<table>
<thead>
<tr>
<th>Penile Cancer Panel Members</th>
<th>Details</th>
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<tbody>
<tr>
<td>Peter E. Clark, MD ٭</td>
<td>Chair</td>
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<tr>
<td>Vanderbilt-Ingram Cancer Center</td>
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<tr>
<td>Philippe E. Spiess, MD, MS ٭</td>
<td>Vice chair</td>
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<tr>
<td>Moffitt Cancer Center</td>
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<tr>
<td>Neeraj Agarwal, MD ‡  †</td>
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<tr>
<td>Huntsman Cancer Institute</td>
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<td>at the University of Utah</td>
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<td>Stephen A. Boorjian, MD ٭</td>
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<tr>
<td>Mayo Clinic Cancer Center</td>
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<tr>
<td>Mark K. Buuyounouski, MD, MS §</td>
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<tr>
<td>Stanford Cancer Institute</td>
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<tr>
<td>Jason A. Efstathiou, MD, DPhil §</td>
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<tr>
<td>Massachusetts General Hospital Cancer Center</td>
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<tr>
<td>Thomas W. Flaig, MD  †</td>
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<td>University of Colorado Cancer Center</td>
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<td>Terence Friedlander, MD  †</td>
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<td>UCSF Helen Diller Family</td>
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<td>Comprehensive Cancer Center</td>
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<td>Richard E. Greenberg, MD ٭</td>
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<td>Khurshid A. Guru, MD ٭</td>
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<td>Roswell Park Cancer Institute</td>
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<tr>
<td>Noah Hahn, MD  †</td>
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<tr>
<td>The Sidney Kimmel Comprehensive Cancer Center at Johns Hopkins</td>
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<td>Harry W. Herr, MD ٭</td>
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<td>Christopher Hoimes, MD  †</td>
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<td>Case Comprehensive Cancer Center</td>
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<td>University Hospitals Seidman Cancer Center</td>
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<td>Cleveland Clinic Taussig Cancer Institute</td>
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**Discussion**

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<tr>
<th>Authors</th>
<th>Institutions</th>
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<tr>
<td>Brant A. Inman, MD, MSc ٭</td>
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<tr>
<td>Duke Cancer Institute</td>
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<td>Fred &amp; Pamela Buffett Cancer Center</td>
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<td>Lance C. Pagliaro, MD  †</td>
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<td>Anthony Patterson, MD ٭</td>
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<td>Kamal S. Pohar, MD ٭</td>
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<tr>
<td>The Ohio State University Comprehensive Cancer Center - James Cancer Hospital and Solove Research Institute</td>
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<td>Michael P. Porter, MD, MS ٭</td>
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<td>Fred Hutchinson Cancer Research Center/Seattle Cancer Care Alliance</td>
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<td>Wade J. Sexton, MD ٭</td>
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<td>Arlene O. Siefker-Radtke, MD  †</td>
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<td>The University of Texas</td>
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<td>Geoffrey Wile, MD  φ</td>
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<td>NCCN</td>
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<td>Mary Dwyer, MS</td>
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<td>Courtney Smith, PhD</td>
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**Panel Disclosures**

- ٭ Urology
-  † Medical oncology
- ‡ Hematology/Hematology oncology
- § Radiotherapy/Radiation oncology
- φ Diagnostic radiology
- ≠ Pathology
- * Discussion writing committee member

**NCCN Guidelines Panel Disclosures**
NCCN Penile Cancer Panel Members
Summary of the Guidelines Updates

Primary Evaluation, Clinical Diagnosis (PN-1)

Primary Treatment Tis, Ta (PN-1)

Primary Treatment T1, T2 or Greater (PN-2)

Management of Non-Palpable Inguinal Lymph Nodes (PN-3)

Management of Palpable Non-Bulky Inguinal Lymph Nodes (PN-4)

Management of Palpable Bulky Inguinal Lymph Nodes (PN-5)

Management of Enlarged Pelvic Lymph Nodes (PN-6)

Surveillance Schedule (PN-7)

Management of Recurrent Disease (PN-8)

Management of Metastatic Disease (PN-9)

Principles of Surgery (PN-A)

Principles of Radiotherapy (PN-B)

Principles of Chemotherapy (PN-C)

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Summary of the changes in the 2.2016 version of the Guidelines for Penile Cancer from the 1.2016 version include:

**PN-1**
- Footnote “b” was added, “Moh’s surgery is an option.”

**PN-6**
- After treatment with neoadjuvant chemotherapy for positive lymph nodes, imaging was added “Abdominal/pelvic CT or MRI and Chest imaging (x-ray, or CT).”
- Footnotes
  - Footnote “o” was added, “Consider PET-CT scan.”
  - Footnote “p” was added, “If M1 disease identified, see Management of Metastatic Disease (PN-9).”

**PN-9**
- After treatment, imaging was added as, “Abdominal/pelvic CT or MRI and Chest imaging (x-ray, or CT).”
- Footnote “o” was added, “Consider PET-CT scan.”

**MS-1**
- The discussion section was updated to reflect the changes in the algorithm.

Note: All recommendations are category 2A unless otherwise indicated.
Clinical Trials: NCCN believes that the best management of any cancer patient is in a clinical trial. Participation in clinical trials is especially encouraged.
Summary of the changes in the 1.2016 version of the Guidelines for Penile Cancer from the 3.2015 version include:

**General**
- “Radiotherapy with concurrent chemotherapy” was replaced with “chemoradiotherapy” throughout.

**PN-2**
- T2 or greater, “T2 tumors only” qualifier was removed for the primary treatment options of radiotherapy and chemoradiotherapy.

**PN-3**
- Imaging was added after intermediate and high risk as, “Abdominal/pelvic CT or MRI and Chest imaging (x-ray, or CT).”
- Treatment for “Dynamic sentinel node biopsy”, the category was changed from a category 2B to a category 2A. Also for PN-A.
- Footnote “n” was added, “Consider prophylactic EBRT to inguinal lymph nodes in patients who are not surgical candidates or who decline surgical management.”

**PN-4**
- Imaging was added for palpable ILN as “Abdominal/pelvic CT or MRI and Chest imaging (x-ray, or CT).”
- Management of palpable inguinial lymph nodes, the risk stratification groups were clarified as:
  - Unilateral mobile lymph node(s) <4 cm
  - Unilateral lymph nodes ≥4 cm (fixed or mobile) or Bilateral lymph node(s) (fixed or mobile)
  - Enlarged pelvic lymph nodes
- For non-bulky, unilateral mobile lymph node(s) <4 cm, after treatment with ILND, options for pN1 and pN2-3 disease were added.
- Footnotes
  - Footnote “o” was added, “Consider PET-CT scan.”
  - Footnote “p” was added, “If M1 disease identified, see Management of Metastatic Disease (PN-9).”
  - Footnote “q” was added, “The size threshold of 4 cm represents the largest diameter of contiguous inguinal lymph node(s) tissue as measured on either physical examination and/or axial imaging (CT or MR) and suspected of harboring metastatic disease.” Also for PN-5.

**PN-5**
- For bulky, unilateral ≥4 cm (fixed or mobile) or Bilateral (fixed or mobile), the lymph node status was clarified as:
  - Unilateral mobile ≥4 cm and
  - Unilateral fixed lymph nodes ≥4 cm or bilateral lymph node(s) (fixed or mobile)
- For unilateral mobile ≥4 cm, after treatment for a positive node, if ≥2 nodes positive or extranodal extension, “Adjuvant chemotherapy (if not already given)” was added.

**PN-6**
- Pelvic lymph nodes enlarged, “needle biopsy” was added and the option for negative nodes was added.

**PN-8**
- Local recurrence in inguinal region, the options were changed from “Consider systemic chemotherapy and/or Consider external beam radiation therapy (EBRT) and/or Consider surgical resection” to “Chemotherapy followed by surgical resection or Chemoradiotherapy or Chemotherapy alone.”

**Principles of Radiotherapy**

**PN-B**
- Primary radiation therapy, if tumor <4 cm, “Consider prophylactisc EBRT to inguinal lymph nodes in patients who are not surgical candidates or who decline surgical management” was added to EBRT.

**Principles of Chemotherapy**

**PN-C 2 of 2**
- TIP, “Repeat every 21 days” was clarified as, “Repeat every 3 to 4 weeks.”
**Primary Evaluation**

- **H&P**
  - Risk factors
    - balanitis, chronic inflammation, penile trauma, lack of neonatal circumcision, tobacco use, lichen sclerosus, poor hygiene, sexually transmitted disease
  - Lesion characteristics
    - diameter, location, number of lesions, morphology (papillary, nodular, ulcerous, or flat), relationship to other structures (submucosal, corpora spongiosa, and/or cavernosa, urethra)

- **Cytology or histologic diagnosis**
  - Punch, excisional, or incisional biopsy

**Clinical Diagnosis**

- Tis or Ta
- T1

**Primary Treatment**

- Topical therapy\(^a\)
  - or
- Wide local excision\(^b\) including circumcision
  - or
- Laser therapy (category 2B)
  - or
- Complete glansectomy (category 2B)

*See Management of Non-Palpable Inguinal Lymph Nodes (PN-3) or Palpable Inguinal Lymph Nodes (PN-4)*

*If recurrent disease, see PN-8 or if metastatic disease, see PN-9*

---

\(^{a}\)Topical therapy may include topical imiquimod (5%) or 5-fluorouracil (5-FU) cream.

\(^{b}\)Moh's surgery is an option.

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**PATHOLOGIC DIAGNOSIS**

- **T1**
  - Grade 1-2
  - Grade 3-4

- **T2 or greater**

**PRIMARY TREATMENT**

- **Grade 1-2**
  - Wide local excision,^b,d^ possible STSG or FTSG or Laser therapy (category 2B) or Radiotherapy^e^ (category 2B)

- **Grade 3-4**
  - Wide local excision^d,f^ or Glansectomy^g^ or Partial penectomy^f,h^ or Total penectomy^f,h^ or Radiotherapy^e^ (category 2B) or Chemoradiotherapy^e,i^ (category 3)

- **T2 or greater**
  - Partial penectomy^f,h^ or Total penectomy^f,h^ or Radiotherapy^e^ (category 2B) or Chemoradiotherapy^e,i^ (category 3)

---

^b^Moh’s surgery is an option.

^c^See Principles of Surgery (PN-A).

^d^Complete excision of the skin with a wide negative margin with skin grafting is needed. STSG = split-thickness skin graft; FTSG = full-thickness skin graft.

^e^See Principles of Radiotherapy (PN-B).

^f^Recommend intraoperative frozen sections to achieve negative margins.

^g^Appropriate with proven negative margins for tumors involving the glans only.

^h^When it is necessary to dissect into the corpora cavernosum to achieve a negative margin, a partial or total penectomy is performed.

^i^See Principles of Chemotherapy (PN-C).

**Note:** All recommendations are category 2A unless otherwise indicated. Clinical Trials: NCCN believes that the best management of any cancer patient is in a clinical trial. Participation in clinical trials is especially encouraged.
### Management of Non-Palpable Inguinal Lymph Nodes

<table>
<thead>
<tr>
<th>Nodal Status</th>
<th>Risk Stratification Based on Primary Lesion</th>
<th>Imaging</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low risk (Tis, Ta, ( T_{1a} ))</td>
<td>Surveillance (See PN-7) or Dynamic sentinel node biopsy (DSNB)(^k,^l)</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>Intermediate risk (T1b, Grade 1 or 2) or High risk (T1b, Grade 3 or 4; Any T2 or greater)</td>
<td>Abdominal/pelvic CT or MRI and Chest imaging (x-ray, or CT)</td>
<td>Inguinal lymph node dissection (ILND)(^m,^n) or DSNB(^l)</td>
<td></td>
</tr>
</tbody>
</table>

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\(^1\)Ta verrucous carcinoma is by definition a well-differentiated tumor and would require surveillance alone of inguinal lymph nodes.

\(^k\)DSNB is recommended provided the treating physician has experience with this modality.

\(^l\)If positive lymph nodes are found on DSNB, ILND is recommended.

\(^m\)A modified/superficial inguinal dissection with intraoperative frozen section is an acceptable alternative to stage the inguinal lymph nodes.

\(^n\)Consider prophylactic EBRT to inguinal lymph nodes in patients who are not surgical candidates or who decline surgical management.

---

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Clinical Trials: NCCN believes that the best management of any cancer patient is in a clinical trial. Participation in clinical trials is especially encouraged.
MANAGEMENT OF PALPABLE NON-BULKY INGUINAL LYMPH NODES

NODAL STATUS IMAGING RISK STRATIFICATION BASED ON PHYSICAL/IMAGING FINDINGS TREATMENT

Unilateral mobile lymph node(s) <4 cm\(^q\)

Low-risk primary lesion \implies\ Lymph node biopsy

Negative \implies\ Excisional biopsy or Surveillance

Positive \implies\ ILND

pN1 \implies\ Consider adjuvant radiotherapy\(^e\) or Chemoradiotherapy\(^e,i\) or Chemotherapy\(^i\)

Unilateral lymph nodes ≥4 cm\(^q\) (fixed or mobile) or Bilateral lymph node(s) (fixed or mobile)

High-risk primary lesion\(^r\)

Management of Palpable Bulky Inguinal Lymph Nodes (PN-5)

Enlarged pelvic lymph nodes

Management of Enlarged Pelvic Lymph Nodes (PN-6)

\(^e\)See Principles of Radiotherapy (PN-B)
\(^i\)See Principles of Chemotherapy (PN-C).
\(^q\)Consider PET-CT scan.
\(^r\)If M1 disease identified, see Management of Metastatic Disease (PN-9).
\(^q\)The size threshold of 4 cm represents the largest diameter of contiguous inguinal lymph node(s) tissue as measured on either physical examination and/or axial imaging (CT or MR) and suspected of harboring metastatic disease.
\(^r\)High-risk primary lesion: T1, high-grade, LVI, >50% poorly undifferentiated.

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MANAGEMENT OF PALPABLE BULKY INGUINAL LYMPH NODES

<table>
<thead>
<tr>
<th>NODE STATUS</th>
<th>LYMPH NODES</th>
<th>TREATMENT</th>
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<tbody>
<tr>
<td>Palpable bulky inguinal lymph node(s):</td>
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<tr>
<td>Unilateral ≥4 cm (fixed or mobile) or</td>
<td>Unilateral mobile ≥4 cm&lt;sup&gt;q&lt;/sup&gt;</td>
<td>Needle biopsy</td>
</tr>
<tr>
<td>Bilateral (fixed or mobile)</td>
<td>Unilateral fixed lymph nodes ≥4 cm&lt;sup&gt;q&lt;/sup&gt; or bilateral lymph node(s) (fixed or mobile)</td>
<td>Needle biopsy</td>
</tr>
<tr>
<td></td>
<td>Negative</td>
<td>ILND&lt;sup&gt;c&lt;/sup&gt; or Consider neoadjuvant chemotherapy&lt;sup&gt;i&lt;/sup&gt; followed by ILND</td>
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<tr>
<td></td>
<td>Positive</td>
<td>0–1 positive nodes with viable disease&lt;sup&gt;s&lt;/sup&gt;</td>
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<td>≤2 nodes positive or extranodal extension</td>
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<td></td>
<td>Negative</td>
<td>Adjuvant chemotherapy&lt;sup&gt;i&lt;/sup&gt; (if not already given) and/or Pelvic lymph node dissection (PLND)&lt;sup&gt;c,t,u&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>Positive</td>
<td>Neoadjuvant chemotherapy&lt;sup&gt;i&lt;/sup&gt; and PLND&lt;sup&gt;c,t,u&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>Excisional biopsy</td>
<td>See Surveillance (PN-7)&lt;sup&gt;t&lt;/sup&gt;</td>
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</table>

<sup>c</sup>See Principles of Surgery (PN-A).
<sup>i</sup>See Principles of Chemotherapy (PN-C).
<sup>q</sup>The size threshold of 4 cm represents the largest diameter of contiguous inguinal lymph node(s) tissue as measured on either physical examination and/or axial imaging (CT or MR) and suspected of harboring metastatic disease.
<sup>s</sup>For viable disease post-chemotherapy, consider PLND.
<sup>t</sup>Consider postoperative radiotherapy or chemoradiotherapy.

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### MANAGEMENT OF ENLARGED PELVIC LYMPH NODES

<table>
<thead>
<tr>
<th>NODE STATUS</th>
<th>LYMPH NODES</th>
<th>TREATMENT</th>
</tr>
</thead>
</table>
| Negative    | Potentially resectable | Neoadjuvant chemotherapy
|             | Abdominal/pelvic CT or MRI and chest imaging (x-ray, or CT) |
|             | Stable or clinical response |
|             | Consolidation surgery |
| Positive    | Non-surgical candidate | Chemoradiotherapy
|             | Additional systemic chemotherapy with consideration of radiation therapy for local control or Clinical trial |
|             | See Surveillance (PN-7) |

- **Negative**:
  - Needle biopsy
  - Potentially resectable
  - Neoadjuvant chemotherapy
  - Abdominal/pelvic CT or MRI and chest imaging (x-ray, or CT)
  - Stable or clinical response
  - Consolidation surgery
  - See Surveillance (PN-7)

- **Positive**:
  - Needle biopsy
  - Non-surgical candidate
  - Chemoradiotherapy
  - Additional systemic chemotherapy with consideration of radiation therapy for local control or Clinical trial
  - See Surveillance (PN-7)

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**Note**: All recommendations are category 2A unless otherwise indicated.

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See Principles of Radiotherapy (PN-B).

See Principles of Chemotherapy (PN-C).

Consider PET-CT scan.

If M1 disease identified, see Management of Metastatic Disease (PN-9).

Consider postoperative radiotherapy or chemoradiotherapy.

Consolidation surgery consists of bilateral superficial and deep ILND and unilateral/bilateral PLND.

On CT or MRI, not pathologic stage.
## Penile Cancer

### SURVEILLANCE SCHEDULE

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<tr>
<th>ANATOMIC SITE</th>
<th>INITIAL TREATMENT</th>
<th>SURVEILLANCE</th>
</tr>
</thead>
</table>
| Primary lesion | • Topical therapy  
• Laser therapy  
• Radiation therapy  
• Wide local excision including circumcision  
• Partial penectomy  
• Total penectomy | Clinical exam: \(^y\)  
year 1–2, every 3 mo then  
year 3–5, every 6 mo then  
year 5–10, every 12 mo  
Clinical exam: \(^y\)  
year 1–2, every 6 mo then  
year 3–5, every 12 mo  
Clinical exam: \(^y\)  
year 1–2, every 3 mo then  
year 3–5, every 6 mo | For patients with recurrence at either local or distant sites, see Management of Recurrent Disease (PN-8) |
| Lymph nodes   | N0, N1            | Clinical exam: \(^y\)  
year 1–2, every 6 mo then  
year 3–5, every 12 mo  
Clinical exam: \(^y\)  
year 1–2, every 3-6 mo then  
year 3–5, every 6–12 mo  
Imaging:  
• Chest (CT or x-ray)  
  ▶ year 1–2, every 6 mo  
• Abdominopelvic (CT or MRI)  
  ▶ year 1, every 3 mo then  
  ▶ year 2, every 6 mo | |
|               | N2, N3            | Clinical exam: \(^y\)  
year 1–2, every 3-6 mo then  
year 3–5, every 6–12 mo  |

\(^x\)Patients on active surveillance of clinically negative nodes and at low risk for inguinal metastases.  
\(^y\)Clinical exam includes examination of the penis and inguinal region.  
\(^z\)If an abnormal clinical exam, obese patient, or prior inguinal surgery, then ultrasound, CT, or MRI of the inguinal region can be considered.

---

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MANAGEMENT OF RECURRENT DISEASE

Recurrence of penile lesion after initial treatment

<table>
<thead>
<tr>
<th>Invasion of corpora cavernosa</th>
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<tbody>
<tr>
<td>Absent</td>
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<tr>
<td>Partial penectomy</td>
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<tr>
<td>or</td>
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<tr>
<td>Total penectomy</td>
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<tr>
<td>or</td>
</tr>
<tr>
<td>Repeat penile-sparing treatment (category 2B)</td>
</tr>
</tbody>
</table>

| Present                       |
| Partial penectomy             |
| or                            |
| Total penectomy               |

Local recurrence in inguinal region

| Chemotherapy\textsuperscript{i} followed by surgical resection\textsuperscript{aa} |
| or                                                                                      |
| Chemoradiotherapy\textsuperscript{e,i}                                                |
| or                                                                                      |
| Chemotherapy alone\textsuperscript{i}                                                 |

\textsuperscript{e}See Principles of Radiotherapy (PN-B).
\textsuperscript{i}See Principles of Chemotherapy (PN-C).

Note: All recommendations are category 2A unless otherwise indicated. Clinical Trials: NCCN believes that the best management of any cancer patient is in a clinical trial. Participation in clinical trials is especially encouraged.
MANAGEMENT OF METASTATIC DISEASE

Metastatic penile cancer

- Systemic chemotherapy
- Radiotherapy
- Chemoradiotherapy

Abdominal/pelvic CT or MRI and chest imaging (x-ray, or CT)

Complete/partial response or stable

Consolidation surgery

Consider salvage systemic chemotherapy
- Consider radiotherapy for local control
- Best supportive care/clinical trial

No response/Disease progression

See Surveillance (PN-7)

See Principles of Radiotherapy (PN-B).
See Principles of Chemotherapy (PN-C).
Consider PET-CT scan.
Consolidation surgery consists of bilateral superficial and deep ILND and possible bilateral PLND.

Note: All recommendations are category 2A unless otherwise indicated. Clinical Trials: NCCN believes that the best management of any cancer patient is in a clinical trial. Participation in clinical trials is especially encouraged.
PRINCIPLES OF SURGERY¹

- Tis, Ta penile cancer lesions may be amenable to conservative penile organ-sparing approaches, including: topical therapy, local excision, circumcision, laser therapy (category 2B), or glansectomy (category 2B).

- Partial penectomy should be considered the standard for high-grade primary penile tumors, provided a functional penile stump can be preserved and negative margins are obtained.

- Standard or modified ILND or DSNB is indicated in patients with penile cancer in the absence of palpable inguinal adenopathy if high-risk features for nodal metastasis are seen in the primary penile tumor:
  - Lymphovascular invasion
  - ≥pT1G3 or ≥T2, any grade
  - >50% poorly differentiated

- PLND should be considered at the time of ILND in patients with ≥2 inguinal nodes (on frozen section) on the ipsilateral ILND site or in a delayed procedure in patients with extranodal extension.

¹See Discussion for further details regarding ILND and PLND.
**Primary Radiation Therapy (category 2B) (Penile Preservation)**

**T1-2, N0**

If tumor <4 cm

- Circumcision followed by either:
  - Brachytherapy alone (preferred approach)\(^1,2\) (should be performed with interstitial implant);
  - or
  - EBRT with or without concurrent chemotherapy:\(^3\) Total dose 65–70 Gy with conventional fractionation using appropriate bolus to primary penile lesion with 2-cm margins. Consider prophylactic EBRT to inguinal lymph nodes in patients who are not surgical candidates or who decline surgical management.

If tumor ≥4 cm

- Circumcision followed by either:
  - EBRT with concurrent chemotherapy:\(^3\) 45–50.4 Gy to a portion of or whole penile shaft depending on bulk and extent of lesion plus pelvic/inguinal nodes, then boost primary lesion with 2-cm margins (total dose 60–70 Gy);
  - or
  - Brachytherapy (in select cases and with careful post-treatment surveillance)

**T3-4 or N+ (surgically unresectable)**

- Circumcision followed by:
  - EBRT with concurrent chemotherapy:\(^3\) 45–50.4 Gy to whole penile shaft, pelvic lymph nodes, and bilateral inguinal lymph nodes, then boost primary lesion with 2-cm margins and gross lymph nodes (total dose 60–70 Gy).

**Postoperative Adjuvant Radiotherapy**

- Inguinal Lymph Node Positive
  - Inguinal and pelvic lymph node EBRT to 45–50.4 Gy (strongly consider concurrent chemotherapy\(^3\)).
  - Boost gross nodes and areas of extracapsular extension to a total dose of 60–70 Gy.
  - Treat primary site of disease if positive margin.

- Primary Site Margin Positive
  - Primary site of disease and surgical scar EBRT to 60–70 Gy (for close margin consider radiation treatment vs. observation).
  - Treat bilateral inguinal lymph nodes and pelvic lymph nodes if no or inadequate lymph node dissection.
  - Brachytherapy (in select cases)


\(^3\)For potential radiosensitizing agents and combinations, see Principles of Chemotherapy (PN-C 2 of 2).
PRINCIPLES OF CHEMOTHERAPY

Neoadjuvant
• Neoadjuvant, cisplatin-based chemotherapy should be considered the standard (prior to ILND) in patients with ≥4 cm inguinal lymph nodes (fixed or mobile), if FNA is positive for metastatic penile cancer.¹
  ▶ Patients with initially unresectable (T4) primary tumors may be downstaged by response to chemotherapy.

• A T1xN2-3, M0 penile cancer can receive 4 courses of neoadjuvant paclitaxel, ifosfamide, and cisplatin (TIP).² Stable or responding disease should then undergo consolidative surgery with curative intent.
  ▶ The phase II response rate was 50% in the neoadjuvant setting.
  ▶ The estimated rate of long-term progression-free survival for intent to treat was 36.7%.
  ▶ Improved progression-free and overall survival times were associated with objective response to chemotherapy.

Adjuvant
• There are no sufficient data to form conclusions about the use of adjuvant chemotherapy. By extrapolation from the neoadjuvant data, it is reasonable to give 4 courses of TIP in the adjuvant setting if it was not given preoperatively and the pathology shows high-risk features. (See Management of Palpable Bulky Inguinal Lymph Nodes, PN-5) Adjuvant EBRT or chemoradiotherapy can also be considered for patients with high-risk features, which include any of the following:
  ▶ Pelvic lymph node metastases
  ▶ Extranodal extension
  ▶ Bilateral inguinal lymph nodes involved
  ▶ 4 cm tumor in lymph nodes

Metastatic/Recurrent
• TIP is a reasonable first-line treatment for patients with metastatic penile cancer, including palliative treatment of patients with distant metastases.²
  ▶ 5-FU + cisplatin has been used historically for metastatic penile cancer and can be considered as an alternative to TIP.³ It appears to be effective for some patients, although the toxicities may be limiting and require dose reductions.⁴
  ▶ Bleomycin-containing regimens are associated with unacceptable toxicity⁵ and are no longer recommended.
  ▶ There are no randomized clinical trials due to the rarity of penile cancer in industrialized countries.

Second-line
• No standard second-line systemic therapy exists.
  ▶ A clinical trial is preferred. The evidence to support the palliative use of second-line therapy is limited.⁶ In select patients, paclitaxel⁷ or cetuximab⁸ may be considered, especially if not previously treated with a similar class of agent.

Note: All recommendations are category 2A unless otherwise indicated. Clinical Trials: NCCN believes that the best management of any cancer patient is in a clinical trial. Participation in clinical trials is especially encouraged.
PRINCIPLES OF CHEMOTHERAPY

Preferred combination chemotherapy regimens

TIP²

Paclitaxel 175 mg/m² IV over 3 hours on Day 1
Ifosfamide 1200 mg/m² IV over 2 hours on Days 1–3
Cisplatin 25 mg/m² IV over 2 hours on Days 1–3
Repeat every 3 to 4 weeks

5-FU + cisplatin⁴ (category 2B)
Continuous infusion 5-FU 1000 mg/m²/d IV on Days 1–5
Cisplatin 100 mg/m² IV on Day 1
Repeat every 3 to 4 weeks

Radiosensitizing agents and combinations⁹ (Chemoradiotherapy)

• Preferred
  ▶ Cisplatin alone, or in combination with 5-FU

• Alternate options
  ▶ Mitomycin C in combination with 5-FU
  ▶ Capecitabine (for palliation)

References


Note: All recommendations are category 2A unless otherwise indicated. Clinical Trials: NCCN believes that the best management of any cancer patient is in a clinical trial. Participation in clinical trials is especially encouraged.
# Table 1

**American Joint Committee on Cancer (AJCC)**

**TNM Staging System for Penile Cancer (7th ed., 2010)**

<table>
<thead>
<tr>
<th>Primary Tumor (T)</th>
<th>ANATOMIC STAGE/PROGNOSTIC GROUPS</th>
</tr>
</thead>
<tbody>
<tr>
<td>TX</td>
<td>Stage 0</td>
</tr>
<tr>
<td>T0</td>
<td>Tis N0 M0</td>
</tr>
<tr>
<td>Ta</td>
<td>Ta N0 M0</td>
</tr>
<tr>
<td>T1a</td>
<td>T1a N0 M0</td>
</tr>
<tr>
<td>Tis</td>
<td>T1a N0 M0</td>
</tr>
<tr>
<td>T1b</td>
<td>T1b N0 M0</td>
</tr>
<tr>
<td>T2</td>
<td>T1b N0 M0</td>
</tr>
<tr>
<td>T3</td>
<td>T1b N0 M0</td>
</tr>
<tr>
<td>T4</td>
<td>T1b N0 M0</td>
</tr>
</tbody>
</table>

*Note: Broad pushing penetration (invasion) is permitted; destructive invasion is against the diagnosis*

<table>
<thead>
<tr>
<th>Regional Lymph Nodes (N)</th>
<th>ANATOMIC STAGE/PROGNOSTIC GROUPS</th>
</tr>
</thead>
<tbody>
<tr>
<td>cNX</td>
<td>Stage 0</td>
</tr>
<tr>
<td>cN0</td>
<td>Tis N0 M0</td>
</tr>
<tr>
<td>cN1</td>
<td>T1a N0 M0</td>
</tr>
<tr>
<td>cN2</td>
<td>T1b N0 M0</td>
</tr>
<tr>
<td>cN3</td>
<td>T1b N0 M0</td>
</tr>
</tbody>
</table>

*Note: Pathologic stage definition based on biopsy or surgical excision.*

<table>
<thead>
<tr>
<th>Distant Metastasis (M)</th>
<th>ANATOMIC STAGE/PROGNOSTIC GROUPS</th>
</tr>
</thead>
<tbody>
<tr>
<td>M0</td>
<td>Stage 0</td>
</tr>
<tr>
<td>M1</td>
<td>Stage 0</td>
</tr>
</tbody>
</table>

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ST-1
## Discussion

### NCCN Categories of Evidence and Consensus

- **Category 1:** Based upon high-level evidence, there is uniform NCCN consensus that the intervention is appropriate.
- **Category 2A:** Based upon lower-level evidence, there is uniform NCCN consensus that the intervention is appropriate.
- **Category 2B:** Based upon lower-level evidence, there is NCCN consensus that the intervention is appropriate.
- **Category 3:** Based upon any level of evidence, there is major NCCN disagreement that the intervention is appropriate.

All recommendations are category 2A unless otherwise noted.

### Table of Contents

- **Overview** ............................................................... MS-2
- **Risk Factors** .......................................................... MS-2
- **Clinical Presentation** .............................................. MS-3
- **Characterization and Clinical Staging** ...................... MS-3
- **Management of Primary Lesions** ............................... MS-4
  - Diagnosis ..................................................................... MS-4
  - NCCN Recommendations ........................................ MS-4
  - T1G1-2 ....................................................................... MS-5
  - T1G3-4 or T≥2 .......................................................... MS-5
- **Management of Regional Lymph Nodes** ..................... MS-6
  - Evaluation and Risk Stratification .............................. MS-6
  - Dynamic Sentinel Node Biopsy ................................ MS-7
  - Inguinal Lymph Node Dissection ................................ MS-8
  - Modified Template Lymphadenectomy ...................... MS-8
  - Delayed Inguinal Lymphadenectomy ........................... MS-9
  - Unilateral Versus Bilateral Lymphadenectomy ............. MS-9
  - Pelvic Lymphadenectomy .......................................... MS-9
  - Chemotherapy ........................................................ MS-10
  - Chemoradiotherapy .................................................. MS-10
  - Non-Palpable Nodes .................................................. MS-11
  - Unilateral Palpable Nodes <4 cm (mobile) ................ MS-11
  - Unilateral Palpable Nodes ≥4 cm (mobile) ................. MS-11
  - Unilateral Palpable Nodes ≥4 cm (fixed) or Bilateral Palpable Nodes (fixed or mobile) MS-11
  - Enlarged Pelvic Lymph Nodes ................................... MS-12
- **Surveillance** .......................................................... MS-12
- **Recurrence** ........................................................... MS-12
- **Metastatic Disease** ................................................ MS-13
- **Summary** .............................................................. MS-13
- **References** ........................................................... MS-14
Overview

Squamous cell carcinoma (SCC) of the penis is a rare disease, representing 0.4% to 0.6% of all malignant neoplasms among men in the United States and Europe. In 2016, the estimated number of new cases of penile and other male genital cancers in the United States is 2030, with 340 predicted cancer-specific deaths. Incidence is higher (up to 10%) among men in the developing countries of Asia, Africa, and South America. The most common age of presentation is between 50 and 70 years. Early diagnosis is of utmost importance, as this is a disease that can result in devastating disfigurement and has a 5-year survival rate of approximately 50% (over 85% for patients with negative lymph nodes and 29%–40% for patients with positive nodes, with the lowest survival rates at 0% for patients with pelvic lymph node [PLN] involvement). As the rarity of this disease makes it difficult to perform prospective, randomized trials, the NCCN Panel relied on the experience of penile cancer experts and the best currently available evidence-based data to collectively lay down a foundation to help standardize the management of the malignancy.

Literature Search Criteria and Guidelines Update Methodology

Prior to the update of this version of the NCCN Guidelines for Penile Cancer, an electronic search of the PubMed database was performed to obtain key literature published between August 20, 2014 and September 8, 2015, using the following search term: penile cancer or squamous cell carcinoma of the penis. The PubMed database was chosen as it remains the most widely used resource for medical literature and indexes only peer-reviewed biomedical literature.

The search results were narrowed by selecting studies in humans published in English. Results were confined to the following article types: Clinical Trial; Guideline; Meta-Analysis; Randomized Controlled Trial; Systematic Reviews; and Validation Studies.

The PubMed search resulted in 9 citations and their potential relevance was examined. The data from key PubMed articles as well as articles from additional sources deemed as relevant to these guidelines and discussed by the panel have been included in this version of the Discussion section (eg, e-publications ahead of print, meeting abstracts). Recommendations for which high-level evidence is lacking are based on the panel’s review of lower-level evidence and expert opinion.

The complete details of the Development and Update of the NCCN Guidelines are available on the NCCN webpage.

Risk Factors

In the United States the median age of diagnosis is 68 years, with an increase in risk for males older than 50 years. Early detection is assisted by the ability to do a good physical exam. Phimosis may hinder the capability to properly inspect the areas of highest incidence—the glans, inner preputial layer, coronal sulcus, and shaft. Men with phimosis carry an increased risk for penile cancer of 25% to 60%. A review of penile SCC in the United States showed that 34.5% of patients had the primary lesion on the glans, 13.2% on the prepuce, and 5.3% in the shaft, with 4.5% overlapping and 42.5% unspecified. Other risk factors include balanitis, chronic inflammation, penile trauma, tobacco use, lichen sclerosus, poor hygiene, and a history of sexually transmitted disease(s), especially HIV and HPV. Overall, about 45% to 80% of penile cancers are related to HPV, with a strong correlation with types 16 and 18. There is an 8-fold increased risk for patients with HIV, which may correspond to a higher incidence of HPV among males with HIV. Cigarette smokers are noted to be 3 to 4.5 times more likely...
to develop penile cancer.\textsuperscript{11,14} Patients with lichen sclerosus are noted to have a 2\% to 9\% risk of developing penile carcinoma.\textsuperscript{15-17} Psoriasis patients undergoing psoralen plus ultraviolet A (PUVA) treatment have an increased penile cancer incidence of 286 times compared to the general population. Therefore, they should be shielded during treatment and any penile lesion should be closely monitored.\textsuperscript{18} A study of men with advanced penile SCC receiving systemic therapy identified visceral metastases and an ECOG performance score greater than or equal to 1 as poor prognostic factors for both overall survival (OS) and progression-free survival.\textsuperscript{19} Studies remain limited on predictive factors of prognosis in this patient population.

**Clinical Presentation**

Most often penile SCC presents as a palpable, visible lesion on the penis, which may be associated with penile pain, discharge, bleeding, or a foul odor if the patient delays seeking medical treatment. The lesion may be characterized as nodular, ulcerative, or fungating, and may be obscured by phimosis. The patient may exhibit signs of more advanced disease, including palpable nodes and/or constitutional symptoms (eg, fatigue, weight loss).

**Characterization and Clinical Staging**

Approximately 95\% of penile cancers originate in squamous epithelial cells and are further categorized as either SCC or penile intraepithelial neoplasia (PIN).\textsuperscript{20} PIN is a premalignant condition at high risk of developing into SCC of the penis and includes the clinical entities bowenoid papulosis, erythroplasia of Queyrat, and Bowen's disease.\textsuperscript{20} The AJCC recognizes four subtypes of SCC: verrucous, papillary squamous, warty, and basaloid.\textsuperscript{21} The verrucous subtype is felt to be of low malignant potential, while other variants reported—adenosquamous and sarcomatoid variants—carry a worse prognosis.\textsuperscript{22,23} The primary lesion is further characterized by its growth pattern with superficial spread, nodular or vertical-phase growth, and verrucous pattern. In addition to the penile lesion, evaluation of lymph nodes is also critical, as involvement of the inguinal lymph nodes (ILNs), the number and site of positive nodes, and extracapsular nodal involvement provide the strongest prognostic factors of survival.\textsuperscript{6,24}

The AJCC Tumor, Node, and Metastasis (TNM) system for penile carcinoma has been used for staging, with the most recent update published in 2010. It was initially introduced in 1968 and was subsequently revised in 1978, 1987, and 2002.\textsuperscript{21,25-28} In the 2010 update, the AJCC has made the distinction between clinical and pathologic staging while eliminating the difference between superficial and deep inguinal metastatic nodes.\textsuperscript{21} Other changes to the 2010 TNM system include: subdivision of T1 into T1a and T1b as determined by the presence or absence of lymphovascular invasion or poorly differentiated cancers; limitation of the T3 category to urethral invasion and T4 prostatic invasion; and grouping of stage II to include T1b N0M0 as well as T2-3 N0M0 (see staging tables in the algorithm). A grading system for SCC of the penis based on degree of cell anaplasia is defined as: grade 1, well differentiated (no evidence of anaplasia); grade 2, moderately differentiated (<50\% anaplasia); and grade 3, poorly differentiated (>50\% anaplastic cells).\textsuperscript{29} According to the AJCC, if no grading system is specified, a general system should be followed: GX, grade cannot be assessed; G1-3 as previously mentioned above; and G4, undifferentiated.\textsuperscript{21} The overall degree of cellular differentiation with high-risk, poorly differentiated tumors is an important predictive factor for metastatic nodal involvement.\textsuperscript{30} The AJCC also recommends collection of site-specific factors, including: the distinction between corpus spongiosum and corpus cavernosum involvement, the percentage of tumor that is poorly differentiated, the depth of invasion in
Management of Primary Lesions

Diagnosis

Evaluation of the primary lesion, regional lymph nodes, and distant metastasis will dictate the appropriate and adequate management of SCC of the penis, beginning with the first evaluation at presentation and then throughout follow-up. Vital to the initial management is a good physical exam of the penile lesion(s) that remarks on the diameter of the lesion(s) or suspicious areas; location(s) on the penis; number of lesions; morphology of the lesion(s); whether the lesion(s) are papillary, nodular, ulcerous, or flat; and relationship with other structures including submucosal, urethra, corpora spongiosa, and/or corpora cavernosa. To complete the initial evaluation, a histologic diagnosis with a punch, excisional, or incisional biopsy is paramount in determining the treatment algorithm based on a pathologic diagnosis. This will provide information on the grade of the tumor, and will assist in the risk stratification of the patient for regional lymph node involvement. MRI or ultrasound can be used to evaluate the depth of tumor invasion. For the evaluation of lymph nodes, see Management of Regional Lymph Nodes.

NCCN Recommendations

**Tis or Ta**

For patients with penile carcinoma in situ or noninvasive verrucous carcinoma, penis-preserving techniques may be utilized, including topical imiquimod (5%) or 5-fluorouracil (5-FU) cream, circumcision and wide local excision such as Mohs surgery, laser therapy (category 2B) using carbon dioxide or neodymium:yttrium-aluminum-garnet, and complete glansectomy (category 2B). Among these, topical therapy and excisional organ-sparing surgery are the most widely used. Retrospective studies of laser therapy reported local recurrence rates of around 18%, comparable to that of surgery, with good cosmetic and functional results. Glansectomy, removal of the glans penis, has also been studied with no recurrence observed in some cases.

While topical therapy for the treatment of PIN has been reported in numerous case studies and case reports, the data are limited by the small sample sizes and variation in treatment protocols. Topical therapy is a valuable outpatient treatment due to ease of administration; however, patients should be monitored for adherence to therapy and for toxicity or adverse events. Local skin and application site reactions may occur and are generally mild to moderate, though severe reactions may occur with a higher frequency of application. Modification of the application frequency can resolve these complications. Despite significant response rates, the probability of relapse is higher following topical therapy than with other more aggressive therapies. Therefore, patients who are eligible for topical therapy should be routinely monitored for recurrence.

A retrospective review from a prospective database of patients diagnosed with PIN over a 10-year range identified patients who received either 5-FU as first-line therapy or imiquimod as the second-line topical agent. Topical chemotherapy was given to 45 patients with a mean follow-up of 34 months. Therapy was standardized to 12 hours every 48 hours for 28 days. A complete response (CR) was reported in 25 patients (57%), while a partial response was seen in 6 patients (13.6%); no response was observed in the remaining 13 patients (29.5%). Following application of 5-FU, local toxicity and adverse events occurred in 10% and 12% of patients, respectively. In a more recent study, 5-FU treatment duration ranged from 3 to 7 weeks...
and was determined based on clinical response. Out of 19 patients, 14 (73.7%) had a CR and none of the patients had recurrence at median time of follow-up at 3.5 years. Topical 5-FU for 6 weeks has also reported good response rates at 5 years.

Imiquimod has been investigated as a second-line therapy for PIN. Due to its ability to produce significant inflammation, initiation of imiquimod therapy at a lower frequency (eg, 2x per week) may be beneficial to evaluate for toxicity or adverse events before increasing the frequency of application. Early studies suggested a 100% response to imiquimod (n = 47; 70% CR), though a recent review identified a lower response to therapy with 63% of patients showing a CR and 29% of patients showing no response. The study highlighted that the difference in response may be related to the frequency and duration of application as well as the PIN subtype. In this study, Bowenoid papulosis and Bowen’s disease subtypes responded better to imiquimod than the erythroplasia of Queyrat subtype. Longer, less frequent application (ie, less than 4 times per week for an average of 113 days) was demonstrated to have a better response than a shorter, more frequent application (ie, 4 times or more per week for an average of 53 days) (81% vs. 68%, respectively).

**T1G1-2**

Careful consideration should be given to penile-preserving techniques if the patient is reliable in terms of compliance with close follow-up. These techniques include wide local excision as well as Mohs surgery as an option plus reconstructive surgery (category 2A), laser therapy (category 2B), and radiotherapy (category 2B) delivered as brachytherapy with interstitial implant (preferred) or external beam radiation therapy (EBRT) with or without chemotherapy. For wide local excision, a complete excision of the skin with a wide negative margin with skin grafting is needed. Either a split-thickness skin graft or full-thickness skin graft may be considered. Emphasis is placed again on patient selection and close follow-up, as the 2-year recurrence rate may reach up to 50%. Studies have shown that surgical margins of 5 to 10 mm are as safe as 2-cm surgical margins, and 10- to 20-mm margins provide adequate tumor control. Circumcision should always precede radiation therapy (RT) to prevent radiation-related complications.

**T1G3-4 or T2**

These lesions typically require more extensive surgical intervention with partial or total penectomy depending on the characteristics of the tumor and depth of invasion. Intraoperative frozen sectioning is recommended to achieve negative surgical margins. If the tumor encompasses less than half of the glans and the patient agrees to very close observation, then a more conservative approach such as wide local excision or glansectomy may be considered for patients with T1G3-4 diagnosis. The patient should understand that there is an increased risk for recurrence and potential for a repeat wide local excision should a local recurrence be noted, provided there is no invasion of the corpora cavernosa. A clear and frank discussion should be had with the patient that a partial or total penectomy will likely be required should a larger or more invasive lesion be present.

The tumor size is an important factor when choosing RT as treatment. As the average length of the glans is about 4 cm, this serves as a cutpoint to reduce the risk of under-treating cavernosal lesions. In a study of 144 patients with penile cancer restricted to the glans treated by brachytherapy, larger tumors, especially those larger than 4 cm, are associated with higher risk of recurrence. A high, 10-year, cancer-specific survival rate of 92% was achieved in this series.
There was nonuniform consensus among NCCN panelists on the use of RT as primary therapy due to scant data. RT alone is a category 2B recommendation for T1G3-4 and T2 tumors, while RT with concurrent chemotherapy is a category 3 recommendation. RT should be given after circumcision has been performed.

For T1G3-4 tumors smaller than 4 cm, brachytherapy with interstitial implant is preferred, but EBRT with or without chemotherapy is a viable option. Consider prophylactic ILN irradiation if selecting EBRT.

For tumors 4 cm or larger, EBRT combined with chemotherapy is recommended. Brachytherapy may be appropriate in select cases of tumors 4 cm or larger, but careful monitoring is necessary as the risks of complications and failures increase. Crook and colleagues reported a 10-year cause-specific survival of 84% in 67 patients with T1-2 (select cases of T3) penile lesions treated with primary brachytherapy. Brachytherapy is not recommended following penectomy or partial penectomy but may be considered following wide local excision or excisional biopsy of small lesions. Brachytherapy should only be performed in centers with significant experience using this treatment modality.

Post-surgical RT to the primary tumor site may be considered for positive margins.

Management of Regional Lymph Nodes
Evaluation and Risk Stratification
The presence and extent of regional ILN metastases has been identified as the single most important prognostic indicator in determining long-term survival in men with invasive penile SCC. Evaluation of the groin and pelvis is an essential component of the metastatic workup of a patient. The involvement of the ILN can be clinically evident (ie, palpable vs. non-palpable), adding to the difficulty in management. Clinical exam for ILN involvement should attempt to evaluate and assess for palpability, number of inguinal masses, unilateral or bilateral localization, dimensions, mobility or fixation of nodes or masses, relationship to other structures (eg, skin, Cooper’s ligaments), and edema of the penis, scrotum, and/or legs. Crossover drainage from left to right and vice versa does occur and is reproducible with lymphoscintigraphy. The physical exam should describe the diameter of node(s) or mass(es), unilateral or bilateral localization, number of nodes identified in each inguinal region, and the relationship to other structures, particularly with respect to the mobility or fixation of the node(s) or mass(es) to adjacent structures and/or involvement of the overlying skin. Imaging for palpable disease by CT or MRI may be used to assess the size, extent, location, and structures that are in close proximity to the ILN, as well as the presence of pelvic and retroperitoneal lymph nodes and distant metastasis. CT and MRI are limited in patients with non-palpable disease. While studies have looked at the use of nanoparticle-enhanced MRI, PET/CT, and 18F-fluorodeoxyglucose (FDG) PET/CT, the small sample sizes require validation in larger prospective studies. When considering one imaging modality to evaluate the stage of the primary lesion and lymph node status, MRI appears to be the best choice to enhance the physical exam in patients where the inguinal region is difficult to assess (eg, morbidity, previous chemotherapy/radiotherapy). Consideration needs to be given to whether or not the primary lesion demonstrated any adverse prognostic factors. If one or more of these high-risk features is present, then pathologic ILN staging must be performed. Up to 25% of patients with non-palpable lymph nodes harbor micrometastases. Therefore, several predictive factors have been evaluated for their ability to identify the presence of occult lymph node metastases.
metastasis. Slaton et al concluded that patients with pathologic stage T2 or greater disease were at significant risk (42%–80%) of nodal metastases if they exhibited greater than 50% poorly differentiated cancer and/or vascular invasion, and therefore should be recommended to undergo an inguinal lymph node dissection (ILND). These factors can then further define patients into low-, intermediate-, and high-risk groups for lymph node metastasis. The European Association of Urology determined risk stratification groups for patients with non-palpable ILNs, and validated this in both uni- and multivariate analyses of prognostic factors. Patients can be stratified based on stage and/or grade into risk groups based on the likelihood of harboring occult node-positive disease, with the low-risk group defined as patients with Tis, Ta, or T1a disease; the intermediate group as those with T1b disease (lymphovascular invasion); and the high-risk group as those with T2 or G3/G4 disease.

There is a paucity of data regarding the predictive value of lymph node removal. A singular study suggests that disease-specific survival following radical lymphadenectomy can be predicted by the lymph node count and lymph node density. Removal of greater than or equal to 16 lymph nodes in patients with pathologic negative nodes was associated with a significantly longer disease-specific survival rate (P < .05). Furthermore, the 5-year disease-free survival in patients with pathologic positive nodes was 81.2% in patients with lymph node density greater than 16% compared to 24.4% in patients with less than 16% lymph node density (P < .001). Although this study suggests that lymph node count and density may be useful in predicting disease-specific survival, a larger validation study is necessary to support these preliminary data.

Dynamic Sentinel Node Biopsy

The work by Cabanas used lymphangiograms and anatomic dissections to evaluate the sentinel lymph node drainage for penile cancer with non-palpable ILNs. This technique has been shown to have false-negative rates as high as 25%; therefore, it is no longer recommended. Advancements have been made with the dynamic sentinel node biopsy (DSNB) technique developed for penile cancer by the Netherlands Cancer Institute using lymphoscintigraphy and performed with technetium-99m–labeled nanocolloid and patent blue dye isosulfan blue. Initially, this technique was associated with a low sensitivity and high false-negative rate (16%–43%). Refinement of the technique to include serial sectioning and immunohistochemical staining of pathologic specimens, preoperative ultrasonography with and without fine-needle aspiration (FNA) cytology, and exploration of groins in which no sentinel node is visualized on intraoperative assessment decreased the false-negative rate from 19% to only 5%. Using FNA with ultrasound can increase the diagnostic yield in metastases greater than 2 mm in diameter. Crashaw et al used ultrasound with DSNB and noted improved accuracy in identifying patients with occult lymph node metastases. With modification of the NCI protocol, Hadway et al were able to achieve a similar false-negative rate (5%) with an 11-month follow-up. A recent observational cohort study of 1000 patients treated between 1956 and 2012 suggests that DSNB can improve 5-year survival in patients with clinically node-negative groins. Data in this study showed that patients treated prior to 1994 (the year DSNB was incorporated into treatment) had an 82% 5-year survival compared to the 91% 5-year survival seen in patients treated between 1994 and 2012 (P = .021). However, there are several limitations of this study including the possibility that improved staging resulted in more patients being grouped in a higher risk group. Therefore, incorporation of DSNB into treatment guidelines...
should be limited to centers with experience. Secondary to the technical challenges associated with DSNB, to be accurate and reliable, it is recommended that DSNB be performed at tertiary care referral centers where at least 20 procedures are done per year. It should be noted that DSNB is not recommended in patients with palpable ILNs.

**Inguinal Lymph Node Dissection**

The most frequent sites of metastasis from penile cancer are the ILNs, typically presenting as palpable inguinal lymphadenopathy. The management of ILNs by ILND has been fraught with great fears of surgical morbidity. Early treatment of lymph node involvement has been shown to have a positive impact on survival, except if the patient has bulky nodal spread or other sites of metastases. Palpable lymphadenopathy at the time of diagnosis does not warrant an immediate ILND. Of the patients with palpable disease, 30% to 50% will be secondary to inflammatory lymph node swelling instead of metastatic disease. Although the distinction between reactive lymph nodes and metastatic disease has traditionally been done with a 6-week course of antibiotics, percutaneous needle biopsy is becoming the favored approach among many penile cancer experts for patients with palpable nodes. In this setting, antibiotics are useful if the patient has a suspected underlying cellulitis at the site of palpable inguinal lymphadenopathy and future site of ILND.

The boundaries of the standard, full-template ILND (ie, Daseler’s quadrilateral area) are: superiorly, the inguinal ligament; inferiorly, the fossa ovalis; laterally, the medical border of sartorius muscle; and medially, the lateral edge of adductor longus muscle. Historically, it has been recommended to keep the patient on bed rest for 48 to 72 hours, especially after myocutaneous flaps or repair of large skin defects, although the necessity for this is debatable and not corroborated with rigorous scientific data. Closed suction drains are placed at surgery and are typically removed when drainage is less than 50 to 100 mL per day. Consideration should be given to keeping the patient on a suppressive dose of an oral cephalosporin (or other gram-positive, broad-spectrum antibiotic) for several days to weeks postoperatively in an attempt to decrease the risk of wound-related issues and minimize the risk for overall complications. However, the data supporting this treatment approach are very limited.

**Modified Template Lymphadenectomy**

In attempts to decrease the morbidity associated with standard ILND, a modified template lymphadenectomy has been proposed that uses a shorter skin incision, limiting the field of inguinal dissection by excluding the area lateral to the femoral artery and caudal to the fossa ovalis, with preservation of the saphenous vein and elimination of the need to transpose the sartorius muscle while providing an adequate therapeutic effect. This technique is commonly reserved for patients with a primary tumor that places them at increased risk for inguinal metastasis but with clinically negative groins on examination. The modified technique has shown a decrease in complications. Contemporary modified ILND should include the central and superior zones of the inguinal region, as these sections were not included in the dissection leading to a false-negative rate of 15%. It is important to note that if nodal involvement is detected on frozen section, the surgical procedure should be converted to a standard, full-template lymphadenectomy. A standard full-template lymphadenectomy should be considered in all patients who have resectable inguinal lymphadenopathy. However, studies would favor neoadjuvant chemotherapy prior to proceeding with surgery, particularly in patients with bulky ILN metastases (ie, fixed nodes or nodal diameter ≥3 cm). Generally, ILND is performed within 4 to 6 weeks following the completion of systemic chemotherapy...
to allow patient recovery while minimizing the risk of cancer progression post-chemotherapy.

**Delayed Inguinal Lymphadenectomy**

Since data exist that suggest men with clinically negative groins undergoing immediate ILND have better survival outcomes than men undergoing delayed ILND once their groins are clinically positive, it is recommended that in most circumstances men with high-risk penile tumors should undergo immediate ILND. However, patients with lower-risk tumors who are undergoing active surveillance or high-risk men who refuse immediate ILND may experience an inguinal nodal recurrence at some time point during follow-up. The median time to inguinal recurrence after treatment of the primary penile tumor is approximately 6 months, with 90% occurring by year 3 and 100% by year 5.  

**Unilateral Versus Bilateral Lymphadenectomy**

In patients with high-risk features that do not have palpable lymph nodes, bilateral lymphadenectomy is generally performed, because it is not possible to predict the laterality of inguinal nodal metastasis based on the location of the tumor on the penis. Similarly, in patients who have a unilateral palpable node, about 30% will have contralateral positive nodes that are not palpable. Therefore, bilateral lymphadenectomy should be considered the standard of care in patients undergoing immediate ILND for high-risk penile tumors or because of palpable nodes. When there is a delayed (>1 year after treatment of the primary penile tumor) inguinal recurrence of cancer, it is usually unilateral, and some authors have suggested that ipsilateral ILND is adequate while others have advocated for bilateral ILND in this circumstance.

**Pelvic Lymphadenectomy**

Approximately 20% to 30% of patients with positive ILNs will also have cancer within PLNs. Interestingly, penile tumors do not appear to metastasize to the PLNs without first affecting the inguinal node echelon (ie, no skip lesions). Patients who have only one positive inguinal node have a risk of pelvic nodal involvement of less than 5% as reported by the Netherlands Cancer Institute. The presence of cancer within the PLN is associated with a very poor 5-year survival rate that is typically of less than 10%. Based on these prior reports, pelvic lymphadenectomy (resection of external iliac, internal iliac, and obturator lymph nodes) is recommended in patients with 2 or more positive ILNs and in the clinical context of high-grade cancer within the ILN pathologic specimen. Pelvic lymph node dissection (PLND) can be conducted during the same operative session as the ILND if the intraoperative frozen section is positive in 2 or more of the inguinal nodes (raising the importance of obtaining a lymph node count intraoperatively) or in a delayed staged fashion based on the pathologic features of the ILND specimen.

A recent retrospective analysis evaluated the benefit of adjuvant chemotherapy on OS of patients with positive PLN following lymph node dissection. Less than half of the patients in this multi-institutional study received adjuvant chemotherapy (36 out of 84). These patients were younger, had a less aggressive pathology, were less inclined to receive adjuvant radiation, and demonstrated less bilateral inguinal disease and more inguinal extranodal extension. The median OS was higher for these patients compared to patients who did not receive adjuvant chemotherapy (21.7 months vs. 10.1 months; \( P = .021 \)). Adjuvant chemotherapy was further shown to be an independent factor in the improved OS based on multivariate analysis (HR, 0.40; 95% CI, 0.19–
One area of controversy is whether the PLND should be performed ipsilaterally or bilaterally in patients with unilateral positive ILNs. Data suggest that the number of positive ILNs identified at the time of dissection may direct clinicians to unilateral or bilateral dissection. In a single retrospective study, the presence of 4 or more positive ILNs supported bilateral PLND.\textsuperscript{106} Unilateral PLND was recommended if 3 or fewer ILN metastases were identified and if there was no suspicion of contralateral pelvic lymphadenopathy on preoperative imaging or intraoperatively. Crossover (right to left or left to right) of inguinal to pelvic nodes has not been well studied; hence, both approaches are feasible and left at the discretion of the surgeon based on case-specific characteristics.

**Chemotherapy**

A patient who presents with resectable bulky disease will rarely be cured with a single treatment modality. Consideration should be given to neoadjuvant chemotherapy if ILNs are greater than or equal to 4 cm. Patients who may benefit from surgical consolidation would be those who had stable, partial, or CR following systemic chemotherapy, thus increasing their potential for disease-free survival.\textsuperscript{96,97} Pagliaro et al\textsuperscript{107} performed a phase II clinical trial in 30 patients, with stage N2 or N3 (stage III or stage IV) penile cancer without distant metastases, receiving neoadjuvant chemotherapy with paclitaxel, ifosfamide, and cisplatin. In this series, 50% of patients were noted to have a clinically meaningful response and 22 patients (73.3%) subsequently underwent surgery. There was an improved time to progression and OS associated with chemotherapy responsiveness ($P < .001$ and $P = .001$, respectively), absence of bilateral residual tumor ($P = .002$ and $P = .017$, respectively), and absence of extranodal extension ($P = .001$ and $P = .004$, respectively) or skin involvement ($P = .009$ and $P = .012$, respectively).

**Chemoradiotherapy**

Chemoradiotherapy has demonstrated improved responses in patients with other SCCs, specifically patients with vulvar and anal cancer.\textsuperscript{108-111} Anecdotal data for the use of chemoradiotherapy in patients with penile cancer have been reported with mixed results.\textsuperscript{112-115} Based on the limited data, chemoradiotherapy is a treatment option in select patients.

For patients with T1 or T2 disease, EBRT with concurrent chemotherapy may be considered though brachytherapy is preferred for tumors less than 4 cm. Similarly EBRT with concurrent chemotherapy can be used for T3 or T4 disease or in patients with nodal involvement. Postoperative adjuvant chemotherapy is recommended in ILN-positive patients. Chemoradiotherapy can be considered for patients with high-risk features including PLN metastases, extranodal extension, bilateral ILN involvement, and tumors in lymph nodes larger than 4 cm.

Chemoradiotherapy is a recommended strategy for patients with resistant disease. The use of chemoradiotherapy as primary treatment is a category 3 recommendation due to the limited studies that have investigated its role for treatment of penile cancer. For patients with palpable, non-bulky pN2 or pN3 disease, treatment may entail adjuvant radiotherapy, chemoradiotherapy, or chemotherapy following ILND or PLND. Chemoradiotherapy is recommended for the management of enlarged PLNs in non-surgical candidates or for local recurrence in the inguinal region or metastatic penile cancer.
NCCN Recommendations

Non-Palpable Nodes
Most low-risk patients (Tis, Ta, T1a) are followed with a surveillance protocol, as the probability of occult micrometastases in ILNs is less than 17%.\(^{71,99}\) If positive lymph nodes are found on DSNB, ILND is recommended. For patients at intermediate (T1bG1-2) or high (T1bG3-4, T2 or greater) risk, a modified or radical inguinal lymphadenectomy is strongly recommended as occult metastatic disease ranges between 68% and 73%.\(^{57,71,99}\) If positive nodes are present on the frozen section, then a superficial and deep inguinal lymphadenectomy should be performed (with consideration of a PLND). Prophylactic EBRT to the ILNs should be considered in patients who are unable or unwilling to undergo surgical management. Alternatively, DSNB can be considered.

As DSNB is currently not widely practiced in the United States, it is a category 2B option for examining non-palpable nodes to determine the need for a modified lymphadenectomy in place of predictive factors.\(^ {116,117}\) This technique should be performed in tertiary care referral centers with substantial experience. DSNB is not recommended for Ta tumors, as observation alone of the ILNs is sufficient for these well-differentiated lesions in the absence of palpable adenopathy.

Unilateral Palpable Nodes <4 cm (mobile)
Lymph node biopsy is considered standard for these patients if no risk feature is present in the primary lesion. Risk features include T1 tumors; high grade; lymphovascular invasion; and poor differentiation in more than half of the tumor cells. The NCCN Panel recommends omitting the procedure for patients with high-risk primary lesions to avoid delay of lymphadenectomy. A negative lymph node biopsy should be confirmed with an excisional biopsy. Alternatively, careful surveillance may be considered following a negative lymph node biopsy. Positive findings from either procedure warrant an immediate ILND. If pN2-3 disease, consider adjuvant radiotherapy, chemoradiotherapy, or chemotherapy following the ILND.

Unilateral Palpable Nodes ≥4 cm (mobile)
Large, unilateral, mobile nodes should first be confirmed by needle biopsy. A negative needle biopsy should be confirmed by an excisional biopsy. If results are negative again, the patient should be closely followed. Patients with confirmed nodes are amenable to standard or modified ILND. Neoadjuvant chemotherapy should be considered before surgery. No further treatment is necessary if no viable tumor elements are detected in the surgical specimen. Patients with viable disease in a single node after undergoing systemic chemotherapy can be considered for a PLND, but the evidence supporting this approach is sparse. If 2 or more positive nodes or extranodal extension is detected, adjuvant chemotherapy (if not already given) and/or PLND is recommended. Recent data suggest that in the setting of 4 or more positive ILNs, a bilateral PLND should be performed.\(^ {106}\) Postoperative radiotherapy or chemoradiotherapy may be considered in patients after PLND, particularly in the setting of a positive surgical margin, viable cancer in multiple inguinal or PLNs, and/or presence of extranodal extension on the final pathologic specimen.

Unilateral Palpable Nodes ≥4 cm (fixed) or Bilateral Palpable Nodes (fixed or mobile)
In the case of large, unilateral, fixed nodes or bilateral ILNs, patients should undergo a needle biopsy of the lymph nodes. A negative result should be confirmed with excisional biopsy. If results are again negative, the patient should be closely followed. Patients with a positive aspiration or biopsy should receive neoadjuvant systemic chemotherapy followed by ILND and PLND. Postoperative radiotherapy or chemoradiotherapy may be considered. As previously mentioned, in the
Enlarged Pelvic Lymph Nodes
Patients with abnormal PLNs on imaging (CT or MRI) are stratified by resectability. Nonsurgical candidates should be treated with chemoradiotherapy. Patients with resectable disease should receive neoadjuvant systemic chemotherapy with consideration of a confirmatory percutaneous biopsy or PET/CT. Patients with disease that responds to therapy or that becomes stable should undergo bilateral superficial and deep ILND and unilateral/bilateral PLND if deemed resectable. Postoperative radiotherapy or chemoradiotherapy should be considered. Patients with disease that progresses may receive additional systemic chemotherapy with consideration of local-field radiation or participation in a clinical trial.

Surveillance
Initial treatment of the primary tumor and lymph nodes dictates the follow-up schedule (see algorithm). A large retrospective review of 700 patients found that penile-sparing therapies carry a significantly higher risk of local recurrence (28%) than partial or total penectomy (5%) and thus require closer surveillance. Patients without nodal involvement had a regional recurrence rate of 2% compared to 19% for patients with node-positive disease. Of all recurrences, 92% were detected within 5 years of primary treatment.

Follow-up for all patients includes a clinical exam of the penis and inguinal region. Imaging is not routinely indicated for early disease (except for obese patients or patients who have undergone inguinal surgery since a physical exam may be challenging), but may be used upon abnormal findings. For patients with N2 or N3 disease, imaging of the chest, abdomen, and pelvis area is recommended.

Recurrence
Invasion of the corpora cavernosa is an adverse finding after initial organ-sparing treatment that warrants partial or total penectomy. For primary tumor recurrences without corpora cavernosa infiltration, repeat penile-sparing options can be considered (category 2B).

A recurrence in the inguinal region carries a poor prognosis (median survival, <6 months) and optimal management remains elusive. Possible options include systemic chemotherapy followed by surgical resection, EBRT, or chemotherapy alone. A recent study suggests that ILND may be beneficial in patients with penile cancer with locally recurrent ILN metastases. While potentially curative, patients must be advised of the high incidence of postoperative complications.

Metastatic Disease
Imaging of the abdomen and pelvis should be obtained when metastasis is suspected to evaluate for pelvic and/or retroperitoneal lymph nodes. PLN metastasis is an ominous finding, with a 5-year survival rate of 0% to 66% for all cases and 17% to 54% for microscopic invasion only, with a mean 5-year survival of approximately 10%. In patients with ILN metastases, 20% to 30% will have PLN metastases. This can be further characterized such that if 2 to 3 ILNs are involved, there is a 23% probability of PLN involvement. With involvement of 3 or more ILNs, this probability increases to 56%.

Lughezzani et al identified three independent predictors of PLN metastases that included the number of inguinal metastases (odds ratio [OR], 1.92; \( P < .001 \)), the diameter of the metastases (OR, 1.03; \( P = .001 \)), and extranodal extension (OR, 8.01; \( P < .001 \)). Similar to previous studies, patients with 3 or more ILN metastases had a 4.77-fold higher risk of PLN metastasis. An ILN metastasis diameter of
30 mm or greater correlated with a 2.53-fold higher risk of PLN metastasis. Patients who showed no risk factors had a 0% risk of metastasis, suggesting that this group may not require PLND. Pettaway et al evaluated the treatment options for stage IV penile cancer—clinical stage N3 (deep inguinal nodes or pelvic nodes) or M1 disease (distant metastases)—including chemotherapy, radiotherapy, and inguinal lymphadenectomy. Cisplatin-based regimens (paclitaxel, ifosfamide, and cisplatin or alternatively 5-FU plus cisplatin) are the most active first-line systemic chemotherapy regimens. A retrospective analysis of 30 patients with non-metastatic N2 or N3 penile cancer who received neoadjuvant cisplatin-based chemotherapy as first-line treatment demonstrated a poor response to treatment when disease progressed (median OS < 6 months).

Systemic chemotherapy, radiotherapy, or chemoradiotherapy may be considered for the treatment of metastatic disease. The panel did not recommend regimens containing bleomycin because of high pulmonary-related toxicity. Patients with a proven objective response to systemic chemotherapy are amenable to consolidative ILND with curative potential or palliation. However, surgical consolidation should not be performed on patients with disease that progresses during systemic chemotherapy except for local symptomatic control. Preoperative radiotherapy may also be given to patients who have lymph nodes greater than or equal to 4 cm without skin fixation to improve surgical resectability and decrease local recurrence. For patients with unresectable inguinal or bone metastases, radiotherapy may provide a palliative benefit after chemotherapy. Systemic chemotherapy may also be considered upon disease progression. The NCCN Panel strongly recommends consideration of clinical trial participation as data are limited in the second-line setting. However, in select patients, paclitaxel or cetuximab may be considered, especially if previous treatments did not include a similar class of agent. Best supportive care remains an option for advanced cases or cases refractory to systemic chemotherapy, radiotherapy, or chemoradiotherapy.

Summary
SCC of the penis is a disease that mandates prompt medical/surgical intervention and patient compliance to obtain the most favorable outcomes. A thorough history and physical is the initial step in this process, followed by a biopsy of the primary lesion to establish a pathologic diagnosis. Accurate clinical staging allows for a comprehensive treatment approach to be devised, thus optimizing therapeutic efficacy and minimizing treatment-related morbidity. Prognostic factors help predict if lymph node metastases are suspected in the absence of any palpable inguinal lymphadenopathy. When clinically indicated, an ILND has curative potential, particularly when performed early, with contemporary surgical series demonstrating its reduced morbidity.
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