



National Comprehensive
Cancer Network®

NCCN Clinical Practice Guidelines in Oncology (NCCN Guidelines®)

Basal Cell Skin Cancer

Version 3.2024 — March 1, 2024

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Clinical Trials: NCCN believes that the best management for any patient with cancer is in a clinical trial. Participation in clinical trials is especially encouraged.

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<https://www.nccn.org/home/member-institutions>.

NCCN Categories of Evidence and Consensus: All recommendations are category 2A unless otherwise indicated.

See [NCCN Categories of Evidence and Consensus](#).

The NCCN Guidelines® are a statement of evidence and consensus of the authors regarding their views of currently accepted approaches to treatment. Any clinician seeking to apply or consult the NCCN Guidelines is expected to use independent medical judgment in the context of individual clinical circumstances to determine any patient's care or treatment. The National Comprehensive Cancer Network® (NCCN®) makes no representations or warranties of any kind regarding their content, use or application and disclaims any responsibility for their application or use in any way. The NCCN Guidelines are copyrighted by National Comprehensive Cancer Network®. All rights reserved. The NCCN Guidelines and the illustrations herein may not be reproduced in any form without the express written permission of NCCN. ©2024.



Terminologies in all NCCN Guidelines are being actively modified to advance the goals of equity, inclusion, and representation.

Updates in Version 3.2024 of the NCCN Guidelines for Basal Cell Skin Cancer from Version 2.2024 include:

[BCC-3](#)

- Primary Treatment, top option revised: Mohs or other forms of PDEMA (preferred for *BCCs that are either recurrent, ≥1 cm; in H zone, recurrent, or ≥1 cm with an aggressive histologic subtype*).

September 14, 2023 - Category correction for Version 1.2024[BCC-2](#)

- Primary Treatment, Non-surgical modalities for tumors clinically and histologically consistent with superficial BCC (without dermal extension):
 - ▶ Third bullet, Photodynamic therapy, porfimer sodium was revised from category 2A to *category 2B*.

Updates in Version 2.2024 of the NCCN Guidelines for Basal Cell Skin Cancer from Version 1.2024 include:

[BCC-1](#)

- Footnote g revised: Imaging modality and targeted area should be at the discretion of the treating team based on the suspected extent of disease (ie, local, regional, metastatic). Histologic confirmation is ~~often~~ sufficient to diagnose local recurrence, but MRI *with and without contrast* can be considered to assess extent of local disease. . .

[BCC-2](#)

- Primary Treatment, Non-surgical modalities for tumors clinically and histologically consistent with superficial BCC (without dermal extension), third bullet revised: Photodynamic therapy (eg, *topical* aminolevulinic acid [ALA], porfimer sodium [category 2B]) (Useful in Certain Circumstances).

[BCC-C](#)

- Second bullet revised: Surgical approaches often offer the most effective and efficient means for accomplishing cure, but considerations of function, *cosmesis*, and patient preference may lead to choosing RT/topical therapy/systemic therapy as primary treatment in order to achieve ~~satisfactory~~ *optimal* overall results.
- Fourth bullet revised: In patients with superficial basal cell skin cancer, ~~non-surgical modalities therapies such as topical imiquimod, topical 5-FU, photodynamic therapy (eg, ALA, porfimer sodium), or cryotherapy~~ may be considered. (See *BCC-2*)

[BCC-E 1 of 2](#)

- None added to Preferred Regimens for Locally Advanced Disease - Neoadjuvant, Locally Advanced Disease, Nodal Disease, and Metastatic Disease.

[MS-1](#)

- The Discussion section has been updated to reflect the changes in the algorithm.

Updates in Version 1.2024 of the NCCN Guidelines for Basal Cell Skin Cancer from Version 1.2023 include:

Global changes:

- Extracapsular extension (ECE) changed to extranodal extension (ENE).

[BCC-1](#)

- Preliminary Workup, third bullet revised: Shave ~~excision-removal~~ if applicable.
- Risk Status, new option added: Locally advanced disease.
- New header added: Staging.
 - ▶ Following High risk, option revised: Consider imaging if clinical exam insufficient for ~~following~~ *determining* disease extent.
 - ▶ Following Locally advanced disease, option added: Consider imaging to determine disease extent.

[Continued](#)**UPDATES**



Updates in Version 1.2024 of the NCCN Guidelines for Basal Cell Skin Cancer from Version 1.2023 include:

[BCC-1](#) (continued)

- New header added: Primary Treatment
 - ▶ Following Low risk, option revised: ~~Primary Treatment of Low Risk BCC (BCC-2)~~.
 - ▶ Following High risk, Consider imaging if clinical exam insufficient for determining disease extent, option revised: ~~Primary Treatment of High Risk BCC (BCC-3)~~.
 - ▶ Following Locally advanced disease, Consider imaging to determine disease extent, option added: BCC-4.
 - ▶ Following Initial presentation of regional or distant metastatic disease, Imaging studies of areas of interest as indicated for suspicion of extensive disease, option revised: ~~Treatment for Recurrence or Advanced Disease (BCC-4)~~.
- New footnote c added: Morgan FC, et al. J Am Acad Dermatol 2021;85:582-587.
- Footnotes revised:
 - ▶ Footnote d: Extensive disease includes deep involvement such as bone, named nerves, and deep soft tissue. If disease of named nerve(s) is suspected, MRI with *and without* contrast is preferred. If bone disease is suspected, CT with contrast is preferred unless contraindicated.
 - ▶ Footnote f: For rare cases that present with regional or distant metastatic disease at diagnosis, treat ~~as per nodal or distant metastases~~ *metastatic* pathways on BCC-4.
- Footnote removed: For more information, See American Academy of Dermatology Association.

[BCC-2](#)

- Primary Treatment, options revised: Curettage and Electrodesiccation (C&E) ~~or shave excision removal~~ or *Shave removal (if tumor appears to extend beyond the dermis, surgical ~~or shave~~ excision should generally be performed rather than C&E or shave removal) or Standard excision with 4-mm clinical margins and postoperative margin assessment. Tissue rearrangement (eg, flap reconstruction, extensive undermining) should not be undertaken until clear margins are identified (second intention healing, linear repair, or skin graft are acceptable) or Radiation therapy (RT) for non-surgical candidates or Topical therapy *Non-surgical modalities for tumors clinically and histologically consistent with superficial BCC (without dermal extension): Topical 5-fluorouracil (5-FU) (Useful in Certain Circumstances), Topical imiquimod (preferred), Photodynamic therapy (eg, aminolevulinic acid [ALA], porfimer sodium) (Useful in Certain Circumstances), Cryotherapy.**
- Footnotes revised:
 - ▶ Footnote l: ~~In patients with superficial basal cell skin cancer, therapies such as topical imiquimod, topical 5-fluorouracil, photodynamic therapy, or cryotherapy may be considered, although cCure rates are approximately 10% lower than for surgical treatment modalities. Jansen MHE, et al. J Invest Dermatol 2018;138:527-533. Drew BA, et al. Dermatol Surg 2017;43:1423-1430.~~
 - ▶ Footnote o: As per other appropriate use criteria. Task Force/Committee Members, Vidal CI, Ambrecht EA, Andrea AA, et al. J Am Acad Dermatol 2019;80:189-207. ~~e44~~. (Also pages BCC-3A and BCC-4)
 - ▶ Footnote p: PDEMA with permanent section analysis or intraoperative frozen section analysis is an alternative to Mohs. See Principles of PDEMA Technique (SCC-G) *within the NCCN Guidelines for Squamous Cell Skin Cancer.* (Also pages BCC-3A and BCC-4)
- New footnotes added:
 - ▶ Footnote i: Shave removal (shaving of epidermal or dermal lesion) is a sharp removal by transverse bowl-shaped slicing to remove epidermal and dermal lesions (without including fat) and does not require suture closure. Emmett AJ and Bradbent GD. Plast Reconstr Surg. 1987;80:47-54. Abramson AK, et al. Dermatol Surg. 2013;39:387-392. Wu X, et al. J Am Acad Dermatol. 2015;73:791-798. Dando EE, et al. Dermatol Surg. 2023;49:130-134.
 - ▶ Footnote k: Determination of the appropriateness of RT should be performed by a radiation oncologist. (Also pages BCC-3A and BCC-4)
 - ▶ Footnote m: Afsar FS, et al. Postepy Dermatol Alergol 2015;32:88-93.

[Continued](#)**UPDATES**



Updates in Version 1.2024 of the NCCN Guidelines for Basal Cell Skin Cancer from Version 1.2023 include:

[BCC-3](#)

- Primary Treatment, options revised: Mohs or other forms of PDEMA (*preferred for ≥ 1 cm, H zone, recurrent, or aggressive histologic subtype*) or Standard excision with wider surgical margins and postoperative margin assessment and second intention healing, linear repair, or skin graft or For non-surgical candidates: *consider multidisciplinary consultation and discussion of definitive RT* ~~RT, Systemic therapy if curative RT is not feasible~~ ~~or For patients in whom surgery may cause significant functional damage, neoadjuvant administration of vismodegib followed by PDEMA may be considered (category 2B).~~
- Additional Treatment:
 - ▶ Top column following Positive margins, third bullet revised: ~~Systemic therapy iff curative~~ *If surgery and/or curative RT are not curative.*
 - ◊ Link to Advanced BCC (BCC-4) added to far right.
 - ▶ Bottom Positive margins pathway, following "Standard excision," removed: Positive margins, Mohs or other forms of PDEMA, if feasible or Standard re-excision if PDEMA is not feasible; If residual disease is present, and further surgery is not feasible, consider multidisciplinary consultation to discuss options: RT, Systemic therapy if curative RT is not feasible.
- Footnotes revised:
 - ▶ Footnote q: For clinically diagnosed non-facial BCCs <6 mm in depth on the head, neck, hands, feet, pretibial, and anogenital *area* that are clinically confined to the dermis, C&E or shave ~~excision~~ *removal* may be considered as an alternative primary treatment option if Mohs, resection with PDEMA, and standard excision are difficult to perform due to patient comorbidities (eg, thrombocytopenia, immunosuppression, bleeding diathesis, multiple primary BCCs). See Risk Factors for Recurrence (BCC-B). (Also page BCC-4)
 - ▶ Footnote t: If named nerve involvement is suspected, consider MRI with *and without* contrast of region of interest to evaluate extent and rule out base of skull involvement or intracranial extension in head and neck tumors.
- New footnote r added: Aggressive histologic subtype is defined as: BCC with squamous differentiation, infiltrative, micronodular, morpheaform, sclerodermiform, or sclerosing. van Loo E, et al. Eur J Cancer 2014;50:3011-3020. Fraga SD, et al. Dermatol Surg 2022;48:704-710.
- Footnotes removed:
 - ▶ For locally advanced disease (extensive disease where surgery and/or RT are unlikely to result in a cure or are not feasible), consider multidisciplinary consultation and treatment with hedgehog pathway inhibitors (HHIs) (vismodegib and sonidegib) or programmed cell death protein 1 (PD-1) inhibitor (cemiplimab-rwlc) for patients previously treated with an HHI or for whom an HHI is not appropriate. Feasibility of surgery or radiation should be assessed by a surgeon and radiation oncologist. Principles of Systemic Therapy (BCC-E).
 - ▶ Principles of Systemic Therapy (BCC-E). (Also page BCC-5)
 - ▶ In one study of 55 patients with locally advanced basal cell carcinoma, neoadjuvant administration of vismodegib before planned surgery allowed for a smaller surgical procedure in 71% of patients, although it carried a high (36.4%) recurrence risk. Bertrand N, et al. E Clinical Medicine 2021;35:100844.

[Continued](#)

UPDATES



Updates in Version 1.2024 of the NCCN Guidelines for Basal Cell Skin Cancer from Version 1.2023 include:

[BCC-4](#)

- New page for Advanced BCC.
- New branch points added to left side of the page:
 - ▶ Locally advanced BCC (laBCC) (primary or recurrent extensive disease where surgery and/or RT may not result in a cure or would possibly produce a significant functional limitation).
 - ◊ Primary Treatment, new options added: Multidisciplinary consultation to consider one or more of the following options: Surgery, Consider neoadjuvant systemic therapy (BCC-E), Mohs or other forms of PDEMA, Standard excision with vertical histologic sectioning (if Mohs or PDEMA are not available) or RT or If surgery and/or RT are not feasible then systemic therapy (BCC-E).
 - ▶ Nodal disease.
 - ◊ Primary Treatment, options added: Multidisciplinary consultation to consider one or more of the following options: Surgery or If surgery is not feasible then RT or systemic therapy (BCC-E), or Clinical trial.
 - ▶ Metastatic disease.
 - ◊ Primary treatment, options added: Multidisciplinary consultation to consider: Systemic therapy (BCC-E) or RT or surgery for limited metastatic disease or Palliation and best supportive care.
- New footnote v added: Fraga SD, et al. *Dermatol Surg* 2022;704-710.

[BCC-5](#)

- Top header revised: ~~Recurrence or Advanced Disease.~~
- ▶ First pathway revised: ~~Local recurrence.~~
- ▶ Second pathway revised: ~~Primary or recurrent Advanced disease: Locally advanced, Nodal metastases, Distant metastases.~~
 - ◊ Options revised: ~~Multidisciplinary consultation to consider one or more of the following options: Surgery or If surgery is not feasible then RT or systemic therapy, Hedgehog pathway inhibitor (HHI), Vismodegib, Sonidegib (category 2B), Programmed cell death protein 1 (PD-1) inhibitor (cemiplimab-rwlc), Clinical trial-Follow Primary Treatment pathways for Advanced BCC (BCC-4).~~
- ▶ Pathway removed: Distant metastases.
 - ◊ Options removed: Multidisciplinary consultation to consider: Systemic therapy, HHI, Vismodegib, PD-1 inhibitor (cemiplimab-rwlc) or RT or surgery for limited metastatic disease or Palliation and best supportive care.
- Footnotes revised:
 - ▶ Footnote x: Follow-up with a dermatologist is strongly recommended if any of the following criteria are met: past or imminent solid organ, marrow, or ~~stem-hematopoietic cell transplant~~; one or more cutaneous melanomas in the past 5 years; or four or more non-melanoma skin cancers in the past 5 years.
 - ▶ Footnote y: Imaging modality and targeted area should be at the discretion of the treating team based on the suspected extent of disease (ie, local, regional, metastatic). Histologic confirmation is ~~often~~ sufficient to diagnose local recurrence, but MRI can be considered to assess extent of local disease. . .
- Footnotes removed:
 - ▶ Principles of Radiation Therapy (BCC-D).
 - ▶ Cemiplimab-rwlc is recommended for patients with locally advanced or metastatic basal cell carcinoma (mBCC) previously treated with an HHI or for whom an HHI is not appropriate.
 - ▶ Under highly selective circumstances, in the context of multidisciplinary consultation, resection of limited metastases can be considered.

[Continued](#)

UPDATES



Updates in Version 1.2024 of the NCCN Guidelines for Basal Cell Skin Cancer from Version 1.2023 include:

[BCC-B](#)

- Location/size, second line under High Risk revised: Head, neck, hands, feet, pretibial, and anogenital *area* (any size).

[BCC-C](#)

- Third bullet revised: In certain patients at high risk for multiple primary tumors (eg, basal cell nevus syndrome [Gorlin syndrome], xeroderma pigmentosum, history of RT), increased surveillance and consideration of prophylactic measures may be indicated. ~~Consider~~ **Referring** patients with suspected basal cell nevus syndrome (~~Gorlin syndrome~~) or xeroderma pigmentosum for genetic evaluation.
- Fourth bullet revised: In patients with superficial basal cell skin cancer, therapies such as topical imiquimod, topical 5-fluorouracil 5-FU, photodynamic therapy (eg, aminolevulinic acid [ALA], porfimer sodium), or cryotherapy may be considered, ~~even though the cure rates may be lower than with surgical treatment modalities.~~
- New footnote a added: Cure rates are approximately 10% lower than for surgical treatment modalities. Jansen MHE, Mosterd K, Arits AHMM, et al. Five-year results of a randomized controlled trial comparing effectiveness of photodynamic therapy, topical imiquimod, and topical 5-fluorouracil in patients with superficial basal cell carcinoma. *J Invest Dermatol* 2018;138:527-533. Drew BA, Karia PS, Mora AN, et al. Treatment patterns, outcomes, and patient satisfaction of primary epidermally limited nonmelanoma skin cancer. *Dermatol Surg* 2017;43:1423-1430.

[BCC-D](#)

- General Principles, second bullet revised: RT is contraindicated for genetic conditions predisposing to skin cancer (eg, basal cell nevus syndrome [~~Gorlin syndrome~~]) and relatively contraindicated for patients with connective tissue diseases (eg, scleroderma).
- Regional Disease, Lymph node regions, without lymph node dissection, option removed: Clinically negative, at risk.
 - ▶ RT dosing, option removed: 50 Gy over 5 to 7 weeks.

[BCC-E 1 of 2](#)

- Header revised: Locally Advanced (laBCC), *Nodal* or *Distant* Metastatic Basal Cell Carcinoma (mBCC).
 - ▶ First bullet revised: Systemic therapy may be considered for laBCC ~~and mBCC~~. Locally advanced disease is defined by those that have primary or recurrent extensive disease where surgery and/or RT ~~are unlikely to result in a cure~~ **may not result in a cure or would possibly produce a significant functional limitation.**
 - ▶ Fourth bullet revised: ~~Systemic therapy options include:~~ **Hedgehog pathway inhibitors (HHIs).**
 - ◇ Sub-bullets removed:
 - Hedgehog pathway inhibitors (HHIs) (ie, vismodegib, sonidegib).
 - Cemiplimab-rwlc is recommended for patients with laBCC or mBCC previously treated with an HHI or for whom an HHI is not appropriate.
 - ◇ Second tertiary-bullet moved out to be a sub-bullet and revised: HHIs may be considered for diffuse BCC formation (eg, basal cell nevus syndrome [~~Gorlin syndrome~~] or other genetic forms of multiple BCC). HHIs are not FDA approved for basal cell nevus syndrome (~~Gorlin syndrome~~); however, they may be used off-label and are effective based on a randomized controlled trial.
 - ◇ Tertiary bullet removed: Current FDA-approved HHIs include vismodegib and sonidegib.1 Vismodegib is FDA approved for the treatment of adults with mBCC or laBCC that has recurred following surgery, or those who are not candidates for surgery or RT. Sonidegib1 is FDA approved for the treatment of adults with laBCC that has recurred following surgery or RT, or those who are not candidates for surgery or RT. Sonidegib is not FDA approved for the treatment of adults with mBCC.

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UPDATES



Updates in Version 1.2024 of the NCCN Guidelines for Basal Cell Skin Cancer from Version 1.2023 include:

[BCC-E 1 of 2](#) (continued)

- New preference stratification table added with information from BCC-3, BCC-3A, BCC-5 and old BCC-E:
 - ▶ Locally Advanced Disease - Neoadjuvant.
 - ◊ Other Recommended Regimens, Vismodegib (category 2B).
 - ◊ Useful in Certain Circumstances, new regimen added: Cemiplimab-rwlc (category 2B).
 - ▶ Locally Advanced Disease:
 - ◊ Other Recommended Regimens:
 - Sonidegib
 - Vismodegib
 - ◊ Useful in Certain Circumstances, Cemiplimab-rwlc.
 - ▶ Nodal Disease:
 - ◊ Other Recommended Regimens:
 - Vismodegib
 - Sonidegib (category 2B)
 - ◊ Useful in Certain Circumstances, Cemiplimab-rwlc.
 - ▶ Metastatic Disease
 - ◊ Other Recommended Regimens, Vismodegib.
 - ◊ Useful in Certain Circumstances, Cemiplimab-rwlc.
- New footnotes added:
 - ▶ Footnote a: In one study of 55 patients with laBCC, neoadjuvant administration of vismodegib before planned surgery allowed for a smaller surgical procedure in 71% of patients, although it carried a high (36.4%) recurrence risk.
 - ▶ Footnote b: Cemiplimab-rwlc is FDA approved for patients with laBCC or mBCC previously treated with an HHI or for whom an HHI is not appropriate.
 - ▶ Footnote c: A multinational single-arm phase 2 trial, consisting of 84 patients with laBCC (local invasion precluding complete resection or in locations for which surgery may result in severe disfigurement or dysfunction) whose disease had progressed on or was intolerant to prior HHI therapy, was conducted. Thirty-one percent had an objective response, including 6% with a complete response. See Discussion.

[BCC-E 2 of 2](#)

- New references added:
 - ▶ Bertrand N, Guerreschi P, Basset-Seguín N, et al. Vismodegib in neoadjuvant treatment of locally advanced basal cell carcinoma: First results of a multicenter, open-label, phase 2 trial (VISMONEO study): Neoadjuvant Vismodegib in Locally Advanced Basal Cell Carcinoma. *EClinicalMedicine* 2021;35:100844.
 - ▶ Sekulic A, Migden MR, Oro AE, et al. Efficacy and safety of vismodegib in advanced basal-cell carcinoma. *N Engl J Med* 2012;366:2171-2179.
 - ▶ Dreno B, Basset-Seguín N, Caro I, Yue H, Schadendorf D. Clinical benefit assessment of vismodegib therapy in patients with advanced basal cell carcinoma. *Oncologist* 2014;19:790-796.
 - ▶ Stratigos AJ, Sekulic A, Peris K, et al. Cemiplimab in locally advanced basal cell carcinoma after hedgehog inhibitor therapy: an open-label, multi-centre, single-arm, phase 2 trial. *Lancet Oncol* 2021;22:848-857.

[BCC-F](#)

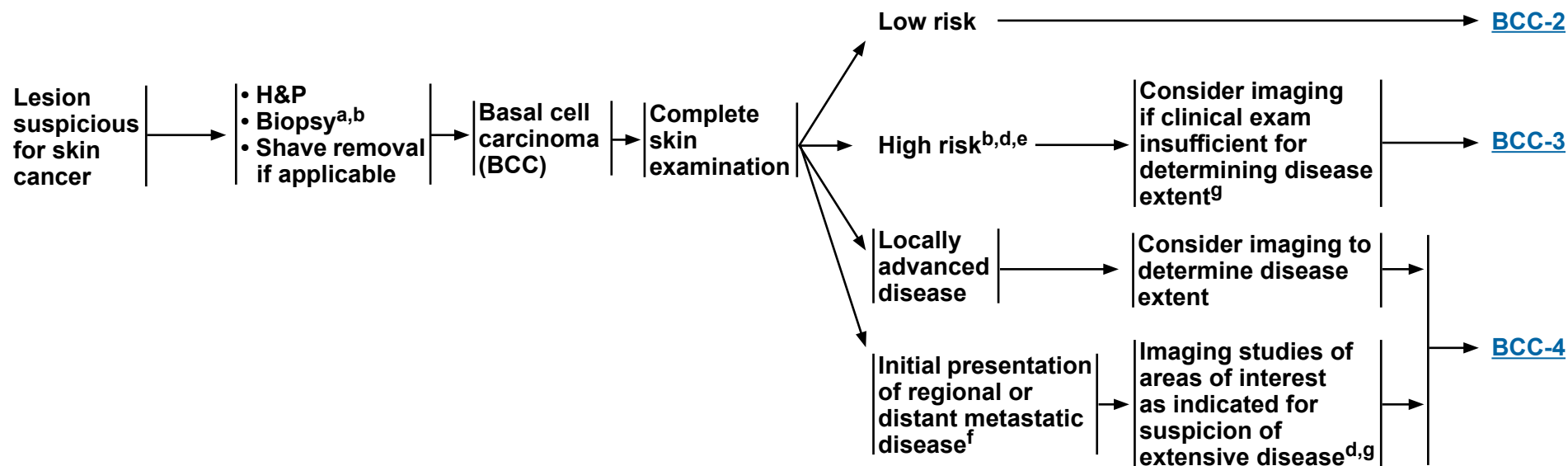
- Second bullet revised: In certain patients at high risk for multiple primary tumors (eg, basal cell nevus syndrome [~~Gorlin syndrome~~], xeroderma pigmentosum, history of RT), increased surveillance and consideration of prophylactic measures may be indicated. . .



NCCN Guidelines Version 3.2024

Basal Cell Skin Cancer

CLINICAL PRESENTATION	PRELIMINARY WORKUP	DIAGNOSIS	ADDITIONAL WORKUP	RISK STATUS ^c	STAGING	PRIMARY TREATMENT
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^a [Principles of Pathology \(BCC-A\)](#).

^b [Risk Factors for Recurrence \(BCC-B\)](#).

^c Morgan FC, et al. J Am Acad Dermatol 2021;85:582-587.

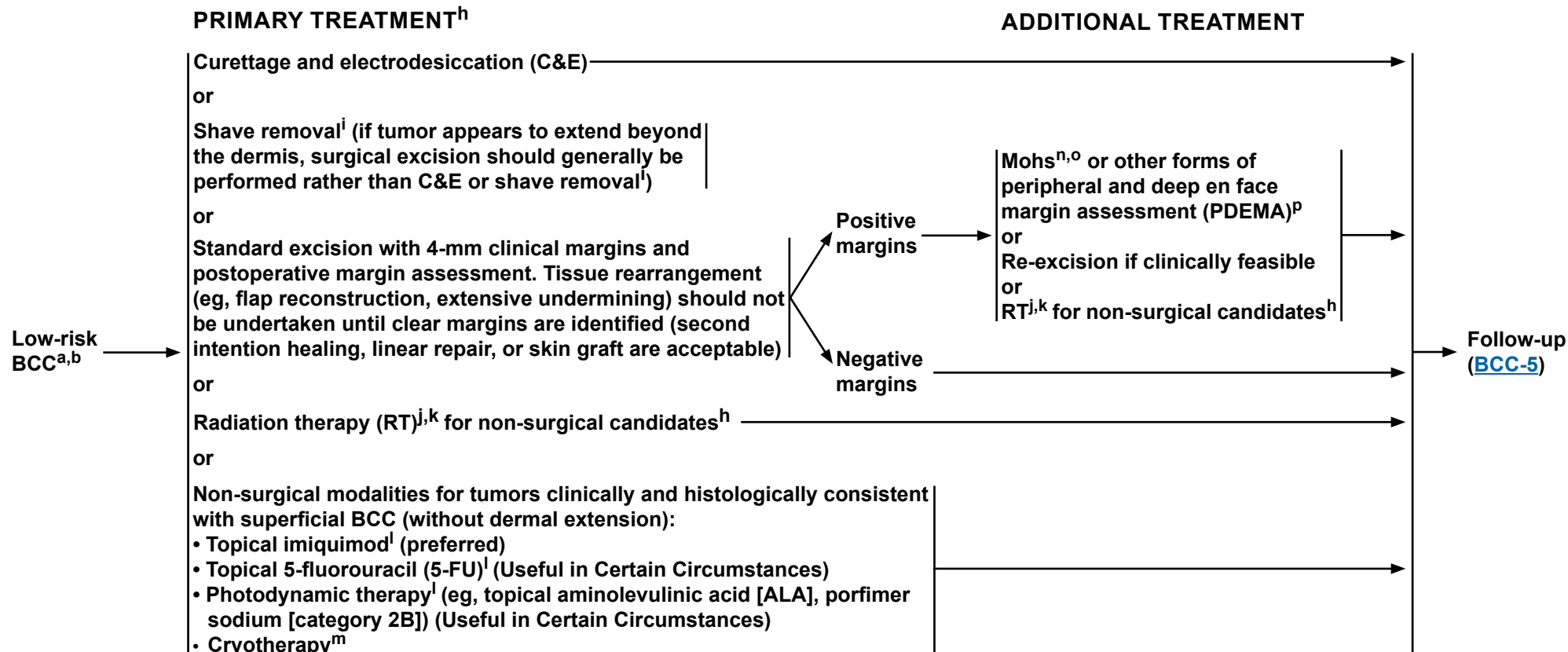
^d Extensive disease includes deep involvement such as bone, named nerves, and deep soft tissue. If disease of named nerve(s) is suspected, MRI with and without contrast is preferred. If bone disease is suspected, CT with contrast is preferred unless contraindicated.

^e Any high-risk factor places the patient in the high-risk category.

^f For rare cases that present with regional or distant metastatic disease at diagnosis, treat per nodal or metastatic pathways on [BCC-4](#).

^g Imaging modality and targeted area should be at the discretion of the treating team based on the suspected extent of disease (ie, local, regional, metastatic). Histologic confirmation is sufficient to diagnose local recurrence, but MRI with and without contrast can be considered to assess extent of local disease. For nodal or distant metastasis, histologic analysis and/or CT imaging can be used for confirmation and to gauge extent of disease.

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



^a [Principles of Pathology \(BCC-A\)](#).

^b [Risk Factors for Recurrence \(BCC-B\)](#).

^h [Principles of Treatment \(BCC-C\)](#).

ⁱ Shave removal (shaving of epidermal or dermal lesion) is a sharp removal by transverse bowl-shaped slicing to remove epidermal and dermal lesions (without including fat) and does not require suture closure. Emmett AJ and Bradbent GD. *Plast Reconstr Surg*. 1987;80:47-54. Abramson AK, et al. *Dermatol Surg*. 2013;39:387-392. Wu X, et al. *J Am Acad Dermatol*. 2015;73:791-798. Dando EE, et al. *Dermatol Surg*. 2023;49:130-134.

^j [Principles of Radiation Therapy \(BCC-D\)](#).

^k Determination of the appropriateness of RT should be performed by a radiation oncologist.

^l Cure rates are approximately 10% lower than for surgical treatment modalities. Jansen MHE, et al. *J Invest Dermatol* 2018;138:527-533. Drew BA, et al. *Dermatol Surg* 2017;43:1423-1430.

^m Afsar FS, et al. *Postepy Dermatol Alergol* 2015;32:88-93.

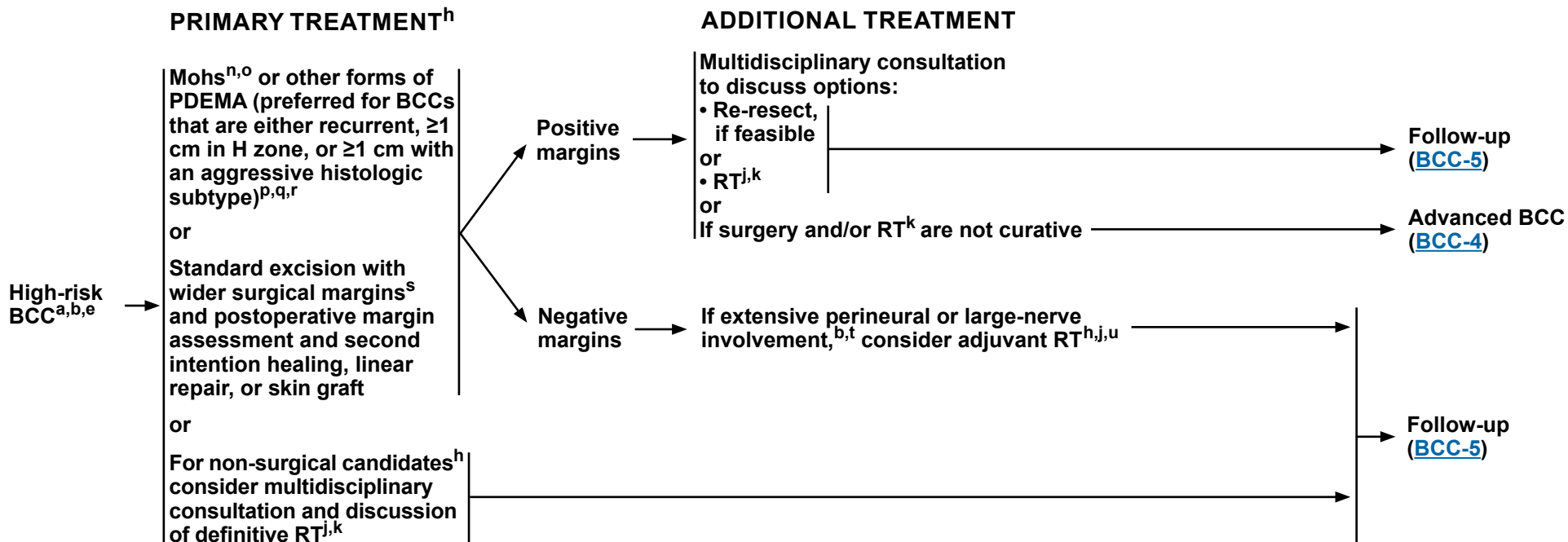
ⁿ Mohs surgery should be performed by dermatologic surgeons who have specialized training and experience in this procedure.

^o As per other appropriate use criteria. Task Force/Committee Members, Vidal CI, et al. *J Am Acad Dermatol* 2019;80:189-207.

^p PDEMA with permanent section analysis or intraoperative frozen section analysis is an alternative to Mohs. See Principles of PDEMA Technique (SCC-G) within the [NCCN Guidelines for Squamous Cell Skin Cancer](#).

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

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^a [Principles of Pathology \(BCC-A\)](#).

^b [Risk Factors for Recurrence \(BCC-B\)](#).

^e Any high-risk factor places the patient in the high-risk category.

^h [Principles of Treatment \(BCC-C\)](#).

^j [Principles of Radiation Therapy \(BCC-D\)](#).

^k Determination of the appropriateness of RT should be performed by a radiation oncologist.

ⁿ Mohs surgery should be performed by dermatologic surgeons who have specialized training and experience in this procedure.

^o As per other appropriate use criteria. Task Force/Committee Members, Vidal CI, et al. *J Am Acad Dermatol* 2019;80:189-207.

^p PDEMA with permanent section analysis or intraoperative frozen section analysis is an alternative to Mohs. See Principles of PDEMA Technique (SCC-G) within the [NCCN Guidelines for Squamous Cell Skin Cancer](#).

^q For clinically diagnosed non-facial BCCs <6 mm in depth on the head, neck, hands, feet, pretibial, and anogenital area that are clinically confined to the dermis, C&E or shave removal may be considered as an alternative primary treatment option if Mohs, resection with PDEMA, and standard excision are difficult to perform due to patient comorbidities (eg, thrombocytopenia, immunosuppression, bleeding diathesis, multiple primary BCCs). See [Risk Factors for Recurrence \(BCC-B\)](#).

^r Aggressive histologic subtype is defined as: BCC with squamous differentiation, infiltrative, micronodular, morpheaform, sclerodermiform, or sclerosing. van Loo E, et al. *Eur J Cancer* 2014;50:3011-3020. Fraga SD, et al. *Dermatol Surg* 2022;48:704-710.

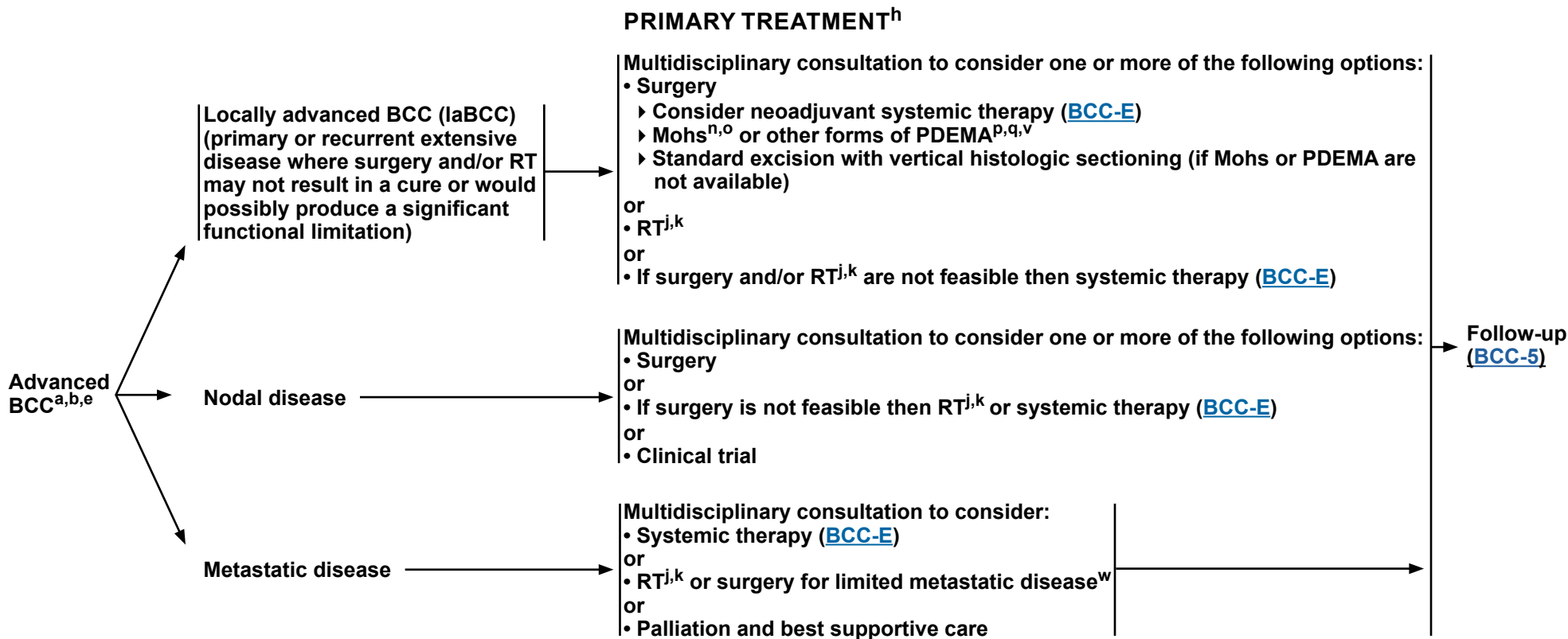
^s Due to the wide variability of clinical characteristics that may define a high-risk tumor, it is not feasible to recommend a defined margin for standard excision of high-risk BCC. Keen awareness of the subclinical extension of BCC is advised when selecting a treatment modality without complete margin assessment for a high-risk tumor. These margins may need to be modified based on tumor- or patient-specific factors.

^t If named nerve involvement is suspected, consider MRI with and without contrast of region of interest to evaluate extent and rule out base of skull involvement or intracranial extension in head and neck tumors.

^u There are conflicting data about the value of adjuvant RT following margin-negative surgical excision, particularly after Mohs.

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

Clinical Trials: NCCN believes that the best management of any patient with cancer is in a clinical trial. Participation in clinical trials is especially encouraged.



^a [Principles of Pathology \(BCC-A\)](#).

^b [Risk Factors for Recurrence \(BCC-B\)](#).

^e Any high-risk factor places the patient in the high-risk category.

^h [Principles of Treatment \(BCC-C\)](#).

^j [Principles of Radiation Therapy \(BCC-D\)](#).

^k Determination of the appropriateness of RT should be performed by a radiation oncologist.

ⁿ Mohs surgery should be performed by dermatologic surgeons who have specialized training and experience in this procedure.

^o As per other appropriate use criteria. Task Force/Committee Members, Vidal CI, et al. *J Am Acad Dermatol* 2019;80:189-207.

^p PDEMA with permanent section analysis or intraoperative frozen section analysis is an alternative to Mohs. See Principles of PDEMA Technique (SCC-G) within the [NCCN Guidelines for Squamous Cell Skin Cancer](#).

^q For clinically diagnosed non-facial BCCs <6 mm in depth on the head, neck, hands, feet, pretibial, and anogenital area that are clinically confined to the dermis, C&E or shave removal may be considered as an alternative primary treatment option if Mohs, resection with PDEMA, and standard excision are difficult to perform due to patient comorbidities (eg, thrombocytopenia, immunosuppression, bleeding diathesis, multiple primary BCCs). See [Risk Factors for Recurrence \(BCC-B\)](#).

^v Fraga SD, et al. *Dermatol Surg* 2022;48:704-710.

^w Under highly selective circumstances, in the context of multidisciplinary consultation, resection of limited metastases can be considered.

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FOLLOW-UP

- H&P
 - ▶ Including complete skin exam every 6–12 mo for the first 5 years, and then at least annually for life^x
- Consider imaging if clinical exam is insufficient for following the disease^y
- Patient education:
 - ▶ Sun protection
 - ▶ Self-examination

RECURRENCE

Local

Follow Primary Treatment pathway for High-risk disease ([BCC-3](#))

Advanced disease:
• Locally advanced
• Nodal metastases
• Distant metastases

Follow Primary Treatment pathways for Advanced BCC ([BCC-4](#))

^x Follow-up with a dermatologist is strongly recommended if any of the following criteria are met: past or imminent solid organ, marrow, or hematopoietic cell transplant; one or more cutaneous melanomas in the past 5 years; or four or more non-melanoma skin cancers in the past 5 years.

^y Imaging modality and targeted area should be at the discretion of the treating team based on the suspected extent of disease (ie, local, regional, metastatic). Histologic confirmation is sufficient to diagnose local recurrence, but MRI can be considered to assess extent of local disease. For nodal or distant metastasis, histologic analysis and/or CT imaging can be used for confirmation and to gauge extent of disease.

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PRINCIPLES OF PATHOLOGY

Principles of Biopsy Reporting:

- The intent of a biopsy is for diagnosis, not to assess the margin status.
- Pathologic evaluation of skin biopsies is ideally performed by a dermatologist, pathologist, dermatopathologist, or Mohs surgeon who is experienced in interpreting cutaneous neoplasms.
- Clinical information to be submitted on biopsy requisition includes patient age and gender, clinical diameter of lesion, anatomic location, and prior treatment of lesion. Additional helpful features to include are immunosuppression and history of RT.
- Pathologic report should include histologic subtype^a and presence and extent of any features that would increase the risk for local recurrence, including invasion of tumor beyond reticular dermis and presence of perineural invasion.¹

Principles of Excision Reporting:

- The intent of excision is to clear the tumor and thus margin status needs to be reported.
- Saucerization specimens intended for definitive surgical therapy should be labeled as such, as they can be histopathologically difficult to distinguish from shave biopsies but must be evaluated for margin status.
- Clinical information to be submitted on excision requisition includes patient age and gender, anatomic location, clinical diameter of lesion, and additional clinical information listed above under Principles of Biopsy Reporting.
- Minimal reporting elements to be reported for all surgical specimens include histologic subtype of BCC,^a invasion of tumor beyond deep reticular dermis, presence of perineural invasion (if involving nerve below dermis or if largest nerve involved is ≥ 0.1 mm in caliber) and angiolymphatic invasion, and peripheral and deep margin status.
- For Mohs excisions, reporting of these elements is also encouraged. Since depth of invasion (in mm) may not be ascertained on tangentially cut Mohs specimens, anatomic level of invasion should be reported. Frozen or permanent section analysis of the clinical tumor specimen may be undertaken if needed for complete reporting of features associated with poor prognosis.²

Footnotes

^a Low-risk histologic subtypes include nodular, superficial, and other non-aggressive growth patterns such as keratotic, infundibulocystic, and fibroepithelioma of Pinkus; high-risk subtypes include basosquamous, infiltrative, sclerosing/morpheaform, micronodular, and BCC with carcinosarcomatous differentiation.

References

¹ Work Group; Invited Reviewers, Kim JYS, et al. Guidelines of care for the management of basal cell carcinoma. J Am Acad Dermatol 2018;78:540-559.

² Morgan FC, Ruiz ES, Karia PS, et al. Brigham and Women's Hospital tumor classification system for basal cell carcinoma identifies patients with risk of metastasis and death. J Am Acad Dermatol 2021;85:582-587.

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STRATIFICATION TO DETERMINE TREATMENT OPTIONS FOR LOCAL BCC BASED ON RISK FACTORS FOR RECURRENCE^a

Risk Group	Low Risk	High Risk
Treatment options	BCC-2	BCC-3
H&P		
Location/size	Trunk, extremities <2 cm	Trunk, extremities ≥2 cm
		Head, neck, hands, feet, pretibial, and anogenital area (any size) ^c
Borders	Well-defined	Poorly defined
Primary vs. recurrent	Primary	Recurrent
Immunosuppression	(-)	(+)
Site of prior RT	(-)	(+)
Pathology (BCC-A)		
Subtype	Nodular, superficial ^b	Aggressive growth pattern ^d
Perineural involvement	(-)	(+)

^a Any high-risk factor places the patient in the high-risk category.

^b Low-risk histologic subtypes include nodular, superficial, and other non-aggressive growth patterns such as keratotic, infundibulocystic, and fibroepithelioma of Pinkus.

^c This area constitutes high risk based on location, independent of size. Narrow excision margins due to anatomic and functional constraints are associated with increased recurrence rates with standard histologic processing. Complete margin assessment such as with Mohs or PDEMA is recommended for optimal tumor clearance and maximal tissue conservation. For tumors <6 mm in size, without other high-risk features, other treatment modalities may be considered if at least 4-mm clinically tumor-free margins can be obtained without significant anatomic or functional distortions.

^d Having basosquamous, infiltrative, sclerosing/morpheaform, micronodular, and BCC with carcinosarcomatous differentiation features in any portion of the tumor. In some cases, basosquamous tumors may be prognostically similar to squamous cell carcinoma (SCC); clinicopathologic correlation is recommended in these cases to further consider prognostic implication.

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PRINCIPLES OF TREATMENT

- **The primary treatment goals of BCC is the complete removal of the tumor and the maximal preservation of function and cosmesis. All treatment decisions should be customized to account for the particular factors present in the individual case and for the patient's preference.**
- **Surgical approaches often offer the most effective and efficient means for accomplishing cure, but considerations of function, cosmesis, and patient preference may lead to choosing RT/topical therapy/systemic therapy as primary treatment in order to achieve optimal overall results.**
- **In certain patients at high risk for multiple primary tumors (eg, basal cell nevus syndrome [Gorlin syndrome], xeroderma pigmentosum, history of RT), increased surveillance and consideration of prophylactic measures may be indicated. Refer patients with suspected basal cell nevus syndrome or xeroderma pigmentosum for genetic evaluation.**
- **In patients with superficial basal cell skin cancer, non-surgical modalities may be considered. (See [BCC-2](#))**
- **When Mohs^b with margin assessment is being performed and the preoperative biopsy is considered insufficient for providing all the staging information required to properly treat the tumor, submission of the central specimen for vertical paraffin-embedded permanent sections or documentation of staging parameters in Mohs report is recommended.**
- **Use of nicotinamide may be effective in reducing the development of basal cell skin cancers.^{1,2}**

Footnotes

^a Cure rates are approximately 10% lower than for surgical treatment modalities. Jansen MHE, Mosterd K, Arits AHMM, et al. Five-year results of a randomized controlled trial comparing effectiveness of photodynamic therapy, topical imiquimod, and topical 5-fluorouracil in patients with superficial basal cell carcinoma. *J Invest Dermatol* 2018;138:527-533. Drew BA, Karia PS, Mora AN, et al. Treatment patterns, outcomes, and patient satisfaction of primary epidermally limited nonmelanoma skin cancer. *Dermatol Surg* 2017;43:1423-1430.

^b Mohs surgery should be performed by dermatologic surgeons who have specialized training and experience in this procedure.

References

¹ Chen AC, Martin AJ, Dalziel RA, et al. A phase II randomized controlled trial of nicotinamide for skin cancer chemoprevention in renal transplant recipients. *Br J Dermatol* 2016;175:1073-1075.

² Chen AC, Martin AJ, Choy B, et al. A phase 3 randomized trial of nicotinamide for skin-cancer chemoprevention. *N Engl J Med* 2015;373:1618-1626.

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Clinical Trials: NCCN believes that the best management of any patient with cancer is in a clinical trial. Participation in clinical trials is especially encouraged.

**PRINCIPLES OF RADIATION THERAPY****General Principles^a**

- Protracted fractionation is associated with improved cosmetic results and should be utilized for poorly vascularized or cartilaginous areas.
- RT is contraindicated for genetic conditions predisposing to skin cancer (eg, basal cell nevus syndrome and relatively contraindicated for patients with connective tissue diseases (eg, scleroderma).
- Given higher complication rates, re-irradiation should not be routinely utilized for recurrent disease within a prior radiation field.
- Isotope-based brachytherapy can be an effective treatment for certain sites of disease, particularly on the head and neck.
- There are insufficient long-term efficacy and safety data to support the routine use of electronic surface brachytherapy.

General Treatment Information

- Radiation treatments should be given by a practicing radiation oncologist with radiation physics support to meet established quality assurance and dosimetric constraints.

Primary Tumor	RT Dosing
Definitive RT	BED10 of 70–93 Gy for conventional fractionation BED10 of 56–88 Gy for hypofractionation
Postoperative adjuvant RT	BED10 of 60–79 Gy for conventional fractionation BED10 of 56–70 Gy for hypofractionation
Regional Disease	
<ul style="list-style-type: none"> • Lymph node regions, after lymph node dissection <ul style="list-style-type: none"> ▶ Negative margins, no extranodal extension (ENE) ▶ Positive margins or ENE 	50–60 Gy over 5 to 6 weeks 60–66 Gy over 6 to 7 weeks
<ul style="list-style-type: none"> • Lymph node regions, without lymph node dissection <ul style="list-style-type: none"> ▶ Clinically positive 	60–70 Gy over 6 to 7 weeks
<ul style="list-style-type: none"> • Clinically at-risk nerves 	50–60 Gy over 5 to 6 weeks

- BED = Biologically effective dose
- Conventionally fractionated radiotherapy consists of five daily treatments per week.
- Hypofractionated radiotherapy consists of daily treatments or two to four treatments per week. Fraction sizes larger than 6 Gy are not routinely recommended outside of the palliative setting.

^a [ASTRO Guideline on Definitive and Postoperative Radiation Therapy for Basal and Squamous Cell Cancers of the Skin.](#)

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PRINCIPLES OF SYSTEMIC THERAPY

Locally Advanced (laBCC), Nodal or Distant Metastatic Basal Cell Carcinoma (mBCC)

- Systemic therapy may be considered for laBCC. Locally advanced disease is defined by those that have primary or recurrent extensive disease where surgery and/or RT may not result in a cure or would possibly produce a significant functional limitation.
- Systemic therapy may be considered for cases of nodal or distant metastatic disease, especially if surgery and RT are not feasible.
- Multidisciplinary consultation may be required to determine the best treatment approach and deem the tumor not amendable to surgery or RT.
- Hedgehog pathway inhibitors (HHIs)
 - ▶ Due to frequency of intolerable side effects associated with HHIs, drug holidays or other alternatives to daily dosing can be used to reduce side effects to improve adherence to therapy and quality of life.
 - ▶ HHIs may be considered for diffuse BCC formation (eg, basal cell nevus syndrome or other genetic forms of multiple BCC). HHIs are not FDA approved for basal cell nevus syndrome; however, they may be used off-label and are effective based on a randomized controlled trial.¹
- The role of adjuvant systemic therapy for resected BCC is unclear and thus, adjuvant systemic therapy is best performed in a clinical trial setting.

	Preferred Regimens	Other Recommended Regimens	Useful in Certain Circumstances
Locally Advanced Disease - Neoadjuvant	• None	• Vismodegib ^{a,2} (category 2B)	• Cemiplimab-rwlc ^b (category 2B)
Locally Advanced Disease	• None	• Sonidegib ³ • Vismodegib ^{4,5}	• Cemiplimab-rwlc ^{b,c,6}
Nodal Disease	• None	• Vismodegib • Sonidegib ³ (category 2B)	• Cemiplimab-rwlc ^b
Metastatic Disease	• None	• Vismodegib ^{4,5}	• Cemiplimab-rwlc ^{b,6}

^a In one study of 55 patients with laBCC, neoadjuvant administration of vismodegib before planned surgery allowed for a smaller surgical procedure in 71% of patients, although it carried a high (36.4%) recurrence risk.²

^b Cemiplimab-rwlc is FDA approved for patients with laBCC or mBCC previously treated with an HHI or for whom an HHI is not appropriate.

^c A multinational single-arm phase 2 trial, consisting of 84 patients with locally advanced BCC (local invasion precluding complete resection or in locations for which surgery may result in severe disfigurement or dysfunction) whose disease had progressed on or was intolerant to prior HHI therapy, was conducted. Thirty-one percent had an objective response, including 6% with a complete response. See [Discussion](#).⁶

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

Clinical Trials: NCCN believes that the best management of any patient with cancer is in a clinical trial. Participation in clinical trials is especially encouraged.

[References on BCC-E \(2 of 2\)](#)



REFERENCES

- ¹ Tang JY, Ally MS, Chanana AM, et al. Inhibition of the hedgehog pathway in patients with basal-cell nevus syndrome: final results from the multicentre, randomised, double-blind, placebo-controlled, phase 2 trial. *Lancet Oncol* 2016;17:1720-1731.
- ² Bertrand N, Guerreschi P, Basset-Seguín N, et al. Vismodegib in neoadjuvant treatment of locally advanced basal cell carcinoma: First results of a multicenter, open-label, phase 2 trial (VISMONEO study): Neoadjuvant Vismodegib in Locally Advanced Basal Cell Carcinoma. *EClinicalMedicine* 2021;35:100844.
- ³ Dummer R, Guminksi A, Gutzmer R, et al. Long-term efficacy and safety of sonidegib in patients with advanced basal cell carcinoma: 42-month analysis of the phase II randomized, double-blind BOLT study. *Br J Dermatol* 2020;182:1369-1378.
- ⁴ Sekulic A, Migden MR, Oro AE, et al. Efficacy and safety of vismodegib in advanced basal-cell carcinoma. *N Engl J Med* 2012;366:2171-2179.
- ⁵ Dreno B, Basset-Seguín N, Caro I, Yue H, Schadendorf D. Clinical benefit assessment of vismodegib therapy in patients with advanced basal cell carcinoma. *Oncologist*. 2014;19:790-796.
- ⁶ Stratigos AJ, Sekulic A, Peris K, et al. Cemiplimab in locally advanced basal cell carcinoma after hedgehog inhibitor therapy: an open-label, multi-centre, single-arm, phase 2 trial. *Lancet Oncol*. 2021;22:848-857.

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PRINCIPLES OF CANCER RISK ASSESSMENT AND COUNSELING

- The decision to offer genetic testing involves three related stages:
 - 1) Pre-test counseling prior to ordering testing;
 - 2) Consideration of the most appropriate testing strategy; and
 - 3) Testing result disclosure and post-test counseling.
- In certain patients at high risk for multiple primary tumors (eg, basal cell nevus syndrome, xeroderma pigmentosum, history of RT), increased surveillance and consideration of prophylactic measures may be indicated. Patients with these conditions should be referred to a cancer center with particular expertise in BCC prevention and prophylaxis.
- It is recommended that a genetic counselor, medical geneticist, endocrinologist, oncologist, surgeon, oncology nurse, or other health professional with expertise and experience in cancer genetics be involved at each stage whenever possible. Clinicians without direct referral access to the appropriate expertise should be aware of the telehealth genetic counseling options available. These resources can be found through the National Society of Genetic Counselors (NSGC) “Find a Genetic Counselor” tool (www.nsgc.org).

See the [NCCN Guidelines for Genetic/Familial High-Risk Assessment: Breast, Ovarian, and Pancreatic](#) for the following:

- Principles of Cancer Risk Assessment and Counseling (EVAL-A)
- Pedigree: First-, Second-, and Third-Degree Relatives of Proband (EVAL-B)
- General Testing Criteria (CRIT-1)

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ABBREVIATIONS

ALA	aminolevulinic acid
BCC	basal cell carcinoma
BED	biologically effective dose
C&E	curettage and electrodesiccation
ENE	extranodal extension
H&P	history and physical
HHI	hedgehog pathway inhibitors
laBCC	locally advanced basal cell carcinoma
mBCC	metastatic basal cell carcinoma
PDEMA	peripheral and deep en face margin assessment
SCC	squamous cell carcinoma

**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 indicated.

NCCN Categories of Preference

Preferred intervention	Interventions that are based on superior efficacy, safety, and evidence; and, when appropriate, affordability.
Other recommended intervention	Other interventions that may be somewhat less efficacious, more toxic, or based on less mature data; or significantly less affordable for similar outcomes.
Useful in certain circumstances	Other interventions that may be used for selected patient populations (defined with recommendation).

All recommendations are considered appropriate.

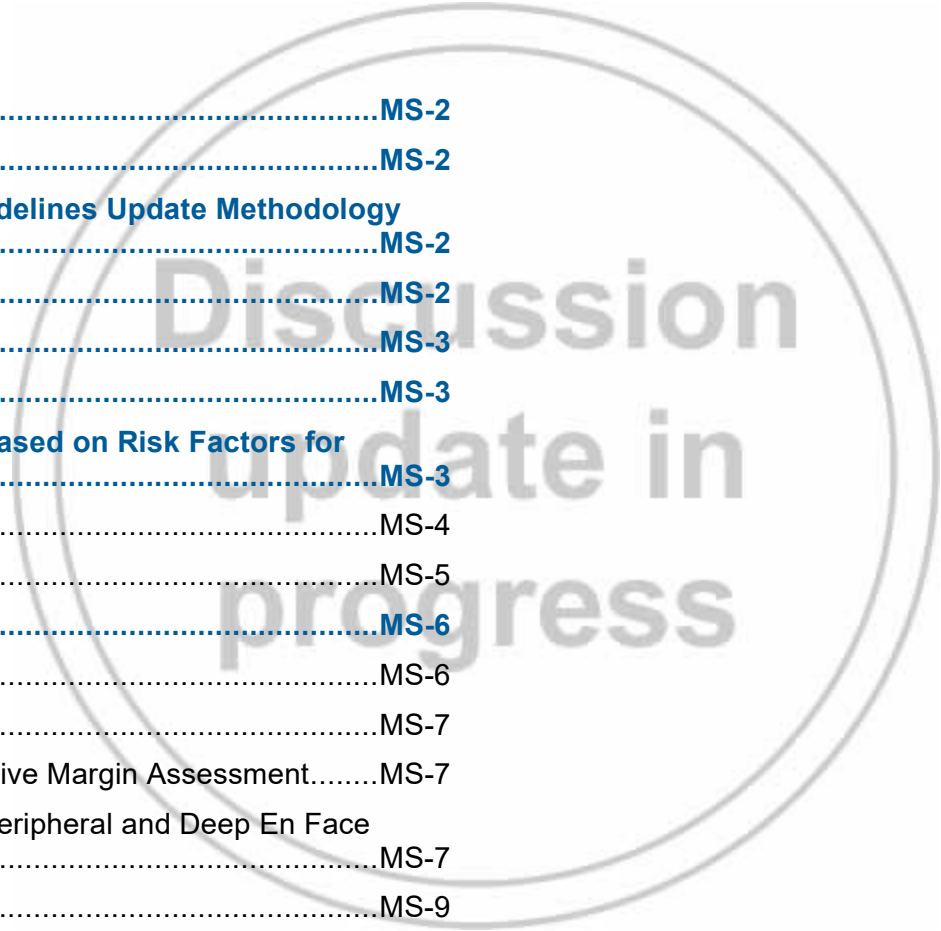


Discussion

This discussion corresponds to the NCCN Guidelines for Basal Cell Skin Cancer. Last updated: September 15, 2023.

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Overview

Basal cell carcinoma (BCC) is the most common cancer in the United States. It is estimated that BCCs occur in 2 million Americans annually, exceeding the incidence of all other cancers combined.¹⁻³ BCCs are at least two times more common than squamous cell carcinomas (SCCs), the second most common type of skin cancer.¹⁻⁶ Furthermore, the incidence of this common malignancy is rising rapidly.^{1,3,6,7} Compared with SCC, BCCs are much less likely to metastasize, with a metastatic rate of <0.1%, and thus generally have a good prognosis.⁸⁻¹⁰ Although rarely metastatic, BCC can produce substantial local destruction along with disfigurement and may involve extensive areas of soft tissue, cartilage, and bone.

A number of risk factors are associated with the development of BCC. The most recognized environmental carcinogen is sunlight. Evidence reveals that the relationship between sun exposure and BCC is complex, and depends on the timing, pattern, and amount of ultraviolet (UV) radiation.¹¹⁻¹⁵ Fair skin, red or blond hair, and light eye color are associated with BCC as independent risk factors due to greater susceptibility to UV damage.^{13,15-22} BCC risk is increased by both UV-A and -B radiation as well as by ionizing radiation. Radiation therapy (RT) for other conditions, especially at a young age, is also associated with an increased risk for developing BCC.²³⁻²⁷ Most BCC tumors develop on skin sites exposed to radiation—either from the sun or from therapy.²³⁻²⁵

Guidelines Update Methodology

The complete details of the Development and Update of the NCCN Guidelines are available at www.NCCN.org.

Literature Search Criteria and Guidelines Update Methodology

Prior to the update of this version of the NCCN Clinical Practice Guidelines in Oncology (NCCN Guidelines®) for Basal Cell Skin Cancer, an electronic search of the PubMed database was performed to obtain key literature using the following search term: basal cell skin carcinoma. The PubMed database was chosen as it remains the most widely used resource for medical literature and indexes peer-reviewed biomedical literature.²⁸

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; Practice Guideline; Randomized Controlled Trial; Systematic Reviews; and Validation Studies.

The data from key PubMed articles as well as articles from additional sources deemed as relevant to these guidelines as discussed by the panel during the Guidelines update have been included in this version of the Discussion section. Recommendations for which high-level evidence is lacking are based on the panel's review of lower-level evidence and expert opinion.

Sensitive/Inclusive Language

NCCN Guidelines strive to use language that advances the goals of equity, inclusion, and representation.²⁹ NCCN Guidelines endeavor to use language that is person-first; not stigmatizing; anti-racist, anti-classist, anti-misogynist, anti-ageist, anti-ableist, and anti-weight biased; and inclusive of individuals of all sexual orientations and gender identities. NCCN Guidelines incorporate non-gendered language, instead focusing on organ-specific recommendations. This language is



both more accurate and more inclusive and can help fully address the needs of individuals of all sexual orientations and gender identities. NCCN Guidelines will continue to use the terms men, women, female, and male when citing statistics, recommendations, or data from organizations or sources that do not use inclusive terms. Most studies do not report how sex and gender data are collected and use these terms interchangeably or inconsistently. If sources do not differentiate gender from sex assigned at birth or organs present, the information is presumed to predominantly represent cisgender individuals. NCCN encourages researchers to collect more specific data in future studies and organizations to use more inclusive and accurate language in their future analyses.

Genetics

Extensive research has led to advances in the understanding of the genetics of BCC. The sonic hedgehog signaling pathway has emerged as playing a pivotal role in the pathogenesis of BCC, and mutations in a number of molecules in this pathway have been implicated in the development of the disease.³⁰⁻³² Mutations in the *PTCH1* (patched 1) gene on chromosome 9q, which codes for the sonic hedgehog receptor, are the underlying cause of nevoid BCC syndrome, and are present in approximately 30% to 90% of sporadic BCCs.³³⁻⁴¹ Specific UV-induced mutations in the tumor suppressor gene *p53* appear to be a common event in BCC development.^{35,38,41,42} Certain genetic syndromes greatly predispose affected individuals to skin cancer formation, including BCC, such as albinism^{43,44} and xeroderma pigmentosum (in which defects exist in UV light-induced unscheduled DNA repair).⁴⁵⁻⁵¹

Clinical Presentation and Workup

On clinical presentation of the patient with lesion suspicious of skin cancer, workup for BCC begins with a history and physical examination, biopsy, and if applicable a shave removal. A skin biopsy is then performed on any suspicious lesion. The biopsy should include deep reticular dermis. This procedure is preferred because an infiltrative histology may sometimes be present only at the deeper, advancing margins of a tumor, and superficial biopsies will frequently miss this component.^{52,53} After BCC diagnosis, a full skin examination is recommended, because individuals with skin cancer often have additional, concurrent precancers or cancers located at other, usually sun-exposed skin sites. These individuals are also at increased risk of developing cutaneous melanoma.⁵⁴

Risk Stratification of Local BCC Based on Risk Factors for Recurrence

After the complete skin examination, a risk assessment should be performed to determine the treatment plan.⁵⁵ The NCCN Panel examined risk factors for BCC associated with recurrence (see *Risk Factors for Recurrence* in the algorithm). Any high-risk factor places the skin lesion in the high-risk category and imaging should be considered if a clinical exam is insufficient to determine disease extent. Skin lesions in populations placed at increased risk may be difficult to assess clinically; therefore, a low threshold for performing skin biopsies in these patients is necessary. Patients with locally advanced disease, which is defined as primary or recurrent extensive disease where surgery and/or RT may not result in a cure or would potentially yield a significant functional limitation, should consider imaging to determine disease extent. For rare cases when patients present with regional or distant metastatic disease at diagnosis, imaging of areas of interest can be



performed when there is suspicion of extensive disease prior to treatment as nodal or distant metastases. Imaging studies may be clinically evident when extensive disease, such as bone involvement, perineural invasion (PNI), or deep soft tissue involvement, is suspected. If perineural disease is suspected, MRI with or without contrast is preferred.^{56,57} If bone disease is suspected, CT with contrast is preferred unless contraindicated. Imaging modality and targeted area should be at the discretion of the treating team based on the suspected extent of disease (ie, local, regional, metastatic). Histologic confirmation is sufficient to diagnose local recurrence, but MRI can be considered to assess extent of local disease. For nodal or distant metastases, histologic analysis and/or CT imaging can be used for confirmation and to gauge the extent of disease.

History & Physical Examination

Location and Size

Anatomic location⁵⁸⁻⁶⁴ and size⁶⁰⁻⁶⁶ have been known to be a risk factor for BCC recurrence and metastasis for many years. In general, BCCs that develop in the head and neck area, which includes the “H zone” or “mask area” of the face, are more likely to recur than those that develop on the trunk and extremities. Based on a 27-year retrospective review of 5755 BCCs, recurrences were significantly more common when tumors in high-risk locations (central face, eyebrows, nose, lips, chin, ear, temple, genitalia, nipples/areola, hands, feet, ankles, and nail units) were greater than or equal to 6 mm in diameter and when tumors in moderate-risk locations (cheeks, forehead, scalp, neck, jawline, pretibial surface) were greater than or equal to 10 mm in diameter.⁶⁷ The American Academy of Dermatology in collaboration with American College of Mohs Surgery, American Society for Dermatologic Surgery Association, and American Society for Mohs Surgery developed an appropriate use criteria document in the treatment of cutaneous

neoplasms based on 270 clinical scenarios including 69 BCCs,⁶⁸ which has been incorporated into *Risk Factors for Recurrence* within the algorithm.

Clinical Borders and Primary Versus Recurrent Disease

The low- and high-risk factors of well-defined versus ill-defined clinical tumor borders⁶⁹⁻⁷¹ and primary versus recurrent disease,^{62,70,72} respectively, have been extensively documented in the literature.

Immunosuppression

Settings of immunosuppression, such as organ transplantation,⁷³⁻⁷⁸ and long-term use of psoralen and UVA (PUVA) light,^{79,80} increase the incidence of BCC. In particular, among patients who have had organ transplants, BCC incidence is approximately 5- to 10-fold higher than in the general population,⁸¹⁻⁸³ occurring in up to half of patients during the 10 years following transplant.⁸⁴⁻⁸⁷ Several large retrospective studies found that BCCs in patients who had received organ transplants were more likely to have the superficial histologic subtype and to occur in extracephalic locations and in younger patients (mean age of onset 15 years lower).⁸⁸⁻⁹⁰ Two of these studies showed similar low recurrence rates for transplant recipients and controls.^{89,90} Nevertheless, because of NCCN Guidelines Panel Members’ own anecdotal experiences, the panel decided to classify BCCs developing in settings of immunosuppression as potentially high-risk tumors.

Site of Prior Radiotherapy

Tumors developing in sites of prior RT refer to primary BCCs arising in areas previously irradiated for unrelated conditions. All recurrent tumors, irrespective of prior therapy, are defined as high risk. Data from a number of studies with large sample sizes support that prior RT for



unrelated, frequently benign conditions is a risk factor for BCC development.^{23-27,91,92}

Pathology

Pathologic Subtypes

Histologic subtyping of BCC as a predictor of risk of recurrence is a well-established concept.^{93,94} The subtypes encompassed by the term “aggressive growth pattern,” including micronodular, infiltrative, sclerosing, and morpheaform (or desmoplastic) patterns, are more likely to recur than the nodular and superficial BCC.^{65,69,70,72,95-99} Non-aggressive subtypes include the keratotic variant, infundibulocystic variant, and fibroepithelioma of Pinkus.

Basosquamous carcinomas are tumors that have the histologic appearance of both a BCC and an SCC. Some basosquamous tumors are the result of a BCC colliding with an adjacent SCC. Others represent truly biphenotypic tumors, many of which may have started as BCC, but have subsequently undergone prominent partial squamous metaplasia.¹⁰⁰ Data suggest that basosquamous carcinomas have a metastatic capacity that is more similar to that of SCC than BCC.¹⁰¹⁻¹⁰³

Perineural Involvement

PNI is uncommon in any nonmelanoma skin cancer (NMSC) (2%–6%), and develops less frequently and is less aggressive in BCC versus SCC.¹⁰⁴⁻¹⁰⁹ BCC with PNI poses a greatly increased risk of recurrence, and is associated with other risk factors including previous recurrent tumors, high grade, larger lesion size, and certain subtypes (infiltrating, morpheaform, and basosquamous).^{108,110,111} If large nerve involvement is suspected, MRI should be considered to evaluate extent and/or rule out skull involvement in those with head and neck tumors.^{57,112-114}

Additionally, in the presence of PNI, a thorough cranial nerve exam is indicated.

Age and Its Effect on BCC Behavior

Whether young age (typically ≤40 years) is an independent risk factor for aggressive BCC behavior is debatable. An analysis of a large database of patients with BCC (N = 3381) documented an increased percentage of BCC with aggressive histologic growth patterns in young persons.¹¹⁵ In contrast, results from other analyses of large databases (N = 1000 to >10,000) indicate that patients presenting with BCC at a young age are more likely to have the superficial subtype.¹¹⁶⁻¹¹⁹ Other analyses report no significant differences in BCC histologic subtype between young versus older patients.¹²⁰⁻¹²² The relationship between tumor location and patient age is also unclear, as several studies showed that younger patients were more likely to present with BCCs on the trunk or extremities,^{116,121,123,124} while another found no significant association.¹²⁰

Most large studies (N = 50-2000) have shown no significant association between age and recurrence rate.^{62,70,120,122} One multivariate analysis, however, showed a positive relationship between increasing age and likelihood of recurrence.¹²⁵ Age has also been evaluated as a risk factor for developing a second or multiple BCCs and many of these studies using fairly large databases (N = 200–2500) found that the risk of developing more than one BCC is associated with increased age.^{65,122,124-130} On the contrary, an analysis of a very large database (N = 71,924) found a significantly higher risk of subsequent NMSC in patients <40 years at the time of their first BCC diagnosis.¹³¹ In addition, an analysis of 100 metastatic BCC cases found that patients with distant metastases tended to be younger than those with only regional metastases.¹³² Consistent with this idea, the Rotterdam Study showed



that while the risk of developing a second BCC increased with age,¹³⁰ the risk of developing multiple BCC lesions was highest in patients who were <65 years at the time of their first BCC diagnosis.¹³³ Taken together, these studies suggest that young age, in and of itself, is not considered a risk factor for aggressive BCC. Nevertheless, there is a small subset of patients who develop BCC at a young age and may have particularly aggressive disease. These patients may benefit from regular follow-up.

Treatment Modalities for BCC

Curettage and Electrodesiccation

Although a fast and cost-effective technique for superficial lesions, curettage and electrodesiccation (C&E) does not allow histologic margin assessment. Studies have reported overall 5-year recurrence rates ranging from 1.2% to 40% in patients with BCC selected for C&E, with high-risk locations and histologically aggressive subtypes reporting higher recurrence rates.^{60,134-143}

This technique is deemed effective for properly selected, low-risk BCC with three caveats.^{60,140} First, C&E should not be used to treat areas with terminal hair growth such as the scalp, pubic and axillary regions, or beard area due to the risk that a tumor extending down follicular structures might not be adequately removed. Second, if the subcutaneous layer is reached during the course of C&E, then surgical removal should generally be performed instead. This change in therapy is necessary as the effectiveness of the C&E technique rests on the ability of the clinician to distinguish between firm, normal dermis, and soft tumor tissue when using a sharp curette. Since subcutaneous adipose is even softer than tumor tissue, the ability of the curette to distinguish and, therefore, selectively and completely remove tumor cells diminishes. Third, if C&E has been performed based only on the appearance of a low-risk tumor, biopsy results of the tissue taken at the time of C&E should be reviewed to make sure that there are no high-risk pathologic features that would require additional therapy. For tumors on the cheeks, forehead, scalp, neck, and pretibial that are less than 6 mm in depth and confined to the dermis, C&E may be considered as an alternative primary treatment option if Mohs micrographic surgery (Mohs) or resection with peripheral and deep en



face margin assessment (PDEMA) and standard excision are not feasible due to patient comorbidities.

Shave Removal

Shave removal, the shaving of epidermal or dermal lesions, is a sharp removal by bowl-shaped slicing of the epidermal and dermal lesions, without including fat, and does not require suture closure.¹⁴⁴ Like C&E, there is concern for inaccurate margin status assessment with shave removal.¹⁴⁵ However, it is a recommended technique for low-risk BCCs located in the trunk or extremities. Shave removal studies have reported 0.5% to 30% rate of recurrence over a 3- to 5-year follow-up, multiple tumors treated in single visits, and a risk for misdiagnosis of only 1%.¹⁴⁴⁻¹⁴⁷

Standard Excision with Postoperative Margin Assessment

Another therapeutic option for BCC is standard surgical excision followed by postoperative pathologic evaluation of margins. This technique has been reported to achieve 5-year recurrence rates of 0.8% to 17.4% for BCC, with lower recurrence rates associated with low-risk tumors and higher recurrence rates associated with high-risk tumors.^{134,136,142,148-150} Studies have reported variable margins required to completely excise 95% of all tumor.¹⁵¹⁻¹⁵⁶ These margins have been suggested to be 2 to 4 mm for low-risk, well-demarcated tumors smaller than 2 cm,¹⁵¹⁻¹⁵⁵ whereas margins of 4 to 6 mm,¹⁵² and in one study, 8 mm¹⁵¹ were suggested for high-risk BCC. Given this wide variability, studies have reported incomplete excision rates after standard excision ranging from 3.2% to 61.5% depending on tumor location, histologic subtype, and medical provider's specialty.¹⁵⁷⁻¹⁶⁶ Therefore, postoperative margin assessment and identification of clear margins are critical to ensure favorable outcomes with standard excision.

The clinical margins chosen by the panel for the primary treatment of low-risk BCC are based on the work of Zitelli et al.¹⁶⁷ Their analysis indicated that for well-circumscribed BCC lesions smaller than 2 cm in diameter, excision with 4-mm clinical margins should result in complete removal in more than 95% of cases. The indications for this approach were also expanded to include re-excision of low-risk primary BCC if positive margins are obtained after an initial excision with postoperative margin assessment. For high-risk BCC, standard excision with wider surgical margins is recommended as the primary treatment. Due to the wide variability of clinical characteristics that may define a high-risk tumor, it is not feasible to recommend a defined margin for standard excision of high-risk BCC. Kean awareness of the subclinical extension of BCC is advised when selecting a treatment modality without complete margin assessment for a high-risk tumor. These margins may need to be modified based on tumor- or patient-specific factors. When standard excision with wider surgical margins yields positive margins, Mohs or other forms of PDEMA or standard re-excision are recommended (if PDEMA is not feasible).

For either low-risk or high-risk BCC, when standard excision is used, tissue rearrangement (eg, flap reconstruction, extensive undermining) should not be undertaken until clear margins are identified. Second intention healing, linear repair, or skin graft are acceptable options.

Mohs Micrographic Surgery and Peripheral and Deep Face Margin Assessment

Mohs is the preferred surgical technique over standard excision for re-excision of low-risk BCC after positive margins with standard excision, as well as the primary surgical technique of choice for high-risk BCC because it allows intraoperative analysis of 100% of the excision margin. Mohs is also recommended when standard excision with wider



surgical margins is unable to achieve negative margins in high-risk BCC. Two meta-analyses published in 1989 associated Mohs with 5-year recurrence rates of 1.0% for primary BCC, and 5.6% for recurrent BCC.^{134,142} In these studies, the recurrence rates for Mohs were lower than those for standard excision (10.1% and 17.4% for primary and recurrent BCC, respectively), and lower than those for any other treatment modality included in the analysis (C&E, cryotherapy, and RT).^{134,142} Studies on the long-term outcomes (~4 years) of Mohs have reported overall recurrence rates of 2.9% to 3.8%,^{168,169} specifically 0% to 6.5% for primary and 4% to 20% for recurrent BCCs.^{94,170-175} The only prospective randomized trial comparing Mohs to standard excision reported fewer 10-year recurrences with Mohs for both primary (2.5% vs. 4.1%; $P = .397$) and recurrent BCC (2.4% vs. 12.1%; $P = .015$), although the difference was only statistically significant for recurrent tumors. Importantly, a large proportion of recurrences occurred more than 5 years after treatment.^{143,176,177} Besides lower recurrence rates, Mohs has also been associated with significant tissue sparing compared with standard excision.^{178,179} It has been demonstrated that H-zone location, recurrent tumor, aggressive subtype, PNI, and tumor size greater than or equal to 11 mm are significantly associated with two or more Mohs stages.^{110,180} However, superficial BCC, despite being generally considered less aggressive, was shown in a Brazilian study to be 9.03 times more likely to require more than one Mohs stage, likely due to “skip areas” and clinically indistinct borders.¹⁸¹

Excision with PDEMA with permanent section analysis or intraoperative frozen section analysis is an acceptable alternative to Mohs provided it includes a complete assessment of all deep and peripheral margins. A 5-year recurrence rate of 0.58% has been reported with slow Mohs using formalin-fixed paraffin-embedded sections and delayed closure in a UK-based prospective study.¹⁸² The descriptive term PDEMA

underscores the panel’s belief that complete histologic assessment of the entire marginal surface is the key to optimal tumor removal. For more information, refer to the [NCCN Guidelines for Squamous Cell Skin Cancer](#) *SCC-G Principles of PDEMA Technique*.



Radiation Therapy

Although surgery is the mainstay of local treatment for BCC, consideration of function and patient preference and other factors may lead to the choice of RT as primary therapy for non-surgical candidates for both low-risk and high-risk as well as patient with advanced BCC (locally advanced, nodal, and metastatic BCC).¹⁸³ The recommendations for RT extend to additional treatment for low-risk BCC after positive margins with standard excision. RT is also recommended for high-risk BCC as additional treatment after standard excision, Mohs, or other forms of PDEMA with positive margins and adjuvant treatment after negative margins in case of extensive perineural or large-nerve involvement.¹⁸⁴ In these patients, local control has been reported to be 50% to 90% with postoperative RT.^{183,185} There are conflicting data about the value of adjuvant RT following margin-negative surgical excision, particularly after Mohs. For high-risk BCC that has undergone multiple resections, and further surgery is not feasible, RT is recommended as part of multidisciplinary consultation if residual disease is present. For specifics about the application of RT, see *Principles of Radiation Therapy* within the algorithm.

Two meta-analyses reported 5-year recurrence rates of 8.7% and 9.8% after RT on primary and recurrent BCC, respectively.^{134,142}

Retrospective analyses of BCC treated with RT have reported 5-year local control, cure, or complete response rates ranging from 93% to 96%,¹⁸⁶⁻¹⁸⁹ and 5-year recurrence rates from 4% to 16%.¹⁹⁰⁻¹⁹² Efficacy of RT was better for BCCs that were less advanced, primary (vs. recurrent), or had smaller diameter or nodular histologic subtype.^{186,187,189-191} A prospective study randomizing 347 patients to receive either surgery (standard excision with free margins ≥ 2 mm from visible borders) or RT as primary treatment of BCC reported higher

recurrence rates with RT than surgery (7.5% vs. 0.7%; $P = .003$),¹⁹³ poorer cosmetic outcomes, and more postoperative complications.¹⁹⁴

A small number of prospective studies have reported high rates of tumor control with specific radiation dose fractionation regimens for small BCCs.^{193,195,196} A systematic review and meta-analysis also reported hypofractionated RT regimens associated with positive cosmetic outcomes.¹⁹⁷ The NCCN Panel recommends ranges of electron beam dose and fractionation that can be used for definite RT and postoperative adjuvant RT. Isotope-based brachytherapy can be an effective treatment for certain sites of disease, particularly on the head and neck.¹⁹⁸⁻²⁰¹ However, there are insufficient long-term efficacy and safety data to support the routine use of electronic surface brachytherapy.^{202,203}

Superficial Therapies

In patients with superficial BCC, therapies such as topical imiquimod, topical 5-fluorouracil (5-FU), or photodynamic therapy (PDT) may be considered, although cure rates are approximately 10% lower than for surgical treatment modalities.²⁰⁴⁻²⁰⁶ Another option for patients with superficial BCC is cryotherapy.²⁰⁷ These options are also recommended for patients where surgery or RT is contraindicated or impractical.

Topical Therapies

Imiquimod was found to be effective for treating nodular and superficial BCC in randomized studies.²⁰⁸⁻²¹³ Two 5-year follow-up studies reported overall treatment success rates of 80.4% and 77.9%, respectively, in patients with superficial BCC treated with imiquimod.^{212,214} Recurrence seems to be associated with tumor thickness.²¹⁵ A phase III randomized trial in patients with superficial or nodular BCC showed that imiquimod



provided an 82.5% clinical success rate.^{216,217} For all of these studies, tumors in the H-zone were excluded. Although the clinical success rate was significantly higher with surgical excision using a 4-mm margin (97.7%; $P < .001$), cosmetic outcomes by dermatologic assessment were significantly better with imiquimod (excellent/good at 3-year follow-up: 61% vs. 36%; $P < .001$). Another topical cream with efficacy against BCC is 5-FU,^{218,219} which has been shown in a large randomized trial to have a 5-year tumor-free survival probability of 70.0%.^{205,220,221} Other studies have reported cure rates of up to 90% with this treatment.²²²⁻²²⁴

Photodynamic Therapy

PDT with photosensitizing agents including 5-aminolevulinic acid (ALA) and porfimer sodium is another option for superficial BCC.²²⁵⁻²²⁷ Multiple randomized trials and a meta-analysis have shown that rates of excellent or good cosmetic outcomes were higher with PDT versus surgery, although surgery was superior to PDT in terms of disease control.^{149,228-235} Data from clinical trials reported cure rates from 60% to 100% by PDT for patients with BCC.^{231,236-241} Most of these studies have focused on the superficial and nodular histologic subtypes, and several have found higher cure rates for superficial versus nodular subtypes in both low- and high-risk locations.^{231,236,241} Ulceration and thickness are associated with lower response to therapy,²⁴¹ and within the nodular subtype, cure rates are better with thinner lesions.²³⁰ Clinical studies have demonstrated PDT activity against “difficult-to-treat” lesions, with a 24-month complete response rate of 78%.^{236,242} Currently, PDT is being used at some NCCN Member Institutions for premalignant or superficial low-risk lesions on any location on the body, although response rates may be higher on the face and scalp.^{243,244}

Cryotherapy

Cryotherapy has been used for many years as a fast and cost-effective means for removal of BCCs.²⁰⁷ Systematic reviews of historical data in primary BCCs have reported recurrence rates for cryotherapy ranging from 0% to 13%, and mean recurrence rates from pooled analyses ranging between 3% and 4%.^{134,136,245,246} In prospective trials, cryotherapy has been shown to result in recurrence rates ranging from 5% to 39%.^{195,247-249} A key limitation of cryotherapy is poorer cosmetic outcomes compared with other treatment options, as demonstrated by prospective randomized trials.²⁴⁸⁻²⁵⁰

Comparisons of Superficial Therapies

Several randomized studies and meta-analyses have compared superficial therapies for BCC (**Table 1**). In summary, these studies indicate that in patients with superficial BCC, PDT has similar efficacy as cryotherapy but much better cosmetic outcomes. Whereas a meta-analysis of 23 randomized and non-randomized trials found no significant difference in efficacy for PDT versus imiquimod,²⁵¹ a randomized trial showed that treatment success was more likely with imiquimod.^{205,221} This study also demonstrates superior imiquimod outcomes compared to 5-FU cream. Exploratory sub-analyses found that treatment success rates were significantly higher with imiquimod for tumors that are large or truncal, while PDT provided significantly better outcomes in older patients with lesions on the lower extremities.²⁵² Safety results showed that while PDT causes moderate to severe pain during treatment administration, imiquimod and 5-FU are more likely to cause moderate to severe local swelling, erosion, crust formation, itching, and wound infections.²²⁰ Both cryotherapy and PDT are associated with pain during and after treatment, and data from a randomized trial indicate a trend toward a higher likelihood of pain with PDT.²⁴⁸



Nicotinamide in Reducing BCC Development

Data from phase II and phase III randomized trials indicated that treatment of actinic keratoses with nicotinamide reduced the occurrence of new BCCs, specifically by 20% at 12-month follow-up.^{253,254} This is supported by data from another study.²⁵⁵ Other agents that might be effective for the prevention of BCC in individuals at high risk for developing NMSCs include celecoxib,²⁵⁶ acitretin,²⁵⁷ capecitabine,²⁵⁸ and tazarotene.²⁵⁹

Systemic Therapy

For advanced BCC, systemic therapy is recommended as a treatment option for locally advanced (laBCC), metastatic (mBCC), and nodal BCC after multidisciplinary consultation. Other options include surgery, RT, and palliation and best supportive care for certain patients. The systemic therapy options for BCC include hedgehog pathway inhibitor (HHI) and immunotherapy. Vismodegib and cemiplimab are currently recommended options for all advanced BCCs while sonidegib is only recommended for nodal and laBCC.

Hedgehog Pathway Inhibitors

Vismodegib is an HHI approved by the U.S. Food and Drug Administration (FDA) for the treatment of adults with laBCC or mBCC that has recurred following surgery, or those who are not candidates for surgery or RT.²⁶⁰ The 9-month follow-up data from the SHH4476g trial, a centrally reported, multicenter, phase I, open-label study had an initial enrollment of 104 patients (laBCC N = 71, mBCC N = 33); however, pathology results excluded eight laBCC patients from the efficacy analysis (N = 63). This trial reported an objective response rate of 30% in the mBCC group and 43% in the laBCC group, with a median duration of response (DOR) of 7.6 months and 9.5-month median progression-free survival (PFS).²⁶¹ A 39-month follow-up to these data

from the ERIVANCE trial, an investigator-reported, multicenter, phase II trial, conveyed an objective response rate of 48.5% in the mBCC group and 60.3% in the laBCC group, with a median DOR of 14.8 months and 26.2 months for each group, respectively.²⁶¹⁻²⁶⁴ Results from these trials for vismodegib in BCC are summarized in **Table 2**. According to these data, nearly all patients treated with vismodegib experienced at least one treatment-emergent adverse event (TEAE), but a significant proportion of these were low grade (grade ≤2).²⁶¹⁻²⁶³ Serious adverse events (SAEs) occurred in 25% to 32% of patients in these studies. The most common adverse events (AEs) included muscle spasms, alopecia, taste loss, weight loss, decreased appetite, fatigue, nausea, and diarrhea.

Vismodegib has also been tested as BCC treatment and prophylaxis in patients with nevoid BCC syndrome. A randomized phase II study in patients with nevoid BCC syndrome and at least 10 operable BCC lesions found that vismodegib significantly reduced incidence of new BCC lesions compared with placebo, and also significantly reduced the size of existing lesions and the number of surgeries needed to remove BCC lesions.²⁶⁵⁻²⁶⁷

Sonidegib is another HHI FDA-approved agent for the treatment of patients with laBCC that has recurred following surgery or RT, or who are not candidates for surgery or RT.²⁶⁸ Sonidegib is FDA approved for laBCC. The 42-month follow-up data from the centrally reported randomized, multicenter, phase II BOLT trial reported similar objective response rates for the 200-mg and 800-mg doses tested among patients with laBCC (56% and 46%, respectively), while there was a 2-fold difference for patients with mBCC (8% and 17%, respectively).²⁶⁹⁻²⁷³ This trial also reported, for each dose and patient group, median DOR and PFS results that are summarized in **Table 2**. The 30-month



investigator-reviewed data for the BOLT trial analyzing only the 200-mg dose showed a higher objective response rate of 71.2% for laBCC and 23.1% for mBCC (**Table 2**).^{271,274} As with vismodegib, nearly all patients experienced at least one AE, and the most common AEs were muscle spasms, dysgeusia, alopecia, nausea, weight decrease, and fatigue. Elevated creatinine kinase was also frequently observed and was one of the most common grade 3–4 AEs, along with elevated lipase.

A key limitation to HHI therapies is that advanced BCC can develop resistance, which limits the DOR. A small investigator-initiated trial in patients with vismodegib-resistant advanced BCC observed no responses during treatment with sonidegib for a median of 6 weeks (range, 3–58 weeks), and in 5 of 9 patients with disease progression.²⁷⁵

Ongoing clinical research is exploring various dosing regimens of vismodegib and sonidegib in a variety of BCC treatment settings, including in the neoadjuvant setting, in patients with multiple BCCs or with radiation-induced multiple BCCs of the scalp, and as maintenance therapy after laBCC complete remission.²⁷⁶⁻²⁸¹ Notably, in the neoadjuvant setting, while one trial reported negative results (unmet predefined complete histologic clearance rate),²⁷⁷ results from two studies indicated vismodegib may reduce surgical defect area and allow for downstaging of the surgical procedure for laBCCs in functionally sensitive locations.^{276,279} The VISMEO trial, a centrally reported, phase II, open-label study, had an enrollment of 55 patients with laBCC. This study reported an objective response rate of 71%, with 36.4% recurrence at the 3-year follow-up.²⁷⁹ Some of these studies included small numbers of patients, and thus their results need to be carefully interpreted.

Other HHIs are also being tested in patients with BCC to see if they can provide higher rates of response, more durable responses, responses in less advanced BCC, or responses in BCC resistant to vismodegib. Results from phase I–II trials with small BCC sample sizes (N < 40) have shown that itraconazole and saridegib can elicit responses in patients with BCC, although not in patients who previously received vismodegib.^{282,283}

Immunotherapy

Cemiplimab-rwlc is an anti-PD-1 immunotherapy FDA-approved for patients with laBCC or mBCC previously treated with an HHI or for whom an HHI is not appropriate.²⁸⁴ Cemiplimab is a recommended treatment option for certain patients with advanced BCC including in the neoadjuvant setting for laBCC. A centrally reported, multicenter, phase II, open-label trial tested cemiplimab-rwlc (N = 84) for patients with laBCC where local invasion precluding complete resection or in locations for which surgery may result in severe disfigurement or dysfunction and whose disease has progressed on or was intolerant to prior HHI therapy.²⁸⁵ This study reported a median follow-up of 15 months, objective response rate of 31%, and grade 3–4 TEAEs in 48% of patients, while SAEs occurred in 35% of patients.²⁸⁵

Due to the rarity of advanced cases, the literature on chemotherapy for BCC is limited to case reports.²⁸⁶⁻²⁹²



Follow-up

Follow-up for BCC should include a history and physical examination, along with a complete skin examination every 6 to 12 months for the first 5 years, and then at least annually for life. Imaging may be considered if clinical examination is insufficient for following the disease. Follow-up with a dermatologist is strongly recommended if any of the following criteria are met: past or imminent solid organ, marrow, or hematopoietic cell transplant; one or more cutaneous melanomas in the past 5 years; or four or more NMSCs within the past 5 years.

Imaging modality and targeted area should be at the discretion of the treating team based on the suspected extent of disease (ie, local, regional, metastatic). Histologic confirmation is sufficient to diagnose local recurrence, but imaging can be considered to assess extent of disease. As part of follow-up, the patients should be educated on sun protection and self-examination. For local recurrence, the primary

treatment pathway for high-risk BCC should be followed. For locally advanced, nodal metastases, and distant metastases, the appropriate path should be followed as found within *Advanced BCC* in the algorithm.

An estimated 30% to 50% of patients with BCC will develop another BCC within 5 years.^{126,129,293-296} This represents a 10-fold increase in risk compared to the general population.²⁹⁴ Patients with a prior BCC are also at increased risk of developing SCC and cutaneous melanoma.^{126,296} A prospective population-based cohort study found that development of a second BCC is most likely during the short-term follow-up period after diagnosis of the first lesion.¹³³ Therefore, close follow-up of patients with BCC in both the short- and long-term is critical.



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Basal Cell Skin Cancer

Table 1. Studies Comparing Superficial Therapies in Patients with Superficial BCC

Study	Histologic Subtype	Tumor Locations	Treatments (n)	Efficacy	Cosmetic Outcome
Phase III randomized trial Wang 2001 ²⁴⁸	Superficial and nodular	Trunk, limb, head, neck	Cryosurgery (39) ALA-PDT (44)	1-year recurrence: 15% } NS 25% }	Excellent: 8% } $P < .001$ 50% }
Randomized trial Basset-Seguín 2008 ²⁴⁹	Superficial	Trunk, limb, head, neck, face	Cryotherapy (58) MAL-PDT (60)	5-year recurrence: 20% } NS 22% }	Excellent: 16% } $P = .00078$ 60% }
Meta-analysis Roozeboom 2012 ²⁵¹	Superficial	Locations depend on individual studies	Imiquimod (1088) PDT (934)	1-year tumor-free survival: 87% } NS 84% }	NR
Randomized, single-blind, non-inferiority trial Jansen 2018 ²⁰⁵	Superficial	Trunk, limb, head, neck	MAL-PDT (202) Imiquimod cream (198) Fluorouracil cream (201)	Treatment success ^a : 63% } $P < .001$ 81% } NS 70% } $P = .04$	Good/excellent: 62% } All comparisons NS 61% } 58% }

MAL, methyl aminolevulinate; NR, not reported; NS, no statistically significant difference; PDT, photodynamic therapy.

^aTreatment success was defined as the product of the percent of patients with clearance at 3 months by the percentage with sustained clearance during the next 9 months.

Table 2. Hedgehog Pathway Inhibitors in Advanced BCC^a

Study	Design	Tx ^b	Patients, n		Objective Response Rate ^d		Duration of Response, Median ^c		Progression-free Survival, Median ^c		Overall Survival, Median ^c	
			laBCC	mBCC	laBCC	mBCC	laBCC	mBCC	laBCC	mBCC	laBCC	mBCC
BOLT – 42-month follow-up NCT01327053 ^{269,270,272,273}	Phase II RDB, CR	Soni 200 mg	66	13	56%	8%	26.1	24.0	22.1	13.1	NR	NR
		Soni 800 mg	128	23	46%	17%	23.3	NE	24.9	11.1	NR	NR
BOLT – 30-month follow-up NCT01327053 ^{271,274}	Phase II RBD, IR	Soni 200mg	66	13	71%	23.1%	15.7	17.7– 18.4	NR	NR	NR	NR
SHH4476g – 9-month follow-up NCT00833417 ²⁶¹	Phase I OL, CR	Vismo	63	33	43%	30%	7.6	7.6	9.5	9.5	NR	NR
ERIVANCE – 39-month follow-up NCT00833417 ²⁶²⁻²⁶⁴	Phase II OL, IR	Vismo	63	33	60%	49%	26.2	14.8	12.9	9.3	NE	33.4

laBCC, locally advanced BCC; mBCC, metastatic BCC; CR, centrally reviewed; IR, investigator reviewed; NE, not reached; NR, not reported; OL, open-label; RDB, randomized double-blind; Soni, sonidegib; Tx, treatment; Vismo, vismodegib.

^aTrials included patients with advanced BCC that was inappropriate for surgery or RT.

^bInhibitors were taken orally once daily. Vismodegib dose was 150 mg.

^cTimes are reported in months.

^dResponse criteria varied between studies.

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Discussion
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