

TeleECHO Series

PRINCIPLES OF TREATMENT:

The Multi-Disciplinary Approach to the Patient
with **Locally Advanced/Metastatic BCC**

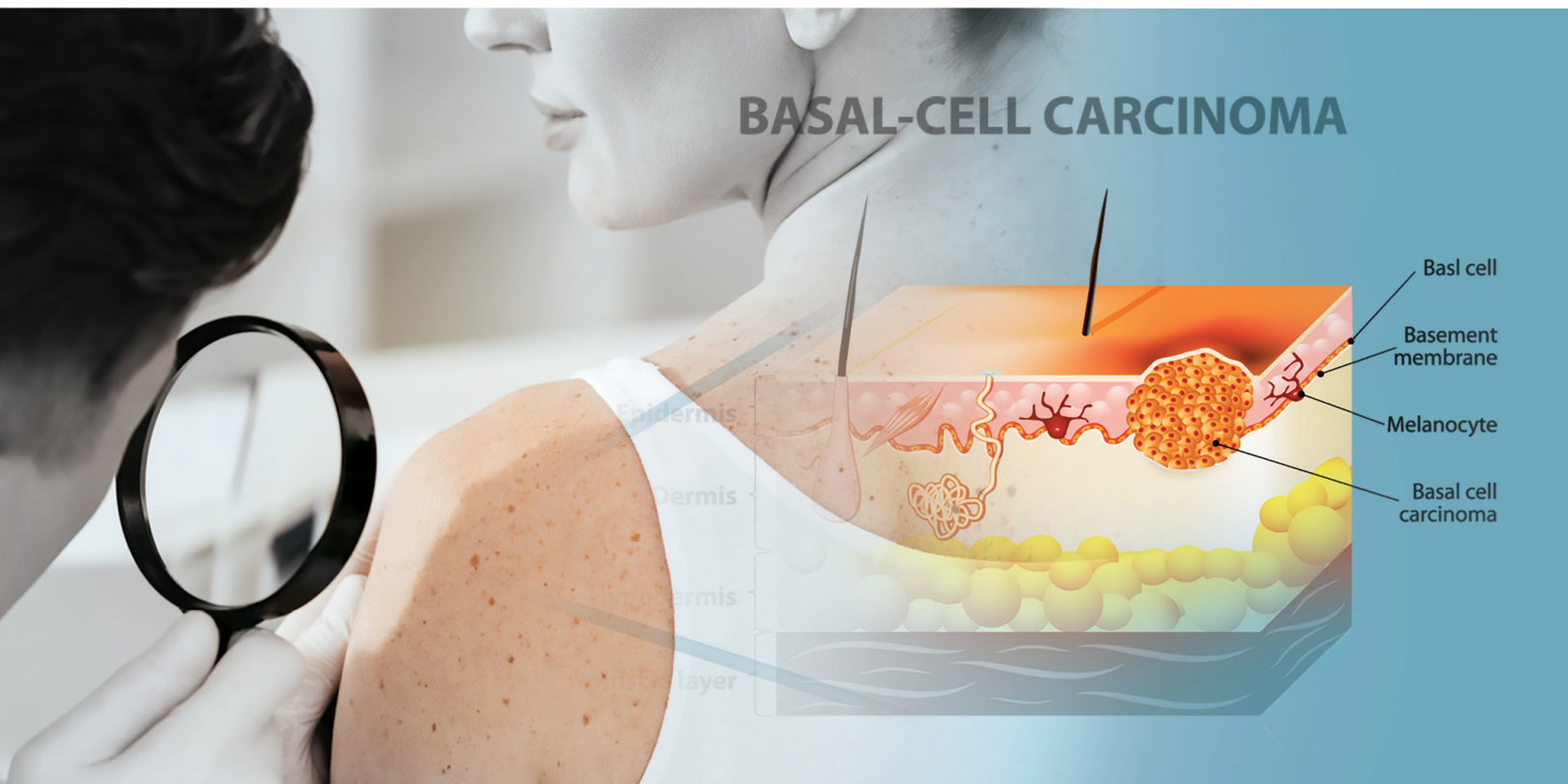
Tuesday, February 9, 2021

FACULTY

Anna Bar, MD

Associate Professor of Dermatology
Co-director of Mohs Micrographic Surgery
Oregon Health & Science University (OHSU)
Portland, OR

BASAL-CELL CARCINOMA





AGENDA

- Basal Cell Carcinoma: Introduction
 - Epidemiology/statistics
 - Prognostic factors
 - Staging
 - NCCN guidelines
 - Considerations for aggressive BCC
- BCC Failing or Not Amenable to Surgery or Radiation
 - Sonic hedgehog pathway
 - Vismodegib and sonidegib
 - Hedgehog side effects
 - Alternate dosing regimens
- When Disease Progresses on Hedgehog Inhibition
 - Immunotherapy and immune checkpoints
 - Cemiplimab, pembrolizumab, etc.
- Case Study Presentation
- Conclusions

***TeleECHO Series: Principles of Treatment:
The Multi-Disciplinary Approach to the Patient With Locally
Advanced/Metastatic BCC***

FACULTY CHAIR

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PROGRAM OVERVIEW

These live virtual TeleECHO® sessions will be a faculty-led didactic and case-based lecture focusing on treatment and management of patients with patients with advanced/metastatic basal cell skin cancer.

TARGET AUDIENCE

This activity is designed to meet the educational needs of medical oncologists, onco-dermatologists, dermatologists, Mohs surgeons, and other healthcare practitioners who care for patients with advanced/metastatic basal cell skin cancer.

LEARNING OBJECTIVES

Upon completion of the program, attendees should be able to:

- Discuss the pathogenesis of metastatic BCC and the attendant involvement of both aberrant pathways and immune dysfunction
- Examine the epidemiology of basal cell carcinoma including prevalence, risk factors, and disease burden
- Apply NCCN practice guidelines for a multi-disciplinary approach to treatment of locally advanced and metastatic BCC
- Explore the clinical trial data for emerging immunotherapy agents in the treatment of advanced or metastatic basal cell skin cancer

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Purpose: This program would be beneficial for nurses involved in the care of patients with advanced/metastatic basal cell skin cancer. **CNE Credits:** 1.0 ANCC Contact Hour.

Accreditation Statement

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Anna Bar, MD receives consulting fees from Regeneron Pharmaceuticals. She received research funding to her institution from Castle, Pelle Pharm and Mavis.

Anokhi Jambusaria, MD, MSCE serves on the advisory board for Regeneron Pharmaceuticals.

Justin Leitenberger, MD has nothing to disclose.

Emily Ruiz, MD, MPH receives consulting fees from Sanofi, Pelle Pharm Inc., and Jounce Therapeutic. She discloses other commercial interests with Leo Pharma, Checkpoint Therapeutics, and Jounce Therapeutic.

Mary L. Stevenson, MD states she is a co-investigator for a Regeneron Pharmaceuticals, Inc. clinical trial and receives consulting fees from Two Sigma

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- Remember to direct all questions to the “co-host.” There is a toggle button above the typing space that allows you to specify the location of your message delivery.

TeleECHO Series: Principles of Treatment: The Multidisciplinary Approach to the Patient With Locally Advanced/Metastatic BCC

Anna Bar, MD

Associate Professor of Dermatology
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Oregon Health & Science University (OHSU)
Portland, OR

Disclosures

- **Dr. Bar** receives consulting fees from Regeneron Pharmaceuticals. She received funding to her institution from Castle, Pelle Pharm and Mavis.
- During the course of this lecture, Dr. Bar may mention the use of medications for both US Food and Drug Administration (FDA)-approved and non-FDA-approved indications
- Case study graphics are not to be copied, reproduced, or distributed in any way

This activity is supported by an independent medical education grant from Regeneron Pharmaceuticals, Inc. and Sanofi Genzyme.

Learning Objectives

- Discuss the pathogenesis of metastatic BCC and the attendant involvement of both aberrant pathways and immune dysfunction
- Examine the epidemiology of BCC including prevalence, risk factors, and disease burden
- Apply practice guidelines for a multidisciplinary approach to the treatment of locally advanced and metastatic BCC
- Explore the clinical trial data for emerging immunotherapy agents in the treatment of advanced or metastatic basal cell skin cancer

Aggressive BCC

Is there such a thing?

Does BCC have a Risk of Metastasis?

- Rates of .0028% to 0.55% reported¹⁻³
- In 1 study of predictive factors: 11,905 BCCs diagnosed between 2000 and 2009 at Brigham and Women's Hospital and Massachusetts General were screened³
- 248 cases \geq 2 cm assessed for risk factors and metastasis/death³
- 248 cases $<$ 2 cm randomly selected for comparison³

1. von Domarus H, Stevens PJ. *J Am Acad Dermatol*. 1984;10(6): 1043-1060. 2. Snow SN, et al. *Cancer*. 1994;73(2):328-335. 3. Morgan F, et al. *J Am Acad Dermatol*. 2020;83:832-838.

Factors predictive of recurrence, metastasis, and death from primary basal cell carcinoma 2 cm or larger in diameter

Frederick C. Morgan, BSPH,^a Emily Stamell Ruiz, MD, MPH,^a Pritesh S. Karia, MPH,^{a,b} Robert J. Besaw, MPH,^a Victor A. Neel, MD, PhD,^c and Chrysalynce D. Schmults, MD, MSCE
Boston, Massachusetts; Baltimore, Maryland; and Providence, Rhode Island

Multivariable logistic regression of ≥ 2 cm BCC tumors

Variable	Local Recurrence		Metastasis and/or Death	
	OR (95% CI)	P	OR (95% CI)	P
Location				
Other	1 (reference)		1 (reference)	
Head and neck	9.7 (3.0-31.3)		5.3 (1.2-23.2)	.026
Diameter				
Other	—	—	1 (reference)	
≥ 4 cm	—		11.9 (2.4-59.4)	.003
Tumor depth				
Other	1 (reference)		1 (reference)	
Beyond fat	3.1 (1.0-9.6)	.049	28.6 (6.7-121.0)	<.001
Treatment modality				
Other	1 (reference)		—	—
Mohs micrographic surgery	0.14 (.04-0.5)	.002	—	—

BCC > 4 cm on the head and neck, and invading beyond subcutaneous fat, may warrant more aggressive follow-up and radiologic imaging.

Morgan F, et al. *J Am Acad Dermatol*. 2020;83:832-838.

BCC Prognostic Factors

- < 2 cm group had no metastasis/death
- In ≥ 2 cm group, 3 significant predictors of metastasis/death:
 - Diameter ≥ 4 cm or larger
 - Depth beyond subcutaneous fat
 - Head/neck location

Morgan F, et al. *J Am Acad Dermatol*. 2019;83:832-838.

Staging BCC

Tumor Staging System	Definition
AJCC 8th Edition T Staging for Cutaneous Carcinoma of the Head and Neck	
T1	< 2 cm in greatest diameter
T2	≥ 2 cm but < 4 cm in greatest diameter
T3	≥ 4 cm in greatest diameter or minor bone invasion or perineural invasion or deep invasion
T4a	Tumor with gross cortical bone and/or marrow invasion
T4b	Tumor with skull bone invasion and/or skull base foramen involvement
BWH T Staging for BCC	
T1	Tumor diameter < 2 cm or tumor diameter ≥ 2 cm with 0-1 risk factors
T2	Tumor diameter ≥ 2 cm with 2-3 risk factors

Morgan F, Schmults C, et al. JAAD, in press.

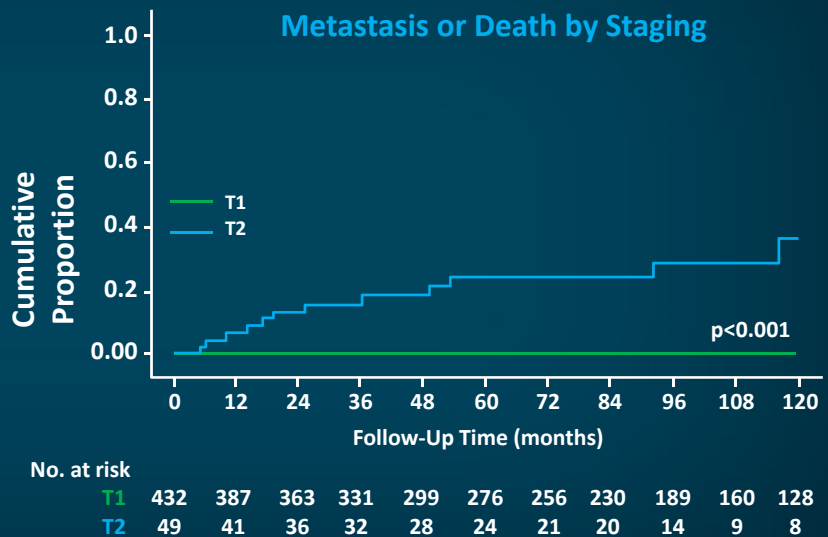
Staging BCC

- BWH T2 BCC:
 - Local recurrence: 47% (28%-70%)
 - Metastasis/death: 37% (21%-60%)

BWH T2:

- ≥ 4 cm + depth beyond fat
- ≥ 4 cm + head/neck
- ≥ 2 cm + depth beyond fat + head/neck

Cumulative Incidence of Metastasis or Death by Staging



Morgan F, Schmults C, et al. JAAD, in press.

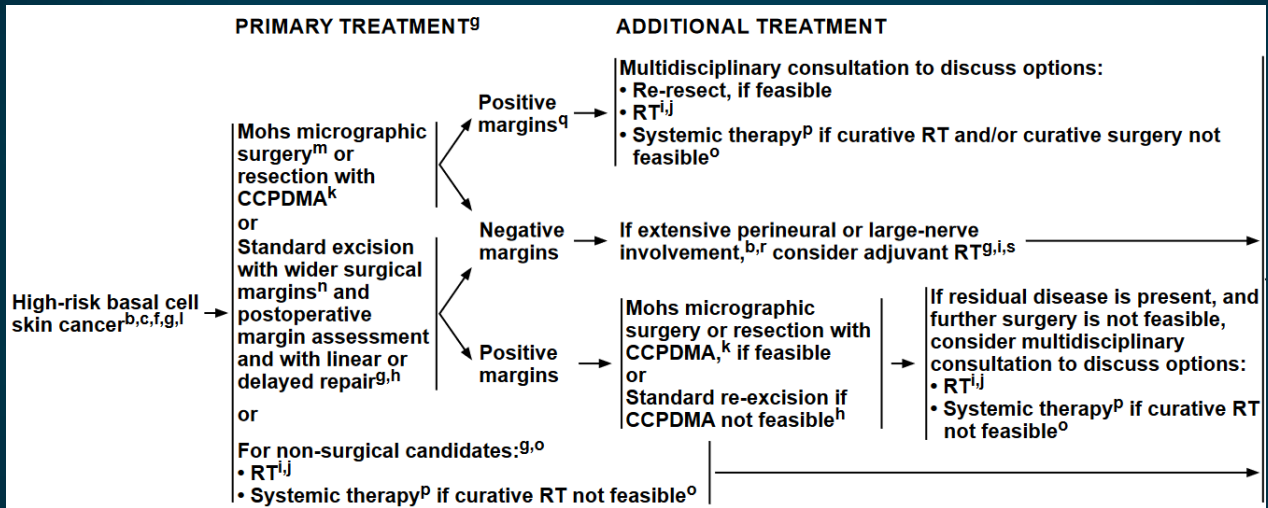
CCPDMA Cuts Recurrence Risk in Half for High-Risk BCC

	Recurrence Risk: Recurrent BCC, n (%)	Recurrence Risk: Aggressive Subtype BCC, n (%)
CCPDMA	132/2911 (4.5%)	97/3149 (3%)
SA	32/263 (12%)	17/234 (7%)
p value	p < .001	P < .001
Risk Ratio	.37 [.26-.54]	.42 [.26-.70]

CCPDMA = complete circumferential peripheral and deep margin assessment; SA = Standard Margin Assessment (vertical sections)

Fraga S, Waldman A. Complete marginal assessment in surgically excised keratinocyte carcinoma: A systematic review. Under review.

NCCN: Algorithm for High-Risk BCC



Schmults C, et al. NCCN Guidelines Version 1.2020: Basal Cell Skin Cancer. Available at: https://www.nccn.org/professionals/physician_gls/pdf/nmsc.pdf. Please see NCCN page above for footnotes and additional information/context for these guidelines.

Lack of Adherence to NCCN for High-Risk BCC/SCC

- Survey study of physicians from the 23 NCCN institutions
- ~50% response, N = 57
- Use of CCPDMA in a majority of NCCN high-risk cases
 - 14/15 Mohs surgeons
 - 10/16 other surgeons, dermatologists, radiation oncologists
 - 2/6 pathologists
- Reasons for not using CCPDMA
 - Surgeons defer to pathologists to determine appropriate margin assessment methods
 - Logistical difficulties (time-consuming, overnight stays, open wounds)

NCCN = National Comprehensive Cancer Network; SCC = squamous cell carcinoma.
Danesh M, et al. *Dermatologic Surgery*. 2020;46:1473-1480.

Conclusions: Aggressive BCC

- BWH T2 BCC may have a risk of local recurrence and metastasis in excess of 20%
 - ≥ 4 cm + depth beyond fat
 - ≥ 4 cm + head/neck
 - ≥ 2 cm + depth beyond fat + head/neck
- Do Mohs surgery (or CCPDMA if needed, OR resection)
- Beware of multiply recurrent BCC
- Radiologic surveillance for recurrence
- Consider off-label adjuvant systemic therapy in extreme or multiply recurrent cases

BCC Failing or Not Amenable to Surgery and Radiation

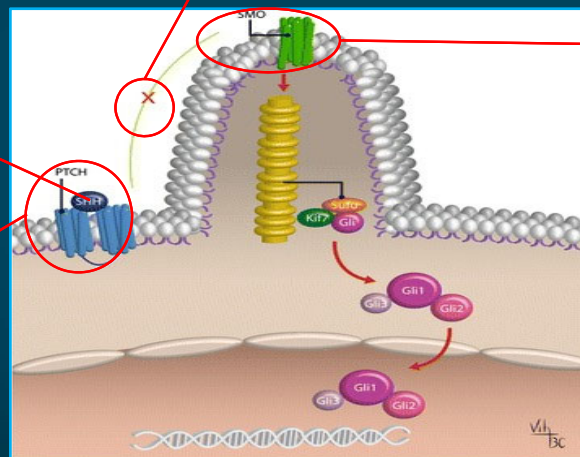
Sonic Hedgehog Pathway

4 Thus, blocking SHH attachment deactivates the carcinogenesis cascade

1 Pathway activation occurs when SHH binds to PTCH at the cell membrane.

2 In response to this binding, PTCH no longer inhibits SMO.

3 SMO accumulates at the primary cilium and initiates the downstream signaling cascade.



SHH = Sonic hedgehog; PTCH = Patched (a multitransmembrane protein); SMO = Smoothened (a key transmembrane protein).
Carballo G, et al. *Cell Commun Signal.* 2018;16:11.

Vismodegib

- Indications
 - Metastatic BCC
 - Locally advanced BCC that has recurred following surgery, or in those who are not candidates for surgery/radiation
- Dosage: oral; 150 mg once daily
- Contraindications: none
- Boxed warning: embryo-fetal toxicity

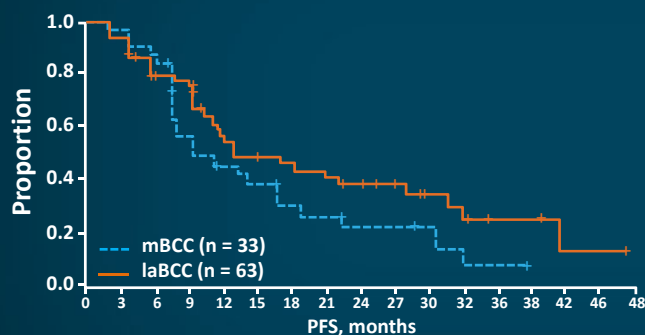
	mBCC (n = 33)	laBCC (n = 63)
ORR	30.3%	42.9%
Complete Response	0%	20.6%
Partial Response	30.3%	22.2%
Median Response Duration	7.6 months	7.6 months

mBCC = metastatic BCC; laBCC = locally advanced BCC; ORR = objective response rate.

Sekulic A, et al. *N Eng J Med.* 2012;366:2171-2179. Vismodegib (Erivedge®) PI 2020 (https://www.gene.com/download/pdf/erivedge_prescribing.pdf). Accessed December 10, 2020.

Vismodegib: ERIVANCE BCC Trial

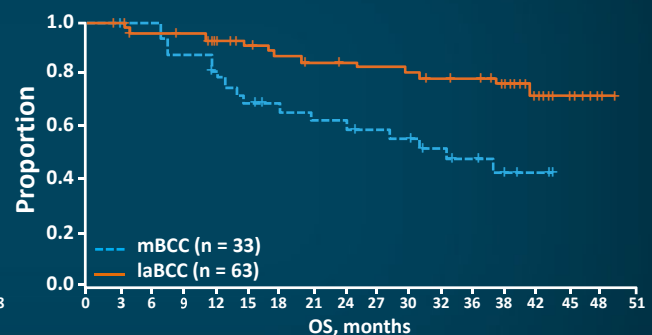
Progression Free Survival (PFS)



No. at risk

	0	3	6	9	12	15	18	21	24	27	30	32	36	38	42	46	48
Metastatic	33	31	26	16	12	10	7	6	4	4	3	1	1	0	0	0	0
Locally Advanced	63	55	42	38	24	19	17	16	13	10	7	5	3	3	1	1	0

Overall Survival (OS)



	0	3	6	9	12	15	18	21	24	27	30	32	36	38	42	45	48	51
Metastatic	33	33	33	29	25	22	20	18	18	16	15	12	10	7	2	0	0	0
Locally Advanced	63	61	58	56	52	48	45	43	42	41	40	38	37	25	13	8	2	0

Sekulic A, et al. *BMC Cancer.* 2017;17:332.

Sonidegib

- Indications
 - Locally advanced BCC that has recurred following surgery, or in those who are not candidates for surgery/radiation
- Dosage: oral; 200 mg once daily
- Contraindications: none
- Boxed warning: embryo-fetal toxicity

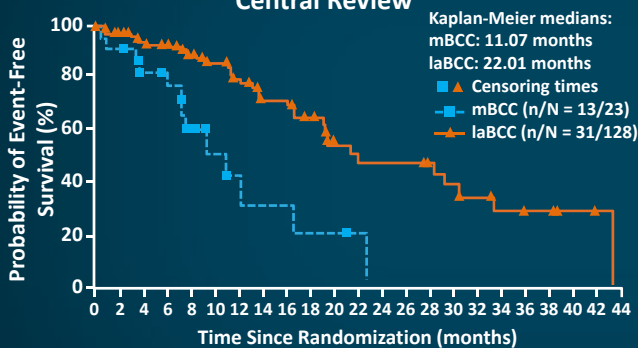
	laBCC (n = 66)
ORR	56.0%
Complete Response	5%
Partial Response	52%
Median Response Duration	26.1 months

Sonidegib (Odomzo®) PI 2019 (<https://www.odomzo.com/themes/custom/odomzo/global/pdfs/pi.pdf>).

Sonidegib: BOLT Study

Progression Free Survival

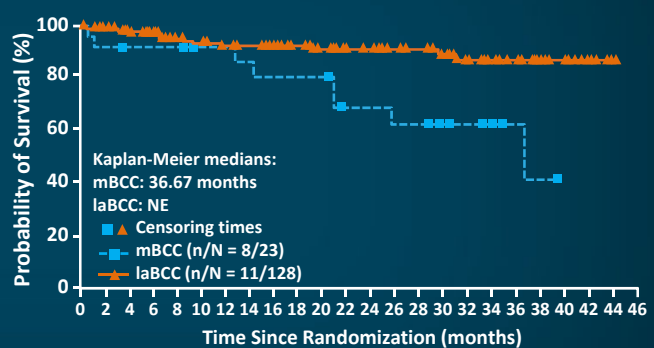
Central Review



No. of patients still at risk

Time (months)	0	2	4	6	8	10	12	14	16	18	20	22	24	26	28	30	32	34	36	38	40	42	44
Metastatic	23	21	16	15	9	6	4	3	2	2	1	0	0	0	0	0	0	0	0	0	0	0	0
Locally	128	98	80	69	57	45	39	30	29	23	15	14	13	13	11	9	7	5	4	4	2	1	0
Advanced																							

Overall Survival



No. of patients still at risk

Time (months)	0	2	4	6	8	10	12	14	16	18	20	22	24	26	28	30	32	34	36	38	40	42	44
Metastatic	23	21	20	20	20	17	16	15	15	15	11	11	10	10	8	6	5	3	2	0	0	0	0
Locally	128	123	113	101	91	83	77	75	71	66	61	55	54	50	47	43	32	28	23	16	11	5	1
Advanced																							

BOLT = Basal Cell Carcinoma Outcomes With LDE225 Treatment.
 Lear JT, et al. *J Eur Acad Dermatol Venereol.* 2018;32(3):372-381.

Efficacy, safety, and comparison of sonic hedgehog inhibitors in basal cell carcinomas: A systematic review and meta-analysis

Pingxing Xie, MD, PhD, and Philippe Lefrançois, MD, PhD
Montreal, Canada

- 16 studies: quantitative meta-analysis of safety and efficacy
- Locally advanced BCC
 - Overall Response Rate: comparable for vismodegib and sonidegib (69% vs 57%)
 - Complete Response Rate: superior for vismodegib (31% vs 3%)
- Metastatic BCC
 - Overall Response Rate: superior for vismodegib (39% vs 15%)
- Side effects (combined prevalence)
 - 67%, 54%, and 58% for muscle spasms, dysgeusia, and alopecia, respectively; comparable for sonidegib and vismodegib
 - Upper GI distress more common in sonidegib than in vismodegib use
- Vismodegib favored over sonidegib in clinical practice

Xie P, Lefrançois P. *J Am Acad Dermatol* 2018;79:1089-1100.

Hedgehog Side Effects

- Hedgehog inhibitor side effects are not life-threatening but are nearly universal and often severe
- High rate of discontinuation in clinical trials (20%-50%)
 - Debilitating muscle pain/cramping
 - Loss of taste leading to major weight loss
 - Hair loss is often severe
- Alternatives to daily dosing are often used (though minimally studied)
 - A dosing regimen of 1 to 2 weeks on, 1 week off, is what we employ
 - Impact on efficacy unknown but such modifications are often needed to remain on treatment

Migden M, et al. *Lancet Oncology*. 2015;16:716-728. Dummer R, et al. *Br J Dermatol*. 2020;182(6):1369-1378. Sekulic A, et al. *BMC Cancer*. 2017;17:332. Sekulic A, et al. *N Eng J Med*. 2012;366:2171-2179. Villani A, et al. *Dermatol Ther*. 2020;10:401-412.

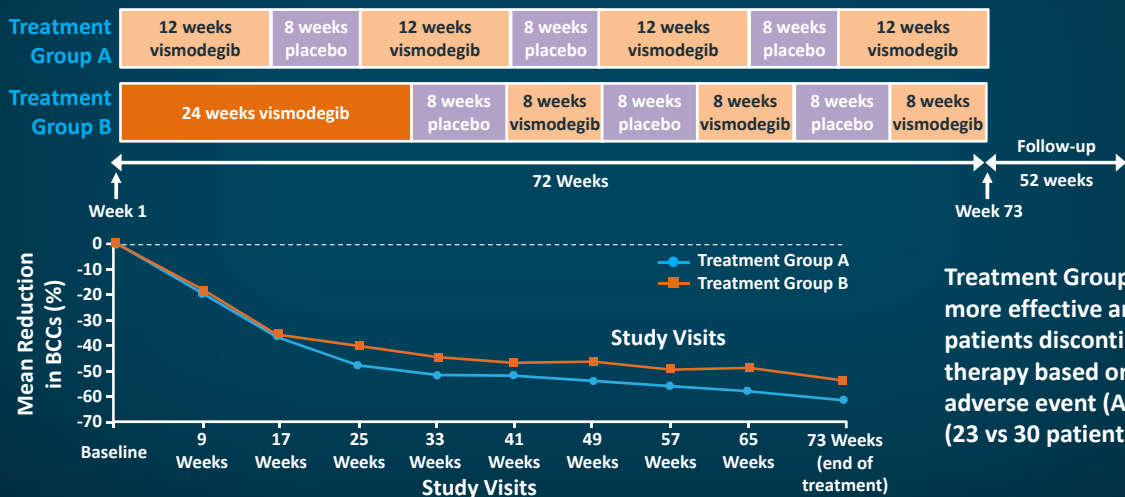
Alternate Dosing Regimens

- More tolerable than continuous therapy, with comparable effectiveness
- Impact on secondary resistance undefined
- BCC managed using hedgehog inhibitor (HHI) is comparable to chronic disease management or palliation
- May take months, weeks, or days off without loss of effectiveness



Two intermittent vismodegib dosing regimens in patients with multiple basal-cell carcinomas (MIKIE): a randomised, regimen-controlled, double-blind, phase 2 trial

Brigitte Dréno, Rainer Kunstfeld, Axel Hauschild, Scott Fosko, David Zloty, Bruno Labelle, Jean-Jacques Grob, Susana Puig, Frank Gilberg, Daniel Bergström, Damian R Page, Gary Rogers, Dirk Schadendorf



Dréno B, et al. *Lancet Oncol.* 2017; 18:404-412.

RESEARCH LETTER

A Novel Alternate Dosing of Vismodegib for Treatment of Patients With Advanced Basal Cell Carcinomas

Vismodegib Dosing Regimen, Response, and Adverse Effects

Patient No./Sex	BCC History	IVT Regimen (with/without medication = no. of cycles)	Length of Follow-Up, mo	BCC Response	Adverse Effects
1/M	>50 BCCs	1 wk/1 wk x 8 cycles; 1 wk/3 wks x 12 cycles	20	Decrease in number and size of lesions	Fatigue, arthralgia, arthritis, and alopecia; resolved after decreasing dose
2/M	Basal cell nevus syndrome	1 wk/2 wks x 10 cycles	12	Decrease in number and size of lesions	Fatigue, dysgeusia, and muscle aches
3/F	Multiple BCCs on the face	1 wk/1 wk x 6 cycles; 1 wk/2 wks x 7 cycles; 1 wk/3 wks x 14 cycles	24	Resolution	Alopecia; resolved with decreased dosage
4/M	Multiple recurrent and aggressive BCCs on face	1 wk/1 wk x 8 cycles; 1 wk/2 wks x 1 cycle; 1 wk/3 wks x 9 cycles	19	Resolution	Fatigue, dysgeusia, muscle spasms, and hand arthritis; improved with lowered dosage
5/F	Large periorbital BCC	1 wk/1 wk x 6 cycles; 1 wk/2 wks x 5 cycles	12	Decrease in BCC size	Fatigue, dysgeusia, muscle aches, and alopecia
6/M	Large BCC on nose	1 wk/1 wk x 6 cycles	7	Decrease in BCC size, followed by MMS	Fatigue, dysgeusia, diarrhea, and constipation
7/M	Incompletely excised periorbital BCC	1 wk/3 wks x 5 cycles	9	Resolution	Fatigue

7 patients: 1 week on/1 to 3 weeks off based on tolerability; 3 BCC clinically resolved, 4 decreased in size
No patients discontinued therapy based on adverse effects.

IVT = intermittent vismodegib therapy; MMS = Mohs micrographic surgery.
Becker LR, et al. *JAMA Dermatol.* 2017; 153:321-322.

Outcomes for Basal Cell Carcinoma Treated With Vismodegib Extended Alternate Day Dosing

- 8 patients
- Vismodegib 150 daily x 3 months, then extended alternate day dosing as tolerated
 - Every other day to once weekly dosing
 - Mean reduction in tumor size: ~50%
 - 4 patients experienced no side effects, 4 experienced mild side effects that
 - improved/resolved with further extended alternate day dosing



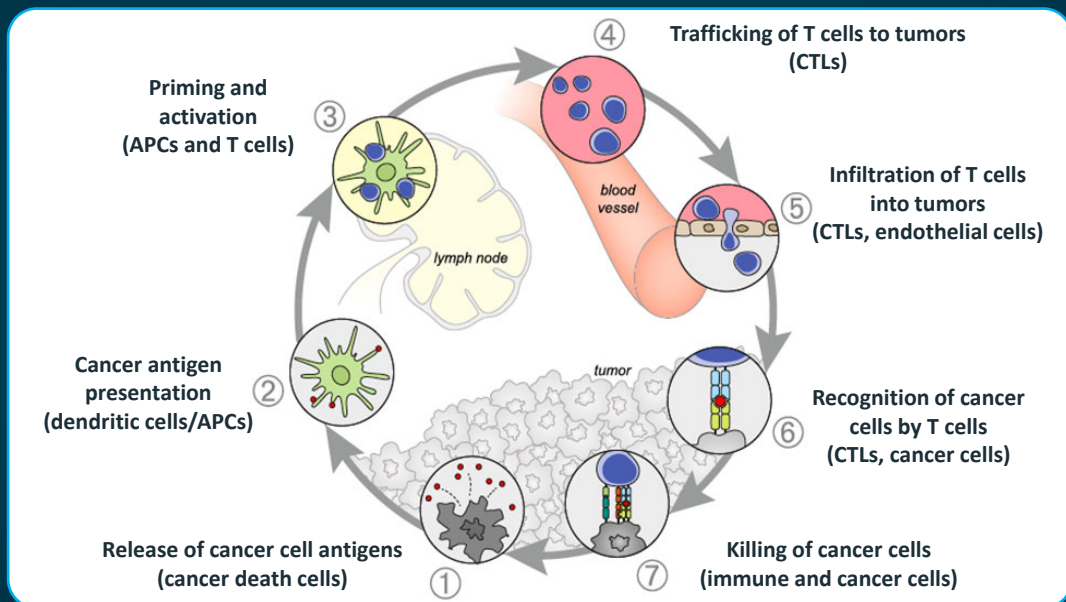
Before

Every 4 days x 10 months

Routt E, Ratner D. *Dermatol Surg.* 2020;46:1109-1112.

When Disease Progresses on Hedgehog Inhibition?

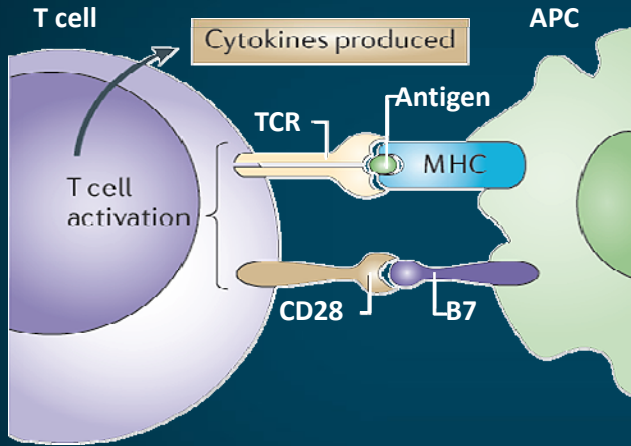
Immunotherapy: The Cancer-Immunity Cycle



APCs = antigen-presenting cells; CTLs = cytotoxic T lymphocytes.
Chen DS, Mellman I. *Immunity*. 2013;39:1-10.

Immune Checkpoints

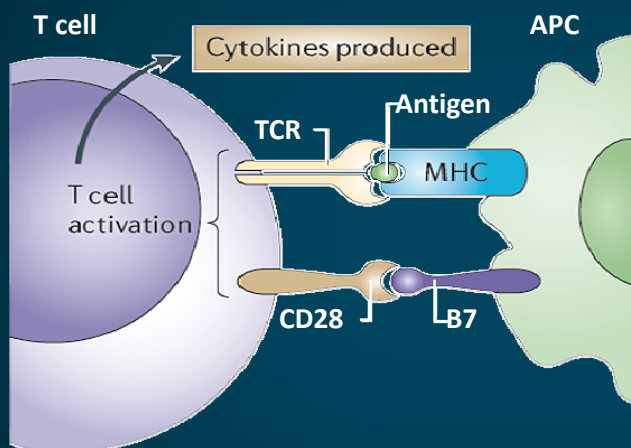
Activation of T lymphocytes



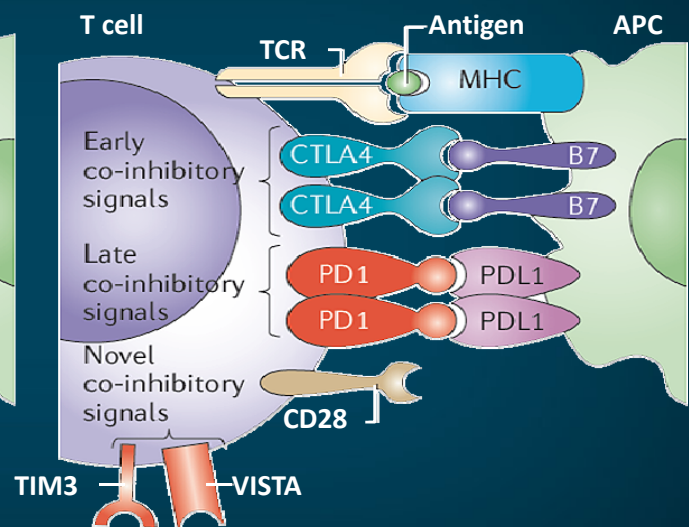
CTLA4 = cytotoxic T lymphocyte-associated protein 4; PD1 = programmed cell death 1; PDL1 = PD1 ligand 1; TIM3 = T cell immunoglobulin and mucin domain-containing protein 3; VISTA = V-domain immunoglobulin suppressor of T cell activation.
Sharma P et al. *Nat Rev Cancer*. 2011;11:805-812.

Immune Checkpoints

Activation of T lymphocytes

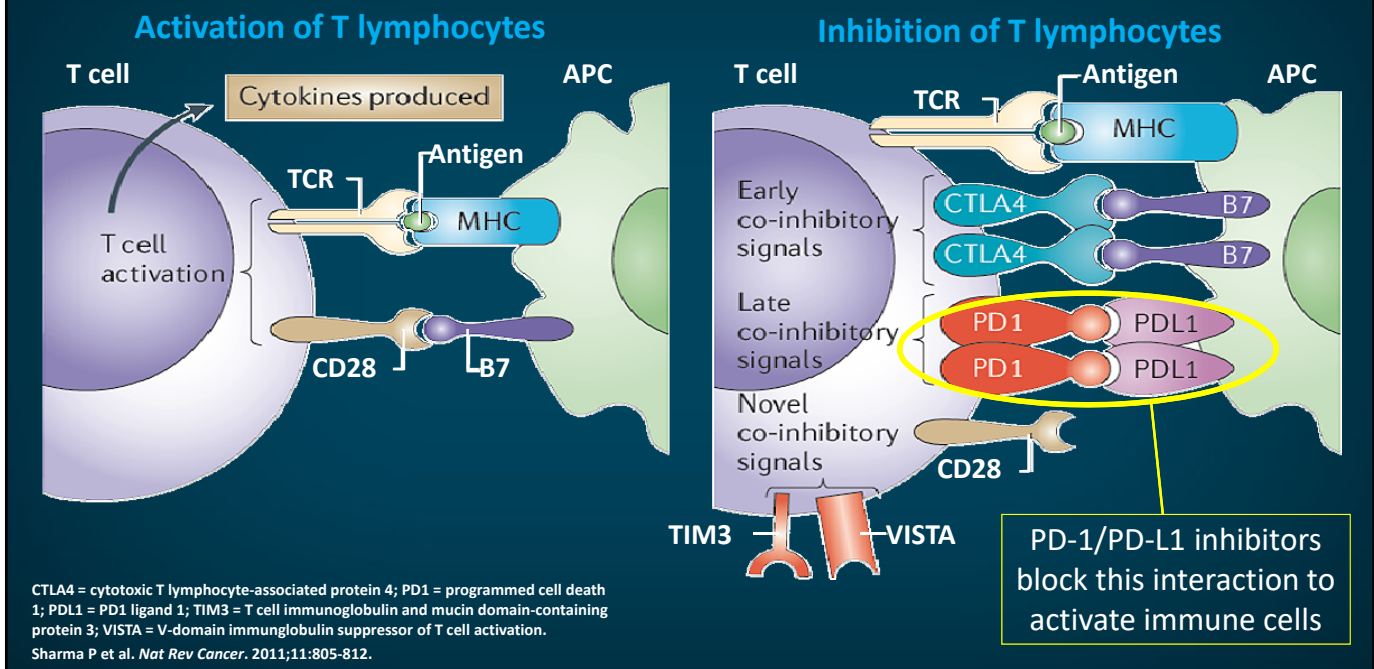


Inhibition of T lymphocytes



CTLA4 = cytotoxic T lymphocyte-associated protein 4; PD1 = programmed cell death 1; PDL1 = PD1 ligand 1; TIM3 = T cell immunoglobulin and mucin domain-containing protein 3; VISTA = V-domain immunoglobulin suppressor of T cell activation.
Sharma P et al. *Nat Rev Cancer*. 2011;11:805-812.

Immune Checkpoints



Cemiplimab: ESMO 2020

- Trial of cemiplimab for those not tolerating or progressing on vismodegib/sonidegib has concluded
 - Submitted to FDA for approval
 - Currently off-label for BCC
 - Approved for metastatic cutaneous SCC and laSCC

	laBCC (n = 84)
ORR	31.0%
Complete Response	6.0%
Partial Response	25.0%
Response	85% responses ongoing at 12 months
Estimated PFS (all patients)	19 months

ESMO = European Society for Medical Oncology.
Stratigos AJ, et al. *Ann Oncology*. 2020;31:S1175-S1176.

Not currently FDA approved for BCC

Cemiplimab: ESMO 2020

- Median baseline TMB: 58.2 (responding; n = 18) and 23.5 (non-responding; n = 38) mutations/Mb
- Responses occurred at all TMB levels
- Downregulation of MHC-I expression identified as an immune evasion mechanism in non-responding BCCs with high TMB
- Adverse events: fatigue (30%), diarrhea (24%) and pruritus (21%)
- 17% of patients discontinued due to AEs

TMB = tumor mutational burden; MHC = major histocompatibility complex.
Stratigos AJ, et al. *Ann Oncology*. 2020;31:S1175-S1176.

Pembrolizumab With/Without Vismodegib for Advanced BCC

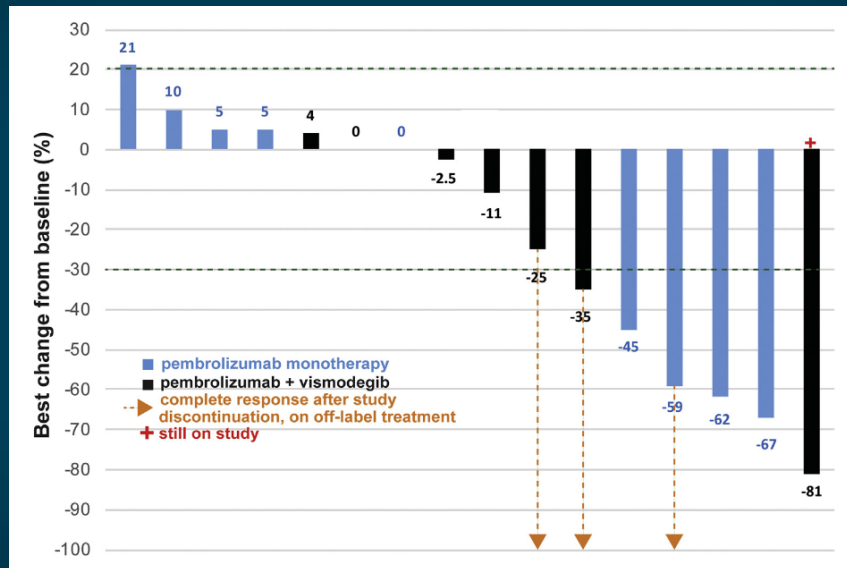
	All Participants (N = 16)	Pembrolizumab (n = 9)	Pembrolizumab + Vismodegib (n = 7)
ORR	38%	44%	29%
1-yr PFS	70%	62%	83%
1-yr OS	94%	89%	100%

- No life-threatening AEs or deaths during the study
- 3 grade 3 AEs occurred out of 98 AEs from 16 participants (1 case of hyponatremia attributed to pembrolizumab)
- 23 immune-related AEs (grade 1/2 dermatitis and fatigue most common)

Chang A, et al. *J Am Acad Dermatol*. 2019;564-566.

Not currently FDA approved for BCC

Pembrolizumab: Best % Change in BCC Diameter From Baseline



Chang A, et al. *J Am Acad Dermatol.* 2019;564-566.

Not currently FDA approved for BCC

Summary

- Advanced BCC is best managed using multidisciplinary consultation
- Diameter > 4 cm, head/neck location, and/or invasion beyond subcutaneous fat significantly increase risk for local recurrence, metastasis, and death
- Vismodegib outperforms sonidegib in a recent meta-analysis on safety and efficacy
- Alternate dosing regimens increase tolerability without compromising efficacy
- Immune checkpoint therapy may be useful in cases of hedgehog inhibitor progression or intolerance

Cases

Case 1

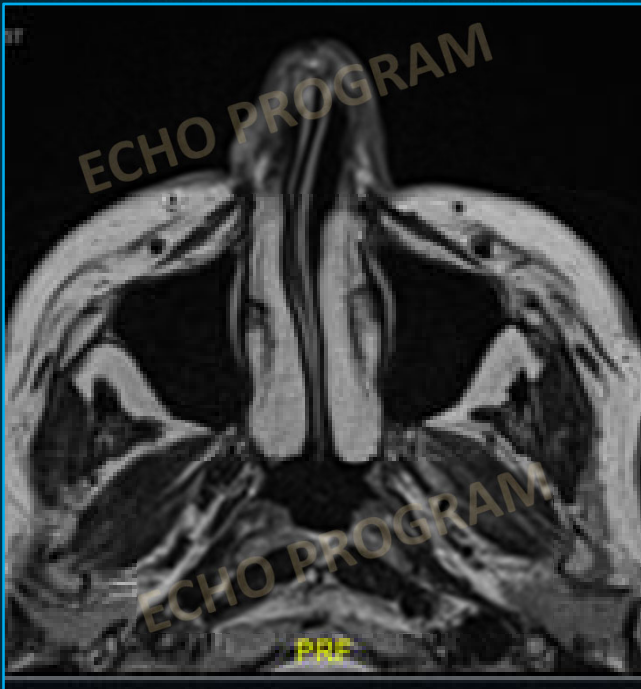
Locally Aggressive BCC: Case 1

Courtesy of Dr Emily Ruiz

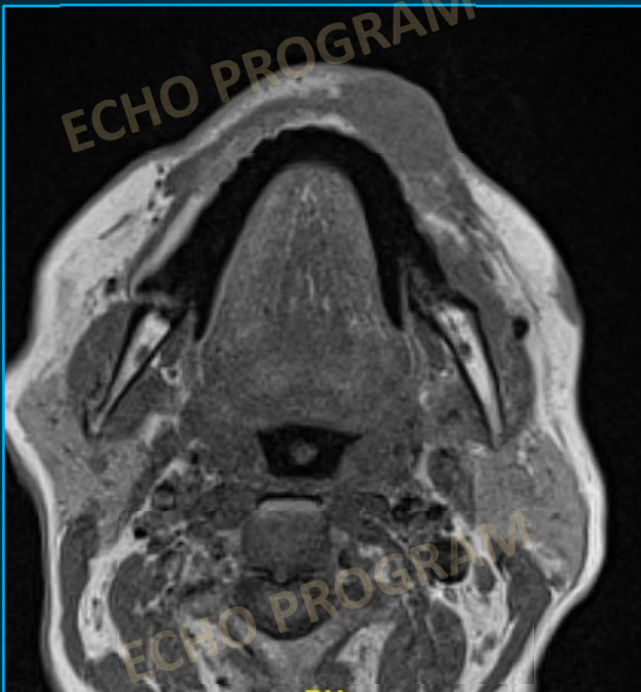


**A 69-year-old
healthy male**

How do you confirm BCC in this patient?



The tumor is inseparable from nasal bones and cartilage.



The tumor involves the mandible and 2 submandibular lymph nodes.

Locally Aggressive BCC: Polling Question

Which of the following would you choose for these BCCs?

1. Radiation
2. Systemic agent
3. Moh's surgery
4. Surgical excision coupled with margin assessment
5. Two different methods



Nose

- Mohs surgery for peripheral and some deep margin
- Bone resection in OR
- Multistage flap

Lip

- Lip resection in OR
- Bilateral neck dissections
- Mandibulectomy
- Fibula free flap
- Tracheostomy
- Skin graft

Surgery

Case 1: Next Steps

Before Vismodegib



S/P 3 Mo Vismodegib



Post Radiation Therapy



- Illustrates success but also imperfect response rate of hedgehog inhibitors
- Radiation may or may not be curative
- Off-label cemiplimab can be considered if he fails radiation

Case 3

Case 3: Locally Aggressive and Metastatic BCC

Courtesy of Dr. Emily Ruiz



- A 60-year-old man, healthy except for ethyl alcohol (EtOH), sometimes undomiciled
- Diagnosis in July 2017
- Histology: infiltrating BCC
- Positron emission tomography (PET) CT: avid tumor, avid left axillary, pectoral, and supraclavicular lymph nodes
- MRI: tumor encasing multiple large nerves

Locally Aggressive BCC: Polling Question

What Do You Offer as Preferred Therapy?

- Observation as progression has been slow over many years
- Excision via forequarter amputation
- Definitive radiation therapy
- Oral hedgehog inhibitor therapy
- Anti-PD1 immunotherapy

Locally Aggressive BCC: Polling Question

What Do You Offer as Preferred Therapy?

- A. Observation as progression has been slow over many years
–No, he's in pain and seeking care
- B. Excision via forequarter amputation
–Nodal metastasis makes this unlikely to control disease
- C. Definitive radiation therapy
–Field is too large
- D. Oral hedgehog inhibitor therapy
- E. Anti-PD1 immunotherapy



- Clinical response to vismodegib (July 2017-May 2018)
- MRI: progression of lymph node metastasis
- July 2018: patient switched to cemiplimab

What do you do now?



- Received 2 infusions of cemiplimab
 - Significant disease progression

- Restarted on vismodegib in September 2018: no response
- Switched to sonidegib in December 2019: no response
- Switched to ipilimumab/nivolumab in March 2019: no response
- Intratumoral flu vaccine in June 2019 (1 injection): no response
- Expired in July 2019 (tumor-related sepsis)



Conclusions

- BCC is the most common form of skin cancer worldwide
- BCCs are slow-growing and rarely metastasize; however, they are locally invasive and can be destructive
- BCC pathogenesis and the implications of aberrant pathways and immune dysfunction continue to be clarified
- Therapy selection depends on the patient's age and gender, along with site, size, and type of lesion
- Current mainstay of BCC treatment involves surgical modalities (excision, electrodesiccation and curettage [EDC], cryosurgery, Mohs surgery)
- Systemics include the HHIs vismodegib and sonidegib; there is no FDA-approved therapeutic option post-HHI use for patients with laBCC; PD1/PDL1 inhibitors are showing promise

Basal Cell Carcinoma: Identification and Management

Resource	Address
Ascierto PA, et al. Immunotherapy in non-melanoma skin cancer: Updates and new perspectives. <i>Drugs Context</i> . 2019;8:212583.	https://www.ncbi.nlm.nih.gov/pmc/articles/PMC6434981/
Carballo GB, et al. A highlight on sonic hedgehog pathway. <i>Cell Commun Signal</i> . 2018;16:11.	https://www.ncbi.nlm.nih.gov/pmc/articles/PMC5861627/
Cameron MC, et al. Basal cell carcinoma: Epidemiology; pathophysiology; clinical and histological subtypes; and disease associations. <i>J Am Acad Dermatol</i> . 2019;80:303-317.	https://pubmed.ncbi.nlm.nih.gov/29782900/
Cameron MC, et al. Basal cell carcinoma: Contemporary approaches to diagnosis, treatment, and prevention. <i>J Am Acad Dermatol</i> . 2019;80:321-339.	https://pubmed.ncbi.nlm.nih.gov/29782901/
Chang ALS, et al. Pembrolizumab for advanced basal cell carcinoma: An investigator-initiated, proof-of-concept study. <i>J Am Acad Dermatol</i> . 2019;80:564-566.	https://www.jaad.org/article/S0190-9622(18)32471-X/pdf
Chen DS, et al. Oncology meets immunology: The cancer-immunity cycle. <i>Immunity</i> . 2013;39:1-10.	https://pubmed.ncbi.nlm.nih.gov/23890059/
Danesh MJ, et al. Adherence to the National Comprehensive Cancer Network Criteria of Complete Circumferential Peripheral and Deep Margin Assessment in Treatment of High-Risk Basal and Squamous Cell Carcinoma. <i>Dermatologic Surg</i> . 2020;46:1473-1480.	https://pubmed.ncbi.nlm.nih.gov/32149872/
Dréno B, et al. Two intermittent vismodegib dosing regimens in patients with multiple basal-cell carcinomas (MIKIE): A randomised, regimen-controlled, double-blind, phase 2 trial. <i>Lancet Oncol</i> . 2017;18:404-412.	https://www.thelancet.com/journals/lanonc/article/PIIS1470-2045(17)30072-4/fulltext

Dummer 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. <i>Br J Dermatol</i> . 2020;182:1369-1378.	https://pubmed.ncbi.nlm.nih.gov/31545507/
McDaniel B, et al. Basal Cell Carcinoma. In: StatPearls [Internet]. Treasure Island, FL: StatPearls Publishing. Updated November 20, 2020.	https://www.ncbi.nlm.nih.gov/books/NBK482439/
Migden MR, et al. Emerging trends in the treatment of advanced basal cell carcinoma. <i>Cancer Treat Rev</i> . 2018;64:1-10.	https://pubmed.ncbi.nlm.nih.gov/29407368/
Morgan FC, et al. Factors predictive of recurrence, metastasis, and death from primary basal cell carcinoma 2 cm or larger in diameter. <i>J Am Acad Dermatol</i> . 2020;83:832-838.	https://pubmed.ncbi.nlm.nih.gov/31600531/
Paulson KG, et al. Immunotherapy for skin cancer. <i>Int Immunol</i> . 2019;31:465-475.	https://pubmed.ncbi.nlm.nih.gov/30753483/
Sekulic A, et al. Long-term safety and efficacy of vismodegib in patients with advanced basal cell carcinoma: final update of the pivotal ERIVANCE BCC study. <i>BMC Cancer</i> . 2017;17:332.	https://bmccancer.biomedcentral.com/articles/10.1186/s12885-017-3286-5
Sekulic A, et al. Efficacy and safety of vismodegib in advanced basal-cell carcinoma. <i>N Eng J Med</i> . 2012;366:2171-2179.	https://www.nejm.org/doi/full/10.1056/nejmoa1113713
Stratigos AJ, et al. LBA47 Primary analysis of phase II results for cemiplimab in patients (pts) with locally advanced basal cell carcinoma (laBCC) who progress on or are intolerant to hedgehog inhibitors (HHIs). <i>Ann Oncology</i> . 2020;31(suppl 4):S1175-S1176.	https://www.annalsofoncology.org/article/S0923-7534(20)42359-2/abstract
Stratigos AJ, et al. European interdisciplinary guideline on invasive squamous cell carcinoma of the skin: Part 2. Treatment. <i>Eur J Cancer</i> . 2020;128:83-102.	https://pubmed.ncbi.nlm.nih.gov/32113942/

Tanese K. Diagnosis and management of basal cell carcinoma. <i>Curr Treat Options Oncol</i> 2019;20:13.	https://pubmed.ncbi.nlm.nih.gov/30741348/
Totonchy M, et al. Emerging concepts and recent advances in basal cell carcinoma. <i>F1000Res</i> . 2017;6:2085.	https://www.ncbi.nlm.nih.gov/pmc/articles/PMC5717469/
Xie P, et al. Efficacy, safety, and comparison of sonic hedgehog inhibitors in basal cell carcinomas: A systematic review and meta-analysis. <i>J Am Acad Dermatol</i> . 2018;79:1089-1100.e17.	https://pubmed.ncbi.nlm.nih.gov/30003981/

Resources and Societies

Resource	Address
American Academy of Dermatology (AAD). <i>Skin Cancer</i> .	https://www.aad.org/media/stats-skin-cancer
American Cancer Society (ACS). <i>Basal and Squamous Cell Skin Cancer</i> .	https://www.cancer.org/cancer/basal-and-squamous-cell-skin-cancer.html
American Society of Clinical Oncology (ASCO).	https://www.asco.org/
Centers for Disease Control and Prevention (CDC). <i>What Is Skin Cancer?</i>	https://www.cdc.gov/cancer/skin/basic_info/what-is-skin-cancer.htm
National Comprehensive Cancer Network. <i>NCCN Clinical Practice Guidelines in Oncology. Basal Cell Skin Cancer. Version 1.2020</i> .	https://www.nccn.org/professionals/physician_gls/pdf/nmsc.pdf
Skin Cancer Foundation. <i>Basal Cell Carcinoma Overview</i> .	https://www.skincancer.org/skin-cancer-information/basal-cell-carcinoma/