic control tissue submitted for TCGA. Note: If normal tissue is being submitted, site matched is preferred.
30	Other Anatomic Site of Normal Tissue		3288189 If the normal control type is normal tissue and the anatomic site is not included in the provided list, specify the site of the non-neoplastic control.
31	Proximity of Normal Tissue to Tumor	Distal (≥ 2 cm) from the primary tumor Adjacent (≤2 cm) from the primary tumor	3088708 If normal tissue is being submitted, confirm that the normal tissue is ≥ 2.0cm from the primary lung tumor. Note: Adjacent and/or tissue of unknown proximity are not accepted for this tissue type.
Date of Norm	al Sample Procurement		
32	Month of Normal Sample Procurement	ПП (ММ)	3288195 Provide the month of the procedure performed to obtain the normal control sample for TCGA.
33	Day of Normal Sample Procurement	□□ (DD)	3288196 Provide the day of the procedure performed to obtain the normal control sample for TCGA.
34	Year of Normal Sample Procurement	□□□ (YYYY)	3288197 Provide the year of the procedure performed to obtain the normal control sample for TCGA.
35	Number of Days from Date of Initial Pathologic diagnosis to Date of Normal Sample Procurement		3288496 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 the procedure that produced the normal control sample submitted for TCGA. 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.
36	Method of Normal Sample Procurement	Blood Draw Cytology Fine Needle Aspiration Biopsy Incisional Biopsy Excisional Biopsy Tumor Resection Other Method (please specify below)	3288147 Indicate the procedure performed to obtain the normal sample submitted for TCGA.
37	Other Method of Normal Sample Procurement		3288151 If the procedure performed to obtain the normal sample is not included in the provided list, specify the procedure.
38	Normal Slide ID #		3288217 If the normal control type is normal tissue, provide the slide ID for the physical top slide OR the digital slide image of the normal control being sent to the BCR.
39	Extracted DNA Quantity		3288185 If the normal control type is extracted DNA from blood, provide the quantity (µg) of the normal control sample sent to the BCR for TCGA.
40	Extracted DNA Quantification Method		3288186 If the normal control type is extracted DNA from blood, provide the quantification method of the normal control sample sent to the BCR for TCGA.
41	Extracted DNA Concentration		3288187 If the normal control type is extracted DNA from blood, provide the concentration (μg/ μL) of the normal control sample sent to the BCR

Question #	Data Element Label	Data Entry Alternatives	CDE ID With Working Instructions
			for TCGA.
			3288188
42	Extracted DNA Volume		If the normal control type is extracted DNA from blood, provide the
			volume (μL) of the normal control sample sent to the BCR for TCGA.
Verification: I	By providing the information below, th	ne Principal Investigator acknowledges that the in	nformation provided by the institution is true and correct and has been
quality contro	lled.	,	,
			3288225
43	Name of Pathologist		Provide the name of the Pathologist that reviewed and prescreened
			the top slide and provided the information for all previous sections.
			3288224
44	Date of Pathologist Review	(MM/DD/YYYY)	Provide the date of the pathology prescreening review performed
		(IVIIVI/DD/TTTT)	by the TSS pathologist above.
			3288497
			Provide the number of days from the date the patient was initially diagnosed pathologically with the disease described on this form to
	Number of Days from Date of		the date of the pathological review performed as part of the
45	Initial Pathologic Diagnosis to		submission process for TCGA.
	Date of Pathological Review		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.
			3288520
4.6	Percent Tumor Nuclei meets	☐ Yes	Confirm that the malignant sample submitted to the BCR meets the
46	TCGA metrics?	□ No	current tumor nuclei metrics for TCGA.
		l No	Check with the BCR to confirm the current acceptable TCGA metrics.
			3288524
47	Percent Tumor Necrosis meets	☐ Yes	Confirm that the malignant sample submitted to the BCR meets the
47	TCGA metrics?	□ No	current necrosis metrics for TCGA.
			Check with the BCR to confirm the current acceptable TCGA metrics.
			3288300
			Confirm that the diagnosis provided on this CQCF for the tumor
			sample being submitted to TCGA is consistent with the diagnosis
			found on the patient's pathology report for the tumor being sent to
			the BCR.
	Is the histologic diagnosis on the		Note: The diagnosis is considered to be consistent if at least one of the following criteria are met:
			Diagnosis on the CQCF is identical to the pathology report for
	CQCF (as determined by the TSS pathology review of the TCGA frozen section top slide) consistent with the histology listed in the final diagnosis on the pathology report?	Yes (skip related question below).	the tumor being sent to the BCR.
48			2) Diagnosis on the CQCF includes at least one of the subtypes
10		│ □ _{No}	listed on the pathology report and all subtypes on the pathology
			report are acceptable for TCGA.
			3) Diagnosis on the CQCF is "histology, NOS" (i.e.
			Adenocarcinoma, NOS) and the pathology report lists a specific
			subtype within the same histological group.
			4) Diagnosis on the CQCF indicates "Mixed Subtype" and the
			pathology report lists two or more acceptable subtypes, provided
			that percent subtype(s) meet applicable TCGA disease-specific
			requirements.
		Macrodissection performed at TSS to	3288315
		select for region containing an acceptable	If the diagnosis provided on this form is not consistent with the diagnosis found on the patient's pathology report for the tumor
		TCGA diagnosis	being submitted for TCGA, specify a reason for this inconsistency.
		–	Note: If a TSS pathology review of the TCGA committed sample
	If the diagnosis on this form is not	Pathology analysis at TSS determined a	resulted in a different histological subtype than what is
49	consistent with the provided	specific histological subtype different from	documented on the pathology report, an amendment to the
	pathology report, indicate the	original pathology report (see note at right)	pathology report should be submitted when the sample is shipped
	reason for the inconsistency.	Pathology review of frozen section for	to the BCR; or in the absence of an amended pathology report, the
		TCGA determined histological subtype	TSS must complete and submit an electronic copy of the "TCGA
		different from the pathology report (see	Pathologic Diagnosis Discrepancy Form." In the case of diagnosis
		note at right)	modifications, institution protocol should be followed for proper
			quality assurance
50	De-Identified Pathology Report	Yes	3288292
	Submitted?		Confirm that a de-identified pathology report will be sent to BCR
	Sabilitieu:	L No	prior to or with the shipment of the physical samples.
	History of Neo-Adjuvant	□ No	3382737
51	Treatment to Tumor Specimen		Indicate whether the patient received therapy for this cancer prior
- -	Submitted for TCGA	Radiation Prior to Sample	to sample procurement of the tumor submitted for TCGA. If the
		Procurement	patient did receive treatment for this cancer prior to procurement,

Question #	Data Element Label	Data Entry Alternatives	CDE ID With Working Instructions
		Pharmaceutical Treatment Prior to Sample Procurement Both Pharmaceutical and Radiation Treatment Prior to Sample Procurement	the TSS should contact the BCR for further instructions. Note: Systemic treatment and certain localized therapies (those administered to the same site as the TCGA submitted tissue) given prior to procurement of the sample submitted for TCGA are exclusionary.
52	Has the Patient Had Any Prior Cancer Diagnosed?	☐ No ☐ History of Prior Malignancy ☐ History of Synchronous / Bilateral Malignancy	3382736 Indicate whether the patient has a history of prior malignancies. Note 1: If this question cannot be answered because the answer is unknown, the case will be excluded from TCGA. Note 2: If the patient has any history of prior malignancies, including synchronous or bilateral malignancies, please complete an "Other Malignancy Form" for each malignancy diagnosed prior to the procurement of the tissue submitted for TCGA. If the patient has a history of multiple diagnoses of basal and/or squamous cell skin cancers, complete an "Other Malignancy Form" for the first diagnosis for each of these types.
53	Consent Status	Consented Deceased Exemption 4 Waiver	3288361 Indicate whether the patient was formally consented, consented by death, or if the case has an exemption or waiver for consent Indicate whether the patient was formally consented, consented by death, or if the case has an exemption or waiver for consent. Note: Either the Date of Consent or the Date of Death must be provided to qualify.
Date of Conse	ent		
54	Month of Consent	□□ (MM)	3081955 If the patient was formally consented, provide the month of consent. Note: Do not answer this question if the patient consented by death only.
55	Day of Consent	(DD)	3081957 If the patient was formally consented, provide the day of consent. Note: Do not answer this question if the patient consented by death only.
56	Year of Consent	□□□□ (YYYY)	3081959 If the patient was formally consented, provide the year of consent. Note: Do not answer this question if the patient consented by death only.
57	Number of Days from Date of Initial Pathologic diagnosis to Date of Consent		3288498 If the patient formally consented, 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 the patient's formal consent. 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.
Date of Death	1		
58	Month of Death	ШШ (ММ)	2897026 If the patient consented by death, provide the month of death. Note: If the patient formally consented, only supply the date the patient consent.
59	Day of Death	□□ (DD)	2897028 If the patient consented by death, provide the day of death Note: If the patient formally consented, only supply the date the patient consent.
60	Year of Death	(YYYY)	2897030 If the patient consented by death, provide the year of death. Note: If the patient formally consented, only supply the date the patient consent.
61	Number of Days from Date of Initial Pathologic diagnosis to Date of Death		3288499 If the patient consented by 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 the patient's death. Note 1: 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. Note 2: If the patient formally consented prior to death, do not

Question #	Data Element Label	Data Entry Alternatives	CDE ID With Working Instructions
			answer this question. Only answer the question above that asks fo
			the number of days between the date of diagnosis and the date of
			the patient consent.
Comments	:		
			
5 · · · · · · · · · · · · · · · · · · ·		D :	
Principal in	vestigator Name:	Principal inv	restigator Signature:
		Date	e Signed (MM/DD/YYYY):