Initial Case Quality Control Form Adrenocortical Carcinoma

| Instructions: This form should be completed for all cases submitted for TCGA, prior to the shipment of samples to the BCR. | | | | |
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| Questions regarding this form should be directed to the Tissue Source Site's primary Clinical Outreach Contact at the BCR. | | | | |
| Tissu | e Source Site (TSS) acknowled | lges that the Biospecimen Core Resource (BCR) may confirm that the diagnosis of the frozen | biospecimen is consistent with the primary diagnosis | |
| repor | ted by the TSS through histop | athology examination in the BCR laboratory. If the BCR identifies a possible discrepancy, th | e TSS authorizes the BCR to report these patient | |
| result | ts to the TSS by means of a for | mal report in confidential email format for the quality assurance program of the TSS to add | Iress. | |
| Tissue | Source Site (TSS): | TSS ID: TSS Unique Patient ID: Interviewer Name: | Interview Date/ / / | |
| # | Question | Entry Alternatives | Working Instructions | |
| Veri | fication: By providing the be | low information, the Principal Investigator acknowledges that the information provided | by the institution is true and correct and has been | |
| | ity controlled. | | | |
| 4 * | Was sample prescreened | | Indicate whether the sample submitted to the BCR was | |
| 1* | at site? | □ Yes. The submitted sample was prescreened. | prescreened at the TSS. <u>3081942</u> | |
| Datk | | | <u>3001742</u> | |
| | lology Review | dage that the Diagnosimon Core Descurse (DCD) may confirm that the diagnosis of the fuga | n biognogimon is consistent with the primary diagnosis | |
| | | dges that the Biospecimen Core Resource (BCR) may confirm that the diagnosis of the frozen | | |
| | | pathology examination in the BCR laboratory. If the BCR identifies a possible discrepancy, th | | |
| resu | its to the 155 by means of a fol | rmal report in confidential email format for the quality assurance program of the TSS to ad | Provide the name of the Pathologist that provided the | |
| 2* | Name of Pathologist | | information for all previous sections. | |
| 2 | Name of Factorogist | | 3288225 | |
| | Data of Dath all stat | | Provide the date of the pathology review performed by the TSS | |
| 3* | Date of Pathologist | | pathologist above. | |
| | Review | | <u>3288224</u> | |
| Prin | cipal Investigator/Authori | zed Designee Confirmation | | |
| | | | Confirm that the malignant sample submitted to the BCR meets | |
| 4* | Percent Tumor Nuclei | | the current tumor nuclei metrics for TCGA. | |
| 4 | meets TCGA metrics? | □ Yes | <u>3288520</u> | |
| | | | Check with the BCR to confirm the current acceptable TCGA metrics. | |
| | | | Confirm that the malignant sample submitted to the BCR meets | |
| 5* | Percent Necrosis meets | □ Yes | the current necrosis metrics for TCGA. 3288524 | |
| | TCGA metrics? | | 3288524 Check with the BCR to confirm the current acceptable TCGA metrics. | |
| | | | Confirm that a de-identified pathology report will be sent to | |
| 6* | De-Identified Pathology | □ Yes | BCR prior to or with the shipment of the physical samples. | |
| Ŭ | Report Submitted? | □ No | 3288292 | |
| | | | Confirm that the diagnosis provided on this CQCF for the tumor | |
| | Is the histologic diagnosis | | sample being submitted to TCGA is consistent with the | |
| | on the CQCF (as | | diagnosis found on the patient's pathology report for the tumor | |
| | determined by the TSS | | being sent to the BCR. 3288300 | |
| | pathology review of the | □ Yes If "yes," skip related question below. | The diagnosis is considered to be consistent if at least one of the | |
| 7* | TCGA frozen section top | | following criteria are met: | |
| , | slide) consistent with the | □ No | 1) Diagnosis on the CQCF is identical to the pathology report for | |
| | histology listed in the | | the tumor being sent to the BCR. | |
| | final diagnosis on the | | 2) Diagnosis on the CQCF includes as least one of the subtypes | |
| | pathology report? | | listed on the pathology report and all subtypes on the | |
| | pathology report: | | pathology report are acceptable for TCGA. 3) Diagnosis on the CQCF is "histology, NOS" (i.e., | |

Initial Case Quality Control Form Adrenocortical Carcinoma

| # | Question | Entry Alternatives | Working Instructions |
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| | | | Adenocarcinoma, NOS) and the pathology report lists a specific subtype within the same histologic 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. |
| †8 | If the diagnosis on this form is not consistent with the provided pathology report, indicate the reason for the inconsistency. | Macrodissection performed at TSS to select for a region containing an acceptable TCGA diagnosis (see note at right) Pathology analysis at TSS determined a specific histological subtype different from original pathology report (see note at right) Pathology review of frozen section for TCGA determined histological subtype different from the pathology report (see note at right) | If the diagnosis provided on this form is not consistent with the diagnosis found on the pathology report provided, specify a reason for this inconsistency. <u>3288315</u> If a TSS pathology review of the TCGA committed sample resulted in a different histological subtype than what is documented on the original pathology report, an amendment to the pathology report should be submitted when the sample is shipped to the BCR; or in the absence of an amended pathology report, the TSS must complete and submit an electronic copy of the "TCGA Pathologic Diagnosis Discrepancy Form". In the case of diagnosis modifications, institution protocol should be followed for proper quality assurance. |
| Patie | ent Information | | |
| 9* | History of Other Malignancy | None History of Prior Malignancy History of Synchronous/ Bilateral Malignancy Both History of Synchronous/ Bilateral and Prior Malignancy | Indicate whether the patient has a history of malignancies. If the patient has any history, 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. <u>3382736</u> 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. |
| 10* | History of Neo-adjuvant Treatment (prior to procurement) of Tumor Submitted for TCGA | ☐ Yes (see note at right) ☐ No | Indicate whether the patient received therapy for this cancer prior to the sample procurement of <i>the tumor submitted for TCGA</i>. If the patient did receive treatment for this cancer prior to procurement, the TSS should contact the BCR for further instruction. <u>3382737</u> Please Note: Systemic therapy and certain localized therapies (those administered to the same site as the TCGA submitted tissue) given prior to the procurement of the sample submitted for TCGA are exclusionary. |
| 11* | Consent Status | Consented Deceased Exemption 4 Waiver | Indicate whether the patient was formally consented, consented by death, or if the case has an exemption or waiver for consent.3288361Please Note:• Exemptions and waivers for consent must be approved by NCI. |
| Date | of Formal Consent Do not an | swer this question if the patient consented by death only. | |
| †12 | Date of Consent | Month Day Year | If the patient was formally consented, provide the date of consent. <u>3081955</u> (Month), <u>3081957</u> (Day), <u>3081959</u> (Year) |

Initial Case Quality Control Form Adrenocortical Carcinoma

| # | Question | Entry Alternatives | Working Instructions |
|------|--|--|--|
| Date | of Death Do not complete d | ate of death, if patient formally consented. | |
| †13 | Date of Death | Month Day Year | If the patient consented by death, provide the date of death. <u>2897026</u> (Month), <u>2897028</u> (Day), <u>2897030</u> (Year) |
| 14* | 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 | Provide the patient's race using the defined categories. 2192199 |
| 15 | Ethnicity | Not Hispanic or Latino A person not meeting the definition of Hispanic or Latino. Hispanic or Latino | Provide the patient's ethnicity using the defined categories. 2192217 |
| Path | ologic/Anatomic Informat | | |
| 16* | Tumor Category | Primary (primary untreated malignant biospecimen) | Indicate the type of tumor submitted for TCGA. <u>3288124</u> This is a biospecimen that has not been treated with chemotherapy or radiation prior to resection. |
| 17* | Histologic Subtype of Tumor Submitted for TCGA | Adrenocortical Carcinoma – Usual Type Adrenocortical Carcinoma – Oncocytic Type Adrenocortical Carcinoma – Myxoid Type | Indicate the confirmed diagnosis of the tumor submitted forTCGA. <u>3081934</u> Note: The listed histologies are the only adrenocortical histologicsubtypes being accepted for this TCGA study.Mixed cases will be excluded from this study. |
| 18* | Anatomic Site of Frozen Biospecimen | □ Adrenal Gland | Indicate the anatomic site of the frozen tumor biospecimen submitted for TCGA. 2735776 |
| 19* | Laterality | □ Right □ Left | Indicate the laterality of the frozen tumor biospecimen submitted for TCGA, if it was located in a paired site. 827 |
| 20* | Date of Cancer Sample Procurement | Month Day Year | Provide the date of the procedure performed to obtain the malignant tissue submitted for TCGA. <u>3008197</u> (Month), <u>3008195</u> (Day), <u>3008199</u> (Year) |

Initial Case Quality Control Form Adrenocortical Carcinoma

| # | Question | Entry Alternatives | Working Instructions |
|--------|--|---|---|
| 21* | Vessel Used | CryovialCassetteOther, specifyBiospecimen Storage BagCryomold | Indicate the type of vessel used to ship the tissue to the Biospecimen Core Resource (BCR) for TCGA. <u>3081940</u> |
| †22 | Other Vessel Used | | If the vessel used to ship the tissue to the BCR is not included in the provided list, specify the vessel used. <u>3288137</u> |
| 23* | Method of Cancer Sample Procurement | □ Surgical Resection | Indicate the procedure performed to obtain the malignant tissue submitted for TCGA. <u>3103514</u> |
| 24* | Weight of Resected Tumor | gm | Using the pathology report, provide the weight (in grams) of the resected adrenal gland tumor. <u>3184957</u> |
| 25* | Maximum Tumor Dimension | cm | Using the pathology report, provide the length (in centimeters) of the largest dimension/diameter of the adrenal tumor. <u>64215</u> |
| 26* | Country Where Cancer Sample was Procured | | Provide the country where the tissue submitted for TCGA was procured. <u>3203072</u> |
| 27* | Is tumor sample being submitted for macrodissection? | □ Yes □ No | Indicate whether the tumor sample submitted to the BCR is intended to undergo macrodissection after the BCR receives the sample. <u>3288488</u> |
| - | or Information | | |
| If the | TSS is submitting multiple p | ieces of the same primary tumor for this case; complete the following information for ea | ch piece of tumor sent to the BCR Provide the TSS unique tumor ID. If multiple pieces of tumor |
| 28* | Tumor Identifier | | are submitted, each tumor needs a unique ID. <u>3288096</u> |
| 29* | Weight of Frozen Tumor | (mg) (0.2cm ³ (0.6cm * 0.6cm * 0.6cm) = ~200mg | Provide the weight of the tumor sample submitted for TCGA. <u>3081946</u> Weight can be estimated based on the size of the tumor submitted. |
| 30* | Tumor Nuclei % | (%) | Provide the percent of tumor nuclei for the sample submitted for TCGA. <u>2841225</u> Check with the BCR to confirm the current acceptable TCGA metrics. |
| 31* | Necrosis % | (%) | Provide the percent of necrosis for the sample submitted for TCGA. <u>2841237</u> Check with the BCR to confirm the current acceptable TCGA metrics. |
| Tum | or Slides Submitted | | |
| 32* | Types of Slides Submitted | Physical Top Slide (Frozen Sample) Digital Top Slide Image (Frozen Sample) Digital FFPE Slide Image | Indicate the type(s) of slide(s) submitted to the BCR. <u>3521909</u> Top Slide Definition: Slide cut directly from frozen biospecimen = mirror image of inked surface |
| 33* | Slide/Digital Image ID # | | Provide the slide ID for each slide (physical and digital image) submitted to the BCR. 2321277 |

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|------|--|--|-------------|---|--|
| # | Question | | Entr | y Alternatives | Working Instructions |
| Norr | nal Information A normal c | ontrol must be present to | | • | |
| 34* | Type(s) of Normal Control Check all that apply | ☐ Whole Blood ☐ Buffy Coat ☐ Lymphocytes | | Extracted DNA from Blood Extracted DNA from Saliva Non-Neoplastic Control Tissue* | Indicate the type of normal control submitted for this case. <u>3081936</u> *Non-neoplastic Control Tissue may only be submitted with NCI approval. |
| Norn | nal Control: Whole Blood, B | uffy Coat, or Lymphocyt | tes | | |
| 35† | Method of Normal Sample Procurement | Blood Draw | | | Indicate the procedure performed to obtain the normal control sample submitted for TCGA. <u>3288147</u> |
| 36† | Date of Normal Sample Procurement | Month | Day | Year | Provide the date of the procedure performed to obtain the normal control submitted for TCGA. <u>3288195</u> (Month), <u>3288196</u> (Day), <u>3288197</u> (Year) |
| 37† | Normal Identifier | | | | Provide the TSS unique normal ID. If multiple normal control samples are submitted, each normal control needs a unique ID. <u>3288138</u> |
| Norn | nal Control: Extracted DNA | | | | |
| 38† | Method of Normal Sample Procurement | Blood Draw Buccal Swab Mouthwash | | | Indicate the procedure performed to obtain the normal control sample submitted for TCGA. <u>3288147</u> |
| 39† | Month of Normal Sample Procurement | Month | Day | Year | Provide the month of the procedure performed to obtain the normal control submitted for TCGA. <u>3288195</u> (Month), <u>3288196</u> (Day), <u>3288197</u> (Year) |
| 40† | Normal Identifier | | | | Provide the TSS unique normal ID. If multiple normal control samples are submitted, each normal control needs a unique ID. <u>3288138</u> |
| 41† | Extracted DNA Quantity | | (µg) | | Provide the quantity (μ g) of the normal control sample sent to the BCR for TCGA. <u>3288185</u> |
| 42† | Extracted DNA Quantification Method | | | | Provide the quantification method of the normal control sample sent to the BCR for TCGA. <u>3288186</u> |
| 43† | Extracted DNA Concentration | | (µg/µL) | | Provide the concentration (μ g/ μ L) of the normal control sample sent to the BCR for TCGA. <u>3288187</u> |
| 44† | Extracted DNA Volume | | (µL) | | Provide the volume (μ L) of the normal control sample sent to the BCR for TCGA. <u>3288188</u> |
| Norn | nal Control: Non-neoplastic | Control Tissue | | | |
| 45† | Method of Normal Sample Procurement | □ Skin Punch □ Surgical Resection □ Other Method (plea | se specify) | | Indicate the procedure performed to obtain the normal control sample submitted for TCGA. <u>3288147</u> |
| 46† | Other Method of Normal Sample Procurement | | | | If the procedure performed to obtain the normal sample is not included in the provided list, specify the procedure. <u>3288151</u> |
| 47† | Date of Normal Sample Procurement | Month | Day | Year | Indicate the date of the procedure performed to obtain the normal control sample submitted for TCGA. <u>3288195</u> (Month), <u>3288196</u> (Day), <u>3288197</u> (Year) |

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| # | Question | Entry Alternatives | Working Instructions | |
|---|---|---|---|--|
| 48† | Normal Identifier | | Provide the TSS unique normal ID. If multiple normal control samples are submitted, each normal control needs a unique ID. <u>3288138</u> | |
| 49† | Anatomic Site of Non- Neoplastic Control Tissue | □ Skin □ Kidney □ Other (please specify)* | If the normal control type is normal tissue, indicate the anatomic site of the non-neoplastic control tissue submitted for TCGA. 3081938 *Adrenal tissue is not acceptable for the normal control. | |
| 50† | Other Site of Non- Neoplastic Control Tissue | | 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. <u>3288189</u> | |
| 51† | Proximity of Normal Tissue to Tumor | □ Distal (> 2cm) from the primary tumor | If the normal control type is normal tissue, confirm that the submitted normal tissue was at least 2cm away from the primary tumor. <u>3088708</u> Adjacent (< 2cm) Normal Tissue is not accepted for this tissue type. Unknown Normal Tissue is not acceptable for this tissue type. | |
| 52† | Normal Slide or Digital Image Identifier | | 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. 3288217 | |
| Time Intervals The following questions are only to be answered if the Tissue Source Site is unable to provide the dates requested on this form. Provided time intervals must begin with the date of initial pathologic diagnosis (i.e., biopsy or resection) Please 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. | | | | |
| i* | Has this TSS received permission from the NCI to provide time intervals as a substitute for requested dates on this form? | □ Yes □ No | Indicate whether the TSS has permission to provide time intervals in lieu of dates. | |
| ii | Number of Days from Date of Initial Pathological Diagnosis to Date of Pathological Review | days | Provide the number of days from the date the patient was initially diagnosed pathologically with the disease to the date of the pathological review performed as part of the submission process <u>3288497</u> | |
| iii | Number of Days from Date of Initial Pathological Diagnosis to Date of Consent | days | If the patient formally consented, provide the number of days from the date the patient was initially diagnosed pathologically with the disease to the date of the patient's formal consent. 3288498 | |
| iv | Number of Days from Date of Initial Pathological Diagnosis to Date of Death | days | If the patient consented by death, provide the number of days from the date the patient was initially diagnosed pathologically with the disease to the date of the patient's death. Note: If the patient formally consented prior to death, do not answer this question only answer the question above that asks for the number of days between the date of diagnosis and the date of the patient consent. 3288499 | |

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Adrenocortical Carcinoma

| # | Question | Entry Alternatives | Working Instructions |
|------|--|--------------------|--|
| v | Number of Days from Date of Initial Pathological Diagnosis to Date of Cancer Sample Procurement | days | Provide the number of days from the date the patient was initially diagnosed pathologically with the disease to the date of the procedure that produced the malignant sample submitted <u>3288495</u> |
| vi | Number of Days from Date of Initial Pathological Diagnosis to Date of Normal Sample Procurement (Whole Blood) | days | Provide the number of days from the date the patient was initially diagnosed pathologically with the disease to the date of the procedure that produced the normal control sample submitted <u>3288496</u> |
| vii | Number of Days from Date of Initial Pathological Diagnosis to Date of Normal Sample Procurement (Buffy Coat/Lymphocytes) | days | Provide the number of days from the date the patient was initially diagnosed pathologically with the disease to the date of the procedure that produced the normal control sample submitted <u>3288496</u> |
| viii | Number of Days from Date of Initial Pathological Diagnosis to Date of Normal Sample Procurement (Extracted DNA) | days | Provide the number of days from the date the patient was initially diagnosed pathologically with the disease to the date of the procedure that produced the normal control sample submitted <u>3288496</u> |
| ix | Number of Days from Date of Initial Pathological Diagnosis to Date of Normal Sample Procurement (Non-Neoplastic Tissue) | days | Provide the number of days from the date the patient was initially diagnosed pathologically with the disease to the date of the procedure that produced the normal control sample submitted <u>3288496</u> |
| | | | // |

Principal Investigator or Designee Signature

Print Name

Date

I acknowledge that the above information provided by my institution is true and correct and has been quality controlled.