

VENTANA PD-L1 (SP263) Assay



07208162001



Rx Only

INTENDED USE

VENTANA PD-L1 (SP263) Assay is a

using rabbit monoclonal anti-PD-L1

clone SP263 intended for use in the

assessment of the PD-L1 protein in

formalin-fixed, paraffin-embedded

(FFPE) urothelial carcinoma tissue

stained with OptiView DAB IHC Detection Kit on a VENTANA

BenchMark ULTRA instrument.

or by the percentage of tumor-

PD-L1 status is determined by the

percentage of tumor cells with any

membrane staining above background

qualitative immunohistochemical assay

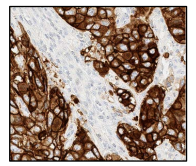


Figure 1. PD-L1 expression in urothelial carcinoma.

associated immune cells with staining (IC+) at any intensity above background. The percent of tumor area occupied by any tumor-associated immune cells (Immune Cells Present, ICP) is used to determine IC+, which is the percent area of ICP exhibiting PD-L1 positive immune cell staining. PD-L1 status is considered High if any of the following are met:

- ≥ 25% of tumor cells exhibit membrane staining; or,
- ICP > 1% and IC+ ≥ 25%; or,
- ICP = 1% and IC+ = 100%.

PD-L1 High status as determined by VENTANA PD-L1 (SP263) Assay was associated with increased objective response rate (ORR) in a single arm study of IMFINZI[™] (durvalumab).

This product is intended for in vitro diagnostic (IVD) use.

SUMMARY AND EXPLANATION

VENTANA PD-L1 (SP263) Assay is an immunohistochemical assay utilizing an anti-PD-L1 rabbit monoclonal primary antibody (VENTANA PD-L1 (SP263)) to recognize the programmed death ligand 1 (PD-L1). It recognizes a transmembrane bound glycoprotein that has a molecular mass of 45-55 kDa. This antibody produces membranous and/or cytoplasmic staining.

Urothelial carcinoma (also known as urothelial cell carcinoma, transitional cell carcinoma of the urinary tract, or urothelial bladder cancer) is the most common cancer of the urinary system worldwide. The majority of urothelial tumors arise in the bladder with the remainder originating in the renal pelvis, urethra, or ureter. Transitional cell carcinoma (TCC) is the most common histologic subtype associated with bladder cancer and accounts for greater than 90% of all urothelial carcinoma cases in the industrialized world; non-urothelial subtypes (e.g., squamous cell carcinoma, adenocarcinoma, small cell carcinoma) are more frequent in other areas of the world.¹

Globally, there were an estimated 429,793 new cases of bladder cancer and 165,084 deaths in 2012.² In Europe alone, for 2012, there were an estimated 151,297 new cases of bladder cancer and 52,411 deaths. In 2015, it was estimated that there would be 74,000 new cases of bladder cancer and 16,000 deaths in the United States.³ Urothelial carcinoma presents as non-muscle-invasive, muscle-invasive, or metastatic disease. The overall 5-year survival rate for metastatic urothelial carcinoma (mUC) is approximately 5.4%.⁴

PD-L1 is a transmembrane protein that downregulates immune responses through binding to its two receptors programmed death-1 (PD-1) and B7-1 (CD80).⁵ PD-1 is an inhibitory receptor expressed on T cells following T-cell activation, which is sustained in states of chronic stimulation such as in chronic infection or cancer.⁶ Binding of PD-L1 with PD-1 inhibits T cell proliferation, cytokine production, and cytolytic activity, leading to the



functional inactivation or exhaustion of T cells. CD80 is a molecule expressed on antigen presenting cells and activated T cells. PD-L1 binding to CD80 on T cells and antigen presenting cells can mediate downregulation of immune responses, including inhibition of T-cell activation and cytokine production.⁷ PD-L1 expression has been observed in immune cells and tumor cells.^{8,9} Aberrant expression of PD-L1 on tumor cells and tumor associated immune cells has been reported to impede anti-tumor immunity, resulting in immune evasion.^{6,9} Therefore, interruption of the PD-L1/PD-1 pathway represents an attractive strategy to reinvigorate tumor-specific T cell immunity suppressed by the expression of PD-L1 in the tumor microenvironment.

PD-L1 is expressed in a broad range of cancers including lung, melanoma, urothelial, ovarian, and colorectal cancer. Prevalence of PD-L1 expression has been reported from 12% to 100% depending on the tumor type, anti PD-L1 clone and cutoff for positivity.¹⁰ IMFINZI (durvalumab) is a human immunoglobulin G1 (IgG1) kappa mAb with immune checkpoint inhibitory and antineoplastic activities that blocks the interaction of PD-L1 (but not programmed cell death ligand-2) with PD-1 on T cells and CD80 on immune cells (IC). IMFINZI (durvalumab) is engineered to reduce antibody dependent cell mediated cytotoxicity and has a calculated molar mass of 146.3 kg/mol.

PRINCIPLE OF THE PROCEDURE

VENTANA PD-L1 (SP263) is a rabbit monoclonal primary antibody which binds to PD-L1 in paraffin-embedded tissue sections. The specific antibody can be localized using a haptenated secondary antibody followed by a multimer anti-hapten-HRP conjugate (OptiView DAB IHC Detection Kit, Cat. No. 760-700 / 06396500001). The specific antibody-enzyme complex is then visualized with a precipitating enzyme reaction product. Each step is incubated for a precise time and temperature. At the end of each incubation step, the VENTANA BenchMark instrument washes the sections to stop the reaction and to remove unbound material that would hinder the desired reaction in subsequent steps. It also applies ULTRA LCS (Cat. No. 650-210 / 05424534001), which minimizes evaporation of the aqueous reagents from the specimen slide.

In addition to staining with VENTANA PD-L1 (SP263), a second slide should be stained with Rabbit Monoclonal Negative Control Ig (Cat. No. 790-4795 / 06683380001). The negative reagent control is used to assess background staining.

REAGENT PROVIDED

VENTANA PD-L1 (SP263) contains sufficient reagent for 50 tests.

One 5 mL dispenser of VENTANA PD-L1 (SP263) contains approximately 8.05 μg of a rabbit monoclonal antibody.

The antibody is diluted in 0.05 M Tris-HCI with 1% carrier protein, and 0.10% ProClin 300, a preservative.

Total protein concentration of the reagent is approximately 10 mg/mL. Specific antibody concentration is approximately 1.61 µg/mL. There is no known non-specific antibody reactivity observed in this product.

VENTANA PD-L1 (SP263) is a recombinant rabbit monoclonal antibody produced as purified cell culture supernatant.

Refer to the appropriate VENTANA detection kit package insert for detailed descriptions of: (1) Principles of the Procedure, (2) Materials and Reagents Needed but Not Provided, (3) Specimen Collection and Preparation for Analysis, (4) Quality Control Procedures, (5) Troubleshooting, (6) Interpretation of Results, and (7) General Limitations.

MATERIALS REQUIRED BUT NOT PROVIDED

Staining reagents, such as VENTANA detection kits and ancillary components, including negative and positive tissue control slides, are not provided.

The following reagents and materials may be required for staining:

- 1. Rabbit Monoclonal Negative Control Ig (Cat. No. 790-4795 / 06683380001)
- 2. Microscope slides, positively charged
- 3. Bar code labels (appropriate for negative reagent control and primary antibody being tested)
- 4. Xylene (Histological grade)
 - Ethanol or reagent alcohol (Histological grade)
 - 100% solution: Undiluted ethanol or reagent alcohol
 - 95% solution: Mix 95 parts of ethanol or reagent alcohol with 5 parts of deionized water
 - 80% solution: Mix 80 parts of ethanol or reagent alcohol with 20 parts of deionized water

5.





- 6. Deionized or distilled water
- 7. OptiView DAB IHC Detection Kit (Cat. No. 760-700 / 06396500001)
- 8. EZ Prep Concentrate (10X) (Cat. No. 950-102 / 05279771001)
- 9. Reaction Buffer Concentrate (10X) (Cat. No. 950-300 / 05353955001)
- 10. ULTRA LCS (Predilute) (Cat. No. 650-210 / 05424534001)
- 11. ULTRA Cell Conditioning Solution (ULTRA CC1) (Cat. No. 950-224 / 05424569001)
- 12. Hematoxylin II counterstain (Cat. No. 790-2208 / 05277965001)
- 13. Bluing Reagent (Cat. No. 760-2037 / 05266769001)
- 14. Permanent mounting medium (Permount Fisher Cat. No. SP15-500 or equivalent)
- 15. Cover glass (sufficient to cover tissue, such as VWR Cat. No. 48393-060)
- 16. Automated coverslipper (such as the Tissue-Tek SCA Automated Coverslipper)
- 17. Light microscope
- 18. Absorbent wipes

Not all products listed in the package insert may be available in all geographies. Consult your local support representative.

STORAGE

Upon receipt and when not in use, store at 2-8°C. Do not freeze.

To ensure proper reagent delivery and stability of the antibody, replace the dispenser cap after every use and immediately place the dispenser in the refrigerator in an upright position.

Every antibody dispenser is expiration dated. When properly stored, the reagent is stable to the date indicated on the label. Do not use reagent beyond the expiration date.

SPECIMEN PREPARATION

Routinely processed, formalin-fixed, paraffin-embedded tissues are suitable for use with this primary antibody when used with the OptiView DAB IHC detection kit and VENTANA BenchMark ULTRA instrument. Based on placenta and tonsil tissues which express PD-L1, the recommended tissue fixative is 10% neutral buffered formalin (NBF) for a period of at least 6 hours up to 72 hours. Acceptable fixatives for use with VENTANA PD-L1 (SP263) Assay are Zinc Formalin and Z-5 fixatives when used with at least 6 hours of fixation time. Other fixatives, including 95% alcohol, AFA and PREFER fixative, are unacceptable for use with the VENTANA PD-L1 (SP263) Assay. The amount used is 15 to 20 times the volume of tissue. Fixation can be performed at room temperature (15-25°C).¹¹ For further information on fixation variables, refer to "Immunohistochemistry Principles and Advances".¹²

WARNINGS AND PRECAUTIONS

- 1. For in vitro diagnostic (IVD) use.
- 2. For professional use only.
- 3. ProClin 300 solution is used as a preservative in this reagent. It is classified as an irritant and may cause sensitization through skin contact. Take reasonable precautions when handling. Avoid contact of reagents with eyes, skin, and mucous membranes. Use protective clothing and gloves.
- 4. Materials of human or animal origin should be handled as biohazardous materials and disposed of with proper precautions.
- Avoid contact of reagents with eyes and mucous membranes. If reagents come in contact with sensitive areas, wash with copious amounts of water.
- 6. Avoid microbial contamination of reagents as it may cause incorrect results.
- Consult local and/or state authorities with regard to recommended method of disposal.
- 8. For supplementary safety information, refer to the product Safety Data Sheet and the Symbol and Hazard Guide located at www.ventana.com.

STAINING PROCEDURE

VENTANA PD-L1 (SP263) Assay has been developed for use on a VENTANA BenchMark ULTRA automated slide stainer in combination with Rabbit Monoclonal Negative Control Ig, OptiView DAB IHC Detection Kit, and ancillary reagents. An assay-specific staining procedure must be used with VENTANA PD-L1 (SP263) Assay. Refer to Table 1 for the recommended staining protocol. Any deviation from recommended test procedures may invalidate expected results. Appropriate controls must be employed and documented. Users who deviate from recommended test procedures must accept responsibility for interpretation of patient results.

This antibody has been optimized for specific incubation times but the user must validate results obtained with this reagent.

The parameters for the automated procedures can be displayed, printed and edited according to the procedure in the instruments Operator's Manual. Refer to the appropriate VENTANA detection kit package insert for more details regarding immunohistochemistry staining procedures.

Table 1. Recommended staining protocol for VENTANA PD-L1 (SP263) Assay and Rabbit Monoclonal Negative Control Ig with OptiView DAB IHC Detection Kit on a BenchMark ULTRA instrument.

Staining Procedure: U VENTANA PD-L1 (SP263) Assay			
Procedure Parameter	Selection		
Deparaffinization	Selected		
Baking	Optional 60°C 12 minutes*		
Cell Conditioning	CC1 Cell Conditioning 64 minutes		
Pre-primary Antibody Peroxidase Selected			
Antibody (Primary)	VENTANA PD-L1 (SP263) Selected*		
	Or		
Negative Control Selected* 16 minutes, 36°C			
OptiView HQ Linker	8 minutes (default)		
OptiView HQ Multimer	8 minutes (default)		
Counterstain	Hematoxylin II, 4 to 8 minutes*		
Post Counterstain Bluing Reagent, 4 minutes			

user-selectable

QUALITY CONTROL PROCEDURES

Rabbit Monoclonal Negative Control Ig

A matched negative reagent control slide must be run for every specimen to aid in the interpretation of results. Rabbit Monoclonal Negative Control Ig, a negative reagent control antibody, is specifically matched for this assay and is used in place of the primary antibody to evaluate nonspecific staining. The staining procedure for the negative reagent control should equal the primary antibody incubation period. Use of a different negative control reagent, or failure to use the recommended negative control reagent, may cause false results.

Placental Tissue Control

A tissue control must be included with each staining run. Qualified normal human term placental tissue is to be used as the control. Control tissue should be fixed as soon as possible and processed in a manner identical to patient tissues. Such tissue may monitor all steps of the analysis, from tissue preparation through staining. Placental tissue contains positive and negative staining elements for the PD-L1 protein and is therefore suitable for use as a tissue control. The positive and negative staining tissue components are used to confirm that the assay functioned properly.

Placental tissue shows moderate to strong uniform staining of the membrane and weak to strong uniform staining of the cytoplasm of trophoblast-lineage cells. Placental stromal tissue and vasculature can be used for assessment of any background staining.

Assay Verification

Prior to initial use of an antibody or staining system in a diagnostic procedure, the specificity of the antibody should be verified by testing it on a series of tissues with known IHC performance characteristics representing PD-L1 positive and negative tissues (refer to the Quality Control Procedures previously outlined in this section of the product insert and to the Quality Control recommendations of the College of American Pathologists Laboratory Accreditation Program, Anatomic Pathology Checklist¹³ or the CLSI Guideline¹⁴). These quality control procedures should be repeated for each new antibody lot, or whenever there is a change in assay parameters. Urothelial carcinoma tissues with known PD-L1 status, and normal human term placental tissue samples, are suitable for assay verification.

STAINING INTERPRETATION / EXPECTED RESULTS

The VENTANA automated immunostaining procedure causes a brown colored DAB reaction product to precipitate at the antigen sites localized by the VENTANA PD-L1





(SP263) Assay. The stained slide(s) are interpreted by a qualified pathologist using light microscopy. A qualified pathologist experienced in IHC procedures must evaluate tissue controls and qualify the stained product before interpreting results.

The cellular staining pattern for VENTANA PD-L1 (SP263) Assay is membranous and/or cytoplasmic staining of tumor cells. Immune cells demonstrate linear membrane, diffuse cytoplasmic, and/or punctate staining.

Positive/Negative System-Level Tissue Controls

The stained positive and negative tissue controls should be examined to ascertain that all reagents are functioning properly. The presence of an appropriately colored reaction product on the PD-L1 positive urothelial tissue consists of membranous and/or cytoplasmic staining of tumor cells, and linear membrane, diffused cytoplasmic, and /or punctate staining of immune cells, and is indicative of positive reactivity.

If the positive or negative tissue controls fail to demonstrate appropriate staining or demonstrate a change in interpretation, any results with the test specimens should be considered invalid.

Placental Tissue Control Evaluation Criteria are described in Table 2. Representative images are provided in the Interpretation Guide for VENTANA PD-L1 (SP263) Assay for Urothelial Carcinoma P/N 1014738US.

 Table 2.
 Placenta Tissue Control Evaluation Criteria for the VENTANA PD-L1 (SP263)
 Assay.

Interpretation	Staining Description	
Acceptable	Moderate to strong uniform membrane staining of trophoblast-lineage cells, and placental stroma and vasculature with no staining.	
No to weak uniform membrane staining of trophoblast lineage cells and/or specific staining within placental st and vascular tissue.		

Negative Reagent Control

Non-specific staining, if present, may have a diffuse appearance and can be evaluated using the negative reagent control slide stained with Rabbit Monoclonal Negative Control Ig. Intact cells should be used for interpretation of staining results, as necrotic or degenerated cells often stain nonspecifically. If background staining is excessive, results from the test specimen should be considered invalid. Examples of background staining for this assay can be found in the interpretation guide (P/N 1014738US).

Patient Tissue

Patient tissue must be evaluated according to the VENTANA PD-L1 (SP263) Assay scoring algorithm provided in Tables 3 and 4. Refer to Interpretation Guide for VENTANA PD-L1 (SP263) Assay Staining of Urothelial Carcinoma P/N 1014738US for representative images and instructions for scoring.

Table 3. VENTANA PD-L1 (SP263) Assay Scoring Algorithm for Urothelial Carcinoma

PD-L1 Interpretation	Staining Description	
PD-L1 status is determined by the percentage of tumor cells with any membrane staining above background or by the percentage of tumor-associated immune cells with staining (IC+) at any intensity above background. The percent of tumor area occupied by any tumor-associated immune cells (Immune Cells Present, ICP) is used to determine IC+, which is the percent area of ICP exhibiting PD-L1 positive immune cell staining is also evaluated.		
High	 PD-L1 Status is considered High if any of the following are met: ≥ 25% of tumor cells exhibit membrane staining; or, ICP > 1% and IC+ ≥ 25%; or, ICP = 1% and IC+ = 100%. 	
Low/negative	PD-L1 Status is considered Low/negative if:none of the criteria for PD-L1 High Status are met.	

Table 4. Non-Specific Background Scoring Criteria for VENTANA PD-L1 (SP263) Assay.

Interpretation	Staining Description	
Acceptable	Background does not interfere with interpretation of PD-L1 status	
Not Acceptable	Background interferes with interpretation of PD-L1 status	

GENERAL LIMITATIONS

- IHC is a multiple step diagnostic process that requires specialized training in the selection of the appropriate reagents, tissue selection, fixation, processing, preparation of the immunohistochemistry slide, and interpretation of the staining results.
- 2. Tissue staining is dependent on the handling and processing of the tissue prior to staining. Improper fixation, freezing, thawing, washing, drying, heating, sectioning, or contamination with other tissues or fluids may produce artifacts, antibody trapping, or false negative results. Inconsistent results may result from variations in fixation and embedding methods, or from inherent irregularities within the tissue.
- 3. Excessive or incomplete counterstaining may compromise proper interpretation of results.
- 4. The clinical interpretation of any positive staining, or its absence, must be evaluated within the context of clinical history, morphology, and other histopathological criteria. The clinical interpretation of any staining, or its absence, must be complemented by morphological studies and system-level controls as well as other diagnostic tests. It is the responsibility of a qualified pathologist to be familiar with the antibodies, reagents, and methods used to interpret the stained preparation. Staining must be performed in a certified licensed laboratory under the supervision of a pathologist who is responsible for reviewing the stained slides and assuring the adequacy of positive and negative controls.
- 5. Ventana Medical Systems, Inc. provides antibodies and reagents at optimal dilution for use when the provided instructions are followed. Any deviation from recommended test procedures may invalidate expected results. Appropriate controls must be employed and documented. Users who deviate from recommended test procedures must accept responsibility for interpretation of patient results.
- 6. This product is not intended for use in flow cytometry, performance characteristics have not been determined.
- Reagents may demonstrate unexpected reactions in previously untested tissues. The possibility of unexpected reactions even in tested tissue groups cannot be completely eliminated because of biological variability of antigen expression in neoplasms, or other pathological tissues.^{15,16}
- Tissues from persons infected with hepatitis B virus and containing hepatitis B surface antigen (HBsAg) may exhibit nonspecific staining with horseradish peroxidase.¹⁷
- 9. False positive results may be seen because of non-immunological binding of proteins or substrate reaction products. They may also be caused by pseudoperoxidase activity (crythrocytes), endogenous peroxidase activity (cytochrome C), or endogenous biotin (example: liver, brain, breast, kidney) depending on the type of immunostain used.¹⁸
- 10. As with any immunohistochemistry test, a negative result means that the antigen was not detected, not that the antigen was absent in the cells or tissue assayed.

SPECIFIC LIMITATIONS

- VENTANA PD-L1 (SP263) Assay has been solely approved on the BenchMark ULTRA instrument with the OptiView DAB IHC Detection Kit and is not approved with any other detection or instruments.
- 2. A patient specimen slide should be stained with Rabbit Monoclonal Negative Control lg. Other negative control reagents are not suitable for this assay.
- This assay has not been validated for use with cytology samples or decalcified bone specimens.Cold ischemia testing of VENTANA PD-L1 (SP263) Assay using a xenograft tissue model did not establish any conditions from zero hours to up to 24 hours that were not favorable with the assay.
- 4. Sections approximately 4-5 µm in thickness should be cut and mounted on positively charged slides. Slides should be stained promptly, as antigenicity of cut tissue sections may diminish over time and may be compromised 6 months after



cutting from the paraffin block for urothelial carcinoma specimens and 9 months for placenta specimens (see the interpretation guide (P/N 1014738US) and the Performance Characteristics section below).

- 5. The clinical associations between VENTANA PD-L1(SP263) Assay and objective response rate have been evaluated in a single arm study of IMFINZI™ (durvalumab) where all patients were treated with durvalumab. The associations observed between PD-L1 status and ORR may be predictive or prognostic.
- 6. There are extremely limited clinical samples which contained ICP = 1% and had 100% of the ICP area stain above background; there were no analytical samples which contained this criteria. Therefore, there is limited analytical or clinical data to support scoring immune cells when ICP = 1%.

PERFORMANCE CHARACTERISTICS

Tests for staining specificity, sensitivity, impact of tissue thickness, repeatability, and intermediate precision, as well as tests for reader precision, inter-laboratory reproducibility, and clinical outcome were conducted and the results are listed in the following section.

Specificity

Arrays containing a variety of normal tissues were stained with VENTANA PD-L1 (SP263) Assay and evaluated for presence of membranous PD-L1 staining as listed in Table 5. Additional staining, such as cytoplasmic or immune cell staining, is also noted (see Table 5 footnote).

Table 5. VENTANA PD-L1 (SP263) Assay staining of formalin-fixed, paraffin-embedded	
normal tissues.	

Tissue	# positive / total cases	Tissue	# positive / total cases
Adrenal gland	0/3*	Mesothelium	0/3†
Bladder	0/3	Myeloid (bone marrow)	0/4*,†
Breast	0/3	Nerve (sparse)	0/3
Cerebellum	0/3	Ovary	0/3
Cerebrum	0/3	Pancreas	0/3*
Cervix	0/3	Parathyroid gland	0/4
Colon	0/3†	Prostate	0/3
Endometrium	0/3	Salivary gland	0/3†
Esophagus	1/3*,†	Skeletal muscle	0/3
Heart	0/3	Skin	0/4§
Hypophysis	0/3*,†	Spleen	0/3†
Intestine, small	0/3†	Stomach	0/3*,†
Kidney	0/3†	Testis	0/3
Larynx	0/3†	Thymus gland	0/3†
Liver	0/3	Thyroid	0/3*,†
Lung	0/3†	Tonsil	3/3†
Lymph node	0/3†		

Additional staining observed: * Cytoplasmic staining, † Immune cell staining, § Melanocyte staining. Percent of IC present above background can not be evaluated in this study because there is no tumor area for which to score tumor infiltrating immune cells.

Sensitivity

Analytical sensitivity was tested on a total of 211 commercially sourced unique cases of urothelial carcinoma FFPE specimens that were tested with manufactured lots. 169/211 tissues were classified as PD-L1 High using the VENTANA PD-L1(SP263) Assay scoring criteria for urothelial carcinoma.

An array of neoplastic tissues was evaluated for TC and IC staining with VENTANA PD-L1 (SP263) Assay as described in Table 6.



 Table 6. VENTANA PD-L1 (SP263) Assay staining of formalin-fixed, paraffin-embedded neoplastic tissues for any tumor cell or immune cell staining.

		# positive / total cases		
Origin	Pathology	Tumor Cells	Immune Cells	
Cerebrum	Glioblastoma	0/1	1/1	
Cerebrum	Atypical meningioma	0/1	0/1	
Cerebrum	Malignant ependymoma	0/1	1/1	
Cerebrum	Oligodendroglioma	0/1	0/1	
Ovary	Serous adenocarcinoma	0/1	1/1	
Ovary	Adenocarcinoma	0/1	0/1	
Pancreas	Islet cell carcinoma	1/1	0/1	
Pancreas	Adenocarcinoma	0/1	1/1	
Testis	Seminoma	0/1	0/1	
Testis	Embryonal carcinoma	0/1	0/1	
Thyroid	Medullary carcinoma	0/1	0/1	
Thyroid	Papillary carcinoma	1/1	0/1	
Breast	Intraductal carcinoma	0/1	1/1	
Breast	Invasive ductal carcinoma	0/2	0/2	
Spleen	Diffuse B-cell lymphoma	0/1	1/1	
Lung	Small cell undifferentiated carcinoma	1/1	1/1	
Lung	Squamous cell carcinoma	1/1	1/1	
Lung	Adenocarcinoma	0/1	0/1	
Esophagus	Neuroendocrine carcinoma	0/1	0/1	
Esophagus	Adenocarcinoma	0/1	0/1	
Stomach	Signet-ring cell carcinoma	0/1	0/1	
Intestine	Adenocarcinoma	0/1	0/1	
Intestine	Stromal sarcoma	0/1	0/1	
Colon	Adenocarcinoma	0/1	1/1	
Colon	Interstitialoma	0/1	0/1	
Rectum	Adenocarcinoma	0/1	0/1	
Rectum	Moderate malignant interstitialoma	0/1	0/1	
Liver	Hepatocellular carcinoma	0/1	0/1	
Liver	Hepatoblastoma	0/1	0/1	
Kidney	Clear cell carcinoma	0/1	0/1	
Prostate	Adenocarcinoma	0/2	0/2	
Uterus	Leiomyoma	0/1	0/1	
Uterus	Adenocarcinoma	0/1	0/1	





Oninin	Dethelesur	# positive / total cases	
Origin	Pathology	Tumor Cells	Immune Cells
Uterus	Clear cell carcinoma of endometrium	0/1	0/1
Uterine cervix	Squamous cell carcinoma	0/2	2/2
Striated muscle	Embryonal rhabdomyosarcoma	0/1	0/1
Rectum	Malignant melanoma	0/1	0/1
Skin	Basal cell carcinoma	0/1	0/1
Skin	Squamous cell carcinoma	0/1	0/1
Back	Neurofibroma	0/1	1/1
Retroperitoneal	Neuroblastoma	0/1	0/1
Abdominal cavity	Malignant mesothelioma	0/1	0/1
Mediastinum	Diffuse B-cell lymphoma	1/1	1/1
Lymph node	Hodgkin's lymphoma	1/1	1/1
Lymph node	Diffuse B-cell lymphoma	1/1	1/1
Pelvic cavity	Anaplastic large cell lymphoma	1/1	1/1
Bladder	Low grade malignant leiomyosarcoma	0/1	0/1
Bone	Osteosarcoma	0/1	1/1
Retroperitoneum	Spindle cell rhabdomyosarcoma	0/1	0/1
Smooth muscle	Moderate malignant leiomyosarcoma	0/1	0/1
Bladder	Transitional cell carcinoma (bladder)	1/1	1/1

Tissue Thickness

Tissue thickness was evaluated using 7 unique cases of human urothelial carcinoma (4 with tumor positivity \geq 25% and 3 with tumor positivity <25%; 4 with IC+ \geq 25% and 3 with IC+ <25% for IC). Tissues were sectioned and tested in duplicate at 2, 3, 4, 5, 6, and 7 microns. All tissue thicknesses from 3 to 7 microns demonstrated appropriate specific staining and acceptable background levels with VENTANA PD-L1 (SP263) Assay. Ventana recommends that specimens be cut at 4-5 microns for the assay.

Repeatability and Intermediate Precision Studies

Repeatability studies for VENTANA PD-L1 (SP263) Assay staining of urothelial carcinoma specimens were completed to demonstrate:

- Intra-day Repeatability Five replicate slides each from 13 unique urothelial carcinoma specimens were stained with VENTANA PD-L1 (SP263) Assay on a single BenchMark ULTRA instrument within one day.
- Inter-day Precision Two replicate slides each from 13 unique urothelial carcinoma specimens were stained with VENTANA PD-L1 (SP263) Assay on a single Benchmark ULTRA instrument across 5 non-consecutive days spanning at least a 20 day period.
- Inter-instrument and Inter-lot Precision 27 replicate slides each from 22 unique urothelial carcinoma specimens were stained with VENTANA PD-L1 (SP263) Assay using three lots of VENTANA PD-L1 (SP263) antibody and three lots of OptiView DAB IHC Detection Kit, on three BenchMark ULTRA instruments.

The 13 unique cases used for the Intra-day Repeatability and Inter-day Precision Studies had an overlapping case distribution of 9 PD-L1 High and 4 PD-L1 Low/negative. The 22 unique cases used for the Inter-instrument and Inter-lot Precision Study had an overlapping case distribution of 13 PD-L1 High and 9 PD-L1 Low/negative.

All slides were blinded and randomized, and then evaluated using the VENTANA PD-L1 (SP263) Assay scoring algorithm as it pertains to tumor cells and tumor associated immune cells (Tables 3 and 4). Results are summarized in Table 7.

 Table 7. Repeatability and Intermediate Precision of VENTANA PD-L1 (SP263) Assay staining of urothelial carcinoma specimens.

Repeatability/Intermediate precision parameter	Positive Percent Agreement (95% CI)*	Negative Percent Agreement (95% CI)*	Overall Percent Agreement (95% CI)*
Intra-day Repeatability	100.0%	95.0%	98.5%
(within a single day)	(45/45)	(19/20)	(64/65)
n = 65 observations	(92.1 – 100%)	(76.4 – 99.1%)	(91.8 - 99.7%)
Inter-day Precision	100.0%	100.0%	100.0%
(5 non-consecutive days)	(90/90)	(40/40)	(130/130)
n = 130 observations	(95.9 – 100%)	(91.2 – 100.0%)	(97.1 – 100.0%)
Inter-instrument and Inter-lot Precision (3 instruments, 3 antibody lots, and 3 detection kit lots) n = 594 observations	98.3% (345/351) (96.3 – 99.2%)	99.6% (242/243) (97.7 –99.9%)	98.8% (587/594) (97.6 – 99.4%)

*Confidence intervals were calculated using the Wilson Score method

Inter- and Intra-Reader Precision Studies

To assess Inter- and Intra-reader Precision, three trained pathologists evaluated an overlapping subset of approximately 50 unique urothelial carcinoma specimens representing the dynamic range of the VENTANA PD-L1 (SP263) Assay with case distribution of 22 PD-L1 High and 28 PD-L1 Low/negative.

Specimens were blinded and randomized prior to evaluation for PD-L1 status using the VENTANA PD-L1 (SP263) Assay scoring algorithm as it pertains to tumor cells as well as tumor associated immune cells (Tables 3 and 4). Readers scored all specimens twice, with a minimum of two weeks between reads. The agreement rates are summarized in Table 8.

Table 8. Inter- and Intra-reader Precision of VENTANA PD-L1 (SP263) Assay staining of urothelial carcinoma specimens.

Reader Precision	Positive Percent Agreement (95% CI)*	Negative Percent Agreement (95% CI)*	Overall Percent Agreement (95% CI)*
Inter-reader Precision (average of all three reader pairwise comparison for the first read) n =143 pairs	92.8% (128/138) (85.5-98.3%)	93.2% (138/148) (86.5-98.5%)	93.0% (133/143) (87.1-98.6%)
Intra-reader Precision (average of all three readers' agreement rates between first and second reads) n =145 pairs	92.1% (128/139) (85.4-96.7%)	92.7% (140/151) (87.1-96.8%)	92.4% (134/145) (87.2-96.6%)

*Percentile Bootstrap confidence intervals were constructed from the 2.5th and 97.5th percentiles using 2,000 replicates when sampling at the case level.

Inter-laboratory Reproducibility Study

An Inter-laboratory Reproducibility Study for VENTANA PD-L1 (SP263) Assay was conducted to demonstrate reproducibility of the assay in determining PD-L1 expression in





urothelial carcinoma cases, using tissue specimens run across 5 non-consecutive days over a 20-day period at three external laboratories.

The Inter-laboratory Reproducibility Study analyzed 26 urothelial carcinoma specimens with a case distribution of 18 PD-L1 High and 8 PD-L1 Low/negative.

The specimens were blinded, randomized and evaluated by a total of 6 readers (2 readers/site). Results are summarized in Table 9.

 Table 9. Inter-laboratory Reproducibility of VENTANA PD-L1 (SP263) Assay staining of urothelial carcinoma specimens.

Inter-laboratory Reproducibility	Positive Percent Agreement (95% CI)	Negative Percent Agreement (95% CI)	Overall Percent Agreement (95% CI)
Overall agreement (across sites, days and readers) n =778 observations	95.2% (512/538) (93.0-96.7%)*	85.8% (206/240) (80.9-89.7%)*	92.3% (718/778) (90.2-94.0%)*
Inter-site agreement (average of site-to-site pairwise comparisons) n =7760 pairs	91.1% (84.2-95.8%)**	79.1% (61.9-89.1%)**	87.5% (86.8-88.2%)**
Inter-reader agreement (average of reader-to-reader pairwise comparisons within each site) n =389 pairs	90.1% (492/546) (82.3-95.7%)**	76.7% (178/232) (58.2-88.4%)**	86.1% (335/389) (82.3-89.2%)**

* Confidence intervals were calculated using the Wilson Score method.

** Percentile Bootstrap confidence intervals were constructed from the 2.5th and 97.5th percentiles using 2,000 replicates when sampling at the case level.

Clinical Outcome Study

The clinical performance of VENTANA PD-L1(SP263) Assay was evaluated in Study 1, a single arm, multicenter, open-label clinical trial of IMFINZI (durvalumab) on patients with locally advanced or metastatic urothelial carcinoma. Tumor specimens from 332 patients were evaluated prospectively for PD-L1 expression at a central laboratory using the VENTANA PD-L1 (SP263) Assay. PD-L1 status was not evaluable for 31 patients due to insufficient tumor cells present in sample (n=25), inappropriate tissue being submitted (n=3), tissue folding (n=1), dye trapping (n=1) or tissue washing of the slide (n=3). There were no instances where a sample was non-evaluable for PD-L1 status due to assay failure.

In Study 1, 182 patients with locally advanced or metastatic urothelial carcinoma were enrolled. The patients that were enrolled had progressed while on or after a platinumbased therapy, including those who progressed within 12 months of receiving therapy in a neo-adjuvant or adjuvant setting. These patients had initiated durvalumab therapy at least 13 weeks prior to the data cut-off date. The trial excluded patients with a history of immunodeficiency; medical conditions that required systemic immunosuppression (not to exceed 10 mg/day of prednisone or equivalent); history of severe autoimmune disease; untreated CNS metastases; HIV; active tuberculosis, or hepatitis B or C infection. All patients received IMFINZI 10 mg/kg via intravenous infusion every 2 weeks for up to 12 months or until unacceptable toxicity or disease progression. Tumor assessments were performed at Weeks 6, 12 and 16, then every 8 weeks for the first year and every 12 weeks thereafter. The major efficacy outcome measures were confirmed Objective Response Rate (ORR) according to RECIST v1.1 as assessed by Blinded Independent Central Review (BICR), and duration of response (DoR).

In Study 1, the median age was 67 years (range: 34 to 88), 72% were male, 64% were Caucasian. Sixty-six percent (66%) had visceral metastasis (bone, liver, or lung), including 34% with liver metastasis. Lymph node only metastasis were present in 13% of patients. Sixty-six percent (66%) of patients had ECOG score of 1 and 41% of patients had a baseline creatinine clearance of <60 mL/min. The Bellmunt risk score (which includes ECOG score, baseline hemoglobin, and liver metastases) was 0 in 23%, 1 in 38%, 2 in 29%, and 3 in 9% of patients. Twenty percent (20%) of patients had disease

progression following platinum-containing neo-adjuvant or adjuvant chemotherapy as their only prior line of therapy. Seventy percent (70%) of patients received prior cisplatin, 30% prior carboplatin and 35% received \geq 2 prior lines of systemic therapy. The median followup time was 5.6 months. A partial or complete response was observed in 31/182 patients (17.0%; 95% CI = 11.9 – 23.3%) following treatment with IMFINZI.

Of the 182 patients that were enrolled in Study 1, 128 were enrolled without regard to PD-L1 status and after the VENTANA PD-L1 (SP263) Assay scoring algorithm was finalized. Of these 128 patients, 58 were classified as PD-L1 High, 56 as PD-L1 Low/negative, and samples for 14 patients were non-evaluable. The efficacy results are summarized in Table 10. The median follow-up time for these patients was 4.9 months.

Table 10. Efficacy Results for a 128 patient subset from Study 1; PD-L1 expression in patients with urothelial carcinoma.

Efficacy Parameter*	PD-L1 High (N=58)	PD-L1 Low/Negative (N=56)	PD-L1 Non-Evaluable (N=14)
Number of confirmed responders by BICR	11	2	3
Objective Response Rate (95% CI)	19.0% (9.9% - 31.4%)	3.6% (0.4% - 12.3%)	21.4% (4.7% - 50.8%)
CR, n (%)	2 (3.4%)	0	1 (7.1%)
PR, n (%)	9 (15.5%)	2 (3.6%)	2 (14.3%)
Median (DoR), months, (range)	4.24 (0.9+ - 4.2)	NR (1.9+ - 4.2+)	NR (2.3+ - 2.6+)

*Objective Response Rate and Duration of Response determined by RECIST v1.1

BICR=Blinded Independent Central Review

CR=Complete Response

CI=Confidence Interval

DoR=Duration of Response NR=Not Reached

PR=Partial Response

(refer to Drugs@FDA for the most recent therapeutic product labeling)



TROUBLESHOOTING

Troubleshooting guidance is provided in Table11. If a problem cannot be attributed to any of these causes, or if the suggested corrective action fails to resolve the problem, consult your local support representative.

Problem	Probable Cause	Suggested Action
Light or no staining of slides	Incorrect staining protocol selected	Verify that U VENTANA PD-L1 (SP263) Assay procedure was used.
		Verify that VENTANA PD-L1 (SP263) was selected for Primary Antibody
	Degradation of tissue	Verify tissue was stained within the recommended time frame following sectioning.
	Dispenser malfunction	Verify nozzle cap is removed.
		Ensure dispenser is primed
		Check the priming chamber for foreign materials or particulates, such as fibers or precipitate
		Refer to inline dispenser package insert associated with P/N 740-4907 / 07208162001 located at www.ventana.com
		Ensure that only recommended fixatives and fixation times are used.
	Incorrect or missing bulk reagent	Ensure bulk reagents are correctly filled.
Excessive background staining of slides	Incorrect staining protocol selected	Verify that U VENTANA PD-L1 (SP263) Assay procedure was used.
	Incorrect or missing bulk reagent	Ensure bulk reagents are correctly filled.
	Inappropriate fixation method used	Ensure that only recommended fixatives and fixation times are used.
Tissue detached from slides	Use of incorrect microscope slides	Ensure positively charged microscope slides are used.

Table 11. Troubleshooting guidance for VENTANA PD-L1 (SP263) Assay.

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CONTACT INFORMATION



Ventana Medical Systems, Inc. 1910 E. Innovation Park Drive Tucson, Arizona 85755 USA

+1 520 887 2155 +1 800 227 2155 (USA)





VENTANA PD-L1 (SP263) Assay Staining in Urothelial Carcinoma

Interpretation Guide

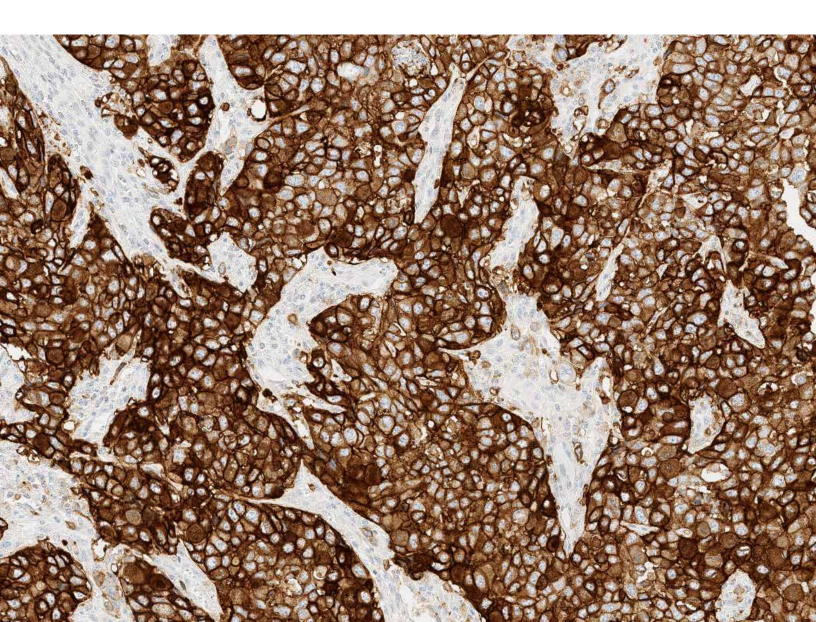


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Introduction

Urothelial carcinoma (also known as urothelial cell carcinoma, transitional cell carcinoma of the urinary tract, or urothelial bladder cancer) is the most common cancer of the urinary system worldwide. The majority of urothelial tumors arise in the bladder with the remainder originating in the renal pelvis, urethra, or ureter. Transitional cell carcinoma (TCC) is the most common histologic subtype associated with bladder cancer and accounts for greater than 90% of all urothelial carcinoma cases in the industrialized world; non-urothelial subtypes (e.g., squamous cell, adenocarcinoma, small cell carcinoma) are more frequent in other areas of the world.¹

Globally, there were an estimated 429,793 new cases of bladder cancer and 165,084 deaths in 2012.² In Europe alone, for 2012, there were an estimated 151,297 new cases of bladder cancer and 52,411 deaths. In 2015, it was estimated that there would be 74,000 new cases of bladder cancer and 16,000 deaths in the United States.³ Urothelial carcinoma presents as non-muscle-invasive, muscle-invasive, or metastatic disease. The overall 5-year survival rate for metastatic urothelial carcinoma (mUC) is approximately 5.4%.⁴

PD-L1 is a transmembrane protein that downregulates immune responses through binding to its two receptors programmed death-1 (PD-1) and B7-1 (CD80).⁵ PD-1 is an inhibitory receptor expressed on T cells following T-cell activation, which is sustained in states of chronic stimulation such as in chronic infection or cancer.⁶ Binding of PD-L1 with PD-1 inhibits T cell proliferation, cytokine production, and cytolytic activity, leading to the functional inactivation or exhaustion of T cells. CD80 is a molecule expressed on antigen presenting cells and activated T cells. PD-L1 binding to CD80 on T cells and antigen presenting cells can mediate downregulation of immune responses, including inhibition of T-cell activation and cytokine production.7 PD-L1 expression has been observed in immune cells and tumor cells.^{8,9} Aberrant expression of PD-L1 on tumor cells and or tumor-associated immune cells has been reported to impede anti-tumor immunity, resulting in immune evasion.^{6,9} Therefore, interruption of the PD-L1/ PD-1 pathway represents an attractive strategy to reinvigorate tumorspecific T cell immunity suppressed by the expression of PD-L1 in the tumor microenvironment. PD-L1 is expressed in a broad range of cancers including lung, melanoma, urothelial, ovarian, and colorectal cancer. Prevalence of PD-L1 expression has been reported from 12% to 100% depending on the tumor type, anti PD-L1 clone and cutoff for positivity.10

Intended Use of Product

VENTANA PD-L1 (SP263) Assay is a qualitative immunohistochemical assay using rabbit monoclonal anti-PD-L1 clone SP263 intended for use in the assessment of the PD-L1 protein in formalin-fixed, paraffin-embedded (FFPE) urothelial carcinoma tissue stained with OptiView DAB IHC Detection Kit on a VENTANA BenchMark ULTRA instrument. PD-L1 status is determined by the percentage of tumor cells with any membrane staining above background or by the percentage of tumor-associated immune cells with staining (IC+) at any intensity above background. The percent of tumor area occupied by any tumor-associated immune cells (Immune Cells Present, ICP) is used to determine IC+, which is the percent area of ICP exhibiting PD-L1 positive immune cell staining. PD-L1 status is considered High if any of the following are met:

- $\geq 25\%$ of tumor cells exhibit membrane staining; or,
- ICP > 1% and IC+ \geq 25%; or,
- ICP = 1% and IC+ = 100%.

PD-L1 High status as determined by VENTANA PD-L1 (SP263) Assay was associated with increased objective response rate (ORR) in a single arm study of IMFINZI[™] (durvalumab).

This product is intended for *in vitro* diagnostic (IVD) use.

Purpose of Interpretation Guide

This guide is intended to aid pathologists in the clinical evaluation of formalin-fixed, paraffin-embedded (FFPE) urothelial carcinoma sections stained with the Assay using the Assay Scoring Algorithms in accordance with the proposed product labeling. Specifically, this guide:

- Provides pathologists with a tool to facilitate the evaluation of formalin-fixed, paraffin-embedded (FFPE) urothelial carcinoma sections stained with the Assay using the Assay Scoring Algorithms in accordance with the proposed product labeling.
- Provides photographic images that illustrate the staining patterns and intensities that may result from staining of urothelial carcinoma tissue samples with the Assay.
- Provides guidance in the use of placenta tissue as a system-level control when stained with the Assay.
- Provides photographic images of internal controls.
- Provides a reference for relating staining patterns to specific PD-L1 scores.
- Provides example images of challenging cases to provide guidance in their evaluation.

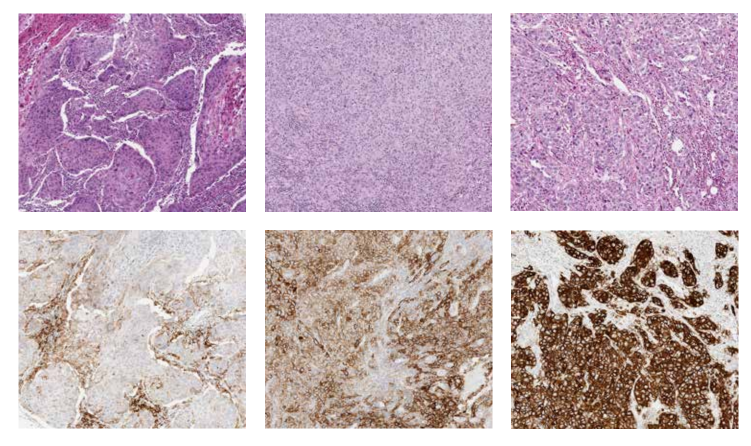
Clinical Evaluation

Evaluating Staining Patterns and Intensities

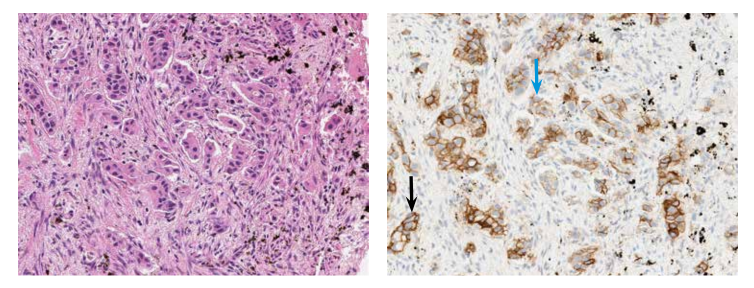
Urothelial carcinoma cases stained with the Assay are assessed for both the percentage of tumor cells with membrane staining and the percentage of tumor-associated immune cells with membrane, cytoplasm, or punctate staining.

Tumor Cell Staining:

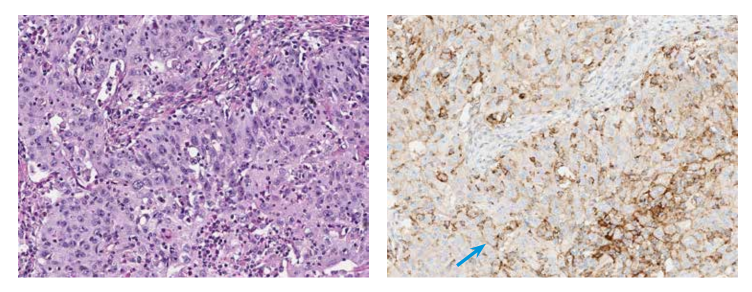
Urothelial carcinoma neoplastic cells labeled with the Assay are evaluated for the percent of the tumor cells with membrane staining at any intensity of the diaminobenzidine (DAB) signal. The immunohistochemical staining in urothelial carcinoma is membranous and/or cytoplasmic, and may be expressed homogeneously or heterogeneously throughout the neoplasm. Membrane staining can have a partial or complete circumferential pattern. Cytoplasmic staining is generally diffuse with some cases displaying a finely granular quality. Tumor cell cytoplasmic staining is disregarded for determining PD-L1 expression. The total percentage of membrane signal intensities is visually estimated and used to generate the PD-L1 expression level. An isotype-matched negative control antibody is used to evaluate the presence of background in test samples and establish a staining intensity baseline.



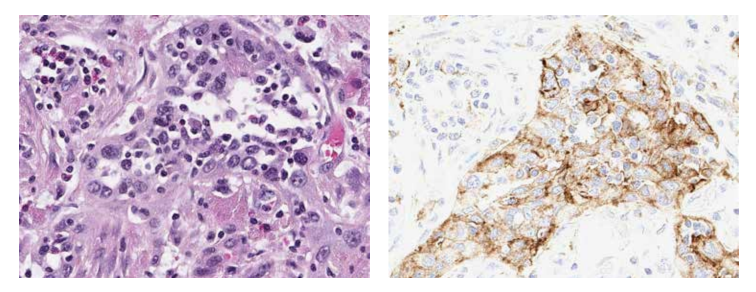
Urothelial carcinoma, H&E and Assay: Various cases demonstrating the range of membrane and cytoplasmic staining in tumor cells (10X)



Urothelial carcinoma, H&E and Assay: Tumor cells with circumferential (black arrow) and partial (blue arrow) membrane staining pattern (20X)



Urothelial carcinoma, H&E and Assay: Tumor cells with weak (blue arrow) membrane staining pattern (20X)



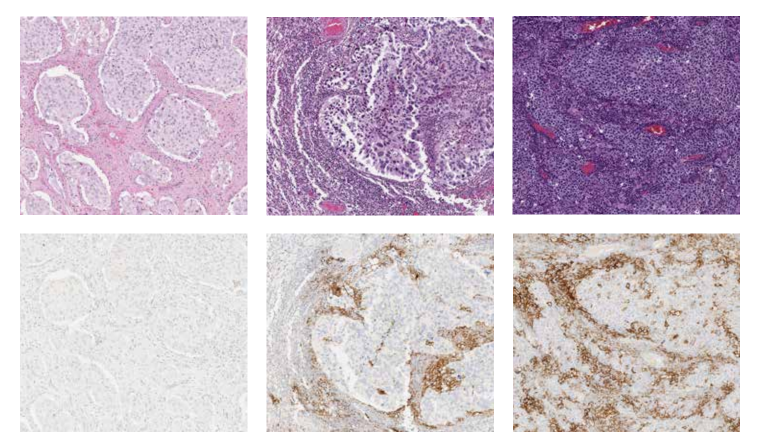
Urothelial carcinoma, H&E and Assay: Tumor cells show membranous and weak granular cytoplasmic staining (40X)

Tumor-Associated Immune Cell Staining:

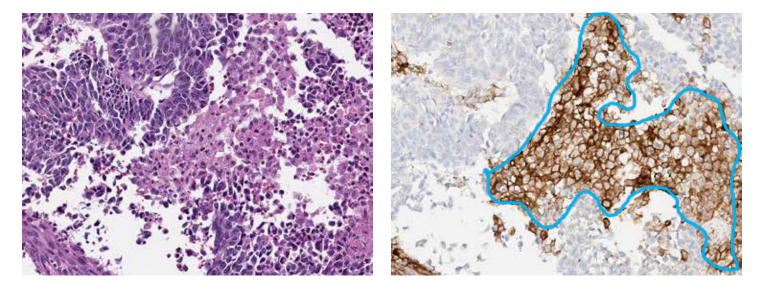
Immune cells exhibit a range of staining intensities and patterns: negative to weak diffuse cytoplasmic and/or weak to strong membranous signal. A punctate pattern of staining may be seen in association with lymphocytes. PD-L1 expression has been observed in lymphocytes, macrophages, histiocytes, plasma cells, and neutrophils.

The percentage of tumor-associated immune cells with staining in urothelial carcinoma cases is evaluated in addition to tumor cell

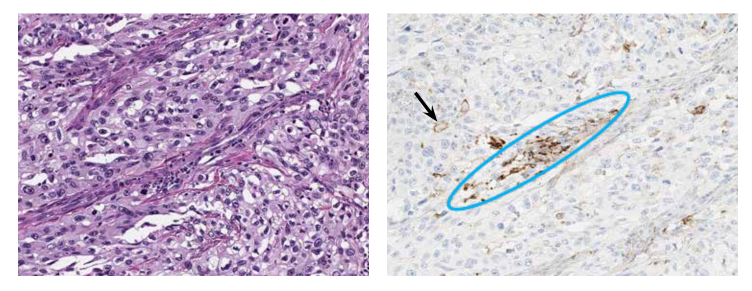
staining. Immune cell staining is assessed by initially reviewing the entire tumor area and determining the percentage of the tumor area occupied by immune cells (ICP). Next the percentage of immune cells demonstrating any pattern of PD-L1 expression within the tumor area is visually estimated (IC+).



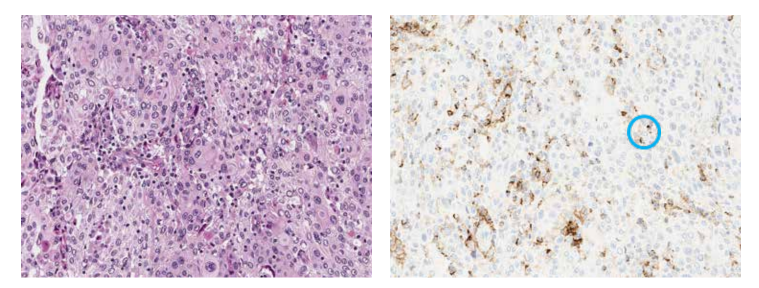
Urothelial carcinoma, H&E and Assay: Various cases demonstrating the range and patterns of immune cells staining (10X)



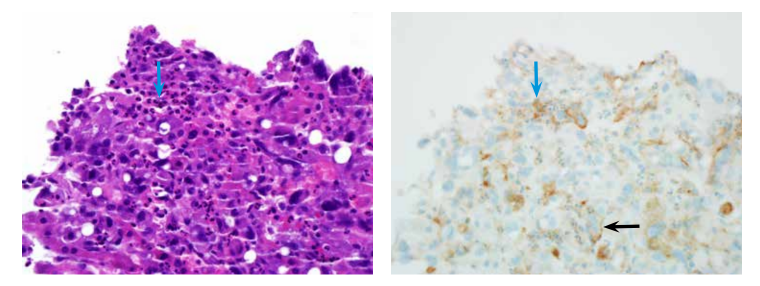
Urothelial carcinoma, H&E and Assay: Positive macrophage staining (blue outline) with adjacent PD-L1 negative tumor (20X)



Urothelial carcinoma, H&E and Assay: Punctate (blue outline) and membranous (black arrow) immune cell staining with adjacent PD-L1 negative tumor (20X)



Urothelial carcinoma, H&E and Assay: Scattered immune cells with punctate staining infiltrating throughout tumor (single example circled in blue) (20X)



NSCLC, H&E and Assay: Case with rare neutrophil-only infiltrate (blue arrow) within the tumor demonstrating PD-L1 staining. Tumor cell membrane staining also present (black arrow) (40X)

9

Tissue Requirements

The VENTANA PD-L1 (SP263) Assay requires one serial tissue section for hematoxylin and eosin (H&E) staining, a second serial tissue section for negative control antibody staining, and a third serial tissue section for staining with the VENTANA PD-L1 (SP263) Assay. In addition, normal human term placenta tissue can be used as a control for the PD-L1 (SP263) Assay. This tissue shows moderate to strong uniform staining of the membrane and weak to strong uniform staining of the cytoplasm of trophoblast-lineage cells. Placental stromal tissue and vasculature can be used for assessment of any background staining. If H&E evaluation indicates that the patient specimen is inadequate then a new specimen should be obtained. Repeat staining of a specimen should be carried out on unstained slides if (1) the tissue run control slide does not exhibit acceptable staining; (2) the negative control case slide does not exhibit acceptable staining; or (3) the VENTANA PD-L1 (SP263) Assay stained case slide (the PD-L1 IHC slide) is not evaluable. If the last of these slides is not interpretable due to artifacts, edge effects, necrosis, lack of tissue, or any other reason, then the slide cannot be used for clinical evaluation. If controls are acceptable and the VENTANA PD-L1 (SP263) Assay stained slide is evaluable, the slide can be evaluated by a trained pathologist as described in the Scoring Criteria.

Morphology and Background Acceptability Criteria

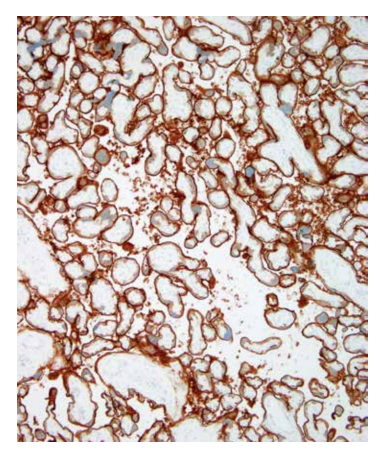
Tissue morphology and background acceptability are assessed for each patient case using the criteria described in Tables 1 and 2.

Table 1: Morphology Acceptability Criteria		
Interpretation	Microscope Observation	
Acceptable	Cellular elements of interest are visualized allowing interpretation of the stain.	
Not Acceptable	Cellular elements of interest are not visualized compromising interpretation of the stain.	

Table 2: Background Acceptability Criteria		
Interpretation	Microscope Observation	
Acceptable	Non-specific staining that is not obtrusive to interpretation of specific staining.	
Not Acceptable	Non-specific staining that is obtrusive to interpretation of specific staining.	

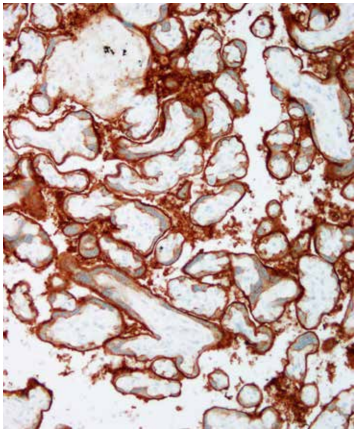
Positive Tissue Control

A known positive control tissue fixed and processed in the same manner as the patient specimens should be run for each set of test conditions and with every VENTANA PD-L1 (SP263) Assay staining procedure performed. The control tissue (an index case) should be a fresh autopsy, biopsy, surgical specimen prepared and fixed as soon as possible in a manner identical to patient specimens. This tissue may be used to monitor all steps of specimen processing and staining. A tissue section fixed or processed differently from the test specimen can be used as a control for reagents and staining but not for fixation or tissue preparation. A positive urothelial carcinoma case with moderate staining is more suitable for quality control than one that stains strongly; it can be used to detect minor levels of reagent



Placenta, VENTANA PD-L1 (SP263) Assay: Strong uniform membrane staining and moderate cytoplasmic staining of trophoblast-lineage cells (10X)

degradation or out-of-specification issues that might be instrumentrelated. Positive membrane staining of neoplastic cells in the control tissue confirms that the VENTANA PD-L1 (SP263) antibody was applied and the instrument functioned properly. The positive tissue control should be used only to monitor performance; it should not be used to aid the clinical diagnosis of patient samples. Additionally, the VENTANA PD-L1 (SP263) Assay can utilize as a positive control human term placental tissue, which shows moderate to strong uniform staining of the membrane and weak to strong uniform staining of the cytoplasm of trophoblast-lineage cells. Placental stromal tissue and vasculature can be used for assessment of any background staining. Please refer to Table 3.



Placenta, VENTANA PD-L1 (SP263) Assay: Stroma and vasculature within villi show no PD-L1 staining (20X)

Table 3: Placenta Tissue Control Evaluation Criteria for the Ventana PD-L1 (SP263) Assay		
Acceptable	Moderate to strong uniform membrane staining of trophoblast-lineage cells, and placental stroma and vasculature with no staining.	
Not Acceptable	No to weak uniform membrane staining of trophoblast lineage cells and/or specific staining within placental stromal and vascular tissue.	

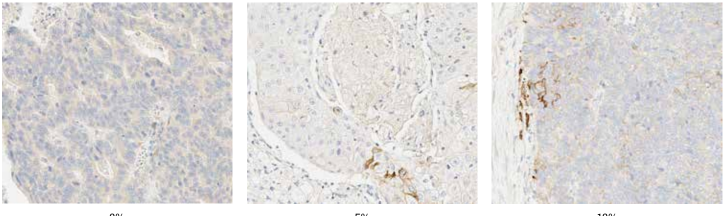
VENTANA PD-L1 (SP263) Assay Scoring Algorithm for Urothelial Carcinoma Tissue

PD-L1 status and expression level is assigned by a trained pathologist based on their evaluation of the percentage of specific staining for both tumor and tumor-associated immune cells. PD-L1 status is based on the total percent of tumor cells with membrane staining or the total percent of tumor-associated immune cells with staining (IC+) at any intensity. Specifically, this guide highlights an expression level of greater than or equal to 25% of tumor cells with membrane staining or of tumor-associated immune cells with staining. In cases where the percent of tumor-associated immune cells in the tumor area (ICP) is 1%, IC+ is scored as either 0%, <100% or 100% due to the difficulties in estimating the percent staining in small volumes of immune cells in low measures. The small amount of PD-L1 staining observed in cases with < 100% IC+, should be considered as < 25% PD-L1 expression. Please refer to Table 4.

Interpretation of urothelial carcinoma cases stained with the Assay is based on the criteria noted in the table below. Images of various expression level staining patterns are provided in the subsequent sections. Please refer to Table 4.

Table 4: VENTANA PD-L1 (SP263) Assay Scoring Algorithm for Urothelial Carcinoma		
PD-L1 Interpretation	Staining Description	
PD-L1 status is determined by the percentage of tumor cells with any membrane staining above background or by the percentage of tumor-associated immune cells with staining (IC+) at any intensity above background. Percent of tumor area occupied by any tumor-associated immune cells (ICP) is used to determine IC+, which is the percent area of ICP exhibiting PD-L1 positive immune cell staining.		
High	 PD-L1 Status is considered high if any of the following are met: ≥ 25% of tumor cells exhibit membrane staining; or, ICP > 1% and IC+ ≥ 25%; or, ICP = 1% and IC+ = 100%. 	
Low/negative	 PD-L1 Status is considered low/negative if: none of the criteria for PD-L1 High Status are met. 	

PD-L1 LOW/NEGATIVE STATUS: TUMOR CELL EXPRESSION < 25%

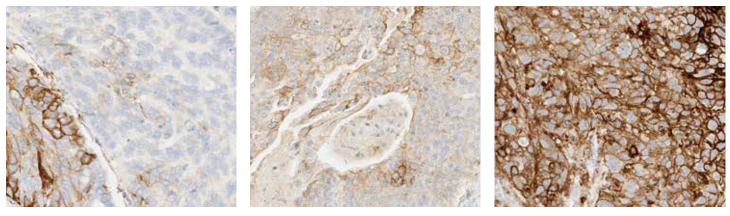


0%

5%

10%

PD-L1 HIGH STATUS: TUMOR CELL EXPRESSION $\ge 25\%$

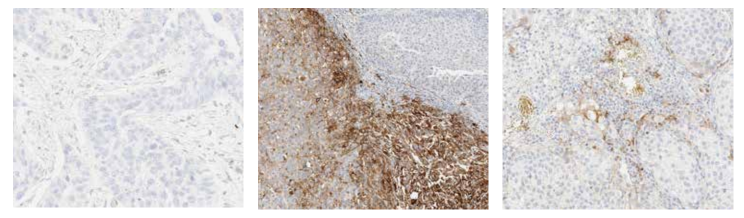


25%

50%

100%

PD-L1 LOW/NEGATIVE STATUS: TUMOR-ASSOCIATED IMMUNE CELL EXPRESSION < 25%



0% IC+

10% IC+

20% IC+

PD-L1 HIGH STATUS: TUMOR-ASSOCIATED IMMUNE CELL EXPRESSION $\geq 25\%$



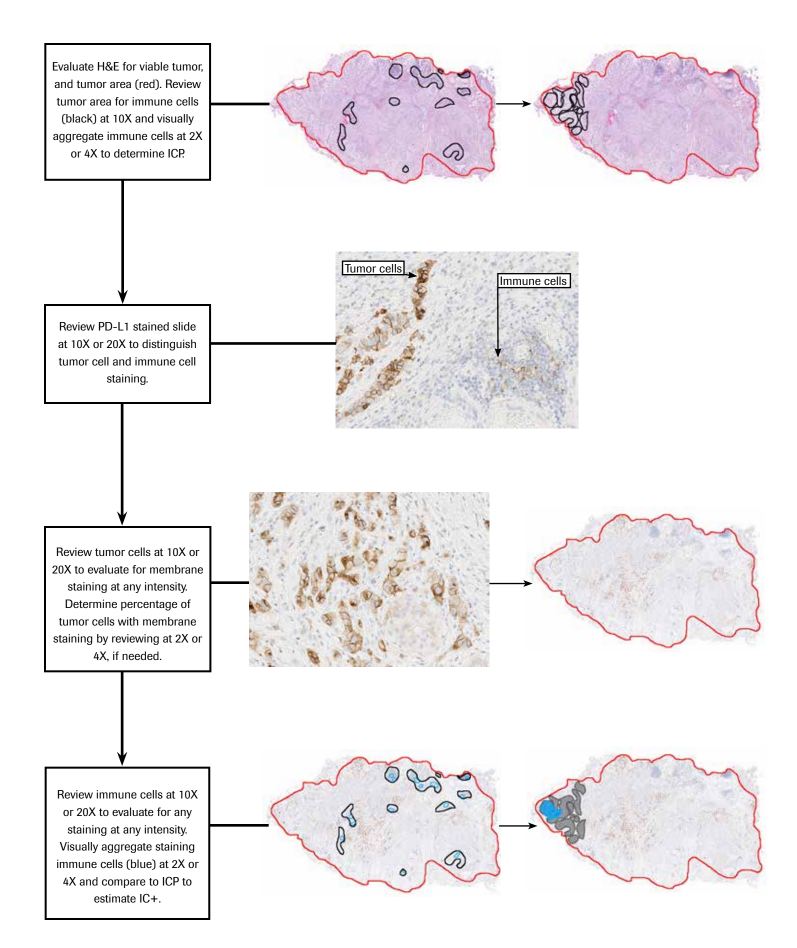
30% IC+

60% IC+

80% IC+

- 1. H&E slide is reviewed for viable tumor, tumor-associated immune cells and tumor area.
 - a. Tumor area encompasses the tumor proper, associated desmoplastic stroma and immune cells infiltrating the tumor and contained within the desmoplasia.
- 2. Visually, tumor-associated immune cells are densely aggregated (with no intervening stroma) within the tumor area when estimating Immune Cells Present (ICP).
 - a. Tumor associated immune cells include those present within the tumor reactive stroma, between the tumor islands and those invading the tumor proper.
- 3. The ICP percentage is scored as 0%, 1%, 5%, and deciles and quartiles (10, 20, 25, 30, 40, 50, 60, 70, 75, 80, 90, 100).
 - a. If a raw percentage falls between a decile or quartile, standard mathematical rounding is used to score to the nearest decile or quartile.
 - When differentiating between 1% and 5% ICP, a raw percentage estimate of 2% is rounded to 1% and an estimate of 3% to 4% is rounded to 5%.
- 4. PD-L1 stained slide is reviewed for tumor cell and immune cell staining.
- 5. Estimate the percentage of tumor cells with partial or complete membrane staining at any intensity.
- 6. The percentage of immune cells (within the ICP) expressing PD-L1 (IC+) is estimated.
 - a. Immune cells with any staining (membranous, cytoplasmic, or punctate) at any intensity are visually aggregated within the ICP to estimate IC+.
 - b. IC+ is scored as deciles and quartiles (0, 10, 20, 25, 30, 40, 50, 60, 70, 75, 80, 90, 100).
- For cases with 1% ICP, the percentage estimate for IC+ is reported as 0%, <100%, or 100%.

Overview of PD-L1 (SP263) Scoring Algorithm for Urothelial Carcinoma



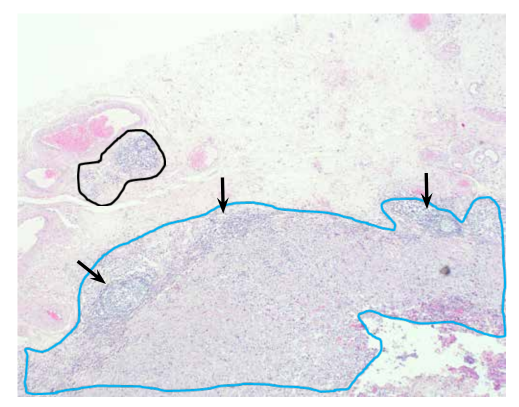
Evaluation of Immune Cell Staining

A variety of immune cells display staining with the Assay, and include lymphocytes, macrophages, histiocytes, reticular dendritic cells, plasma cells and neutrophils. The H&E-stained slide is initially examined to determine the total percentage of the tumor area (tumor cells and any desmoplastic stroma) involved by immune cells with no intervening stroma (ICP). Areas not considered part of the tumor area include non-viable tumor such as those with cautery or crush artifacts, and extensive necrosis. Normal lymphoid tissue uninvolved by the neoplasm, as seen in lymph nodes with metastatic tumor, is not considered as a part of the tumor area or immune cells involving the tumor. The PD-L1 IHC slide is then scored for the percentage of tumorassociated immune cells staining for PD-L1 (IC+). Membranous, cytoplasmic, and punctate PD-L1 immune cell staining are all included in the estimation. In cases where positively-staining immune cells are intermixed with positively-staining tumor cells, it can be difficult to quantify the amount of staining for each component. Examples of immune cell interpretation are given in the following photo sets.

The following steps describe the estimation of tumor-associated immune cell staining percentage.

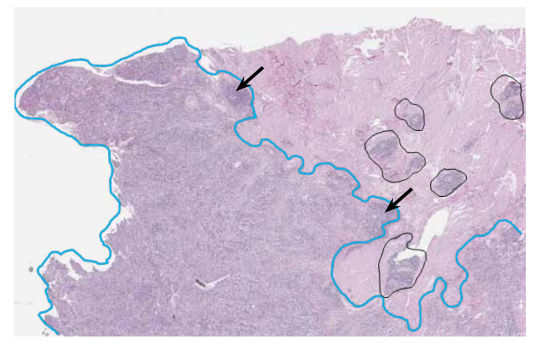
Tumor-associated immune cell staining percentage estimation:

1. Review case H&E to assess viable tumor with attached desmoplastic stroma containing tumor-associated immune cells. Tumor associated immune cells include those present within the tumor reactive stroma, between the tumor islands and those invading the tumor proper. Tumor within lymphatics is not included within the tumor area.



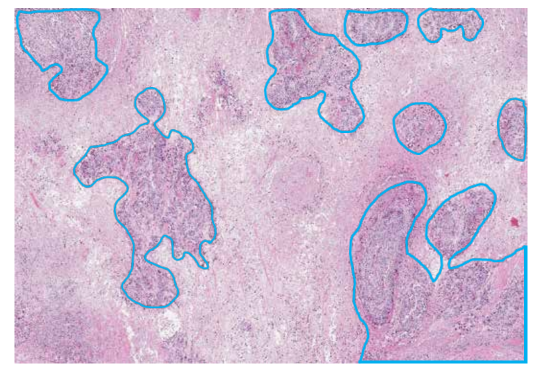
Immune Cell scoring 1: Tumor-associated immune cells (black arrows) within the tumor area (blue outline) are present at the leading edge of the tumor and are included in ICP estimation. A separate aggregate of lymphocytes is not within the tumor area (black outline) (4X)

2. Lower magnification review is recommended to perform the percentage estimate of immune cells within and adjacent to the tumor area (ICP). Immune cells located within blood vessels and lymphatics are disregarded.



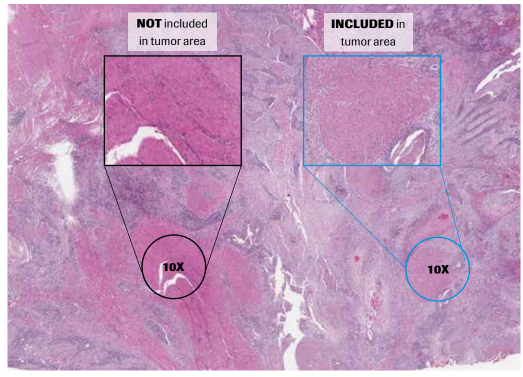
Immune Cell scoring 2: Tumor area containing tumor cells, desmoplastic stroma and tumor-associated immune cells is outlined in blue, which includes immune cell aggregates identified by black arrows. Immune cells outside of the tumor area are also present (black outlines) (1X)

3. Exclude non-neoplastic areas not involved by tumor, areas with necrotic tumor, crush and cautery artifacts.



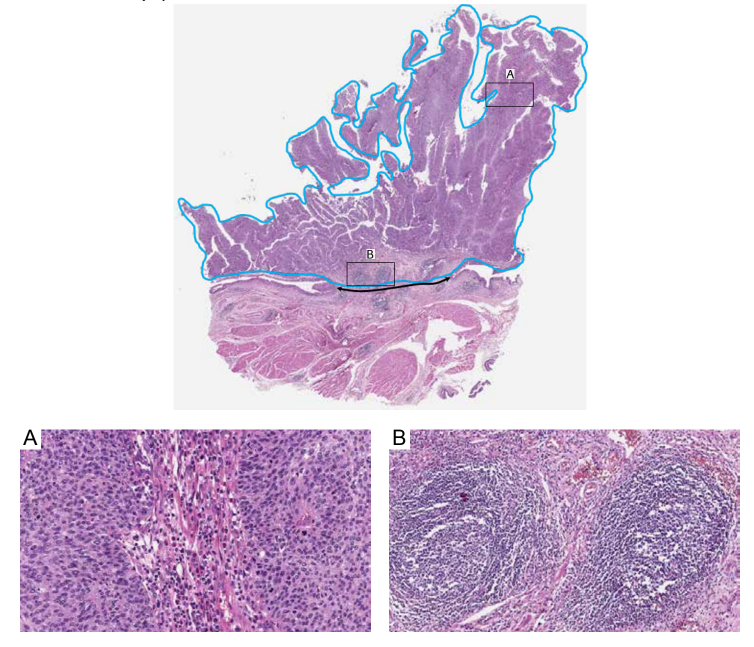
Immune Cell scoring 3: Tumor area consists of only viable areas of tumor and desmoplasia (areas outlined in blue). Any necrosis, cautery artifact, crush artifact or large areas of non-neoplastic tissue are disregarded (3X)

4. If tumor islands are separated by muscle or stroma, they are included as part of the tumor area if the tumor borders on both sides within a 10X field.



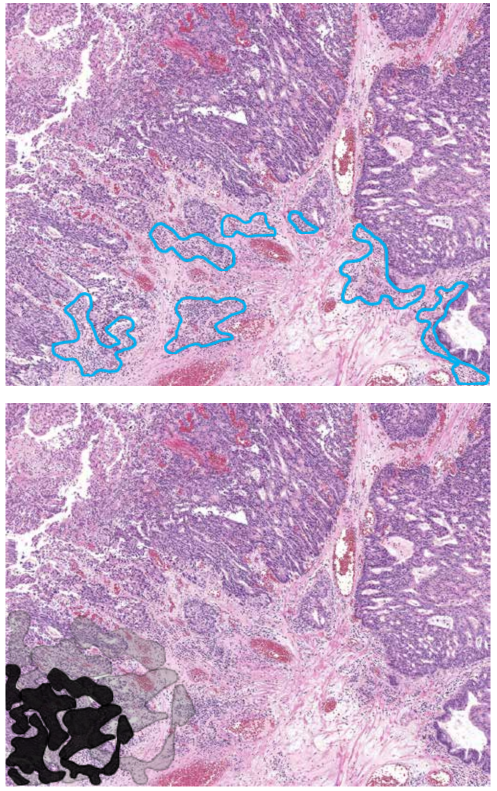
Immune Cell scoring 4: The circled areas show large areas of fibromuscular stroma uninvolved by tumor. When viewed at higher power (10X), the stroma with tumor on both sides within a single field of view (blue) is included in the overall tumor area. When the large bundle of fibromuscular stroma is not bordered by tumor within a single 10X field (black), it is not included in the overall tumor area. (1X)

5. For noninvasive urothelial carcinoma (papillary carcinoma), include the immune cells within the fibrovascular cores as well as within the immediately adjacent base of the frond/stalk.

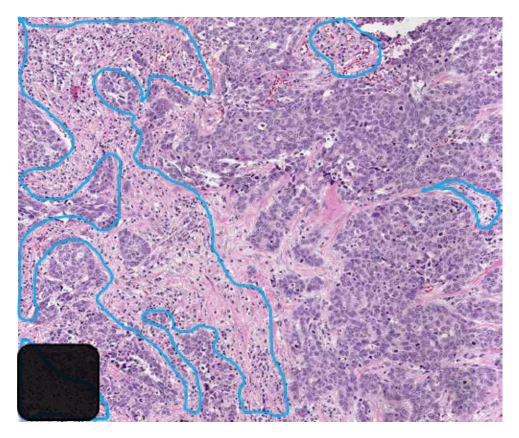


Immune Cell scoring 5: Double sided arrow (black) shows base of papillary tumor with the tumor area for this noninvasive urothelial carcinoma outlined (blue). Immune cells from the intratumoral fibrovascular core (A) (10X) and adjacent base of the stalk (B) (5X) are included in estimation of ICP (0.8X)

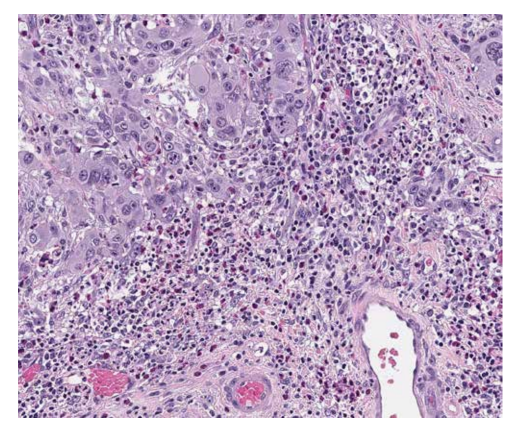
6. Visually estimate the area occupied by the tumor-associated immune cells relative to the total tumor area (ICP = percent of tumor area occupied by immune cells).



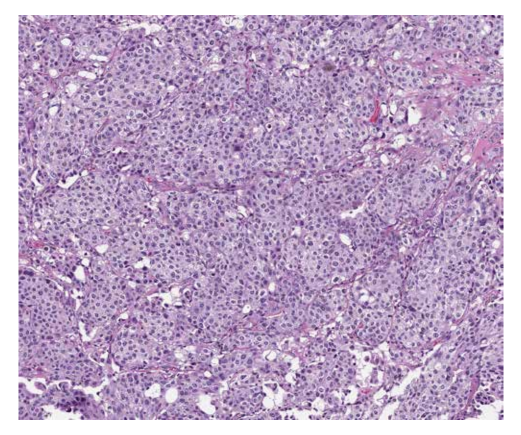
Immune Cell scoring 6: The regions occupied by immune cells within the tumor area in the upper image (blue outlines) are identified in the lower image (gray outlines) and then densely aggregated (illustrated by the smaller black areas) in order to estimate the percentage of the tumor area with immune cells, or ICP – in this example 10% (black areas together relative to the tumor area) (5X)



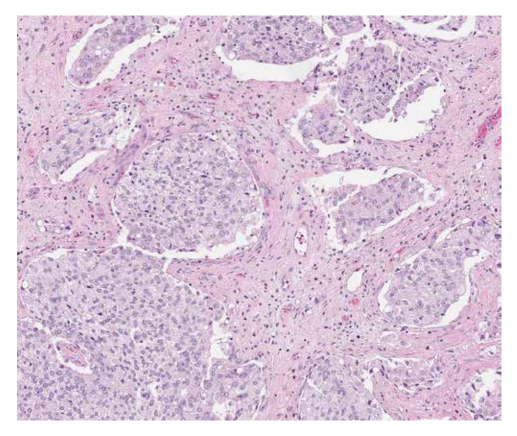
Immune Cell scoring 7: The stromal regions occupied by widely dispersed immune cells (blue outlines) are identified. When the immune cells are densely aggregated (represented by the black area) in order to estimate the percentage of the tumor area with immune cells, the ICP in this example is 5% (10X)



Immune Cell scoring 8: The immune cell infiltrates with neutrophils and eosinophils, are included in the ICP estimation. The ICP in this example is 25% (10X)

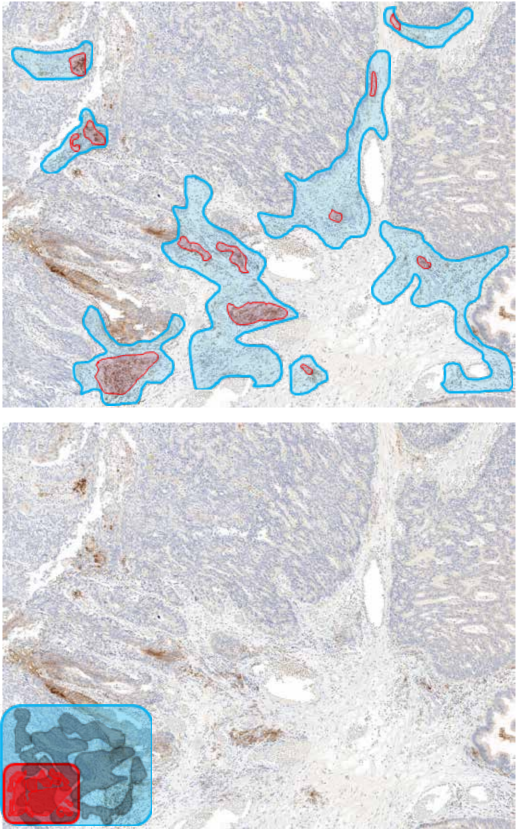


Immune Cell scoring 9: In this field of view, the ICP for the small amount of immune cells in the upper right corner is 1% (10X)



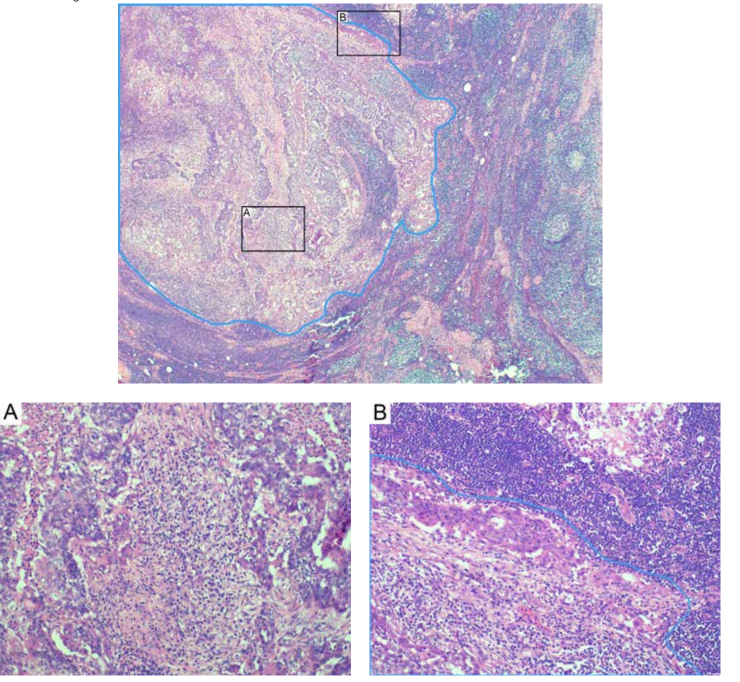
Immune Cell scoring 10: The ICP for this field of view when densely aggregated is 5% (10X)

 Review PD-L1-stained slide and estimate the percentage of tumor-associated immune cells which demonstrate PD-L1 expression (IC+) including diffuse cytoplasmic, linear membrane and punctate immune cell staining.



Immune Cell scoring 11: The immune cells comprising ICP are outlined in blue (upper image) and can be seen densely aggregated in the lower image. The immune cells with PD-L1 staining (IC+) are highlighted in red (upper image) and when aggregated constitute 25% of the ICP (lower image) (5X). Please refer to the associated H&E image contained within this guide, "Immune Cell scoring 6."

8. When evaluating lymph node metastases the reactive stroma generated by the tumor is included as part of the tumor area when determining ICP. In cases where the tumor does not generate a stromal response, the tumor area is limited to the tumor nests and adjacent immune cells in direct contact with the tumor only. Any immune cells part of the uninvolved lymphoid tissue are disregarded for ICP.



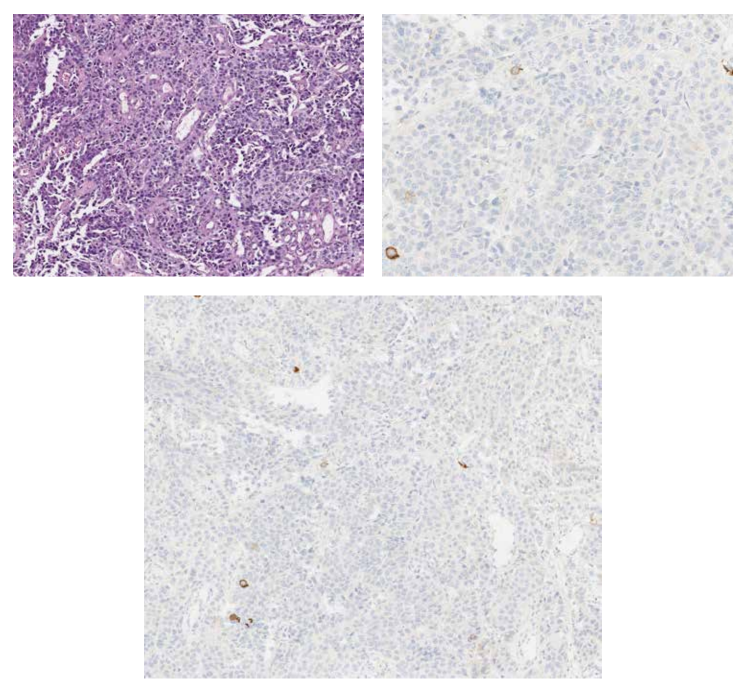
Immune Cell scoring 12: In this metastatic tumor to a lymph node, the tumor area containing tumor cells, desmoplastic stroma and immune cells is outlined in blue (4X). Higher power shows immune cells that are included as part of ICP within the reactive stroma (inset A (20X)). Along the periphery of the metastatic nodule only immune cells immediately adjacent to the tumor cells outlined in blue are included as part of ICP (inset B (20X)).

9. Only in cases where the ICP equals 1%, the IC+ is only scored as 0%, <100% or 100% due to the difficult nature of quantifying expression is such small amounts of cells.

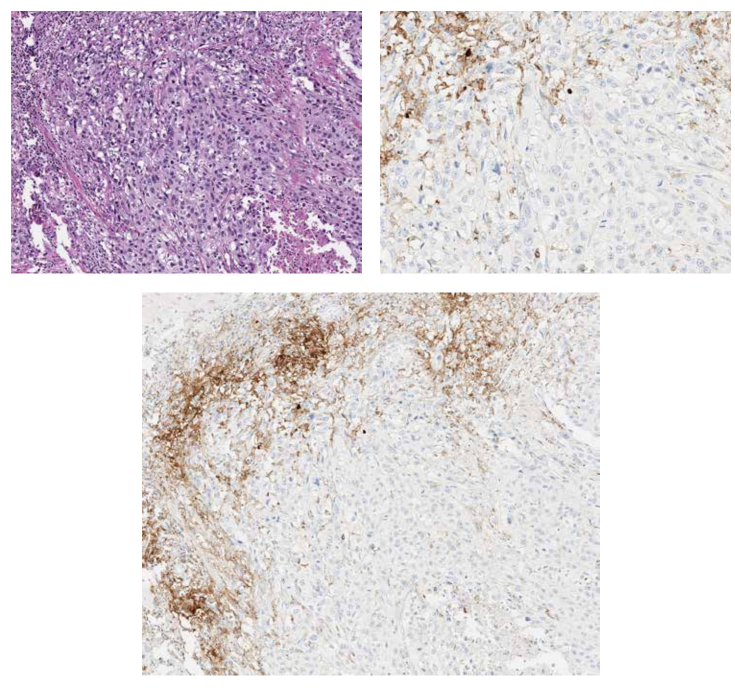
10. If the PD-L1-stained slide shows more immune cell staining than the initial ICP estimation from the H&E slide, refer back to the H&E slide and revise the ICP percentage, if necessary.

PD-L1 Expression Atlas in Urothelial Carcinoma

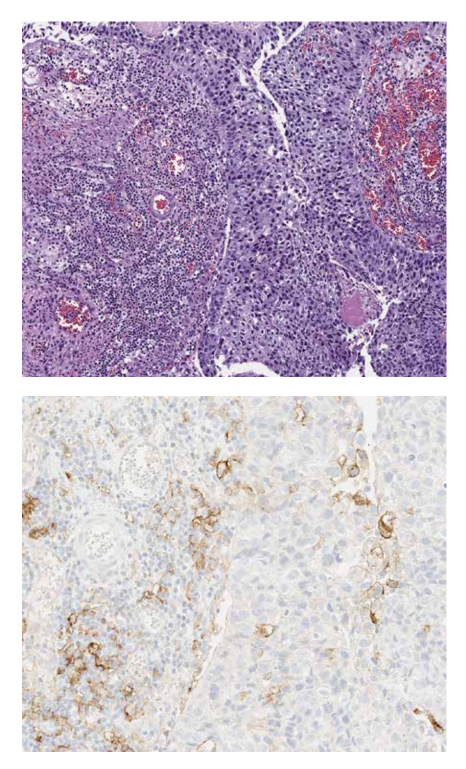
Tumor Cell Cases



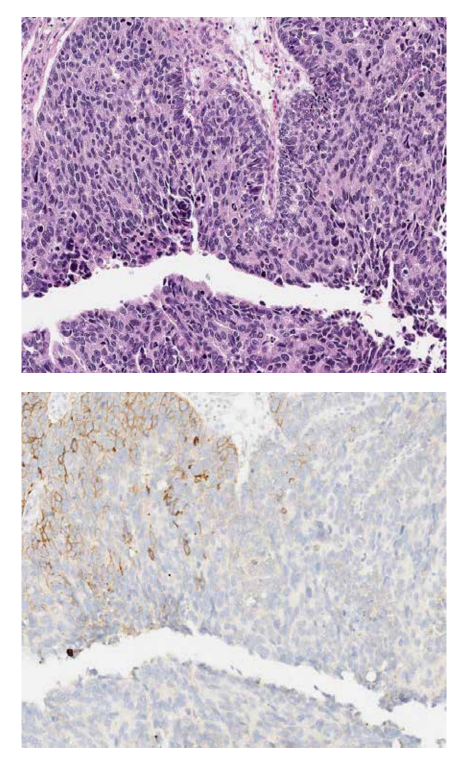
Tumor Cell Case 1: Tumor cells with 0% staining, Immune cells with 0% staining. (IC+) (Top row H&E 10X, PD-L1 20X. Bottom row PD-L1 10X.)



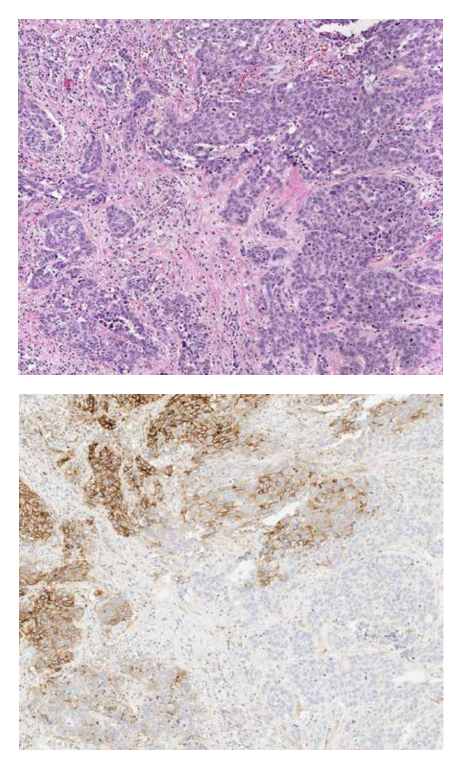
Tumor Cell Case 2: Tumor cells with 0% staining, Immune cells with 80% staining. (IC+) (Top row H&E 10X, PD-L1 20X. Bottom row PD-L1 10X.)



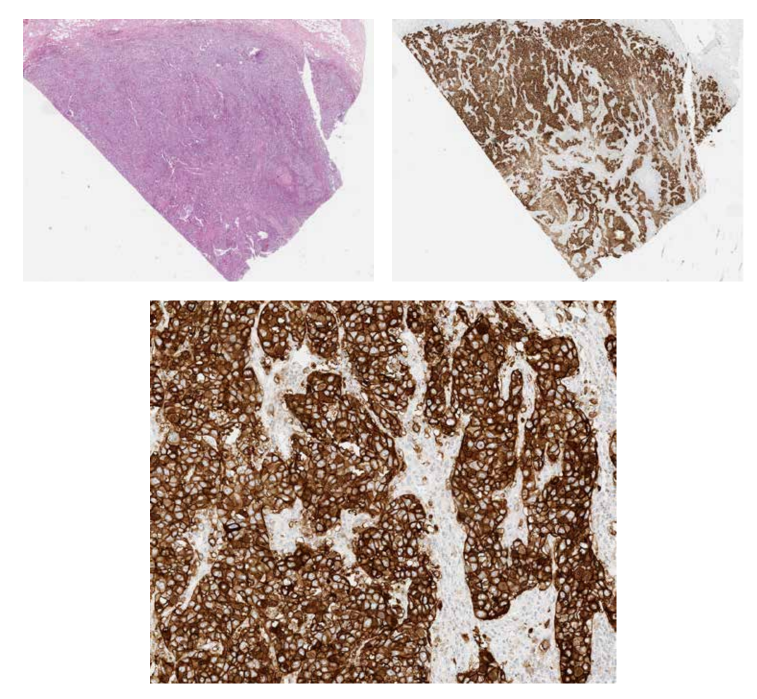
Tumor Cell Case 3: Tumor cells with 10% staining, Immune cells with 40% staining (IC+) (10X)



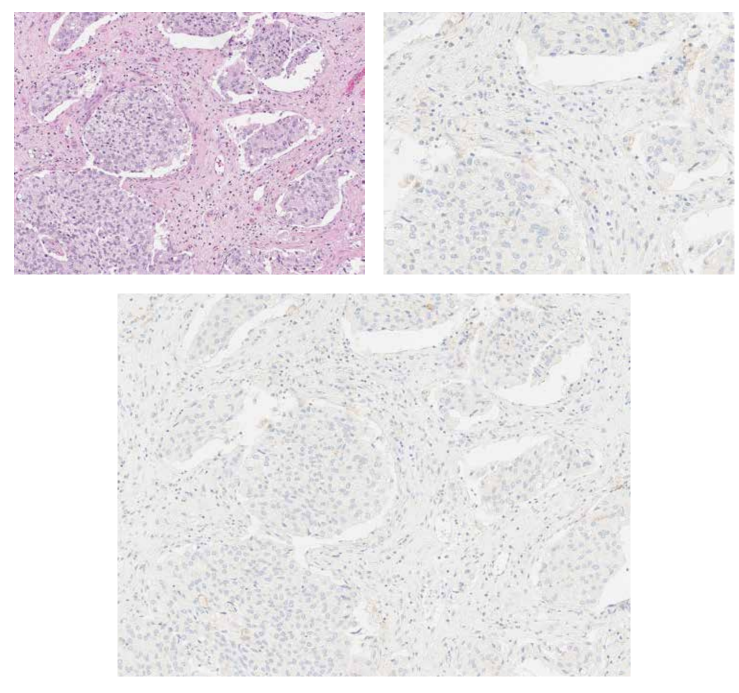
Tumor Cell Case 4: Tumor cells with 25% staining, Immune cells with 10% staining (IC+) (10X)



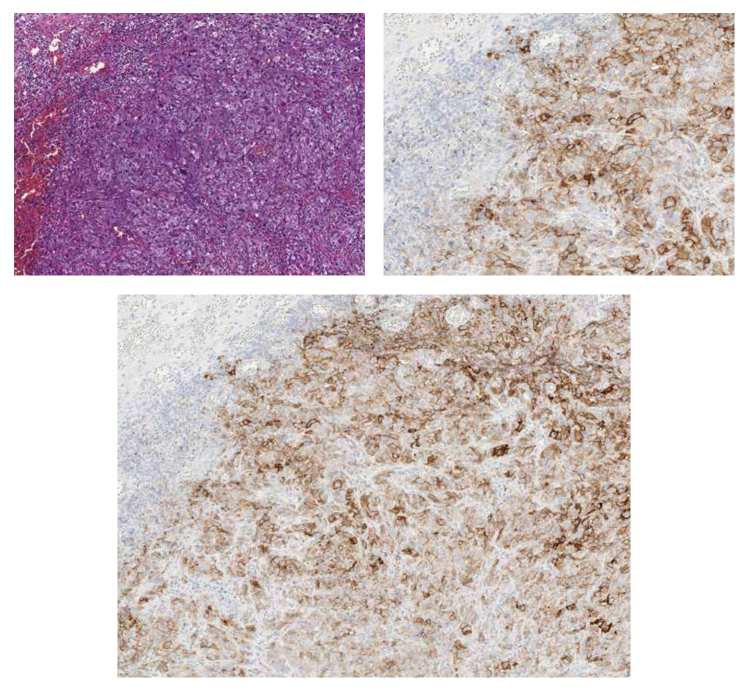
Tumor Cell Case 5: Tumor cells with 50% staining, Immune cells with 20% staining (IC+) (10X)



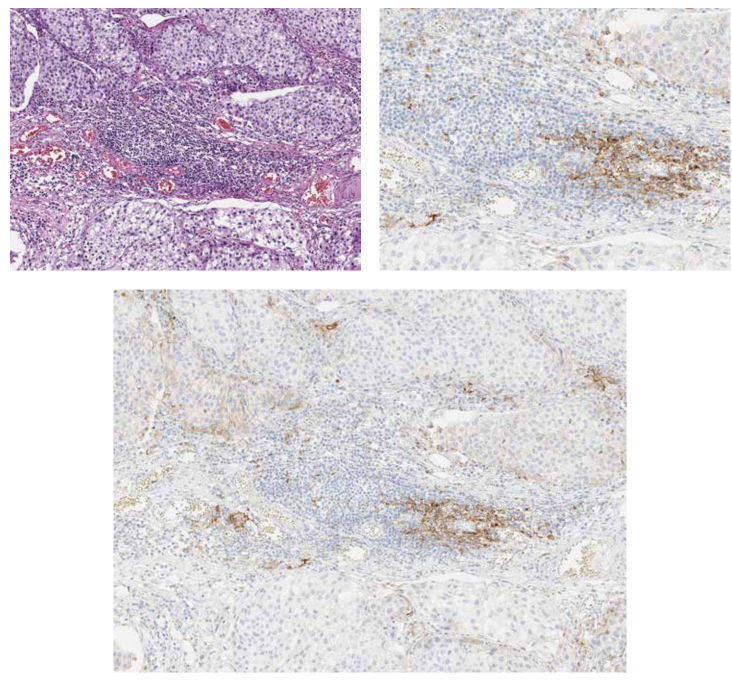
Tumor Cell Case 6: Tumor cells with 100% staining, Immune cells with 20% staining. (IC+) (Top row H&E 10X, PD-L1 20X. Bottom row PD-L1 10X.)



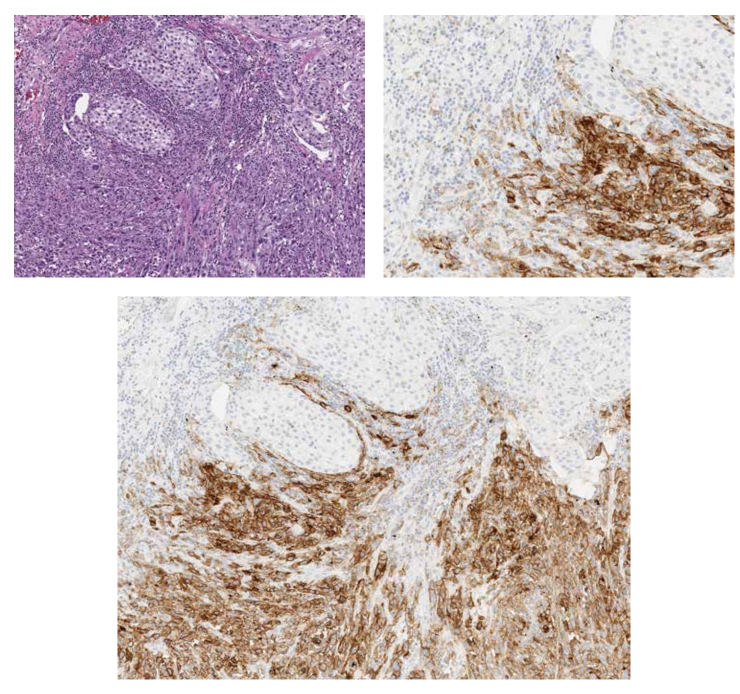
Immune Cell Case 1: Immune cells with 0% staining (IC+), Tumor cells with 0% staining. (Top row H&E 10X, PD-L1 20X. Bottom row PD-L1 10X.)



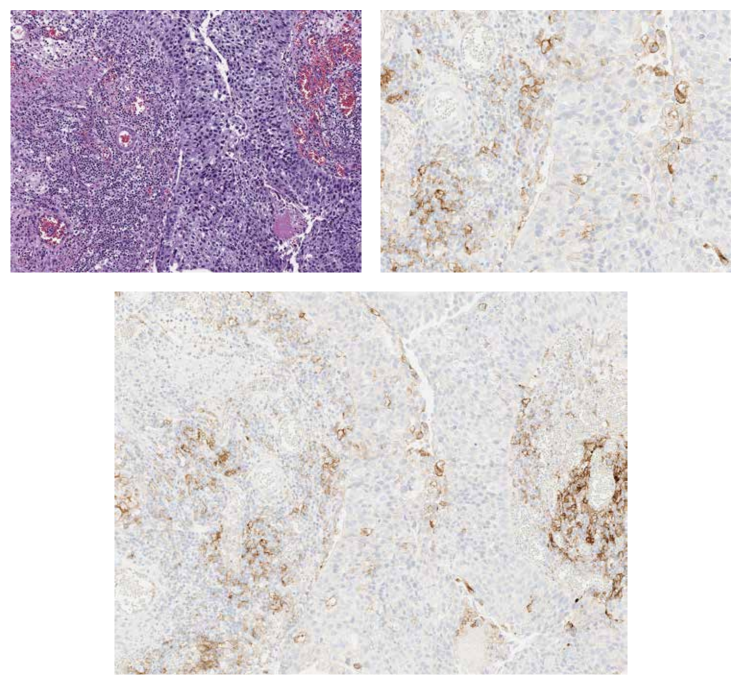
Immune Cell Case 2: Immune cells with 10% staining (IC+), Tumor cells with 80% staining. (Top row H&E 10X, PD-L1 20X. Bottom row PD-L1 10X.)



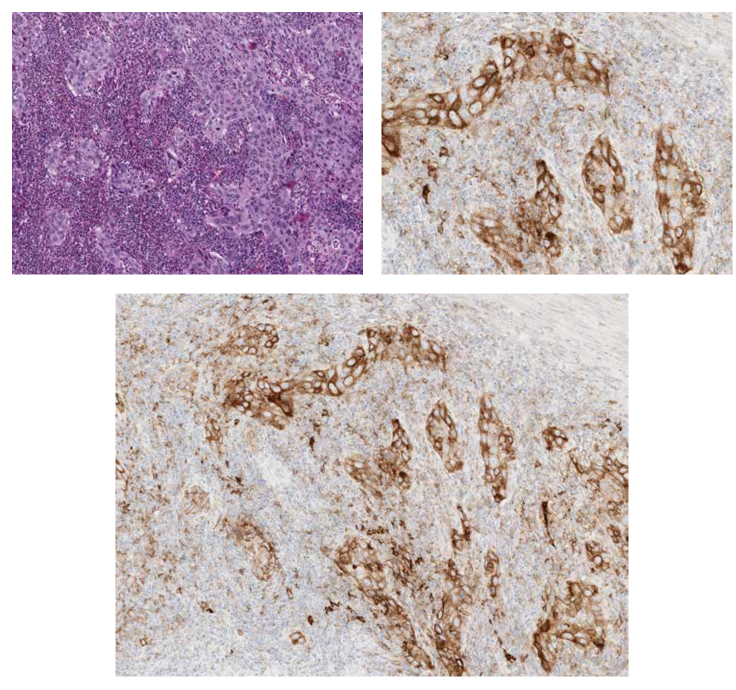
Immune Cell Case 3: Immune cells with 30% staining (IC+), Tumor cells with 10% staining. (Top row H&E 10X, PD-L1 20X. Bottom row PD-L1 10X.)



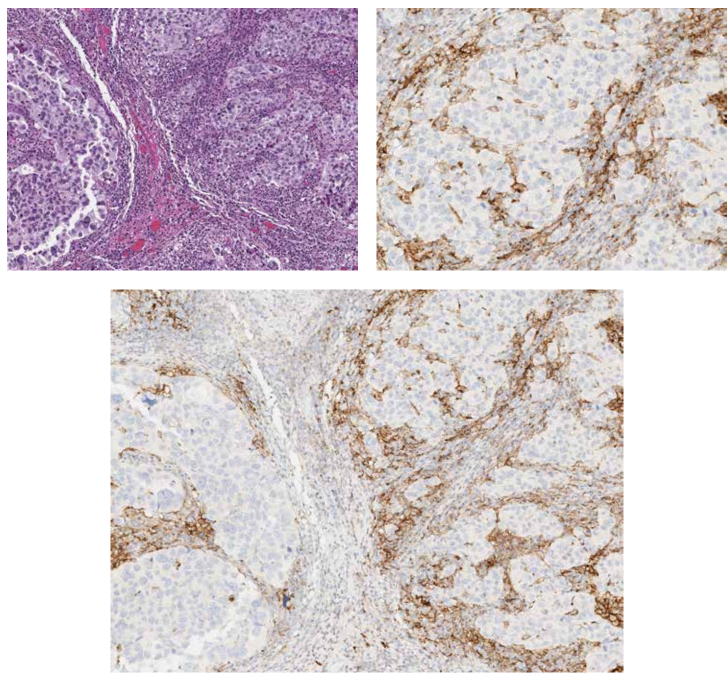
Immune Cell Case 4: Immune cells with 25% staining (IC+), Tumor cells with 100% staining. (Top row H&E 10X, PD-L1 20X. Bottom row PD-L1 10X.)



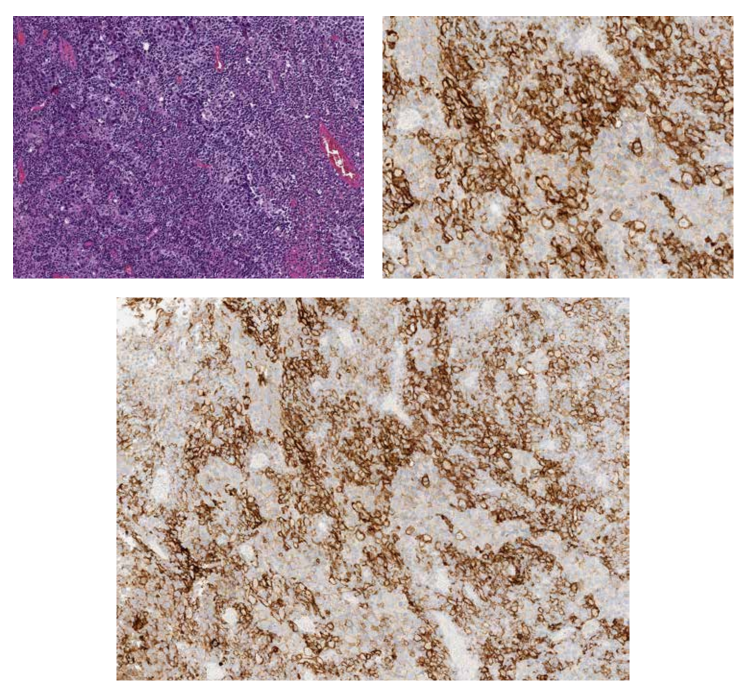
Immune Cell Case 5: Immune cells with 40% staining (IC+), Tumor cells with 10% staining. (Top row H&E 10X, PD-L1 20X. Bottom row PD-L1 10X.)



Immune Cell Case 6: Immune cells with 50% staining (IC+), Tumor cells with 100% staining. (Top row H&E 10X, PD-L1 20X. Bottom row PD-L1 10X.)



Immune Cell Case 7: Immune cells with 60% staining (IC+), Tumor cells with 0% staining. (Top row H&E 10X, PD-L1 20X. Bottom row PD-L1 10X.)



Immune Cell Case 8: Immune cells with 90% staining (IC+), Tumor cells with 100% staining. (Top row H&E 10X, PD-L1 20X. Bottom row PD-L1 10X.)

Challenging Cases

Cases are given a PD-L1 score according to percentage of tumor cells with membrane staining and tumor-associated immune cells. Various staining patterns and morphologic features may make interpretation and quantification of tumor membrane staining difficult.

Some cases may be particularly challenging due to the following issues:

Weak Tumor Cell Membrane vs Weak Tumor Cell Cytoplasmic Staining

Some specimens may exhibit weak cytoplasmic staining of the tumor cells that may be confused at low power with weak tumor cell membrane staining. For this reason when evaluating PD-L1 IHC stained slides, weak staining should be confirmed with examination at higher powers to distinguish between tumor cell membranous and cytoplasmic staining.

Strong Immune Cell Staining Overlapping with Tumor Cell Staining

Some tumors may contain an extensive inflammatory component both surrounding the tumor and infiltrating within the tumor. In instances when significant staining is seen for both tumor and immune cells, it can be challenging to differentiate and quantify the PD-L1 IHC staining between the two cell populations. The presence of immune cells infiltrating the tumor should be confirmed using the H&E slide. The pattern of PD-L1 staining and nuclear morphology are utilized to help attribute expression to immune cells (punctate staining, small uniform nucleus) and tumor cells (linear membrane staining, large irregular nucleus).

Borderline PD-L1 (SP263) Status

Some cases are near the cut-off between a PD-L1 High and Low/negative status. These cases are particularly challenging to estimate the percent of target cells with staining. For these borderline cases it may be helpful to view the case at a magnification that enables the entire tumor area to be assessed. Consultation with a trained colleague may also be helpful.

Cases with Multiple Tissue Fragments

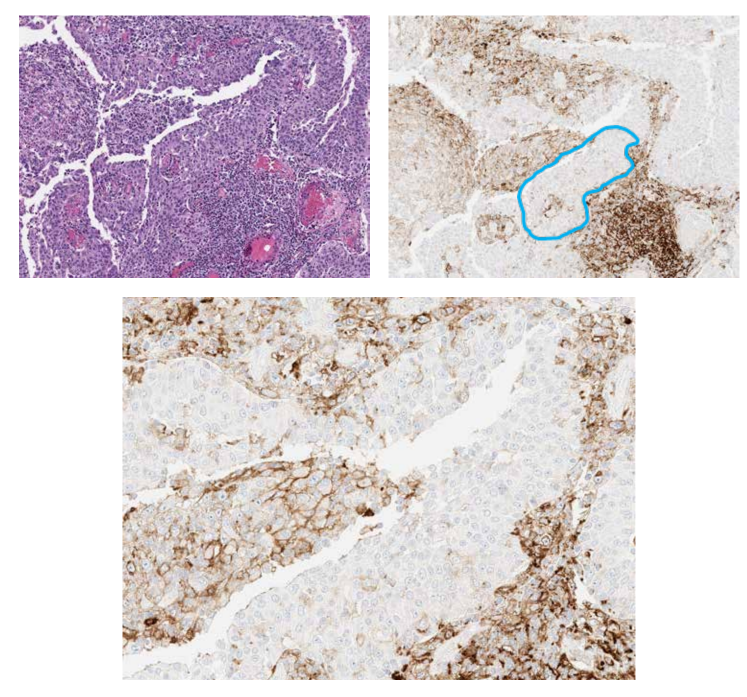
Urothelial carcinoma samples resulting from transurethral resections of the bladder (TURB) can result in challenging cases complicated by multiple fragments of tissue scattered across the slide with varying amounts of necrosis, crush/cautery artifact, viable tumor cells and/or tumor-associated immune cells. A fragment-by-fragment examination approach with individual percent staining estimates followed by an overall average can help in the evaluation.

Obscuring Endogenous Material

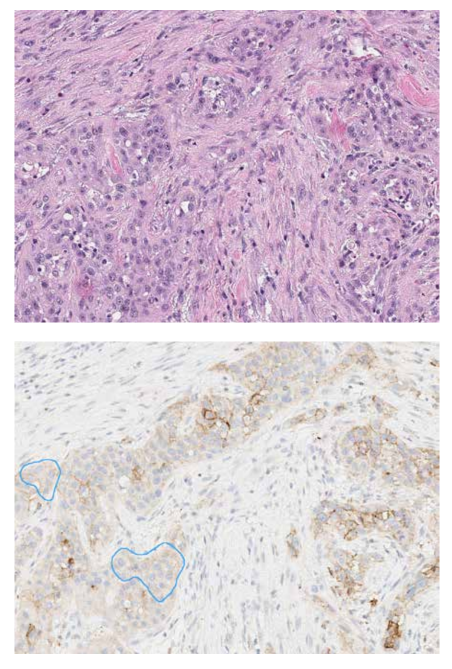
Occasionally in urothelial carcinoma samples endogenous material, such as melanin pigment or hemosiderin, may obscure and interfere with interpretation of PD-L1 IHC staining of tumor and immune cells. Comparison of the negative isotype control slide with the PD-L1 stained slide can aid in differentiating between biomarker staining and endogenous material. Non-specific staining can also be seen on the PD-L1 stained slide in areas of necrosis.

Some challenging cases are shown.

Challenging cases: Weak Tumor Membrane vs Cytoplasmic Staining

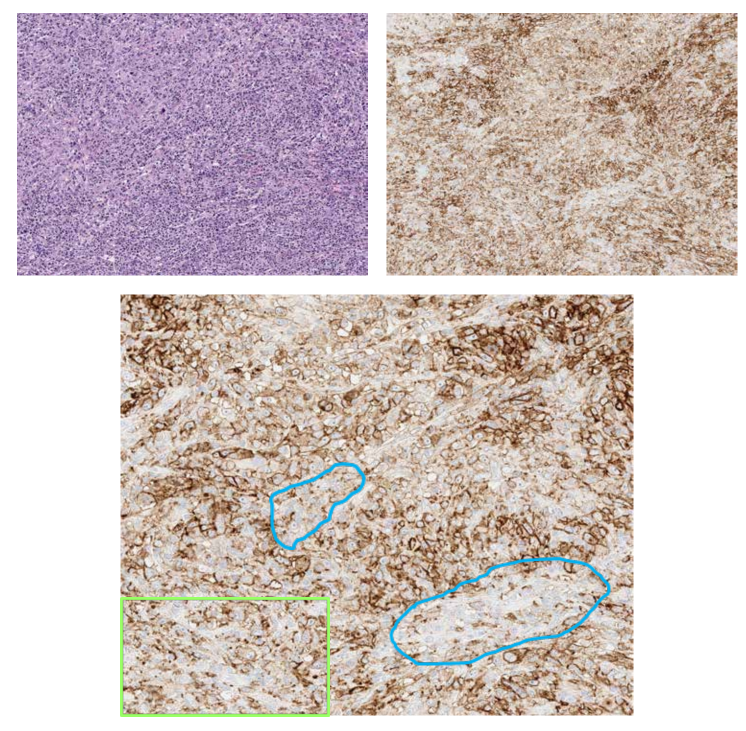


Challenging Case 1: The area outlined in blue (10X PD-L1 image, upper right) contains weak tumor cell staining that is difficult to distinguish. Examination at higher power (20X) shows there is scattered weak tumor membrane staining. For 10X image: Tumor cells with 40% staining, Immune cells with 80% staining. (IC+) (Top row H&E 10X, PD-L1 10X. Bottom row PD-L1 20X.)

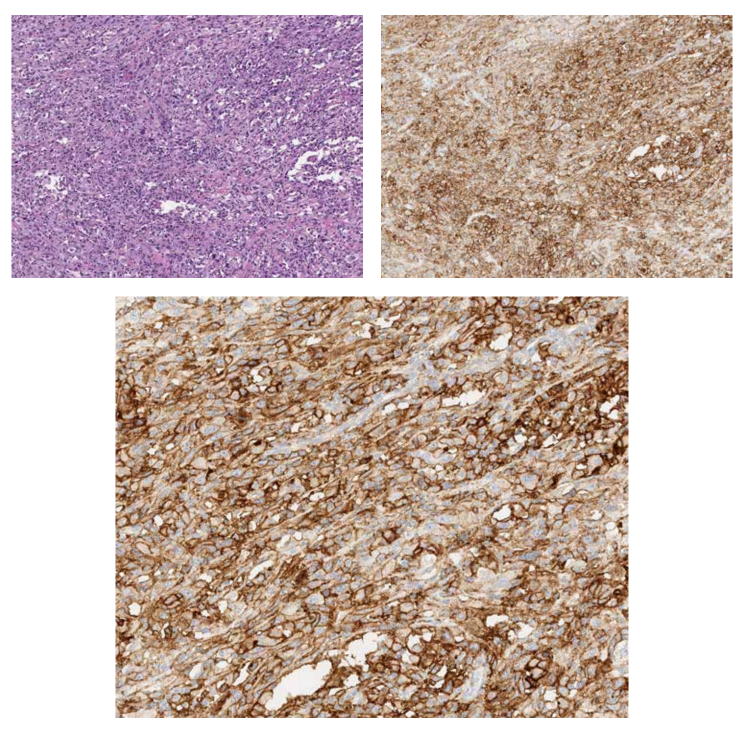


Challenging Case 2: Tumor cells with moderate or stronger membrane staining are readily identified within the right half of this field. Careful examination of the tumor demonstrates cells with weak membrane expression (blue outlines) that should also be counted towards the total percentage of membrane staining.

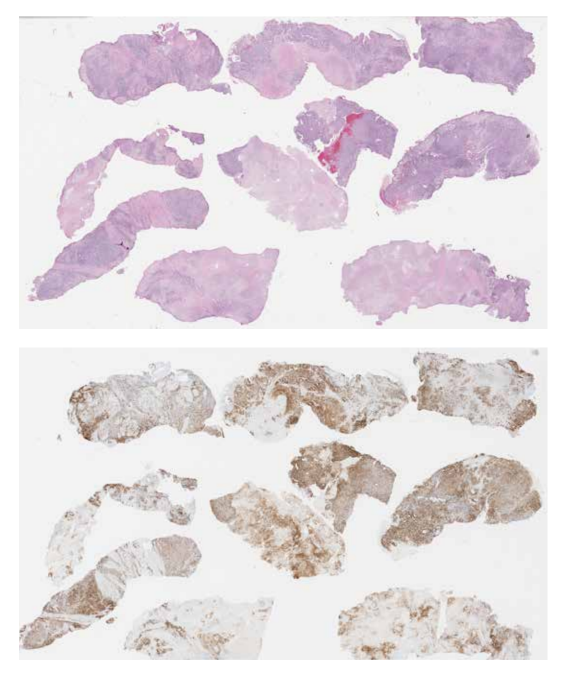
Challenging cases: Strong Immune Cell Staining Overlapping with Tumor Cell Staining



Challenging Case 3: TC and IC intermixed together with both expressing PD-L1. In areas with strong TC staining, IC can only be evaluated where discernible from tumor cell staining. The different patterns of IC staining (punctate pattern) as well as nuclear morphology can be used to identify PD-L1 expression. The blue outlines contain areas with discernible immune cell staining with the green outlined area having scattered punctate immune cell staining. Tumor cells with 100% staining, Immune cells with 25% staining (IC+) (Top row H&E 10X, PD-L1 10X. Bottom row PD-L1 20X.)

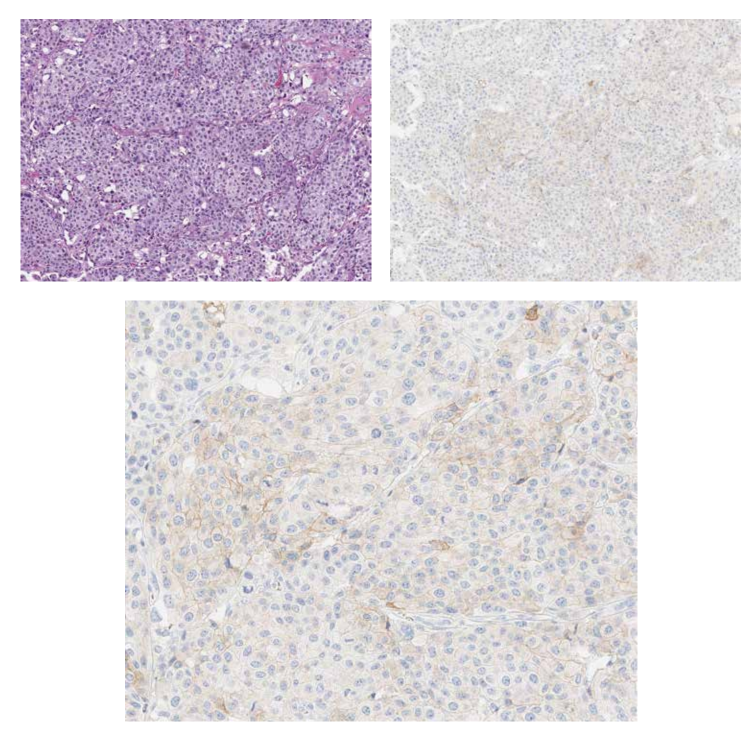


Challenging Case 4: TC and IC intermixed together with TC membrane staining more distinct and IC difficult to evaluate and can only be scored along the periphery of the tumor. Tumor cells with 100% staining. (Top row H&E 10X, PD-L1 10X. Bottom row PD-L1 20X.)

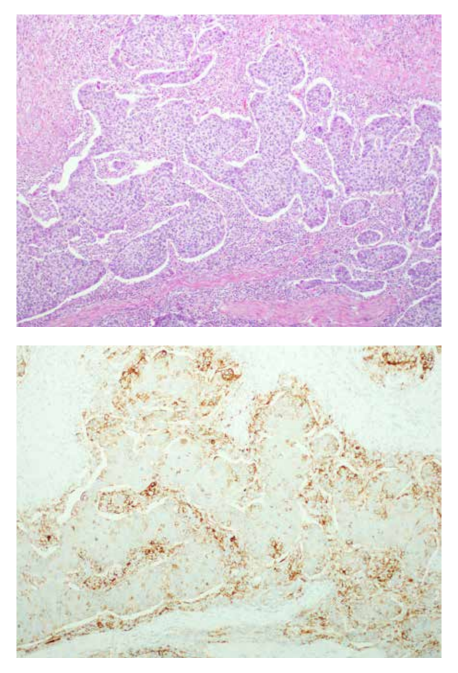


Challenging Case 5: TURB specimen contains multiple pieces with crush artifact, blood clot and necrosis which shows non-specific staining on the IHC slide(1X).

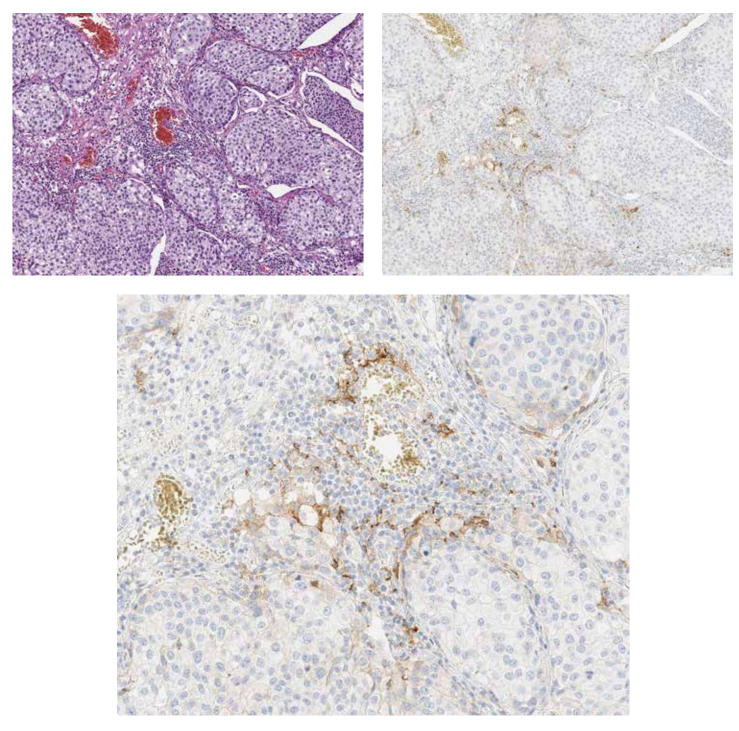
Challenging cases: Borderline PD-L1 Status



Challenging Case 6: Weak tumor cell staining seen at 10X magnification is difficult to distinguish between membrane and cytoplasmic staining. Examination at higher power (20X) reveals tumor cells with 20% weak, delicate membranous staining.

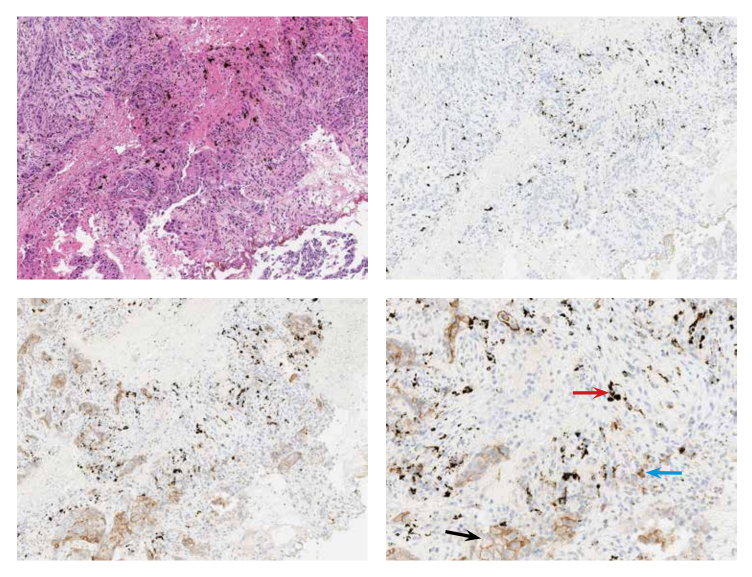


Challenging Case 7: Borderline TC PD-L1 expression with 25% tumor cell membrane staining (10X)

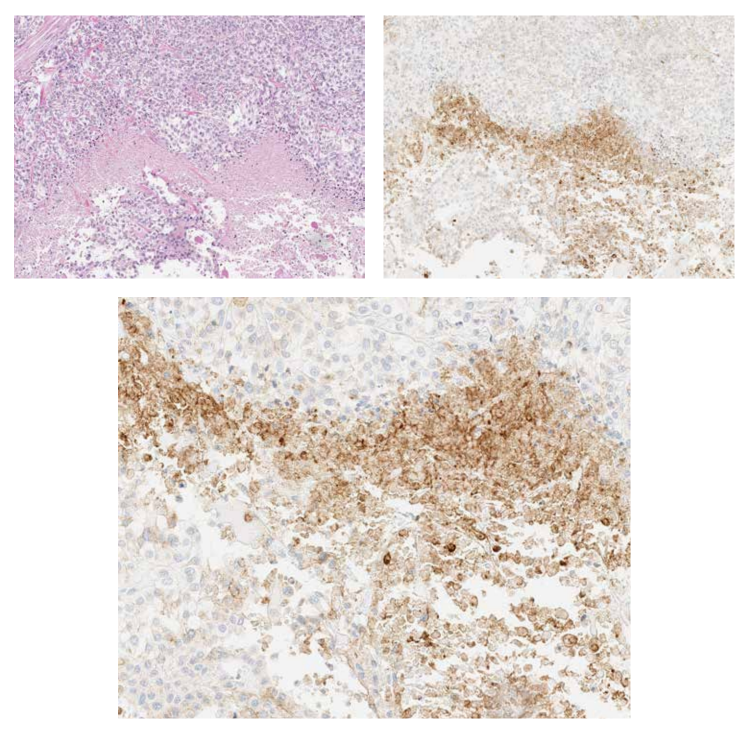


Challenging Case 8: Borderline IC PD-L1 expression with an IC+ of 20% (20X). (Top row H&E 10X, PD-L1 10X. Bottom row PD-L1 20X.)

Challenging cases: Obscuring Endogenous Material



Challenging Case 9: Case contains anthracotic pigment (red arrow) overlapping with punctate immune cell staining (blue arrow) as well as tumor cell membrane staining (black arrow) (20X). (Top row H&E 10X, Neg Rb 10X. Bottom row PD-L1 10X.)



Challenging Case 10: Large necrotic areas can have non-specific DAB staining which may be confused with immune cell staining. (Top row H&E 10X, PD-L1 10X. Bottom row PD-L1 20X.)

Impact of Pre-Analytical Conditions on VENTANA PD-L1 (SP263) Assay Staining

Fixative Recommendations to Achieve Optimal Staining Results with the VENTANA PD-L1 (SP263) Assay

Ventana recommends fixation in 10% NBF for 6-72 hours. Acceptable fixatives and fixation times are outlined in blue.

Time	VENTANA PD-L1 (SP263) Assay Staining of Placenta Tissue Across Fixatives and Fixation Times Fixative							
Point (Hrs)	10% NBF	Zinc Formalin	PREFER fixative** AFA**		Alcoholic Formalin**	95% Ethanol**		
1*		100 00						
6								
12								
24								
48								
72								

NOTES

*One hour fixation time is not recommended for all fixative types

**Use of PREFER fixative and 95% Ethanol (weaker staining) or alcoholic fixatives (high background) is not recommended. See additional higher magnification images to the right.



Impact of Tissue Thickness on Assay Staining

Ventana recommends tissue thickness of 4-5 microns for use with the Assay.

Cut Slide Stability

Sections approximately 4-5 microns in thickness should be cut and mounted on positively charged glass slides. Slides should be stained within 6 months of sectioning.

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Ventana Medical Systems, Inc. 1910 E. Innovation Park Drive Tucson, Arizona 85755 USA +1 520 887 2155 +1 800 227 2155 (USA)

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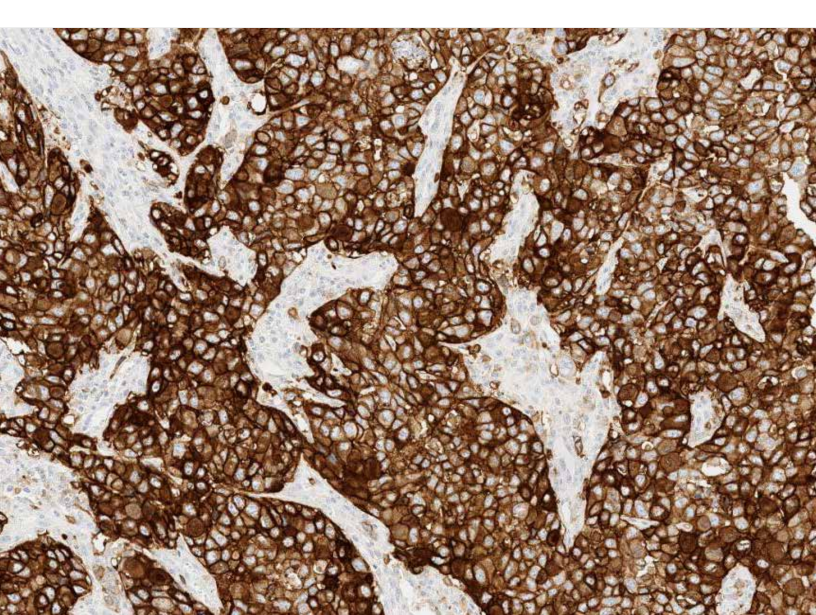
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NOTES:

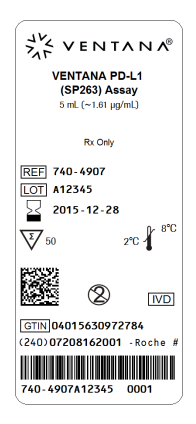
- 1. Text: Black. Background: White.
- 2. Make from: Ventana P/N 1442800 blank label stock (DMS Mat. No. 05249368001), approximately 3.00 x 1.75".
- 3. The individual label file can be located in the Document Control System (DCS) under 07447710001c.btw.
- 4. Outside border line is shown here for illustrative purposes only and does not print.
- 5. Label image is shown here at an approximate size. Label dimensions are contained in the drawing of the label stock found on the bill of materials.
- 6. Lot Number, Expiry Date, and 2D barcode are shown here for illustrative purposes only. Actual data will vary by job.



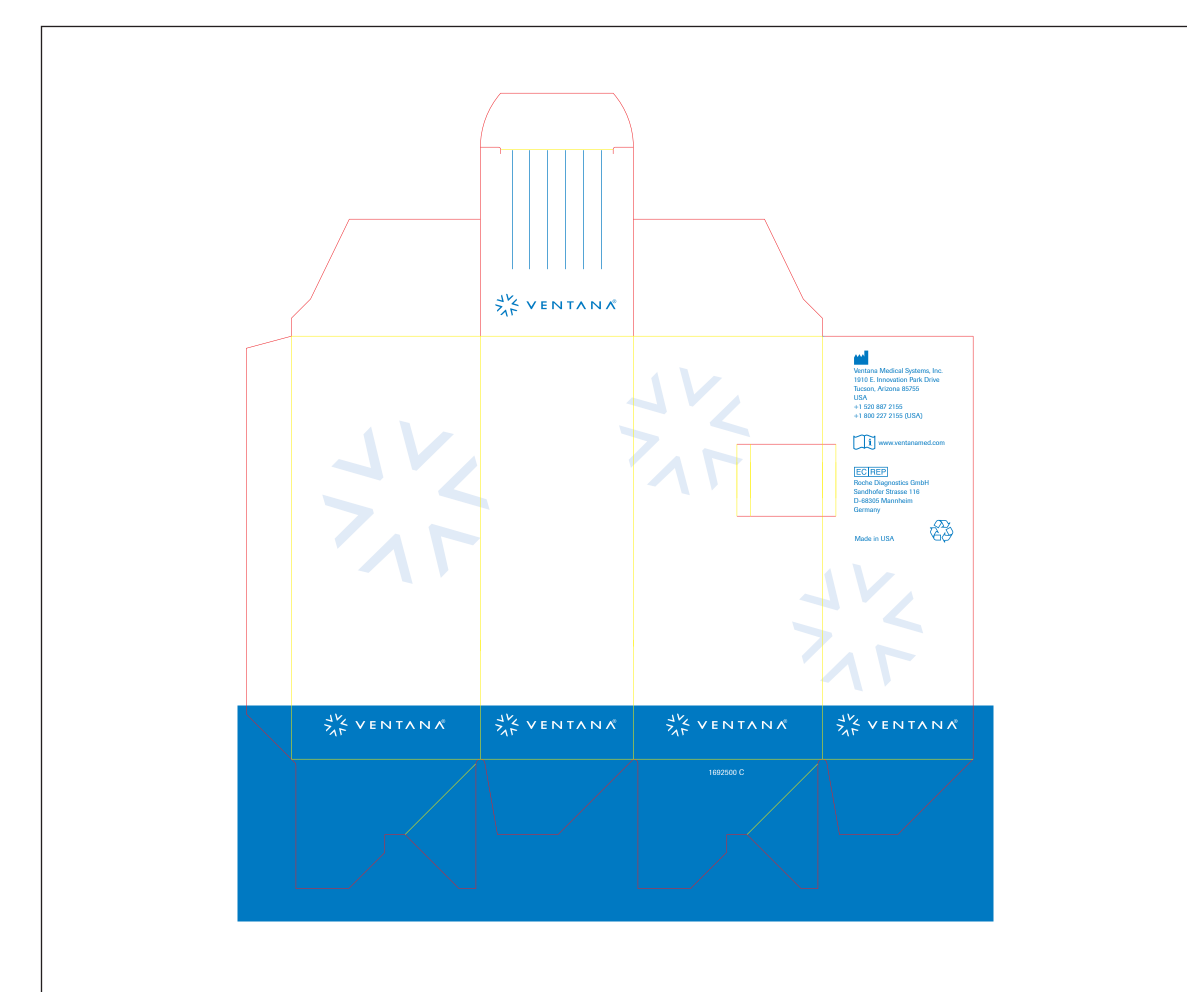
С	Change product name, ad	See Agile	CS	C057844		
REV	DESCRIPTION			EFFECTIVE DATE	RP	CO
TITLE: LBL, 1 DSPNR BRL, 2DBC, 740-4907						
This document contains confidential, proprietary information of Ventana Medical Systems, Inc. It may not be copied or reproduced without prior written permission from Ventana Medical Systems, Inc.					REV: C	
Ventan	<u>a Medical Systems, Inc.</u> Tucson, Arizona	PART NUMBER: 07447710001		RIAL NUMBER: 7710001	SHEET 1 OF 1	
FT0710-300	Rev A					

NOTES

- 1. Text: Black. Background: White.
- 2. Make from: Ventana P/N 07184603001, approximately 1.75 x 4.0".
- 3. The individual label file can be located in the Document Control System (DCS) under 07447396001c.btw.
- 4. Outside border line is shown here for illustrative purposes only and does not print.
- 5. Label image is shown here at an approximate size. Label dimensions are contained in the drawing of the label stock found on the bill of materials.
- Lot Number, Expiry Date, barcodes, and barcode-related human readable text are shown here for illustrative purposes only. Actual data will vary by job.



С	Change product name, add	See Agile	CS	C057844				
REV	DESCRIPTION EFFECTIVE DATE RF				CO			
LBL, 1 DSPNR BOX, 2DBC, FG 740-4907								
This document contains confidential, proprietary information of Ventana I It may not be copied or reproduced without prior written permission from Vent					REV: C			
Ventan	a Medical Systems, Inc. Tucson, Arizona	PART NUMBER: 07447396001	DMS MATERIAL NUMBER: 07447396001		SHEET 1 OF 1			



С	UPDA	ATE REV FOR SP	SEE AGILE	GV	C007663			
В	UPDATE ARTWORK WITH BLUE BORDER				SEE AGILE	PM	C003389	
Α	RELEASED DRAWING				SEE AGILE	GA	10682	
REV	DESCRIPTION				Effective Date	RP	CO	
$\frac{2V_{2}}{V_{AF}}$ VENTAN $\overset{\text{Ventana Medical Systems Inc.}}{\text{Tucson, AZ}}$								
ARTWORK, SINGLE DISPENSER, BOX PN 1692500								
SCA	ALE	SHEET	SIZE	DWG NUMBER			REV	
DO I SCA		1 OF 1	В	000014900				

NOTES:

1. COLOR SWATCHES a.PMS 300