

Indolent and Mantle Cell Lymphoma: Updating the Toolbox, Including and Beyond Anti-CD20 Antibodies

Sonali M. Smith, MD FASCO
Elwood V. Jensen Professor of Medicine
Chief, Section of Hematology/Oncology
Co-Leader, Cancer Service Line
Co-Director, Lymphoma Program
The University of Chicago

Overview

- Current landscape of treatment options for iNHL and MCL
- Recently approved treatments/modalities
- Emerging/future therapies

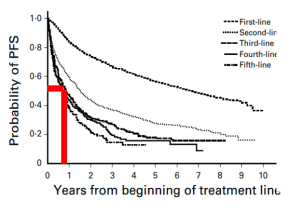


Follicular lymphoma



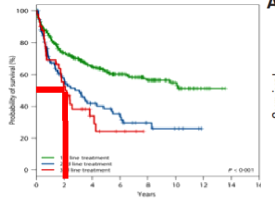
Follicular lymphoma snapshot: outcome after relapse

Median survival for follicular lymphoma approaches 20 years, but...



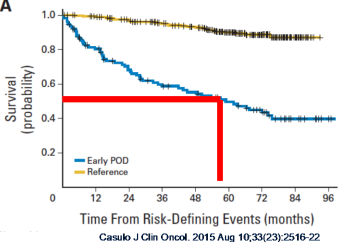
Link et al. BJH, 2018; 184: 660-63

PFS declines with each subsequent relapse



Rivas-Delgado et al. BJH 2018; 184: 753-59

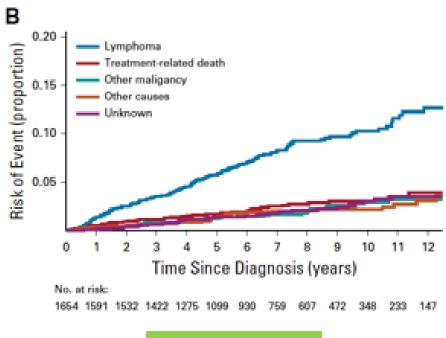
OS declines with each subsequent relapse



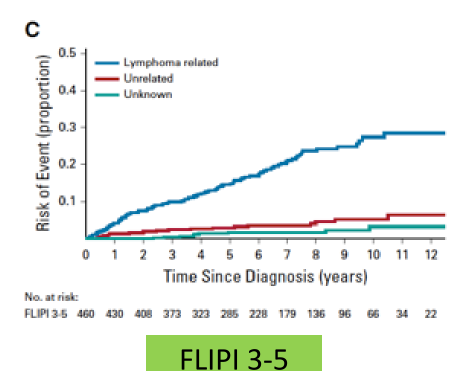
Early relapse (POD24) predicts 5y OS of 50%



FL remains important cause of death

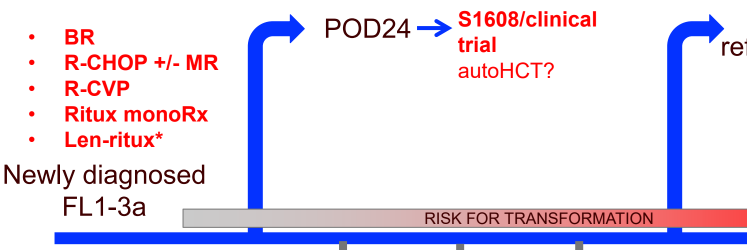








FL: Clinical categories and treatment options



Doublerefractory FL

- PI3Ki
- CAR-T?
- AlloHCT?

Biopsy critical at relapse to r/o transformation



HTB vs. LTB, symptomatic vs. asymptomatic

Kelap Relap

- ASCT
- Benda + Obinu or Ritux
- Len-rituximab
- PI3Ki

- Radiation therapy
 Radioimmunotherapy
- Ritux monoRx

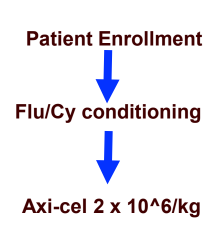
Tazemetostat

*not approved HTB = high tumor burden

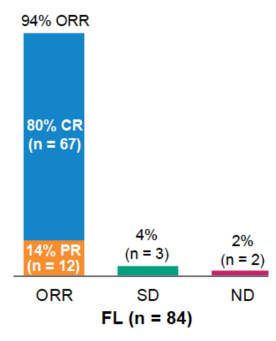
LTB = low tumor burden

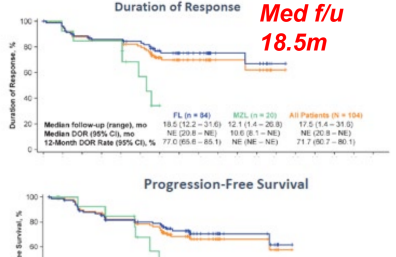
POD24 = early progression of disease w/i 24m of initial treatment

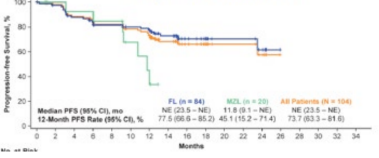
March 2021: Axi-cel approved for r/r FL with ≥ 2 lines of therapy (ZUMA-5)



Primary endpoint: ORR





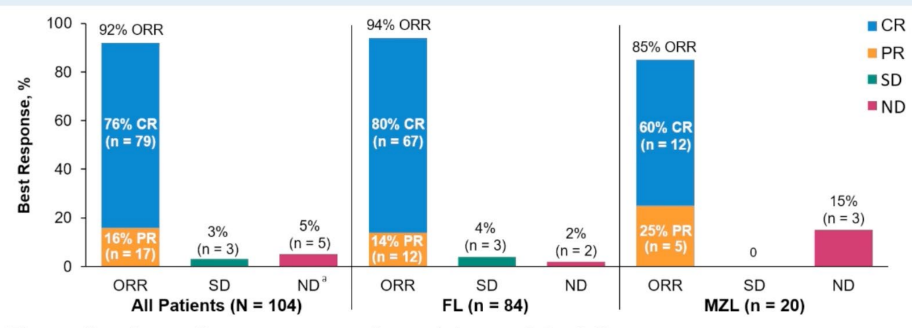


Figures and data courtesy of Caron Jacobsen, MD from ASH 2020



ZUMA-5: Response for FL and MZL

ORR by IRRC Assessment Was 92% (95% CI, 85 – 97); CR Rate Was 76% (95% CI, 67 - 84)

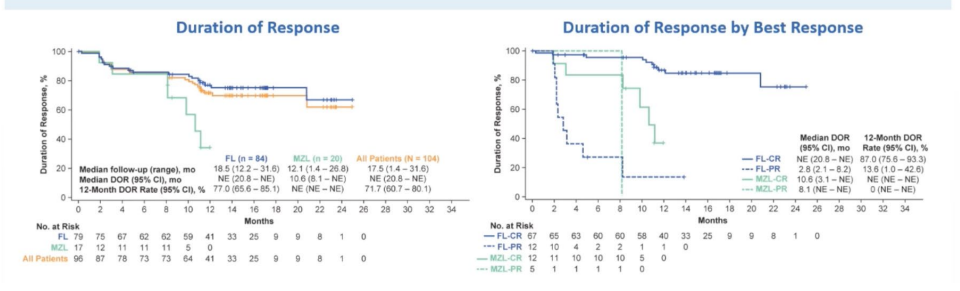


- The median time to first response was 1 month (range, 0.8 3.1)
- Among the 25 patients with FL who initially had a PR, 13 (52%) subsequently converted to a CR after a median of 2.2 months (range, 1.9 - 11.2)



ZUMA-5: Duration of response for FL and MZL

Duration of Response



Median f/u 17.5m
All 151 pts received intended treatment

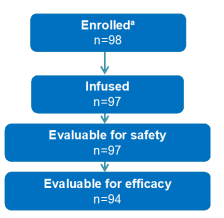


ZUMA-5: CRS and ICANS for FL and MZL

Parameter	FL (n = 124)	MZL (n = 22)	All Patients (N = 146)		
CRS, n (%) ^a					
Any grade	97 (78)	22 (100)	119 (82)		
Grade ≥ 3	8 (6)	2 (9)	10 (7)		
Most common symptoms of any grade, n/n (%)					
Pyrexia	94/97 (97)	20/22 (91)	114/119 (96)		
Hypotension	39/97 (40)	10/22 (45)	49/119 (41)		
Median time to onset (range), days	4 (1 – 15)	4 (1 – 9)	4 (1 – 15)		
Median duration of events (range), days	6 (1 – 27)	6 (2 – 14)	6 (1 – 27)		
Patients with resolved events, n/n (%)	96/97 (99) ^b	22/22 (100)	118/119 (99) ^b		
Neurologic events, n (%) ^a					
Any grade	70 (56)	17 (77)	87 (60)		
Grade ≥ 3	19 (15)	9 (41)	28 (19)		
Most common events of any grade, n/n (%)					
Tremor	36/70 (51)	9/17 (53)	45/87 (52)		
Confusional state	28/70 (40)	7/17 (41)	35/87 (40)		
Median time to onset (range), days	7 (1 – 177)	7 (3 – 19)	7 (1 – 177)		
Median duration of events (range), days	14 (1 – 452)	10 (2 – 81)	14 (1 – 452)		
Patients with resolved events, n/n (%)	67/70 (96)	14/17 (82)	81/87 (93)		



ELARA: Tisa-cel in r/r FL



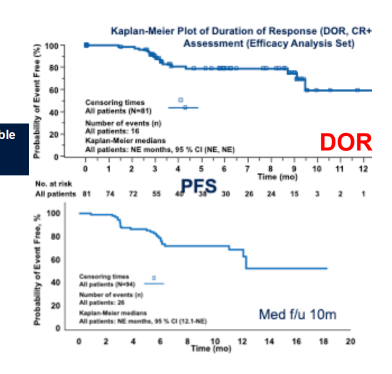
Vast majority of patients received cellular product

Lymphodepleting therapy could be Flu/Cy or bendamustine

18% of patients received tisagenlecleucel infusion in outpatient setting

Response Rate, %	Patients Evaluab for Efficacy ^b (n=94)
CR	66-0 ^b
PR	20.2
ORR (CR+PR)	86.2

Primary endpoint: CR





ELARA: phase 2 international trial of tisa-cel in FL (n=98)

Adverse Events, n (%)	Treated Patients N=97
Any AE (all grade)	92 (94.8)
AEs suspected to be drug-related	71 (73.2)
Any SAE	37 (38.1)
Suspected to be drug-related	26 (26.8)
Any grade 3/4 AE	68 (70.1)
Suspected to be drug-related	37 (38.1)
Death	3 (3.1)
Deaths due to study indication	3 (3.1)
Deaths within 30 days post infusion	0

	Treated Patients N=97		
AESI (within 8 weeks of infusion)	All grades, %	Grade ≥3, %	
Cytokine release syndrome ^a	48.5	0	
Serious neurological adverse reactions	9.3	1.0	
Infections	18.6	4.1	
Tumor lysis syndrome	1.0	0	
Prolonged depletion of B cells/ agammaglobulinemia	9.3	0	
Hematologic disorders including cytopenias			
Neutropenia ^{b,c}	28.9	24.7	
Anemia ^b	22.7	12.4	
Thrombocytopenia ^b	15.5	8.2	

Promising safety profile



Comparing cytokine release syndrome and neurotoxicity

	All grades <u>></u> Gr 3				
	ELARA				
CRS	48.5%	0%			
NT/ICANS	9.3% 1%				
Time to onset	8 days				
ZUMA-5					
CRS	78%	6%			
NT/ICANS	56% 15%				
Time to onset	4 days				

In ELARA trial:

Most CRS (75%) and all neurotoxicity (100%) occurred in patients with bulky disease



Key patient characteristics in ELARA and ZUMA-5 trials

ELARA

	All Patients (N=97)
Median age (range), y ≥65 y, n (%)	57.0 (29-73) 24 (24.7)
ECOG PS, n (%) 0 1	56 (57.7) 37 (38.1)
Bulky disease at study entry,° n (%)	63 (64.9)
CLIDI 23 et atruly cotor a (9/)	50 (50.0)
Median no. of prior therapies (range)	4 (2-13)
POD24 from first anti-CD20 mAb-containing therapy, ^d n (%)	58 (59.8)
Discould be 1907 (1)	05 (00.4)
Refractory to ≥2 regimens, f n (%)	74 (76.3)
Double reliactory,- If (78)	07 (09.1)
Prior therapy Anti-CD20 mAb and alkylating agents, h n (%) PI3K inhibitors, n (%) Lenalidomide and rituximab, n (%)	63 (64.9) 20 (20.6) 16 (16.5)

ZUMA-5

Characteristic	FL (n = 124)
Median age (range), years	60 (34 – 79)
≥ 65 years, n (%)	38 (31)
Male, n (%)	15 (39)
ECOG 1, n (%)	46 (37)
Stage III-IV disease, n (%)	106 (85)
≥ 3 FLIPI, n (%)	54 (44)
High tumor bulk (GELF criteria), n (%)*	64 (52)
Median no. of prior therapies (range)	3 (1 – 10)
≥ 3, n (%)	78 (63)
Prior PI3Ki therapy n (%)	39 (27)
Refractory disease, n (%) ^c	84 (68)
POD24 from first anti-CD20 mAb-containing therapy, n (%) ⁶	68 (55)
Prior autologous SCT, n (%)	30 (24)



Predictors of Response and Toxicity





POST-TREATMENT

Improved Response

- Low tumor burden, low LDH
- Low pretreatment inflammatory markers
- · Absence of medical comorbidities
- Lack of need for bridging therapy
- Proportion of CCR7+ and other early memory T-cells in the CAR product
- Faster doubling time in vitro
- Higher CAR T-cell peak to tumor burden ratio
- Absence of CD58 mutations
- Low tumor MDSCs
- High TILs
- Absence of MYC overexpression

Increased Toxicity

- High tumor burden, pretreatment LDH
- High pretreatment inflammatory markers
- ? High pretreatment monocyte levels
- High peak CAR T-cell levels
- High peak cytokine levels
- Markers of DIC (including fibrinogen levels)
- Early CRS

16



Which indolent lymphoma patients should be considered for CAR-T?

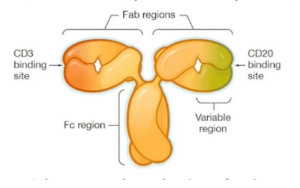
Patient identification is key! The vast majority of patients with FL do well without aggressive treatments □ Disease characteristics: early POD, double refractory, multiple prior regimens with sequentially shorter PFS Patient characteristics: ■ No upper age limit, adequate cardiac/renal/pulmonary/neurologic reserve ☐ Bulky disease and need for bridging therapy are poor prognostic factors No clear difference between axi-cel and tisa-cel in terms of efficacy in FL Marginal zone lymphoma needs more data

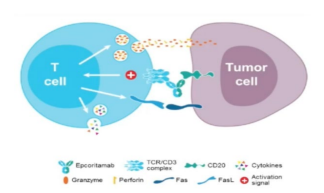


Very short follow up so far: is this a cure??

Emerging class of agents: bispecific antibodies

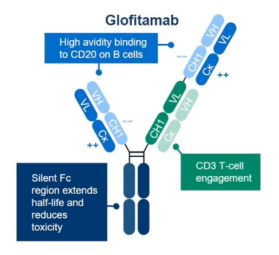
Odronextamab bispecific antibody structure

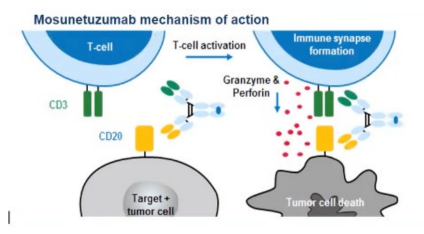




Distinct CD20 epitope





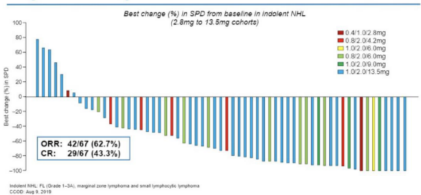


Olszewski ASH 2020 #401 Matasar ASH 2020 #2096 Philips ASH 2020 #1184 REGN1979 Bannerji ASH 2020 #400; Glofitamab Hutchings ASH 2020 #403 Epcoritamab Hutchings ASH 2020 #402

Bispecific antibodies in indolent lymphomas

Mosunetuzumab

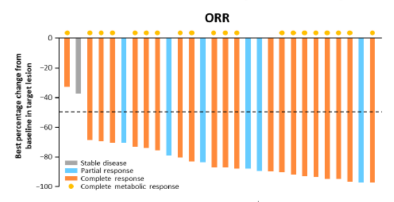
Objective response rate in indolent NHL

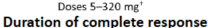


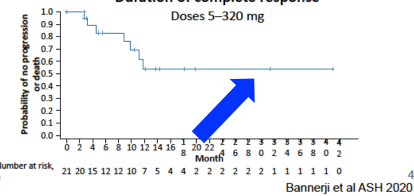
Schuster et al ASH 2019



Odronextamab (REGN1979)







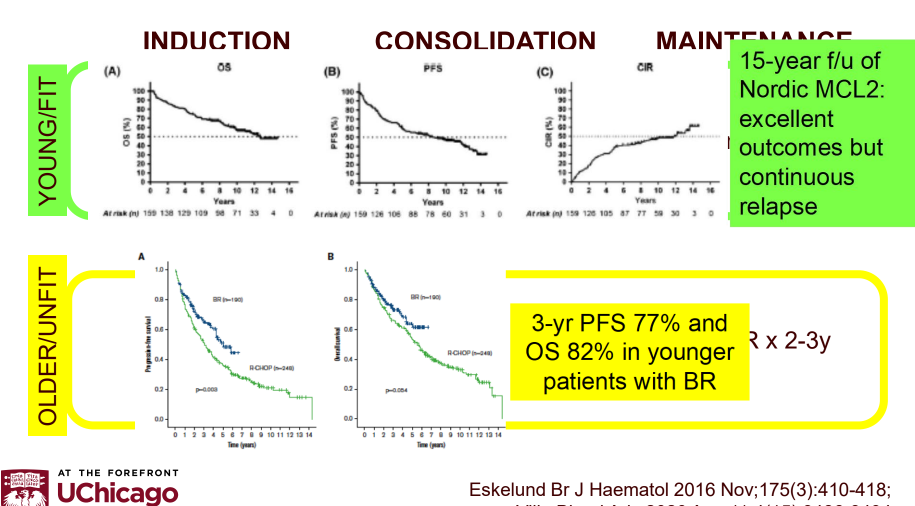


Mantle cell lymphoma

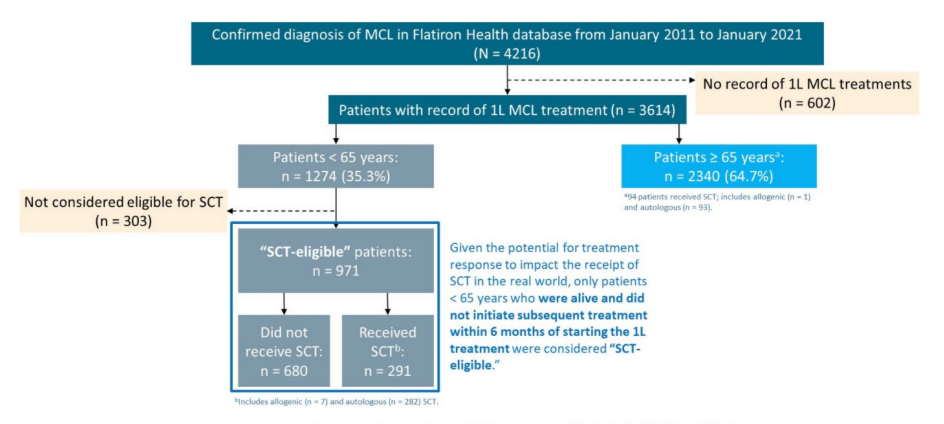


Mantle cell lymphoma: initial treatment approach

Medicine



Does the real-world experience match the data?



ASCO 2021, Martin P, et al.

Additional information can be viewed by accessing this link: https://www.oncologysciencehub.com/OncologyAM2021/ibrutinib/Martin/
Copies of this oral obtained through Quick Response (QR) Code are for personal use only and may not be reproduced without permission from ASCO® and the author of this oral.



Martin ASCO 2020

Real-world (rw) results: mantle cell lymphoma (n=3600)

- BR was most commonly used 1L treatment
 - Only one-third of pts < 65 received cytarabinecontaining regimen
 - Only 23% underwent SCT
- Med rwTTNT was 28m in pts <65y and 22m in pts ≥
 65y
 - Worse than reported in trials
- Despite lower use of SCT, there was no clear rwTTNT or rwOS benefit among SCT-eligible patients

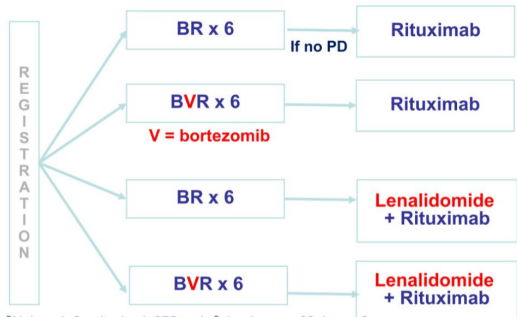


Implication for clinical trial development!!

Martin ASCO 2020

BR as a backbone for new regimens: ECOG-ACRIN E1411





Induction:

BR = bendamustine 90 mg/m²/d days 1, 2 + rituximab 375 mg/m² day 1, every 28 days x 6
BVR = BR + bortezomib 1.3 mg/m² days 1, 4, 8, 11 (later amended to 1.6 mg/m² days 1, 8), IV or SQ
Consolidation:

Rituximab 375 mg/m² every 8 weeks x 12 doses ± Lenalidomide 15 mg/d 21/28 days x 24 cycles

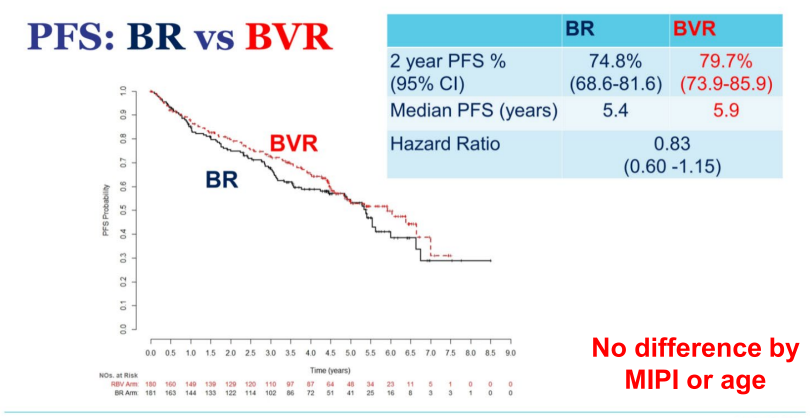
Presented By:

#ASCO21 | Content of this presentation is the property of the author, licensed by ASCO. Permission required for reuse.





Addition of bortezomib to BR does not improve outcomes







#ASCO21 | Content of this presentation is the property of the author, licensed by ASCO. Permission required for reuse.



Smith M ASCO 2021

Treatment approach in rel/ref MCL

- No data on sequencing
- List of options:
 - BTK inhibitors
 - Lenalidomide-rituximab
 - Venetoclax



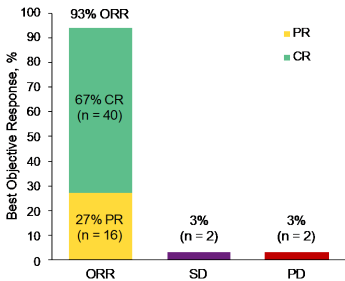
- Chemoimmunotherapy (if long duration of response to prior treatment)
- Bortezomib-based treatment
- Potential role of delayed autoHCT
- Allogeneic HCT
- CAR-T * (not yet FDA-approved) APPROVED!

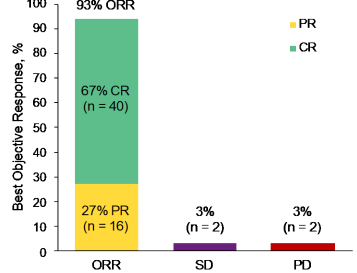


ZUMA-2: CAR-T in rel/ref MCL

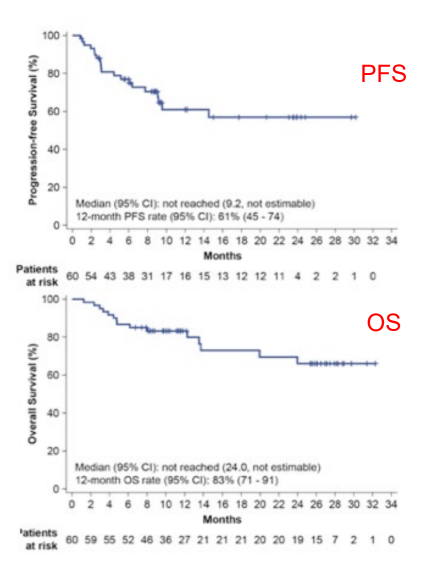
Key patient features:

- N=74 enrolled patients
- Med age 65 y
- $Ki-67 \ge 50\% 69\%$
- TP53 mut 17%
- Blastoid 25%









ZUMA-2: ASH 2020 multicenter phase 2 trial of KTE-X19 in r/r MCL (N=60)

Ongoing Response Rate Was Consistent Across Adverse Prognostic Subgroups

Overall 60 29 ————————————————————————————————————			Evaluable Patients	Responding Patie	nts	(95% CI)
 < 65 Years ≥ 65 Years 32 17 0.43 (0.24 - 0.63) ≥ 65 Years 32 17 0.53 (0.35 - 0.71) Sex Male 51 25 0.44 (0.14 - 0.79) Morphological characteristics Classical 35 16 0.46 (0.29 - 0.63) Pleomorphic 4 3 0.55 (0.19 - 0.99) Blastoid 14 5 0.50 (0.10 - 0.65) Ki-67 proliferation index ≥ 30% 40 21 0.53 (0.36 - 0.68) ≥ 50% 34 19 0.55 (0.38 - 0.73) Disease stage II-II 2 1 0.50 (0.01 - 0.99) III-IV 58 28 0.48 (0.35 - 0.62) S-MIPI score Low risk 25 11 0.44 (0.24 - 0.65) Intermediate or high risk 33 16 0.44 (0.24 - 0.65) Intermediate or high risk 33 16 0.50 (0.12 - 0.88) Mutation detected 3 0.50 (0.12 - 0.88) Mutation detected 30 17 0.0 0.1 0.2 0.3 0.4 0.5 0.6 0.7 0.8 0.9 1.0 0.57 (0.37 - 0.75) 0.07 (0.37 - 0.75) 0.09 0.1 0.2 0.3 0.4 0.5 0.6 0.7 0.8 0.9 1.0 0.57 (0.37 - 0.75) 0.57 (0.37 -		Overall	60	29	—	0.48 (0.35 - 0.62)
≥ 65 Years 32 17		Age at baseline				
Sex Male 51 25		< 65 Years	28	12	├	0.43 (0.24 - 0.63)
Male 51 25 4 0.49 (0.35 - 0.63) Female 9 4 0.44 (0.14 - 0.79) Morphological characteristics Classical 35 16 0.46 (0.29 - 0.63) Pleomorphic 4 3 0.75 (0.19 - 0.99) Blastoid 14 5 0.36 (0.13 - 0.65) Ki-67 proliferation index 230% 40 21 0.53 (0.36 - 0.68) ≥ 50% 34 19 0.50 (0.01 - 0.99) III-IV 58 28 0.40 (0.24 - 0.65) III-IV 58 28 0.44 (0.24 - 0.65) Intermediate or high risk 33 16 0.44 (0.24 - 0.65) Intermediate or high risk 33 16 0.44 (0.24 - 0.65) Mutation detected 6 3 0.50 (0.12 - 0.88) 0.57 (0.37 - 0.75) Mutation undetected 30 17 0.00 0.1 0.2 0.3 0.4 0.5 0.6 0.7 0.8 0.9 1.0 0.57 (0.37 - 0.75) MIPL, simplified Mantle Cell Lymphoma International Prognostic Index. Wang et al ASH 2020 Abstract 1120		≥ 65 Years	32	17	· ·	0.53 (0.35 - 0.71)
Female 9 4		Sex				
Morphological characteristics Classical 35 16		Male	51	25		0.49 (0.35 - 0.63)
Classical 35 16		Female	9	4		0.44 (0.14 - 0.79)
Pleomorphic 4 3		Morphological characteristic	cs		1	
Blastoid 14 5		Classical	35	16	· · · · · · · · · · · · · · · · · · ·	0.46 (0.29 - 0.63)
Ki-67 proliferation index 230%		Pleomorphic	4	3	<u> </u>	0.75 (0.19 – 0.99)
≥ 30% 40 21 0.53 (0.36 − 0.68) ≥ 50% 34 19 0.56 (0.38 − 0.73) Disease stage I-II 2 1 0.50 (0.01 − 0.99) III-IV 58 28 0.48 (0.35 − 0.62) S-MIPI score Low risk 25 11 0.44 (0.24 − 0.65) Intermediate or high risk 33 16 0.48 (0.31 − 0.66) TP53 mutation Mutation detected 6 3 0.50 (0.12 − 0.88) Mutation undetected 30 17 0.50 (0.12 − 0.88) Mutation undetected 30 17 0.50 (0.12 − 0.88) Mutation undetected 30 17 0.50 (0.12 − 0.88) MIPI, simplified Mantle Cell Lymphoma International Prognostic Index. Wang et al ASH 2020 Abstract 1120		Blastoid	14	5	· -	0.36 (0.13 - 0.65)
Disease stage I-II 2 1		Ki-67 proliferation index				
Disease stage		≥ 30%	40	21	<u> </u>	0.53 (0.36 - 0.68)
I-II		≥ 50%	34	19	├	0.56 (0.38 - 0.73)
III-IV 58 28		Disease stage				
S-MIPI score Low risk 25 11		1-11	2	1		0.50 (0.01 – 0.99)
Low risk 25 11 0.44 (0.24 – 0.65) Intermediate or high risk 33 16 0.48 (0.31 – 0.66) TP53 mutation Mutation detected 6 3 0.50 (0.12 – 0.88) 0.57 (0.37 – 0.75) Mutation undetected 30 17 0.00 0.1 0.2 0.3 0.4 0.5 0.6 0.7 0.8 0.9 1.0 0.57 (0.37 – 0.75) MIPI, simplified Mantle Cell Lymphoma International Prognostic Index. Wang et al ASH 2020 Abstract 1120		III-IV	58	28	├	0.48 (0.35 - 0.62)
Intermediate or high risk 33 16 0.48 (0.31 – 0.66) TP53 mutation Mutation detected 6 3 0.50 (0.12 – 0.88) Mutation undetected 30 17 0.57 (0.37 – 0.75) MIPI, simplified Mantle Cell Lymphoma International Prognostic Index. Wang et al ASH 2020 Abstract 1120		s-MIPI score				
## TP53 mutation Mutation detected 6 3		Low risk	25	11	 • 	0.44 (0.24 - 0.65)
Mutation detected 6 3		Intermediate or high risk	33	16	 	0.48 (0.31 - 0.66)
Mutation undetected 30 17 0.57 (0.37 – 0.75) 0.0 0.1 0.2 0.3 0.4 0.5 0.6 0.7 0.8 0.9 1.0 MIPI, simplified Mantle Cell Lymphoma International Prognostic Index. Wang et al ASH 2020 Abstract 1120		TP53 mutation				
0.0 0.1 0.2 0.3 0.4 0.5 0.6 0.7 0.8 0.9 1.0 MIPI, simplified Mantle Cell Lymphoma International Prognostic Index. Wang et al ASH 2020 Abstract 1120		Mutation detected	6	3	 	0.50 (0.12 - 0.88)
MIPI, simplified Mantle Cell Lymphoma International Prognostic Index. Wang et al ASH 2020 Abstract 1120		Mutation undetected	30	17	⊢	0.57 (0.37 - 0.75)
Wang et al ASH 2020 Abstract 1120					0.0 0.1 0.2 0.3 0.4 0.5 0.6 0.7 0.8 0.	9 1.0
	MIPI, simplified Ma	ntle Cell Lymphoma International Prog	nostic Index.		Ongoing Response Rate	
	L.		W	ang et al AS	SH 2020 Abstract 1120	



Ongoing Response Rate

ZUMA-2: ASH 2020 multicenter phase 2 trial of KTE-X19 in r/r MCL (N=60)

Duration of Response, Progression-Free Survival, and Overall Survival The medians for DOR, PFS, and OS were not reached after a median follow-up of 17.5 months DOR **PFS** OS 100 % Progression-free Survival, 80 Duration of Response, 60 40 0 2 4 6 8 10 12 14 16 18 20 22 24 26 28 30 32 34 36 0 2 4 6 8 10 12 14 16 18 20 22 24 26 28 30 32 34 36 38 Time, months Patients 55 47 43 40 39 35 24 18 13 13 13 12 12 12 11 1 1 1 Patients 60 53 44 42 39 35 30 22 13 13 13 13 12 12 12 4 1 1 0 Patients at risk 60 59 55 52 50 50 50 36 29 21 20 20 19 19 19 14 8 DOR **PFS** OS Median 15-Mo Rate Median 15-Mo Rate 15-Mo Rate Median (95% CI), mo (95% CI), % (95% CI), mo (95% CI), % (95% CI), mo (95% CI), % Evaluable pts(N = 60)NR (14 - NE)a $59(43 - 72)^a$ NR (10 - NE) 59 (45 – 71) NR (NE – NE) 76 (63 – 85) Pts in CR (n = 40)NR (14 - NE) 70(49 - 83)NR (15 - NE) 75(57 - 87)NR (NE - NE) 92(76 - 97)Pts in PR (n = 15)2(1-4)24(6-49)3(2-5)24(6-49)13 (3 - NE) 47(21 - 69)a Of 55 total responding patients. CR, complete response; DOR, duration of response; NE, not evaluable; NR, not reached; PFS, progression-free survival; PR, partial response; pts, patients; OS, overall survival Wang et al **ASH 2020** Abstract 1120



Take home points: indolent and mantle cell lymphomas

- Toolbox is growing (!)
- Advent of cellular therapy for indolent lymphomas
 Patient selection is critical
- Watch for bispecifics in indolent lymphomas
- New regimens for MCL on the horizon



Thank You



