

# Protocol for the Examination of Specimens From Patients With Carcinoma of the Penis

**Version:** 4.1.0.0

Protocol Posting Date: June 2021

CAP Laboratory Accreditation Program Protocol Required Use Date: March 2022

The changes included in this current protocol version affect accreditation requirements. The new deadline for implementing this protocol version is reflected in the above accreditation date.

## For accreditation purposes, this protocol should be used for the following procedures AND tumor types:

Procedure	Description
Penectomy	Includes specimens designated partial penectomy and total penectomy.
Circumcision	Required if margins can be assessed.
Tumor Type	Description
Carcinoma	Includes carcinomas arising from foreskin glands or penile shaft.

## This protocol is NOT required for accreditation purposes for the following:

Procedure				
Biopsy (incisional or excisi	onal)			
Primary resection specime	n with no residual	cancer (eg, following neoa	djuvant therapy)	
Cytologic specimens				

#### The following tumor types should NOT be reported using this protocol:

	ar approximation and representational grand processing
Tumor Type	
Urothelial carcinoma	(consider Urethra protocol)
Lymphoma (consider the Hodgkin or non-Hodgkin Lymphoma protocols)	
Sarcoma (consider th	e Soft Tissue protocol)

#### **Authors**

Antonio L. Cubilla, MD\*; Gladell P. Paner, MD\*; Ming Zhou, MD, PhD\*; Lara R. Harik, MD; Robert Allan, MD; Mahul B. Amin, MD; Jonathan I. Epstein, MD; David J. Grignon, MD; Peter A. Humphrey, MD, PhD; Curtis A. Pettaway, MD; Jason Pettus, MD; Victor E. Reuter, MD; John R. Srigley, MD; Elsa F. Velazquez, MD.

With guidance from the CAP Cancer and CAP Pathology Electronic Reporting Committees.

<sup>\*</sup> Denotes primary author.

## **Accreditation Requirements**

This protocol can be utilized for a variety of procedures and tumor types for clinical care purposes. For accreditation purposes, only the definitive primary cancer resection specimen is required to have the core and conditional data elements reported in a synoptic format.

- <u>Core data elements</u> are required in reports to adequately describe appropriate malignancies. For accreditation purposes, essential data elements must be reported in all instances, even if the response is "not applicable" or "cannot be determined."
- <u>Conditional data elements</u> are only required to be reported if applicable as delineated in the protocol. For instance, the total number of lymph nodes examined must be reported, but only if nodes are present in the specimen.
- Optional data elements are identified with "+" and although not required for CAP accreditation purposes, may be considered for reporting as determined by local practice standards.

The use of this protocol is not required for recurrent tumors or for metastatic tumors that are resected at a different time than the primary tumor. Use of this protocol is also not required for pathology reviews performed at a second institution (ie, secondary consultation, second opinion, or review of outside case at second institution).

#### **Synoptic Reporting**

All core and conditionally required data elements outlined on the surgical case summary from this cancer protocol must be displayed in synoptic report format. Synoptic format is defined as:

- Data element: followed by its answer (response), outline format without the paired Data element: Response format is NOT considered synoptic.
- The data element should be represented in the report as it is listed in the case summary. The response for any data element may be modified from those listed in the case summary, including "Cannot be determined" if appropriate.
- Each diagnostic parameter pair (Data element: Response) is listed on a separate line or in a tabular format to achieve visual separation. The following exceptions are allowed to be listed on one line:
  - o Anatomic site or specimen, laterality, and procedure
  - Pathologic Stage Classification (pTNM) elements
  - Negative margins, as long as all negative margins are specifically enumerated where applicable
- The synoptic portion of the report can appear in the diagnosis section of the pathology report, at the end of the report or in a separate section, but all Data element: Responses must be listed together in one location

Organizations and pathologists may choose to list the required elements in any order, use additional methods in order to enhance or achieve visual separation, or add optional items within the synoptic report. The report may have required elements in a summary format elsewhere in the report IN ADDITION TO but not as replacement for the synoptic report ie, all required elements must be in the synoptic portion of the report in the format defined above.

#### **Summary of Changes**

### v 4.1.0.0

- General Reformatting
- Revised Margins Section
- Revised Lymph Nodes Section
- Added Distant Metastasis Section
- Removed pTX and pNX Staging Classification

## Reporting Template

Select a single response unless otherwise indicated.
CASE SUMMARY: (PENIS) Standard(s): AJCC-UICC 8 This case summary is recommended for reporting biopsy specimens, but is not required for accreditation purposes.
SPECIMEN (Note A)
Procedure Incisional biopsy Excisional biopsy Partial penectomy Total penectomy Circumcision Other (specify): Not specified
Foreskin (presence and type)  Not identified (circumcised)  Present (uncircumcised)  Short  Medium  Long  Phimotic  _ Cannot be determined:
TUMOR
+Tumor Focality Unifocal Multifocal:
Tumor Site (select all that apply)  Glans: Foreskin mucosal surface: Foreskin skin surface: Coronal sulcus (balanopreputial sulcus): Skin of the shaft: Penile urethra: Penis, NOS:
+Tumor Macroscopic Features (select all that apply)  Flat Ulcerated Polypoid Verruciform

Necrosis	
Hemorrhage	
Other (specify):	
Tumor Size	
	cm
Greatest dimension in Centimeters (cm):  +Additional Dimension in Centimeters (cm):	CIII
Cannot be determined (explain):	
Histologic Type (Note B)	
Non-HPV-related squamous cell carcinoma	
Squamous cell carcinoma, usual type	
Pseudohyperplastic carcinoma	
Pseudoglandular carcinoma	
Verrucous carcinoma	
Carcinoma cuniculatum	
Papillary squamous cell carcinoma, NOS	
Adenosquamous carcinoma	
Sarcomatoid squamous cell carcinoma	
HPV-related squamous cell carcinoma	
Basaloid squamous cell carcinoma	
Papillary-basaloid squamous cell carcinoma	
Warty carcinoma	
Warty-basaloid squamous cell carcinoma	
Clear cell squamous cell carcinoma	
Lymphoepithelioma-like carcinoma	
Other Histologic Type	
Paget disease	
Adnexal carcinoma (specify type):	
Carcinoma, type cannot be determined:	
Other histologic type not listed (specify):	
+Histologic Type Comment:	
Histologic Grade (Note C)	
G1, well-differentiated	
G2, moderately differentiated	
G3, poorly differentiated	
Other (and if i)	
GX, cannot be assessed:	
Not applicable:	
Not applicable.	
+Tumor Thickness / Donth of Invesion (Note D)	
+Tumor Thickness / Depth of Invasion (Note D)	
Specify in Millimeters (mm): m	m
Other (specify):	
Cannot be determined:	
+Tumor Deep Borders (Note <u>E</u> ) (select all that apply)	
Pushing (broadly based)	
Infiltrative (jagged)	
Other (specify):	

Tumor Extent (select all that apply)	
Carcinoma in situ	
Noninvasive localized squamous cell ca	arcinoma
Invades lamina propria	
Invades dermis	
Invades dartos fascia	
Invades cornus spondiosum	
Invades corpus cavernosum	
Invades tunica albuginea	
Invades Buck's fascia	
Invades penile (distal) urethra	tum, prostate, pubic bone) (specify):
Invades regional skin (pubis, inguinal)	
Invades adjacent structure(s) (i.e., scro	tum, prostate, pubic bone) (specify):
Invades other structure(s) (specify):	
Cannot be determined (explain):	<del></del>
No evidence of primary tumor	
Lymphovascular Invasion (Note F)	
Not identified	
Present	
Cannot be determined:	
Perineural Invasion (Note G)	
Not identified	
Present	
Cannot be determined:	
4Tumor Comments	
+Tumor Comment:	
MARGINS (Note <u>H</u> )	
Margin Status for Invasive Carcinoma	
All margins negative for invasive carcin	
+Closest Margin(s) to Invasive Carcino	oma (Select all that apply)
Urethral:	prostive tiesus [leveine proprie] compre energiasure. Duelde
	nnective tissue [lamina propria], corpus spongiosum, Buck's
fascia):	
Corpus cavernosum:	<del></del>
Buck's fascia at penile shaft:	
Skin: * For circumcision specimens only	
Coronal sulcus mucosal#:	
Cutaneous#:	<del></del>
Other (specify):	
Cannot be determined (explain):	_
Califict be determined (explain)	
+Distance from Invasive Carcinoma to	Closest Margin
Specify in Millimeters (mm)	
Exact distance:	
Greater than:	nm

At least:	mm		
Less than:			
Less than 1 mm			
Other (specify):			
Cannot be determined	·	_	
Invasive carcinoma prese	_		
Margin(s) Involved by Inv Urethral:	asive Carcinoma (se	elect all that apply)	
	ubepithelial connective	e tissue [lamina propria], corpus spongiosum, Buc	k's
fascia) :			
Corpus cavernosum: _			
Buck's fascia at penile	shaft:	<del></del>	
Skin: # For circumcision specimens only			
Coronal sulcus mucos			
Other (specify):			
Cannot be determined	(explain):	<del></del>	
Other (specify):			
Cannot be determined (e	xplain):		
Not applicable	' /		
Margin Status for Noninvas	ive Carcinoma / Car	cinoma in Situ	
All margins negative for r	on-invasive carcinom	na / carcinoma in situ	
Noninvasive carcinoma /	carcinoma in situ pre	sent at margin	
Margin(s) Involved by No			
Specify involved marg	in(s):		
Cannot be determined	(explain):	<del> </del>	
Other (specify):			
Cannot be determined (e	xplain):		
Not applicable			
+Margin Comment:			
+wargin comment.	<del></del>		
REGIONAL LYMPH NODES	(Note <u>I</u> )		
Danis and Lamant Nada Otata			
Regional Lymph Node Statu		Standard Community	
Not applicable (no region		itted or found)	
Regional lymph nodes pr			
All regional lymph nod			
Tumor present in regio			
Number of Lymph Noc			
	cify):	<del></del>	
At least (specify): _			
Other (specify):	and (avalois):		

Nodal Site(s) with Tumor (select all that apply)
Sentinel:
Inguinal:
Number of Inguinal Lymph Nodes with Tumor
Exact number (specify):
At least (specify):
Other (specify):
Cannot be determined (explain):
Laterality of Inguinal Lymph Node(s) with Tumor
Unilateral
Bilateral
Cannot be determined (explain):
Pelvic:
Pelvic: Other (specify):
Cannot be determined
+Size of Largest Nodal Metastatic Deposit Specify in Centimeters (cm)
Exact size: cm
At least:cm
Greater than: cm
Less than: cm
Other (specify):
Other (specify): Cannot be determined:
+Nodal Site with Largest Metastatic Deposit (specify site):
+Size of Largest Lymph Node with Tumor
Specify in Centimeters (cm)
Exact size: cm
At least: cm
Greater than: cm
Less than: cm
Other (specify):
Cannot be determined
+Largest Lymph Node with Tumor (specify site):
Extranodal Extension
Not identified
Present
Cannot be determined:
<u> </u>
Other (specify):
_ Other (specify): _ Cannot be determined (explain):

From the market of the second	
Exact number (specify):	
At least (specify):	
Other (specify):	
Cannot be determined (explain):	
+Regional Lymph Node Comment:	
DISTANT METASTASIS	
Distant Site(s) Involved, if applicable (select all that apply)	
Not applicable	
Site(s) outside the true pelvis:	
Lung:	
Liver:	
Cutaneous nodules distant from the primary site:	
Bone:	
Other (specify):	
Cannot be determined:	
based upon all pertinent information, including but potentially not limited to this pathology report.  TNM Descriptors (select all that apply)	
Not applicable:	
Not applicable:	
m (multiple primary tumors)	
m (multiple primary tumors) r (recurrent)	
m (multiple primary tumors) r (recurrent) y (post-treatment)	
Not applicable: m (multiple primary tumors) r (recurrent) y (post-treatment)  pT Category  pT not assigned (cannot be determined based on available pathological information)	
m (multiple primary tumors) r (recurrent) y (post-treatment)  pT Category pT not assigned (cannot be determined based on available pathological information)	
m (multiple primary tumors) r (recurrent) y (post-treatment)  pT Category pT not assigned (cannot be determined based on available pathological information) pT0: No evidence of primary tumor	
m (multiple primary tumors) r (recurrent) y (post-treatment)  pT Category pT not assigned (cannot be determined based on available pathological information) pT0: No evidence of primary tumor pTis: Carcinoma *in situ* (Penile intraepithelial neoplasia [PelN])	
m (multiple primary tumors) r (recurrent) y (post-treatment)  pT Category pT not assigned (cannot be determined based on available pathological information) pT0: No evidence of primary tumor	r
m (multiple primary tumors) r (recurrent) y (post-treatment)  pT Category pT not assigned (cannot be determined based on available pathological information) pT0: No evidence of primary tumor pTis: Carcinoma *in situ* (Penile intraepithelial neoplasia [PeIN]) pTa: Noninvasive localized squamous cell carcinoma pT1: (Glans) Tumor invades lamina propria; (Foreskin) Tumor invades dermis, lamina propria, or dartos fascia; (Shaft) Tumor invades connective tissue between epidermis and corpora regardless of location; All sites with or without lymphovascular inv	
m (multiple primary tumors) r (recurrent) y (post-treatment)  pT Category pT not assigned (cannot be determined based on available pathological information) pT0: No evidence of primary tumor pTis: Carcinoma *in situ* (Penile intraepithelial neoplasia [PeIN]) pTa: Noninvasive localized squamous cell carcinoma pT1: (Glans) Tumor invades lamina propria; (Foreskin) Tumor invades dermis, lamina propria, or dartos fascia; (Shaft) Tumo invades connective tissue between epidermis and corpora regardless of location; All sites with or without lymphovascular inv perineural invasion and is or is not high grade	asion o
m (multiple primary tumors) r (recurrent) y (post-treatment)  pT Category pT not assigned (cannot be determined based on available pathological information) pT0: No evidence of primary tumor pTis: Carcinoma *in situ* (Penile intraepithelial neoplasia [PelN]) pTa: Noninvasive localized squamous cell carcinoma pT1: (Glans) Tumor invades lamina propria; (Foreskin) Tumor invades dermis, lamina propria, or dartos fascia; (Shaft) Tumo invades connective tissue between epidermis and corpora regardless of location; All sites with or without lymphovascular invaperineural invasion and is or is not high grade pT1a: Tumor is without lymphovascular invasion or perineural invasion and is not high grade (i.	asion o
m (multiple primary tumors) r (recurrent) y (post-treatment)  pT Category pT not assigned (cannot be determined based on available pathological information) pT0: No evidence of primary tumor pTis: Carcinoma *in situ* (Penile intraepithelial neoplasia [PelN]) pTa: Noninvasive localized squamous cell carcinoma pT1: (Glans) Tumor invades lamina propria; (Foreskin) Tumor invades dermis, lamina propria, or dartos fascia; (Shaft) Tumo invades connective tissue between epidermis and corpora regardless of location; All sites with or without lymphovascular inv perineural invasion and is or is not high grade pT1a: Tumor is without lymphovascular invasion or perineural invasion and is not high grade (i. grade 3 or sarcomatoid)	e.,
m (multiple primary tumors) r (recurrent) y (post-treatment)  PT Category pT not assigned (cannot be determined based on available pathological information) pT0: No evidence of primary tumor pTis: Carcinoma *in situ* (Penile intraepithelial neoplasia [PelN]) pTa: Noninvasive localized squamous cell carcinoma pT1: (Glans) Tumor invades lamina propria; (Foreskin) Tumor invades dermis, lamina propria, or dartos fascia; (Shaft) Tumo invades connective tissue between epidermis and corpora regardless of location; All sites with or without lymphovascular inv perineural invasion and is or is not high grade pT1a: Tumor is without lymphovascular invasion or perineural invasion and is not high grade (i.e. pT1b: Tumor exhibits lymphovascular invasion and / or perineural invasion or is high grade (i.e.	e.,
m (multiple primary tumors) r (recurrent) y (post-treatment)  pT Category pT not assigned (cannot be determined based on available pathological information) pT0: No evidence of primary tumor pTis: Carcinoma *in situ* (Penile intraepithelial neoplasia [PelN]) pTa: Noninvasive localized squamous cell carcinoma pT1: (Glans) Tumor invades lamina propria; (Foreskin) Tumor invades dermis, lamina propria, or dartos fascia; (Shaft) Tumor invades connective tissue between epidermis and corpora regardless of location; All sites with or without lymphovascular invaerineural invasion and is or is not high grade pT1a: Tumor is without lymphovascular invasion or perineural invasion and is not high grade (i. grade 3 or sarcomatoid) pT1b: Tumor exhibits lymphovascular invasion and / or perineural invasion or is high grade (i.e. grade 3 or sarcomatoid)	e.,
m (multiple primary tumors) r (recurrent) y (post-treatment)  pT Category pT not assigned (cannot be determined based on available pathological information) pT0: No evidence of primary tumor pTis: Carcinoma *in situ* (Penile intraepithelial neoplasia [PelN]) pTa: Noninvasive localized squamous cell carcinoma pT1: (Glans) Tumor invades lamina propria; (Foreskin) Tumor invades dermis, lamina propria, or dartos fascia; (Shaft) Tumo invades connective tissue between epidermis and corpora regardless of location; All sites with or without lymphovascular inv perineural invasion and is or is not high grade pT1a: Tumor is without lymphovascular invasion or perineural invasion and is not high grade (i. grade 3 or sarcomatoid) pT1b: Tumor exhibits lymphovascular invasion and / or perineural invasion or is high grade (i.e. grade 3 or sarcomatoid) pT1 (subcategory cannot be determined)	e.,
m (multiple primary tumors) r (recurrent) y (post-treatment)  pT Category pT not assigned (cannot be determined based on available pathological information) pT0: No evidence of primary tumor pTis: Carcinoma *in situ* (Penile intraepithelial neoplasia [PelN]) pTa: Noninvasive localized squamous cell carcinoma pT1: (Glans) Tumor invades lamina propria; (Foreskin) Tumor invades dermis, lamina propria, or dartos fascia; (Shaft) Tumo invades connective tissue between epidermis and corpora regardless of location; All sites with or without lymphovascular inv perineural invasion and is or is not high grade pT1a: Tumor is without lymphovascular invasion or perineural invasion and is not high grade (i. grade 3 or sarcomatoid) pT1b: Tumor exhibits lymphovascular invasion and / or perineural invasion or is high grade (i.e. grade 3 or sarcomatoid) pT1 (subcategory cannot be determined) pT2: Tumor invades into corpus spongiosum (either glans or ventral shaft) with or without ureth	e.,
m (multiple primary tumors) r (recurrent) y (post-treatment)  pT Category pT not assigned (cannot be determined based on available pathological information) pT0: No evidence of primary tumor pTis: Carcinoma *in situ* (Penile intraepithelial neoplasia [PelN]) pTa: Noninvasive localized squamous cell carcinoma pT1: (Glans) Tumor invades lamina propria; (Foreskin) Tumor invades dermis, lamina propria, or dartos fascia; (Shaft) Tumor invades connective tissue between epidermis and corpora regardless of location; All sites with or without lymphovascular invarineural invasion and is or is not high grade pT1a: Tumor is without lymphovascular invasion or perineural invasion and is not high grade (i. grade 3 or sarcomatoid) pT1b: Tumor exhibits lymphovascular invasion and / or perineural invasion or is high grade (i.e. grade 3 or sarcomatoid) pT1 (subcategory cannot be determined) pT2: Tumor invades into corpus spongiosum (either glans or ventral shaft) with or without ureth invasion	e., , ,
m (multiple primary tumors) r (recurrent) y (post-treatment)  pT Category pT not assigned (cannot be determined based on available pathological information) pT0: No evidence of primary tumor pTis: Carcinoma *in situ* (Penile intraepithelial neoplasia [PelN]) pTa: Noninvasive localized squamous cell carcinoma pT1: (Glans) Tumor invades lamina propria; (Foreskin) Tumor invades dermis, lamina propria, or dartos fascia; (Shaft) Tumo invades connective tissue between epidermis and corpora regardless of location; All sites with or without lymphovascular inv perineural invasion and is or is not high grade pT1a: Tumor is without lymphovascular invasion or perineural invasion and is not high grade (i. grade 3 or sarcomatoid) pT1b: Tumor exhibits lymphovascular invasion and / or perineural invasion or is high grade (i.e. grade 3 or sarcomatoid) pT1 (subcategory cannot be determined) pT2: Tumor invades into corpus spongiosum (either glans or ventral shaft) with or without ureth	e., , ,

pN Category  pN not assigned (no nodes submitted or found)  pN not assigned (cannot be determined based on available pathological information)  pN0: No lymph node metastasis  pN1: less than or equal to 2 unilateral inguinal metastases, no extranodal extension  pN2: greater than or equal to 3 unilateral inguinal metastases or bilateral metastases, no ENE  pN3: Extranodal extension of lymph node metastases or pelvic lymph node metastases
pM Category (required only if confirmed pathologically)  Not applicable - pM cannot be determined from the submitted specimen(s)  **Including lymph node metastasis outside the true pelvis, lung, liver, cutaneous nodules distant from the primary site, and bone.  pM1: Distant metastasis present**
ADDITIONAL FINDINGS (Note K)
+Additional Findings (select all that apply)  None identified HPV-related penile intraepithelial neoplasia (PelN), warty type HPV-related penile intraepithelial neoplasia (PelN), basaloid type HPV-related penile intraepithelial neoplasia (PelN), warty-basaloid type Non-HPV-related PelN (differentiated [simplex] penile intraepithelial neoplasia) Pleomorphic PelN Spindle PelN Clear cell PelN Pagetoid PelN Lichen sclerosus Squamous hyperplasia Condyloma acuminatum Other (specify):
SPECIAL STUDIES
+Ancillary Studies Specify: Not performed
COMMENTS
Comment(s):

## **Explanatory Notes**

#### A. Types of Foreskin

There are three foreskin types: in the short foreskin, the preputial orifice is located behind the glans corona; in the medium foreskin, the orifice is between the corona and the meatal orifice; in the long foreskin, the entire glans is covered and the meatus is not identified without retracting the foreskin. Phimotic foreskins are unretractable and long. Phimosis is present in up to one-half of patients with penile carcinoma, and its presence is considered a risk factor for the development of this tumor.

#### References

- Velazquez EF, Bock A, Soskin A, Codas R, Arbo M, Cubilla AL. Preputial variability and preferential association of long phimotic foreskins with penile cancer: an anatomic comparative study of types of foreskin in a general population and cancer patients. Am J Surg Pathol. 2003;27(7):994-998.
- 2. Daling J, Madeleine MM, Johnson LG, et al. Penile cancer: importance of circumcision, human papillomavirus and smoking in in situ and invasive disease. Int J Cancer. 2005;116(4):606-616.
- 3. Tsen HF, Morgenstern H, Mack T, Peters RK. Risk factors for penile cancer: results of a population-based case-control study in Los Angeles County (United States). Cancer Causes Control. 2002;12(3):267-277.
- 4. Madsen BS, van den Brule AJ, Jensen HL, Wholfahrt J, Frisch M. Risk factors for squamous cell carcinoma of the penis: population-based case-control study in Denmark. Cancer Epidemiol Biomarkers Prev. 2008;17(10):2683-2691.

#### B. Histologic Subtype of Squamous Cell Carcinoma

The World Health Organization (WHO) classification of tumors of the penis was recently published.¹ Most penile cancers are squamous cell carcinomas (SCC), and most arise from the epithelium of the distal portion of the penis (including glans, coronal sulcus, and mucosal surface of the prepuce). Squamous cell carcinoma of the usual type (keratinizing SCC) comprises about 50% to 60% of all cases.² There are other SCC variants showing distinctive morphological and outcome features.³ 4.5 The different histological subtypes correlate with different rates of regional/nodal and systemic dissemination. Penile cancer subtypes can be prognostically stratified in three groups. The low-risk group includes verruciform tumors such as verrucous, papillary, and warty/condylomatous carcinomas. More recently described subtypes, such as pseudohyperplastic and carcinoma cuniculatum of the penis, also belong to this category of excellent prognosis. The high-risk category is comprised by basaloid, sarcomatoid, adenosquamous, and poorly differentiated SCC of the usual type. 9.10.11 There is an intermediate category of metastatic risk that includes most SCCs of the usual type, some mixed neoplasms (such as hybrid verrucous carcinomas), and high-grade variants of warty/condylomatous carcinomas.6

- 1. Moch H, Humphrey PA, Ulbright TM, Reuter VE. WHO Classification of Tumours of the Urinary System and Male Genital Organs. Geneva, Switzerland: WHO Press; 2016.
- 2. Epstein JH, Humphrey PA, Cubilla AL. Tumors of the Prostate Gland, Seminal Vesicles, Male Urethra, Penis and Scrotum. Washington, DC: Armed Forces Institute of Pathology; 2011 Atlas of Tumor Pathology.
- 3. Cubilla AL, Dillner J, Schellhammer PF, Horenblas S. Malignant epithelial tumors. In: Eble JN, Sauter G, Epstein J, Sesterhenn I, eds. Pathology and Genetics of Tumors of the Urinary System and Male Genital Organs. Lyon, France: IARC Press; 2004. World Health Organization Classification of Tumours.
- Velazquez EF, Barreto JE, Ayala G, Cubilla AL. Penis. In: Mills SE, Carter D, Greenson JK, et al, eds. Sternberg's Diagnostic Surgical Pathology. 5th ed. Philadelphia, PA: Lippincot Williams & Wilkins; 2009.

- 5. Cubilla AL, Reuter V, Velazquez E, Piris A, Saito S, Young RH. Histologic classification of penile carcinoma and its relation to outcome in 61 patients with primary resection. Int J Surg Pathol. 2001;9(2):111-120.
- 6. Cubilla AL, Velazques EF, Reuter VE, Oliva E, Mihm MC Jr, Young RH. Warty (condylomatous) squamous cell carcinoma of the penis: a report of 11 cases and proposed classification of verruciform tumors penile tumors. Am J Surg Pathol. 2000;24(4):505-512.
- 7. Cubilla AL, Velazquez EF, Young RH. Pseudohyperplastic squamous cell carcinoma of the penis associated with lichen sclerosus an extremely well-differentiated nonverruciform neoplasm that preferentially affects the foreskin and is frequently misdiagnosed: a report of 10 cases of a distinctive clinicopathologic entity. Am J Surg Pathol. 2004;28(7):895-900.
- 8. Barreto JE, Velazquez EF, Ayala E, Torres J, Cubilla AL. Carcinoma cuniculatum of the penis a distinctive variant of penile squamous cell carcinoma: report of 7 cases. Am J Surg Pathol. 2007;31(1):71-75.
- 9. Cubilla AL, Reuter VE, Gregoire L, et al. Basaloid squamous cell carcinoma: a distinctive HPV related penile neoplasm: a report of 20 cases. Am J Surg Pathol. 1998;22(6):751-761.
- 10. Velazquez EF, Melamed J, Barreto JE, Aguero F, Cubilla AL. Sarcomatoid carcinoma of the penis: a clinico-pathological study of 14 cases. Am J Surg Pathol. 2005;29(9):1152-1158.
- 11. Cubilla AL, Ayala MT, Barreto JE, Bellasai JG, Nöel JC. Surface adenosquamous carcinoma of the penis: a report of three cases. Am J Surg Pathol. 1996;20(2):156-160.

#### C. Histologic Grade

Histological grade has been consistently reported as an influential predictive factor of groin metastasis and dissemination of penile cancer. 1.2.3 We recommend a method to grade penile SCCs as follows:

- Grade 1 is an extremely well-differentiated carcinoma, with a minimal deviation from the morphology of normal/hyperplastic squamous epithelium.
- Grade 2 tumors show a more disorganized growth as compared to grade 1 lesions, higher nuclear-to-cytoplasmic ratio, evident mitoses, and, although present, less prominent keratinization.
- Grade 3 are tumors showing any proportion of anaplastic cells, identified as solid sheets or irregular small aggregates, cords or nests of cells with little or no keratinization, high nuclear-tocytoplasmic ratio, thick nuclear membranes, nuclear pleomorphism, clumped chromatin, prominent nucleoli, and numerous mitosis.

A tumor should be graded according to the least differentiated component. Any proportion of grade 3 should be noted in the report. $^4$ 

- 1. Slaton JW, Morgenstern N, Levy DA, et al. Tumor stage, vascular invasion and the percentage of poorly differentiated cancer: independent prognosticators for inguinal node metastasis in penile squamous cancer. J Urol. 2001;165(4):1138-1142.
- 2. Cubilla AL, Velazquez EF, Ayala GE, Chaux A, Torres J, Reuter V. Identification of prognostic pathologic parameters in squamous cell carcinoma of the penis: significance and difficulties. Pathol Case Rev. 2005;10:3-13.
- 3. Velazquez EF, Ayala G, Liu H, et al. Histologic grade and perineural invasion are more important than tumor thickness as predictor of nodal metastasis in penile squamous cell carcinoma invading 5 to 10 mm. Am J Surg Pathol. 2008;32(7):974-979.
- 4. Chaux A, Torres J, Pfannl R, et al. Histologic grade in penile squamous cell carcinoma: visual estimation versus digital measurement of proportions of grades, adverse prognosis with any proportion of grade 3 and correlation of a Gleason-like system with nodal metastasis. Am J Surg Pathol. 2009;33:1042-1048.

#### D. Depth of Invasion

The tumor depth in small lesions is best obtained by perpendicularly sectioning along the tumor central axis. For large glans tumors, it is preferred to section the specimen longitudinally in half, with additional parallel sections of each half, using as an axis the central and ventral penile urethra. The depth of invasion of SCC is defined as a measurement in millimeters from the epithelial-stromal junction of the adjacent nonneoplastic epithelium to the deepest point of invasion. In larger tumors, especially verruciform ones, the previously mentioned system is not applicable, and we measure the thickness from the surface (excluding the keratin layer) to the deepest point of invasion. Depth of invasion and tumor thickness are of equivalent significance. There is a correlation between depth of invasion and outcome in penile cancers. Minimal risk for metastasis was reported for tumors measuring less than 5 mm in thickness. 1.2 Tumors invading deeper into penile anatomical levels are usually associated with a higher risk for nodal involvement. There is also a correlation between deeper infiltration and higher histological grade, although some exceptions do occur.<sup>3</sup> Tumors invading corpus cavernosum are at higher risk for presenting nodal metastases than those invading only corpus spongiosum, 3.4 and the deepest erectile tissue invaded should be clearly stated in the final pathology report. Per AJCC 8th edition, tumor invading into subepithelial connective tissue (lamina propria), Dartos muscle, and Buck's fascia is staged as T1; tumor invading into corpus spongiosum (either glans or ventral shaft) with or without urethral invasion is staged as T2; tumor invading into corpora cavernosum (including tunica albuginea) with or without urethral invasion is staged as T3; and tumor invading into adjacent structures (ie, scrotum, prostate, pubic bone) is staged as T4.

#### References

- 1. Velazquez EF, Ayala G, Liu H, et al. Histologic grade and perineural invasion are more important than tumor thickness as predictor of nodal metastasis in penile squamous cell carcinoma invading 5 to 10 mm. Am J Surg Pathol. 2008;32(7):974-979.
- 1. Emerson RE, Ulbright TM, Eble JN, Geary WA, Eckert GJ, Cheng L. Predicting cancer progression in patients with penile squamous cell carcinoma: the importance of depth of invasion and vascular invasion. Mod Pathol. 2001;14(10):963-968.
- 2. Leijte J, Gallee M, Antonini N, Horenblas S. Evaluation of current TNM classification of penile carcinoma. J Urol. 2008;180(3):933-938.
- Chaux A, Caballero C, Soares F, et al. The prognostic index: a useful pathologic guide for prediction of nodal metastases and survival in penile squamous cell carcinoma. Am J Surg Pathol. 2009;33(7):1049-1057.

#### E. Tumor Base of Infiltration

Two patterns are recognized: infiltrating (invasion in blocks of small solid strands of cell tumors broadly infiltrating the stroma) and pushing infiltration (tumor cells invading in large cell blocks with well-defined tumor-stroma interface). The infiltrating pattern of invasion is associated with a higher risk for nodal involvement.<sup>1</sup>

#### References

1. Guimarães G, Lopes A, Campos RS, et al. Front pattern of invasion in squamous cell carcinoma of the penis: new prognostic factor for predicting risk of lymph node metastases. Urology. 2006;68(1):148-153.

## F. Lymphovascular Invasion

Vascular invasion, lymphatic or venous, adversely affects prognosis of penile cancer. 1.2.3.4.5 The TNM staging classification in the 8th edition of the AJCC Cancer Staging Manual subdivides T1 tumors into T1a and T1b based on the absence or presence of lymphovascular invasion or poorly differentiated tumors. Embolic involvement of lymphatic vascular spaces occurs usually near the invasive tumor front, but it may also be found at a certain distance from the primary tumor in anatomical areas such as the

lamina propria, penile fascia, and especially in the subepithelial connective tissues surrounding penile urethra. Venous invasion indicates a more advanced stage of the disease and is related to the compromise of the specialized erectile venous structures of corpora spongiosa and cavernosa.

#### References

- 1. Lopes A, Hidalgo GS, Kowalski LP, Torloni H, Rossi BM, Fonseca FP. Prognostic factors in carcinoma of the penis: multivariate analysis of 145 patients treated with amputation and lymphadenectomy. J Urol. 1996;156(5):1637-1642.
- 2. Ficarra V, Zattoni F, Cunisco SC, et al. Lymphatic and vascular embolizations are independent predictive variables of inguinal node involvement in patients with squamous cell carcinoma of the penis: Gruppo Uro-Oncologico del Nord Est (Northeast Uro-Oncological Group) Penile Cancer data base data. Cancer. 2005;103(12):2507-2516.
- 3. Ficarra V, Zattoni F, Artibani W, et al; GUONE Penile Cancer Project Members. Nomogram predictive of pathological inguinal lymph node involvement in patients with squamous cell carcinoma of the penis. J Urol. 2006;175(6):1700-1705.
- 4. Kattan MW, Ficarra V, Artibani W, et al; GUONE Penile Cancer Project Members. Nomogram predictive of cancer specific survival in patients undergoing partial or total amputation for squamous cell carcinoma of the penis. J Urol. 2006;175(6):2103-2108.
- 5. Novara G, Galfano A, De Marco V, Artibani W, Ficarra V. Prognostic factors in squamous cell carcinoma of the penis. Nat Clin Pract Urol. 2007;4(3):140-146.
- 6. Amin MB, Edge SB, Greene FL, et al, eds. AJCC Cancer Staging Manual. 8th ed. New York, NY: Springer; 2017.

#### G. Nomograms, Risk Groups, and Perineural Invasion

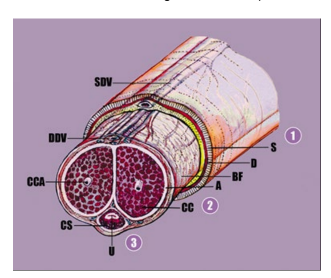
An evaluation of clinical and pathological variables using a nomogram was recently developed.¹ The selected factors were clinical stage of lymph nodes, microscopic growth pattern, grade, vascular invasion, and invasion of corpora spongiosa and cavernosa and urethra. The probability of nodal metastasis as predicted by the nomogram was close to the real incidence of metastasis observed at follow up. A second nomogram to estimate predictions of survival at 5 years with the same clinical and pathological factors gave similar results.² More recently, perineural invasion and histological grade were found to be the strongest independent predictors of mortality in penile tumors 5 to 10 mm thick. A nomogram considering the predictive value of perineural invasion and histological grade was accordingly constructed.³ Risk groups stratification systems are available to predict the likelihood of inguinal nodal involvement and for therapeutic planning and are based on a combination of histological grade and pT stage.⁴.5.6.7 Strongest predictive power results from the combination of histological grade, deepest anatomical level of infiltration, and presence of perineural invasion. These factors are used for constructing the prognostic index.<sup>8</sup>

- 1. Ficarra V, Zattoni F, Artibani W, et al; GUONE Penile Cancer Project Members. Nomogram predictive of pathological inguinal lymph node involvement in patients with squamous cell carcinoma of the penis. J Urol. 2006;175(6):1700-1705.
- 2. Kattan MW, Ficarra V, Artibani W, et al; GUONE Penile Cancer Project Members. Nomogram predictive of cancer specific survival in patients undergoing partial or total amputation for squamous cell carcinoma of the penis. J Urol. 2006;175(6):2103-2108.
- 3. Cubilla AL, Velazquez EF, Ayala GE, Chaux A, Torres J, Reuter V. Identification of prognostic pathologic parameters in squamous cell carcinoma of the penis: significance and difficulties. Pathol Case Rev. 2005;10:3-13.
- 4. Solsona E, Iborra I, Rubio J, Casanova JL, Ricos JV, Calabuig C. Prospective validation of the association of local tumor stage and grade as a predictive factor for occult lymph node micrometastasis in patients with penile carcinoma and clinically negative inguinal lymph nodes. J Urol. 2001;165(5):1506-1509.

- 5. Solsona E, Algaba F, Horenblas S, Pizzocaro G, Windahl T; European Association of Urology. EAU guidelines on penile cancer. Eur Urol. 2004;46(1):1-8.
- 6. Hungerhuber E, Schlenker B, et al. Risk stratification in penile carcinoma: 25-year experience with surgical inguinal lymph node staging. Urology. 2006;68(3):621-625.
- 7. Ornellas AA, Nóbrega BL, Wei Kin Chin E, et al. Prognostic factors in invasive squamous cell carcinoma of the penis: analysis of 196 patients treated at the Brazilian National Cancer Institute. J Urol. 2008;180(4):1354-1359.
- 8. Chaux A, Caballero C, Soares F, et al. The prognostic index: a useful pathologic guide for prediction of nodal metastases and survival in penile squamous cell carcinoma. Am J Surg Pathol. 2009;33(7):1049-1057.

#### H. Resection Margins

Positive margins adversely affect prognosis in patients with penile squamous cell carcinomas. 1.2.3 Important margins to be examined in partial penectomy specimens include: (1) proximal urethra and surrounding periurethral cylinder consisting of epithelium, subepithelial connective tissue (lamina propria), corpus spongiosum, and penile fascia; (2) proximal shaft with corresponding corpora cavernosa separated and surrounded by the tunica albuginea and Buck's fascia; and (3) skin of shaft with underlying corporal dartos 4 (Figure 1). The coronal sulcus mucosal margin and cutaneous margin should be entirely examined when evaluating circumcision specimens.



**Figure 1.** Partial penectomy specimen; anatomical structures of proximal resection margin. The ventral urethra (U) is surrounded by the corpus spongiosum (CS) and a delicate white tunica albuginea (A). The latter is also surrounding the corpora cavernosa (CC). The penile fascia (Buck's fascia) (BF) is located underneath skin (S) and dartos (D). The proximal margin of resection should be cut en face and all the structures including the entire circumference of the urethra with periurethral cylinder should be examined. The 3 important margins to be examined include (1) skin of the shaft with underlying dartos and penile fascia, (2) the corpora cavernosa with surrounding tunica albuginea, and (3) the urethra and periurethral cylinder that includes the lamina propria, corpus spongiosum, albuginea, and penile fascia.

Abbreviations: CCA, cavernous artery; DDV, deep dorsal vein; SDV, superficial dorsal vein.

#### References

 Epstein JH, Humphrey PA, Cubilla AL. Tumors of the Prostate Gland, Seminal Vesicles, Male Urethra, Penis and Scrotum. Washington, DC: Armed Forces Institute of Pathology; 2011 Atlas of Tumor Pathology.

- Velazquez EF, Barreto JE, Ayala G, Cubilla AL. Penis. In: Mills SE, Carter D, Greenson JK, et al, eds. Sternberg's Diagnostic Surgical Pathology. 5th ed. Philadelphia, PA: Lippincot Williams & Wilkins; 2009.
- 3. Velazquez EF, Soskin A, Bock A, Codas R, Barreto JE, Cubilla AL. Positive resection margins in partial penectomies: sites of involvement and proposal of local routes of spread of penile squamous cell carcinoma. Am J Surg Pathol. 2004;28(3):384-389.
- Chaux A, Caballero C, Soares F, et al. The prognostic index: a useful pathologic guide for prediction of nodal metastases and survival in penile squamous cell carcinoma. Am J Surg Pathol. 2009;33(7):1049-1057.

#### I. Number of Involved Lymph Nodes and Extension of the Lymphadenectomy

The presence of more than two positive lymph nodes in one inguinal basin increases the likelihood of contralateral inguinal and ipsilateral pelvic nodal involvement. In such cases, prophylactic contralateral inguinal and ipsilateral pelvic lymphadenectomy is advised. The number and percentage of positive nodes involved also has an impact on survival. 2.3

#### References

- 1. Lont AP, Kroon BK, Gallee MP, van Tinteren H, Moonen LM, Horenblas S. Pelvic lymph node dissection for penile carcinoma: extent of inguinal lymph node involvement as an indicator for pelvic lymph node involvement and survival. J Urol. 2007;177(3):947-952.
- 2. Svatek RS, Munsell M, Kincaid JM, et al. Association between lymph node density and disease specific survival in patients with penile cancer. J Urol. In press.
- 3. Pandey D, Mahajan V, Kannan RR. Prognostic factors in node-positive carcinoma of the penis. J Surg Oncol. 2006;93(2):133-138.

#### J. TNM Staging Classification

The protocol recommends the use of the TNM staging system of the American Joint Committee on Cancer (AJCC) for carcinoma of the penis. By AJCC convention, the designation T refers to a primary tumor that has not been previously treated. The symbol p refers to the pathologic classification of the TNM, as opposed to the clinical classification, and is based on gross and microscopic examination. pT entails a resection of the primary tumor or a biopsy adequate to evaluate the highest pT category, pN entails removal of nodes adequate to validate lymph node metastasis, and pM implies microscopic examination of distant lesion. Pathologic staging is usually performed after surgical resection of the primary tumor. The summary of changes in the TNM staging classification in the 8th edition of the AJCC Cancer Staging Manual is as follows:

Change	Details of Change
Histologic Grade (G)	The 3-tiered World Health Organization (WHO)/International Society of Urological Pathology (ISUP) grading system has been adopted. Any proportion of anaplastic cells is sufficient to categorize a tumor as grade 3.
Definition of Primary Tumor (T)	Ta definition is now broadened to include noninvasive localized squamous carcinoma.
Definition of Primary Tumor (T)	T1a and T1b have been separated by an additional prognostic indicator—the presence or absence or perineural invasion.
Definition of Primary Tumor (T)	T1a or T1b are described by the site where they occur on the penis and are designated glans, foreskin, or shaft. Anatomic layers invaded are described for the three locations.
Definition of Primary Tumor (T)	T2 definition includes corpus spongiosum invasion.
Definition of Primary Tumor (T)	T3 definition now involves corpora cavernosum invasion.

Change	Details of Change
Definition of Regional Lymph	pN1 is defined as ≤2 unilateral inguinal metastases, no extranodal extension.
Nodes (N)	
Definition of Regional Lymph	pN2 is defined as ≥3 unilateral inguinal metastases or bilateral metastases
Nodes (N)	

## **Additional Descriptor**

<u>The m suffix</u> indicates the presence of multiple primary tumors and is recorded in parentheses, eg, pTa(m)N0M0.

## **Anatomic Stage/Prognostic Groups**

Group	T	N	M
Stage 0is	Tis	N0	M0
Stage 0a	Та	N0	M0
Stage I	T1a	N0	M0
Stage IIA	T1b	N0	M0
Stage IIA	T2	N0	M0
Stage IIB	T3	N0	M0
Stage IIIA	T1-3	N1	M0
Stage IIIB	T1-3	N2	M0
Stage IV	T4	Any N	M0
Stage IV	Any T	N3	M0
Stage IV	Any T	Any N	M1

## **Prognostic Factors (Site-Specific Factors)**

Factors required for staging: None.

Clinically significant factors:

- Involvement of corpus spongiosum
- Involvement of corpus cavernosum
- Percentage of tumor that is poorly differentiated
- Verrucous carcinoma depth of invasion
- Size of largest lymph node metastasis
- Extranodal/extracapsular extension
- Human papillomavirus (HPV) status

#### References

1. Amin MB, Edge SB, Greene FL, et al, eds. AJCC Cancer Staging Manual. 8th ed. New York, NY: Springer; 2017.

#### K. Penile Intraepithelial Neoplasia

Penile Intraepithelial Neoplasia (PeIN) may be subclassified as differentiated (simplex), warty, basaloid, and warty/basaloid (mixed). L2 Differentiated PeIN shows parakeratosis, epithelial thickening, elongation of rete ridges, prominent bridges, basal cell atypia, enlarged nuclei, and prominent nucleoli. Differentiated PeIN is frequently associated with lichen sclerosus. It is considered HPV-unrelated, there is no koilocytosis, and p16 immunohistochemical staining results (surrogate of high-risk types of HPV) are usually negative. Basaloid PeIN is characterized by a replacement of the normal epithelium by small, uniform cells with round nuclei and scant cytoplasm. Numerous mitosis and apoptotic cells are usually present. Warty PeIN shows a spiky surface with parakeratosis. The normal epithelium is replaced by

markedly pleomorphic cells showing prominent koilocytosis. Mixed warty-basaloid lesions are not infrequent. Warty and basaloid PeIN are HPV-related lesions and usually overexpress p16.

#### References

- 1. Cubilla AL, Pfannl R, Rodriguez I, et al. Morphological characterization and distribution of penile precancerous lesions using a simplified nomenclature: a study of 198 lesions in 115 patients. Lab Invest. 2008;88:696(A).
- 2. Pfannl R, Hernandez M, Velazquez EF, et al. Expression of p53 and p16 in differentiated and warty/basaloid penile intraepithelial neoplasia (PelN). Lab Invest. 2008;88:807(A).

## L. Handling of the Specimen

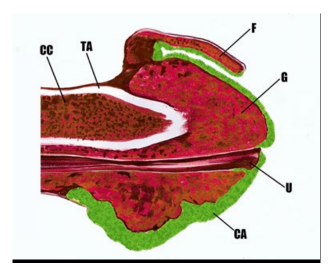
## **Circumcision Specimen**

Take measurements, describe specimen, and identify and describe tumor. Identify and ink the mucosal and cutaneous margins with different colors. Most SCCs arise from the mucosal surface of the foreskin, therefore the coronal sulcus (mucosal) margin is especially important. Lightly stretch and pin the specimen to a cardboard. Fix for several hours in formalin. Cut vertically the whole specimen labeling from 1 to 12, clockwise.

#### **Penectomy Specimen**

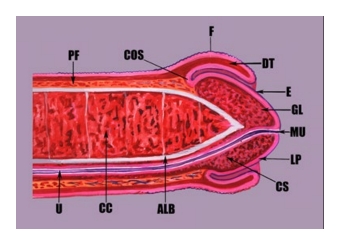
Take measurements, describe specimen, and identify and describe tumor. Most SCCs of the penis arise from the epithelium of the distal portion of the organ (glans, coronal sulcus, and mucosal surface of the prepuce; the tumor may involve one or more of these anatomical compartments).41 If present, classify the foreskin as short, medium, long, and/or phimotic.<sup>2</sup> Cut the proximal margin of resection en face making sure to include the entire circumference of the urethra (Figure 1). If the urethra has been retracted, it is important to identify its resection margin and submit it entirely. The resection margin can be divided in three important areas that need to be analyzed: the skin of the shaft with underlying dartos and penile fascia; corpora cavernosa with albuginea; and urethra with periurethral cylinder that includes subepithelial connective tissue (lamina propria), corpus spongiosum, albuginea, and penile fascia (Figure 1). The urethra and periurethral cylinder can be placed in one cassette. The skin of the shaft with dartos and fascia can be included together with the corpora cavernosa. Because this is a large specimen, it may need to be included in several cassettes to include the entire resection margin. Fix the rest of the specimen overnight. Then, in the fixed state and if the tumor is large and involves most of the glans, cut longitudinally and centrally by using the meatus and the proximal urethra as reference points. Do not probe the urethra. Separate the specimen into halves, left and right (Figures 2 and 3). Then cut two to six serial sections of each half. If tumor is small and asymmetrically located in the dorsal or ventral area, the central portion of the tumor may be used as the axis of sectioning. If the tumor is large involving multiples sites (glans, sulcus and foreskin), it is important not to remove the foreskin leaving the entire specimen intact for sectioning.

In cases of small carcinomas exclusively located in the glans with no foreskin involvement, one may choose to remove the foreskin leaving a 3-mm redundant edge around the sulcus. Proceed cutting the foreskin as indicated for circumcision specimens. If the primary tumor is located in the glans, one should still submit the foreskin serially and in orderly fashion labeled from 1 to 12 clockwise. The rest of the penectomy specimen should be handled as described above.



**Figure 2.** Partial penectomy specimen. After submitting the proximal resection margin, the specimen is cut in half longitudinally. Parallel serial sections will follow.

Abbreviations: CA, carcinoma; CC, corpus cavernosum; F, foreskin; G, glans; TA, tunica albuginea; U, urethra.



**Figure 3.** Longitudinal and central section showing the ventral urethra (U) and the penile main anatomic compartments: glans (GL), coronal sulcus (COS), and foreskin (F). The Buck's (penile) fascia (PF) encases the shaft and inserts into the coronal sulcus.

Abbreviations: ALB, albuginea; CC, corpus cavernosum; CS, corpus spongiosum; DT, dartos; E, epithelium; LP, lamina propria; MU, urethral meatus.

## M. Pathology Report for Penile Squamous Cell Carcinoma

The report should contain the following information: primary tumor: tumor site or sites, size in centimeters, histologic subtype, histologic grade, anatomical level of invasion, tumor thickness in millimeters, and vascular and perineural invasion. In penectomy specimens, the margins of resection to be reported are urethral/periurethral, corporal, and skin of the shaft. In circumcision specimens, margins include coronal sulcus mucosal margin and cutaneous margin. Common associated lesions to be reported are penile intraepithelial neoplasia (differentiated or undifferentiated), lichen sclerosus, and other "inflammatory dermatologic" conditions.

If the specimen is accompanied by inguinal nodes, the number and size of nodes should be described. All nodes should be included for microscopic examination. The number of positive nodes and total number of

nodes examined should be reported as well as the presence of extracapsular extension and the number and site (eg, inguinal versus pelvic) of metastatic nodes. The distinction between superficial and deep inguinal lymph nodes has been eliminated in the seventh edition TNM classification.<sup>2</sup>

- 1. Velazquez EF, Soskin A, Bock A, Codas R, Barreto JE, Cubilla AL. Positive resection margins in partial penectomies: sites of involvement and proposal of local routes of spread of penile squamous cell carcinoma. Am J Surg Pathol. 2004;28(3):384-389.
- 2. Amin MB, Edge SB, Greene FL, et al, eds. AJCC Cancer Staging Manual. 8th ed. New York, NY: Springer; 2017.