

高雄榮民總醫院

皮膚癌(BCC)診療原則

2016年03月21日 第一版

皮膚癌醫療團隊擬定

注意事項：這個診療原則主要作為醫師和其他保健專家診療癌症病人參考之用。假如你是一個癌症病人，直接引用這個診療原則並不恰當，只有你的醫師才能決定給你最恰當的治療。

修訂指引

- 本共識依下列參考資料修改版本
– NCCN 2016版 診療指引

BCC診療指引審視修訂會議討論日期

- 上次會議：2015/09/29
- 本共識經審視後與上一版之差異

上一版：

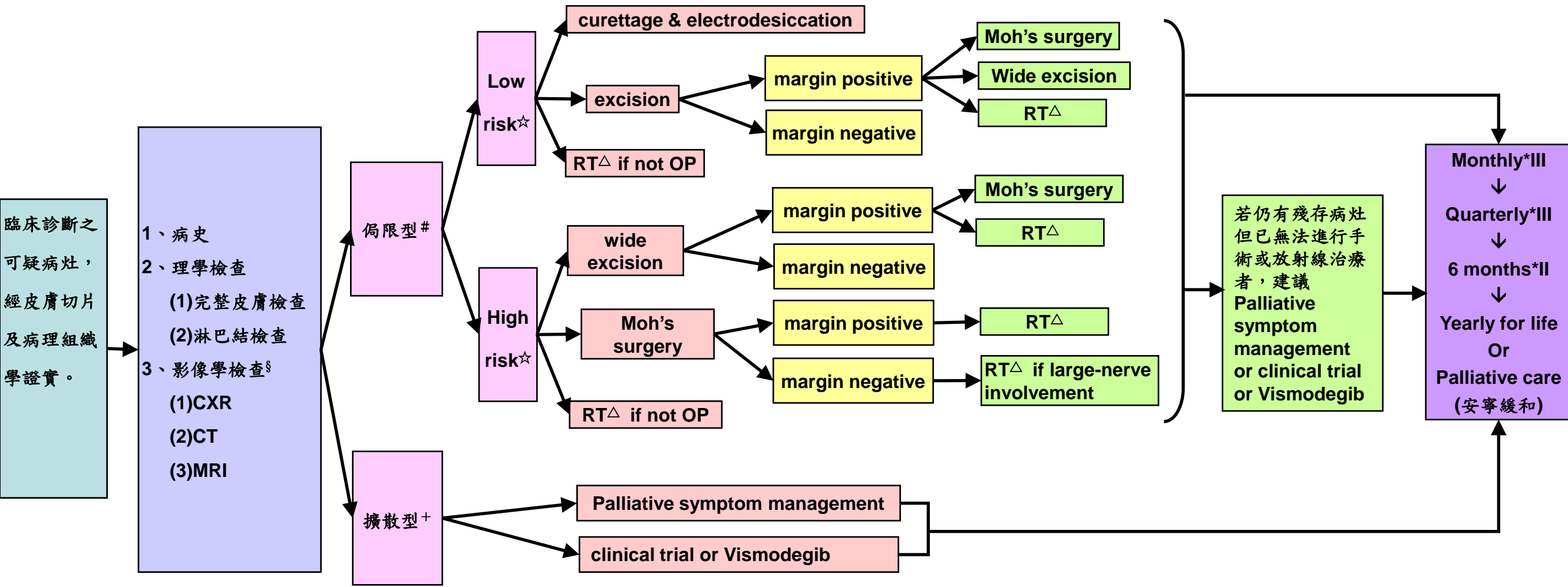
使用NCCN 2015版 診療指引

新版：

更新 NCCN 2016版 診療指引

基底細胞癌(BCC)

診斷	初步評估	分期	初始治療	療效評估	輔助治療	追蹤
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§ : Image studies is indicated for extensive disease (deep structural involvement such as bone, deep soft tissue, perineural disease)

: T any, N0, M0(附件三)

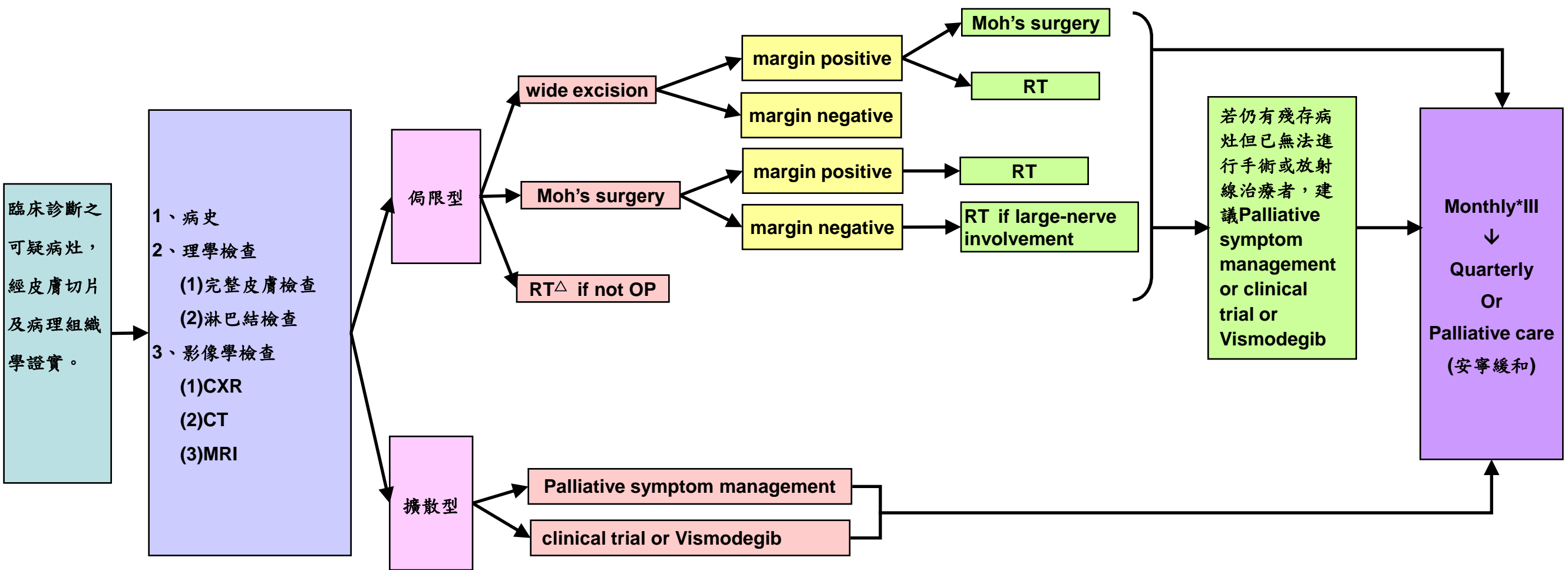
+ : regional or distal metastatic disease(初始皮膚病灶治療同局限型)

☆ : 附件一

△ : 附件二

基底細胞癌(BCC)

復發



基底細胞癌(BCC)

癌症藥物停藥準則

- 根據CTCAE (Common Terminology Criteria for Adverse Events, Version 4.0 Published: May 28, 2009 【v4.03: June 14, 2010】)，出現Grade 3 ~ Grade 4 adverse event。
- 停藥至adverse event回復至Grade 1或Baseline時可再次用藥，但有些患者必須調整用藥劑量。
- 使用BRAF inhibitor時可能產生cutaneous SCC。此現象雖被CTCAE列為Grade 3 toxic effect, 但此現象不必停藥或調整劑量
- 特定藥物治療下疾病仍持續進展，根據追蹤及評估顯示疾病對此特定藥物治療無效 (考慮停止投藥並選擇其他治療方法)。
- 病患要求 (Hospice care或其他因素)
- 病患死亡

基底細胞癌(BCC)

附件一：



NCCN Guidelines Version 1.2016 Basal Cell Skin Cancer

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RISK FACTORS FOR RECURRENCE

<u>H&P</u>	<u>Low Risk</u>	<u>High Risk</u>
Location/size	Area L <20 mm Area M <10 mm ¹ Area H <6 mm ¹	Area L ≥20 mm Area M ≥10 mm Area H ≥6 mm
Borders	Well defined	Poorly defined
Primary vs. Recurrent	Primary	Recurrent
Immunosuppression	(-)	(+)
Site of prior RT	(-)	(+)
<u>Pathology</u>		
Subtype	Nodular, superficial ²	Aggressive growth pattern ³
Perineural involvement	(-)	(+)

Area H = "mask areas" of face (central face, eyelids, eyebrows, periorbital, nose, lips [cutaneous and vermilion], chin, mandible, preauricular and postauricular skin/sulci, temple, ear), genitalia, hands, and feet.
Area M = cheeks, forehead, scalp, neck, and pretibia.
Area L = trunk and extremities (excluding pretibia, hands, feet, nail units, and ankles).

¹Location independent of size may constitute high risk.
²Low risk histologic subtypes include nodular, superficial and other non-aggressive growth patterns such as keratotic, infundibulocystic, and fibroepithelioma of Pinkus.
³Having morpheaform, basosquamous (metatypical), sclerosing, mixed infiltrative, or micronodular features in any portion of the tumor. In some cases basosquamous (metatypical) tumors may be prognostically similar to SCC. Clinicopathologic consultation is recommended.

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.

基底細胞癌(BCC)

附件二：



NCCN Guidelines Version 1.2016 Basal Cell Skin Cancer

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PRINCIPLES OF RADIATION THERAPY FOR BASAL CELL SKIN CANCER

<u>Dose and Field Size</u>		
<u>Tumor Diameter</u>	<u>Margins</u>	<u>Examples of Electron Beam Dose and Fractionation</u>
<2 cm	1–1.5 cm ¹	64 Gy in 32 fractions over 6–6.4 weeks ² 55 Gy in 20 fractions over 4 weeks 50 Gy in 15 fractions over 3 weeks 35 Gy in 5 fractions over 5 days
≥2 cm	1.5–2 cm ¹	66 Gy in 33 fractions over 6–6.6 weeks 55 Gy in 20 fractions over 4 weeks
Postoperative adjuvant		50 Gy in 20 fractions over 4 weeks 60 Gy in 30 fractions over 6 weeks

- Protracted fractionation is associated with improved cosmetic results.
- Radiation therapy is contraindicated in genetic conditions predisposing to skin cancer (eg, basal cell nevus syndrome, xeroderma pigmentosum) and connective tissue diseases (eg, scleroderma)

¹When using electron beam, wider field margins are necessary than with orthovoltage x-rays due to the wider beam penumbra. Tighter field margins can be used with electron beam adjacent to critical structures (eg, the orbit) if lead skin collimation is used. Bolus is necessary when using electron beam to achieve adequate surface dose. An electron beam energy should be chosen which achieves adequate surface dose and encompasses the deep margin of the tumor by at least the distal 90% line. Appropriate medical physics support is essential.

²Electron beam doses are specified at 90% of the maximal depth dose (Dmax). Orthovoltage x-ray doses are specified at Dmax (skin surface) to account for the relative biologic difference between the two modalities of radiation.

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.

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附件三-1:



NCCN Guidelines Version 1.2014 Basal and Squamous Cell Skin Cancers

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Staging

Table 1 American Joint Committee on Cancer (AJCC) TNM Staging Classification for Cutaneous Squamous Cell Carcinoma (cSCC) and Other Cutaneous Carcinomas (7th ed., 2010)		Regional Lymph Nodes (N)	
Primary Tumor (T)*		NX	Regional lymph nodes cannot be assessed
TX	Primary tumor cannot be assessed	N0	No regional lymph node metastases
T0	No evidence of primary tumor	N1	Metastasis in a single ipsilateral lymph node, 3 cm or less in greatest dimension
Tis	Carcinoma in situ	N2	Metastasis in a single ipsilateral lymph node, more than 3 cm but not more than 6 cm in greatest dimension; or in multiple ipsilateral lymph nodes, none more than 6 cm in greatest dimension; or in bilateral or contralateral lymph nodes, none more than 6 cm in greatest dimension
T1	Tumor 2 cm or less in greatest dimension with less than two high-risk features**	N2a	Metastasis in a single ipsilateral lymph node, more than 3 cm but not more than 6 cm in greatest dimension
T2	Tumor greater than 2 cm in greatest dimension or Tumor any size with two or more high-risk feature	N2b	Metastasis in multiple ipsilateral lymph nodes, none more than 6 cm in greatest dimension
T3	Tumor with invasion of maxilla, mandible, orbit, or temporal bone	N2c	Metastasis in bilateral or contralateral lymph nodes, none more than 6 cm in greatest dimension
T4	Tumor with invasion of skeleton (axial or appendicular) or perineural invasion of skull base	N3	Metastasis in a lymph node, more than 6 cm in greatest dimension
*Excludes cSCC of the eyelid		Distant Metastasis (M)	
** High-risk features for the primary tumor (T) staging		M0	No distant metastases
Depth/invasion	> 2 mm thickness Clark level ≥ IV Perineural invasion	M1	Distant metastases
Anatomic location	Primary site ear Primary site non-hair-bearing lip		
Differentiation	Poorly differentiated or undifferentiated		

[Continue](#)

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基底細胞癌(BCC)

附件三-2:



NCCN Guidelines Version 1.2014 Basal and Squamous Cell Skin Cancers

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Table 1 Continued

American Joint Committee on Cancer (AJCC)
TNM Staging Classification for Cutaneous Squamous Cell Carcinoma (cSCC) and Other Cutaneous Carcinomas (7th ed., 2010)

Anatomic Stage/Prognostic Groups

Stage 0	Tis	N0	M0
Stage I	T1	N0	M0
Stage II	T2	N0	M0
Stage III	T3	N0	M0
	T1	N1	M0
	T2	N1	M0
Stage IV	T3	N1	M0
	T1	N2	M0
	T2	N2	M0
	T3	N2	M0
	T Any	N3	M0
	T4	N Any	M0
	T Any	N Any	M1

Histologic Grade (G)

GX	Grade cannot be assessed
G1	Well differentiated
G2	Moderately differentiated
G3	Poorly differentiated
G4	Undifferentiated

Used with the permission of the American Joint Committee on Cancer (AJCC), Chicago, Illinois. The original and primary source for this information is the AJCC Cancer Staging Manual, Seventh Edition (2010) published by Springer Science and Business Media LLC (SBM). (For complete information and data supporting the staging tables, visit www.springer.com.) Any citation or quotation of this material must be credited to the AJCC as its primary source. The inclusion of this information herein does not authorize any reuse or further distribution without the expressed, written permission of Springer SBM, on behalf of the AJCC.

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