



AT THE FOREFRONT

UChicago
Medicine

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

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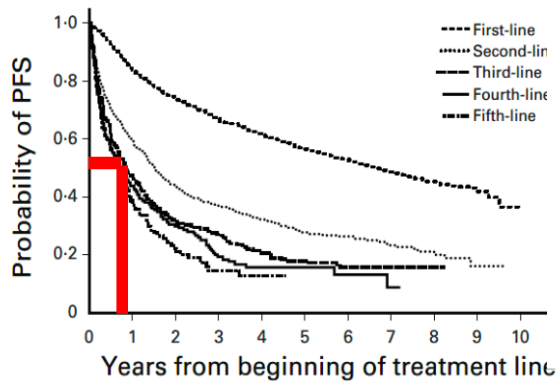
Overview

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

Follicular lymphoma

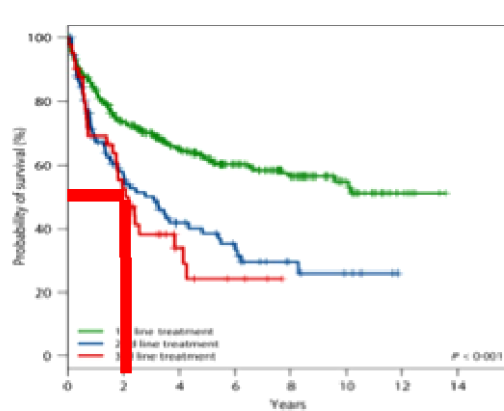
Follicular lymphoma snapshot: outcome after relapse

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



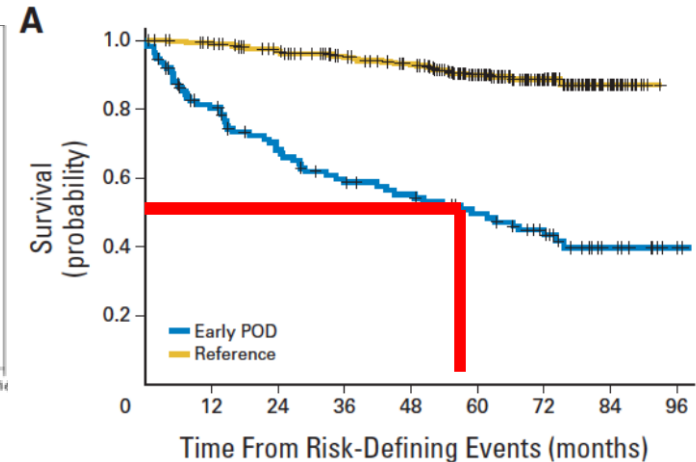
Link et al. *BJH*, 2018; 184: 660-63

PFS declines with each subsequent relapse



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

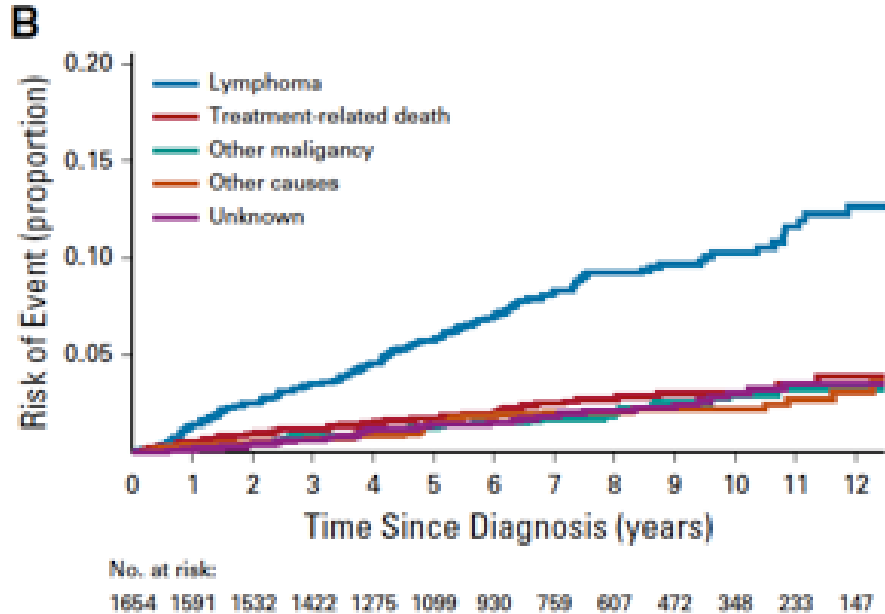
OS declines with each subsequent relapse



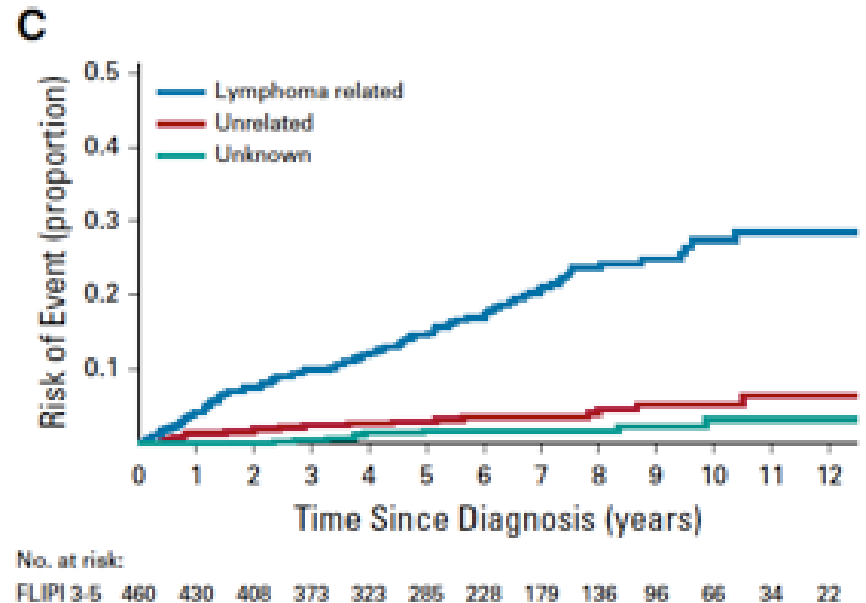
Casulo *J Clin Oncol*. 2015 Aug 10;33(23):2516-22

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

FL remains important cause of death

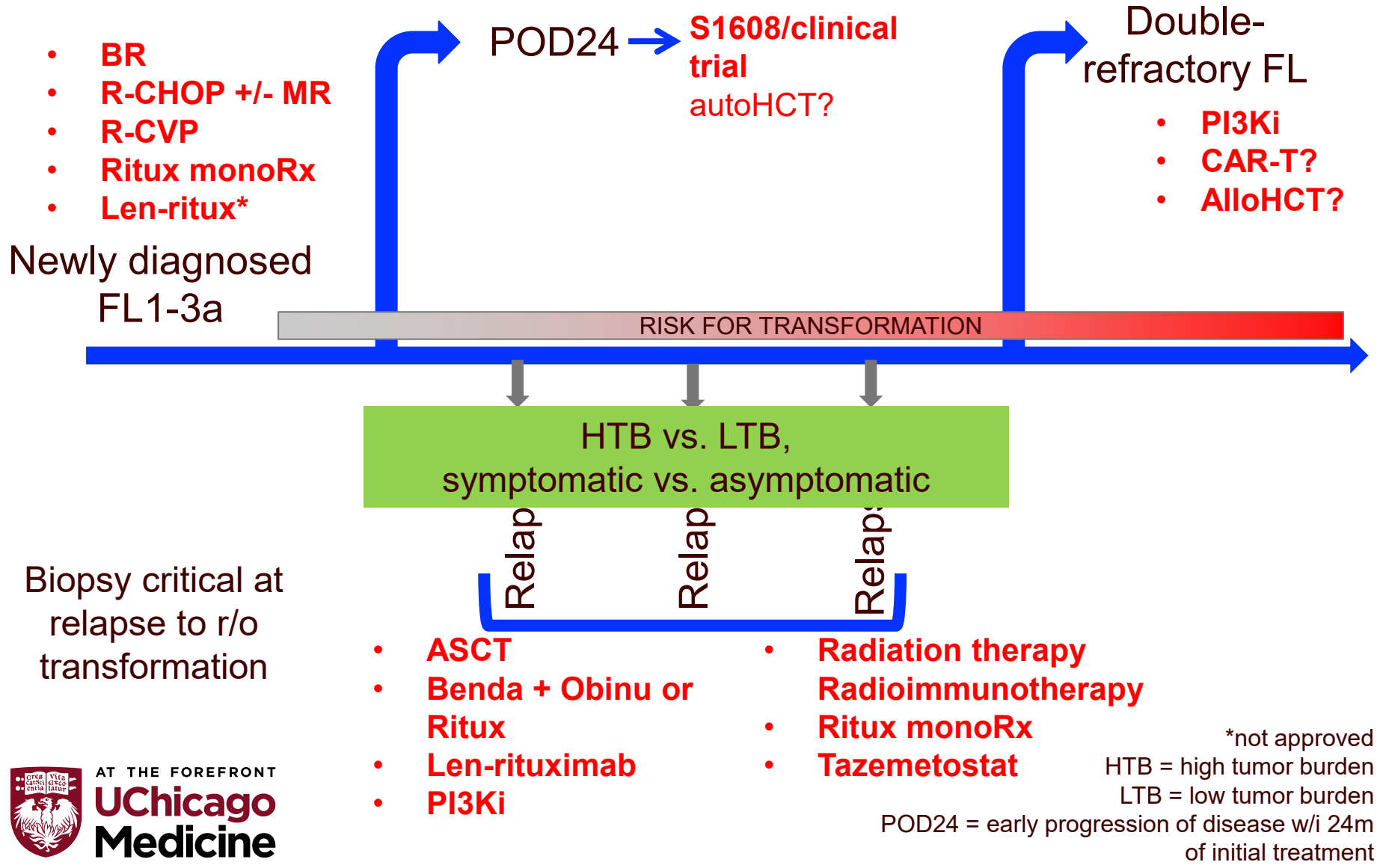


All patients

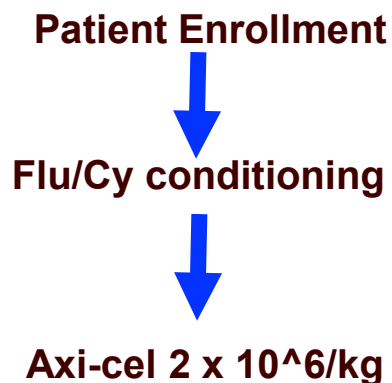


FLIPI 3-5

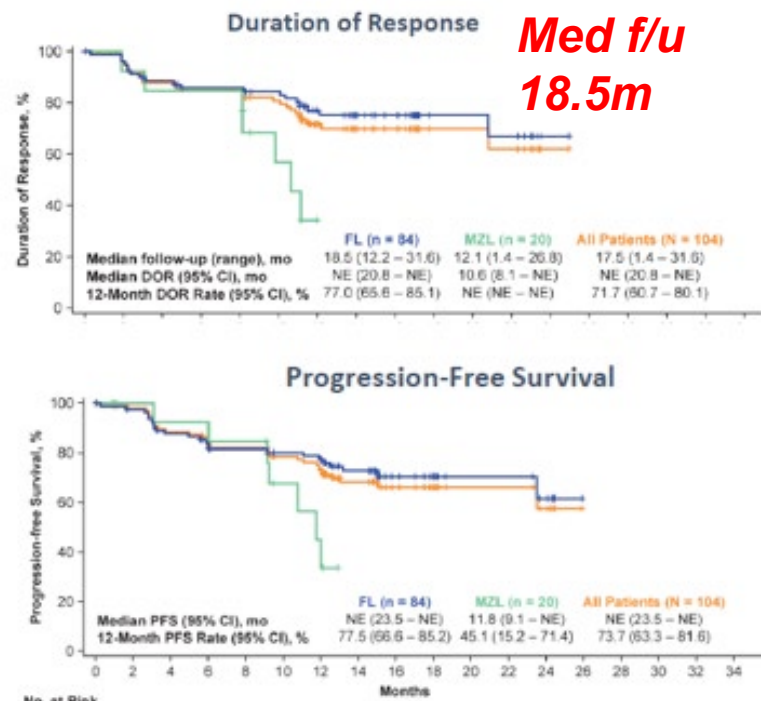
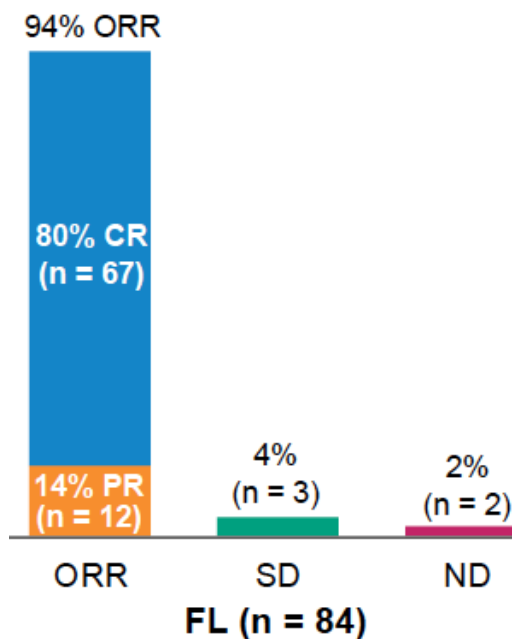
FL: Clinical categories and treatment options



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



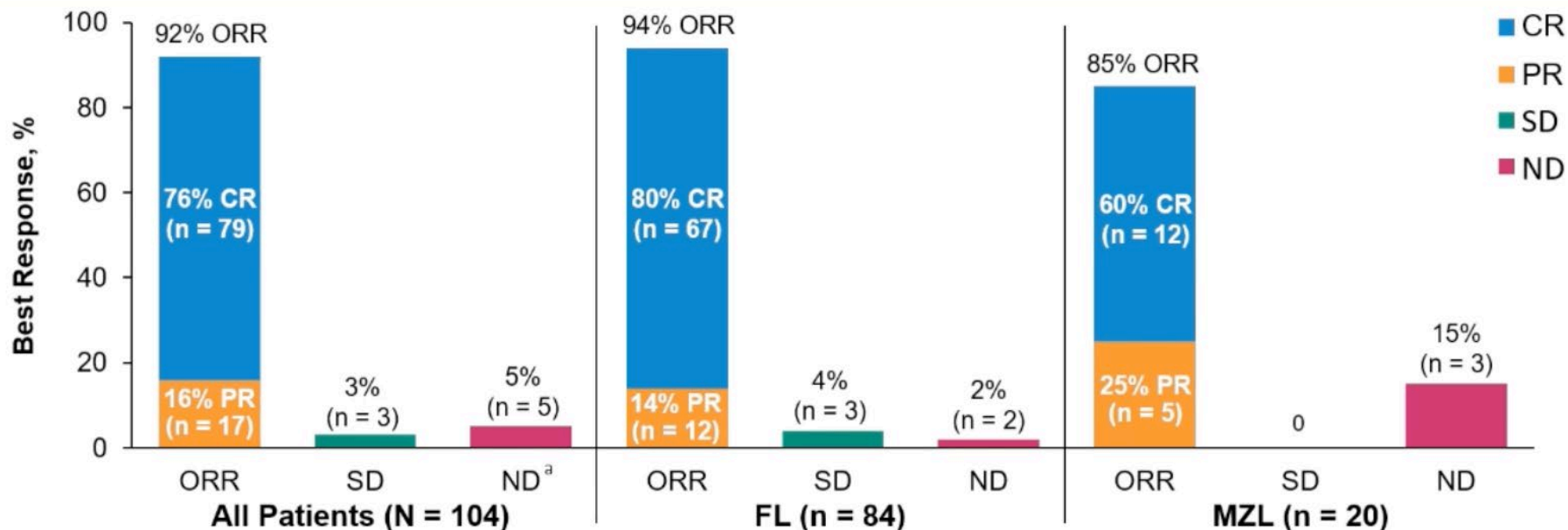
Primary endpoint: ORR



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)

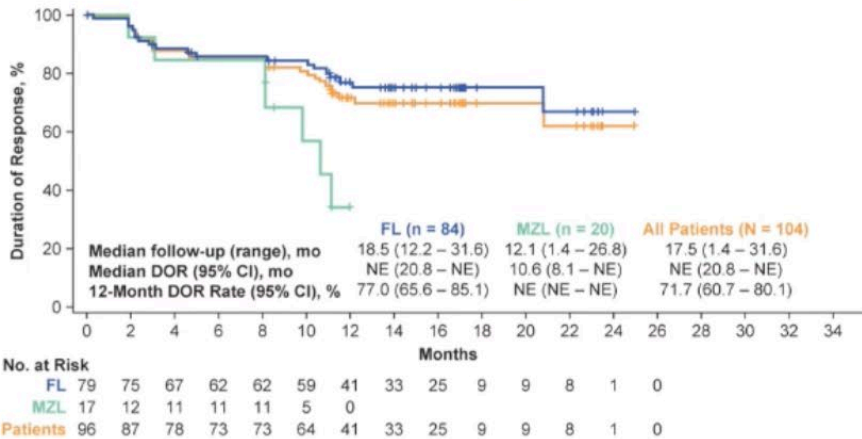


- The median time to first response was 1 month (range, 0.8 – 3.1)
- Among the 25 patients with FL who initially had a PR, 13 (52%) subsequently converted to a CR after a median of 2.2 months (range, 1.9 – 11.2)

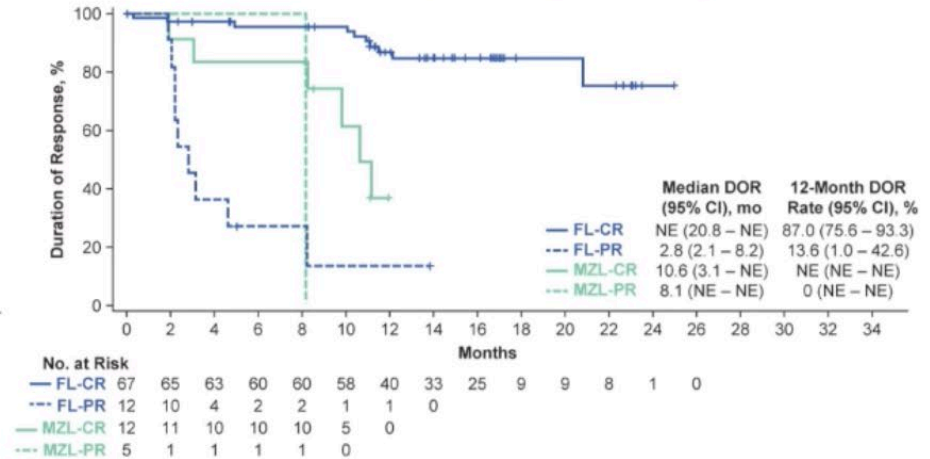
ZUMA-5: Duration of response for FL and MZL

Duration of Response

Duration of Response



Duration of Response by Best Response

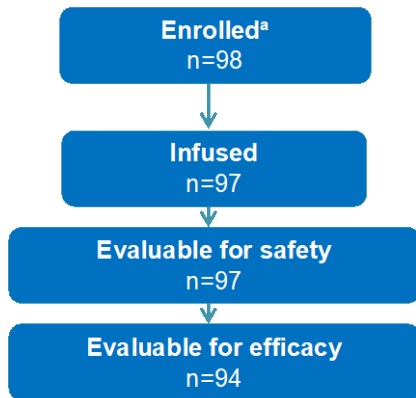


Median f/u 17.5m
All 151 pts received intended treatment

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



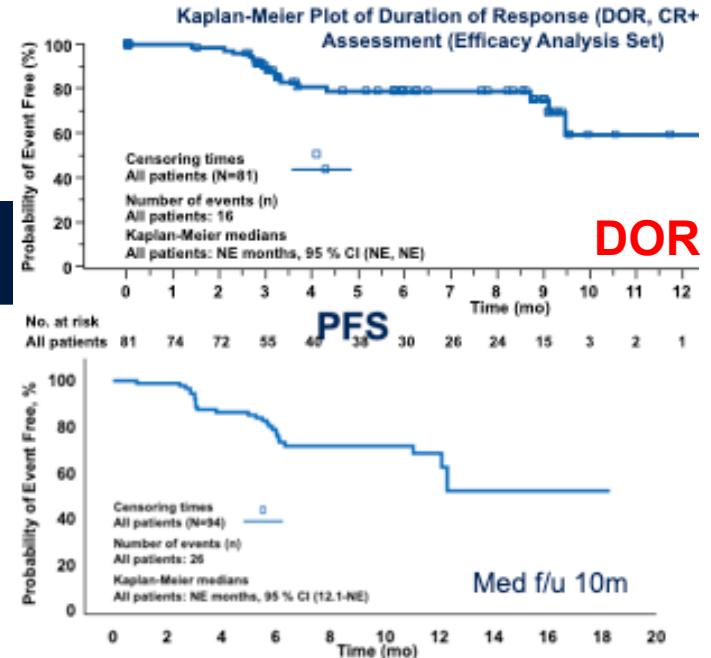
Vast majority of patients received cellular product

Lymphodepleting therapy could be Flu/Cy or bendamustine

18% of patients received tisagenlecleucel infusion in outpatient setting

Response Rate, %	Patients Evaluable 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

AESI (within 8 weeks of infusion)	Treated Patients N=97	
	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	≥ 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	57.0 (29-73)
≥65 y, n (%)	24 (24.7)
ECOG PS, n (%)	
0	56 (57.7)
1	37 (38.1)
2	4 (4.1)
Bulky disease at study entry, ^c n (%)	63 (64.9)
Stage at study entry III-IV, n (%)	82 (84.6)
FLIPI-2 at study entry, n (%)	59 (60.8)
Median no. of prior therapies (range)	4 (2-13)
≥5, n (%)	27 (27.8)
POD24 from first anti-CD20 mAb-containing therapy, ^d n (%)	58 (59.8)
Refractory to ≥2 regimens, ^f n (%)	74 (76.3)
Double refractory, ^f n (%)	67 (69.1)
Prior therapy	
Anti-CD20 mAb and alkylating agents, ^h n (%)	63 (64.9)
PI3K inhibitors, n (%)	20 (20.6)
Lenalidomide and rituximab, n (%)	16 (16.5)

ZUMA-5

Characteristic	FL (n = 124)
Median age (range), years	60 (34 – 79)
≥ 65 years, n (%)	38 (31)
Male, n (%)	73 (59)
ECOG 1, n (%)	46 (37)
Stage III-IV disease, n (%)	106 (85)
≥ 3 FLIPI, n (%)	54 (44)
High tumor bulk (GELF criteria), n (%) ^a	64 (52)
Median no. of prior therapies (range)	3 (1 – 10) ^b
≥ 3, n (%)	78 (63)
Prior PI3K therapy, n (%)	34 (27)
Refractory disease, n (%) ^c	84 (68)
POD24 from first anti-CD20 mAb-containing therapy, n (%) ^d	68 (55)
Prior autologous SCT, n (%)	30 (24)

Predictors of Response and Toxicity

PATIENT

T-CELLS

TUMOR

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

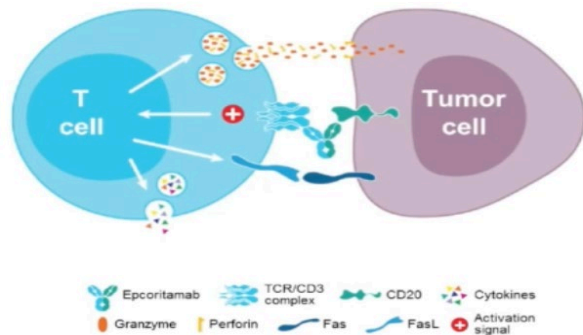
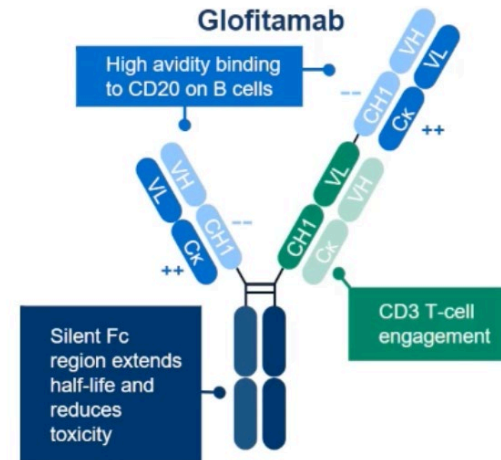
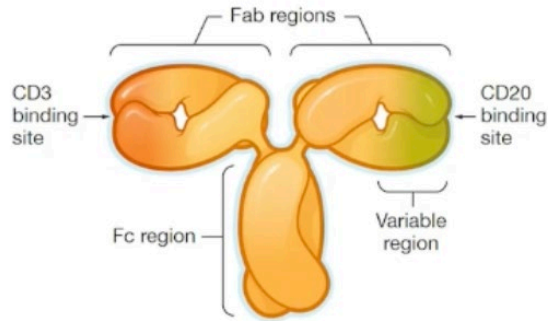
- High tumor burden, pretreatment LDH
- High pretreatment inflammatory markers
- ? High pretreatment monocyte levels
- High peak CAR T-cell levels
- High peak cytokine levels
- Markers of DIC (including fibrinogen levels)
- Early CRS

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

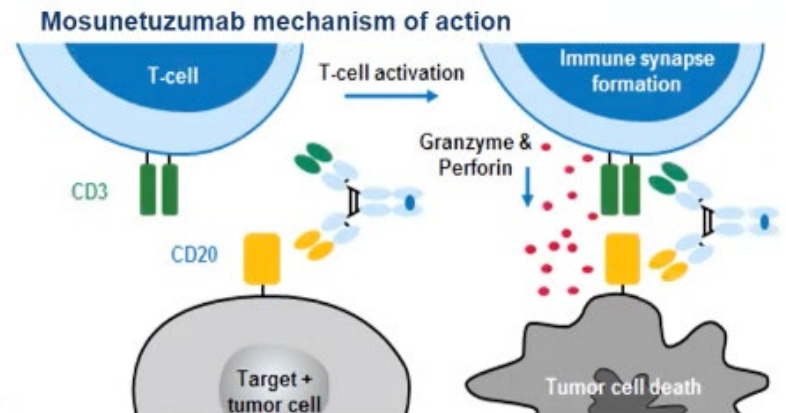
- Patient identification is key!
 - The vast majority of patients with FL do well without aggressive treatments
 - Disease characteristics:**
 - early POD, double refractory, multiple prior regimens with sequentially shorter PFS
 - Patient characteristics:**
 - No upper age limit, adequate cardiac/renal/pulmonary/neurologic 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

Emerging class of agents: bispecific antibodies

Odronextamab bispecific antibody structure



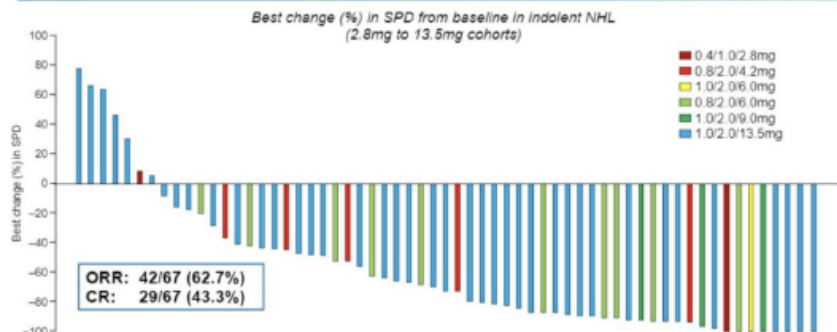
Distinct CD20 epitope



Bispecific antibodies in indolent lymphomas

Mosunetuzumab

Objective response rate in indolent NHL



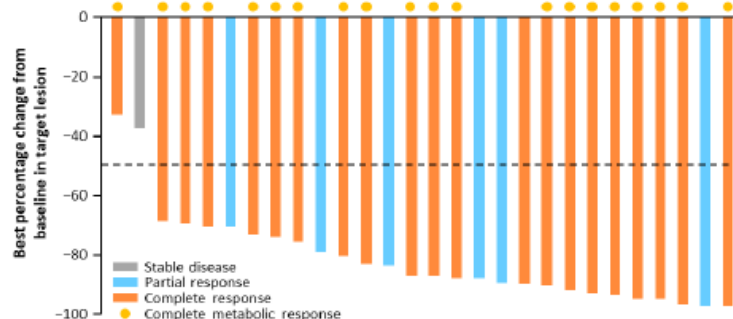
Indolent NHL, FL (Grade 1-3A), marginal zone lymphoma and small lymphocytic lymphoma
CCOD: Aug 9, 2019

Schuster et al ASH 2019



Odronextamab (REGN1979)

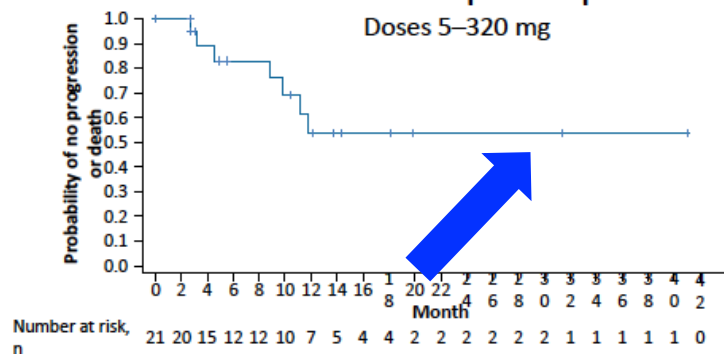
ORR



Doses 5-320 mg[†]

Duration of complete response

Doses 5-320 mg



Bannerji et al ASH 2020

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Mantle cell lymphoma

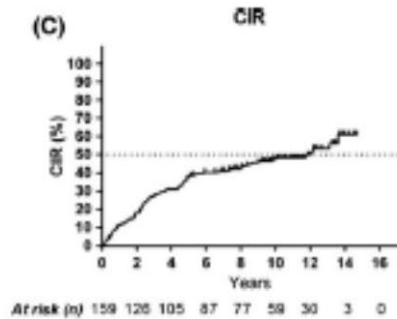
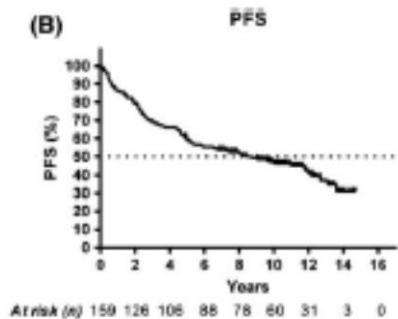
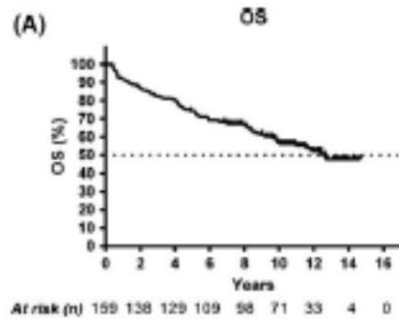
Mantle cell lymphoma: initial treatment approach

INDUCTION

CONSOLIDATION

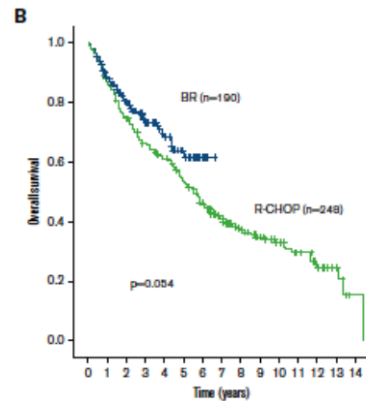
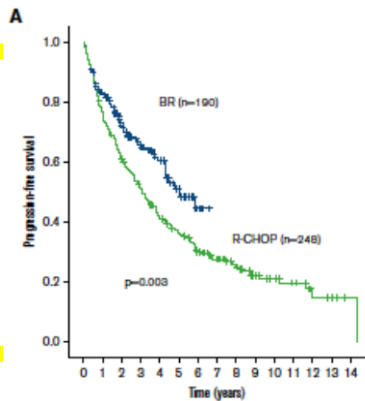
MAINTENANCE

YOUNG/FIT



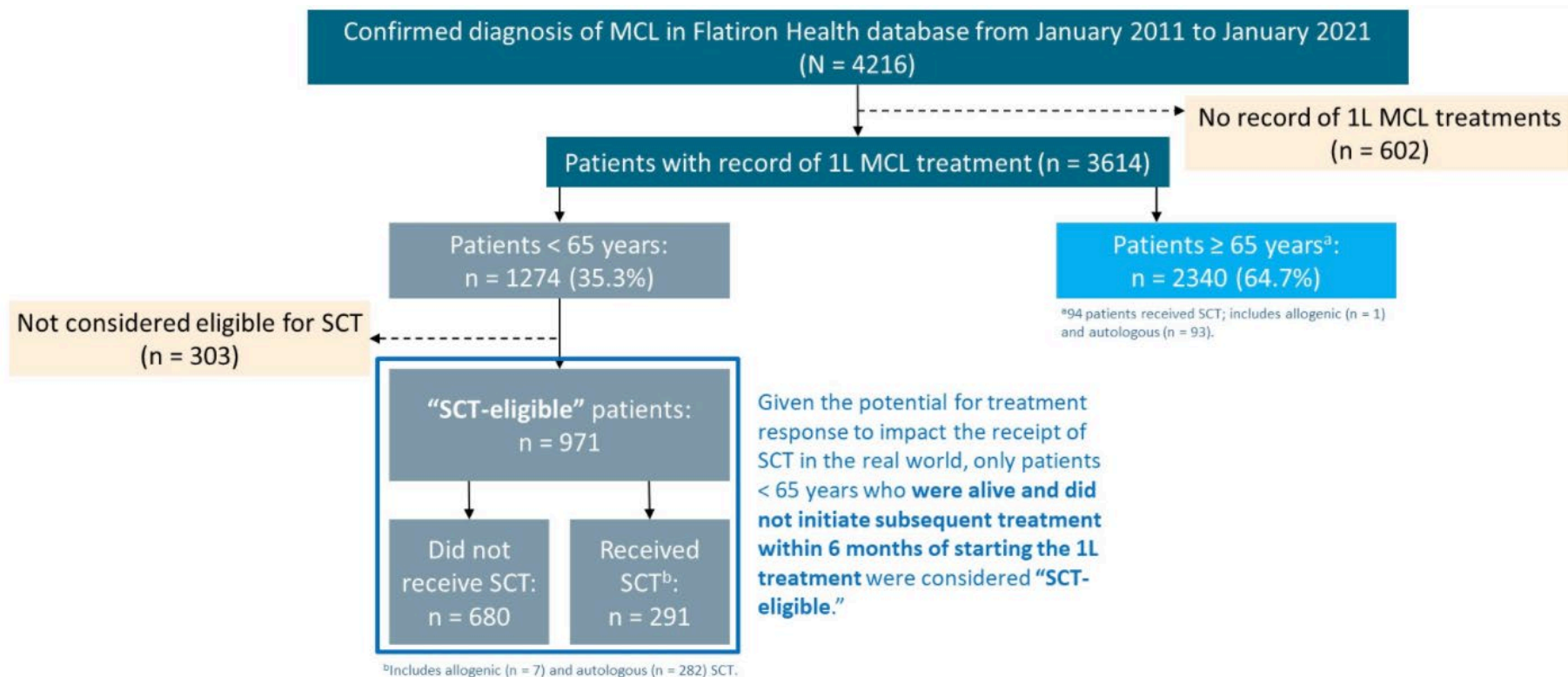
15-year f/u of Nordic MCL2: excellent outcomes but continuous relapse

OLDER/UNFIT



3-yr PFS 77% and OS 82% in younger patients with BR
R x 2-3y

Does the real-world experience match the data?



Given the potential for treatment response to impact the receipt of SCT in the real world, only patients < 65 years who **were alive and did not initiate subsequent treatment within 6 months of starting the 1L treatment** were considered “SCT-eligible.”

Additional information can be viewed by accessing this link: <https://www.oncologysciencehub.com/OncologyAM2021/ibrutinib/Martin/>
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ASCO 2021, Martin P, et al.

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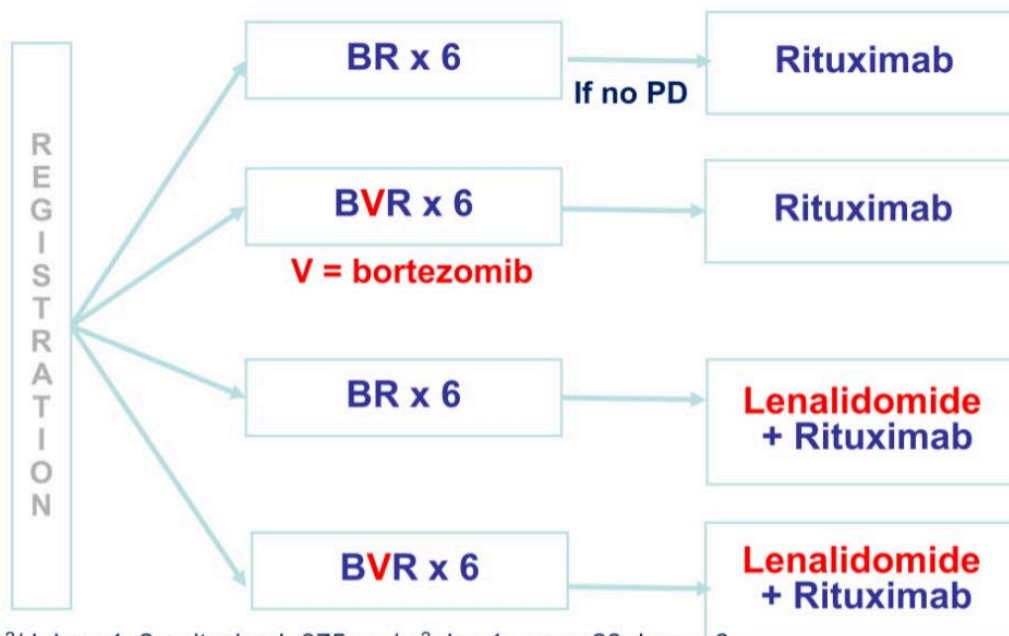
Martin ASCO 2020

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

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

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

E1411 SCHEMA



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

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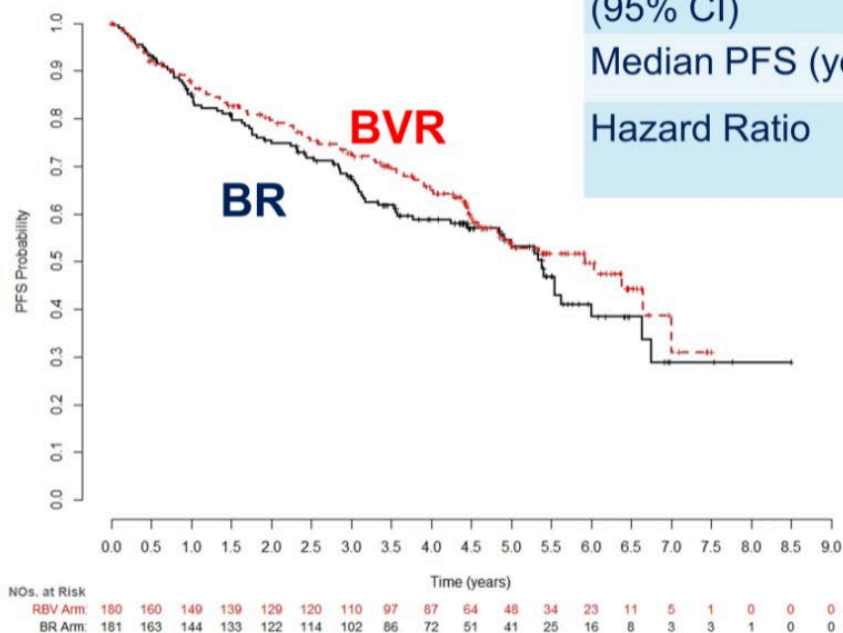
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Addition of bortezomib to BR does not improve outcomes

PFS: BR vs BVR

	BR	BVR
2 year PFS % (95% CI)	74.8% (68.6-81.6)	79.7% (73.9-85.9)
Median PFS (years)	5.4	5.9
Hazard Ratio	0.83 (0.60 -1.15)	



**No difference by
MIPI or age**

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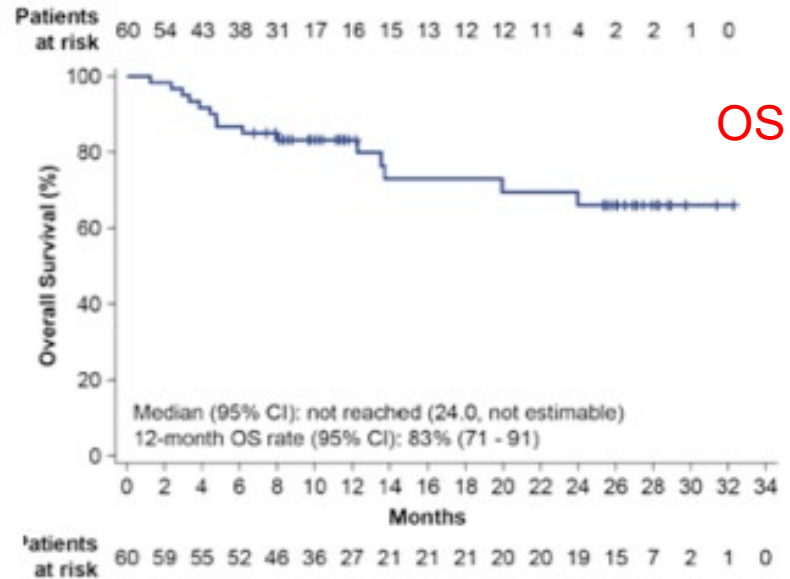
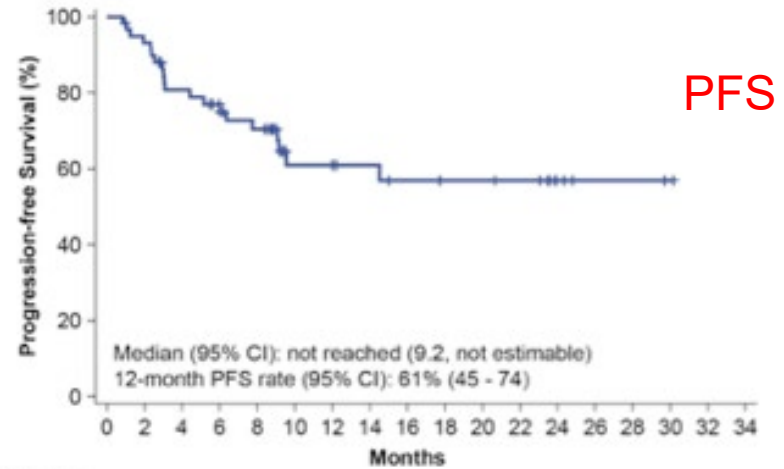
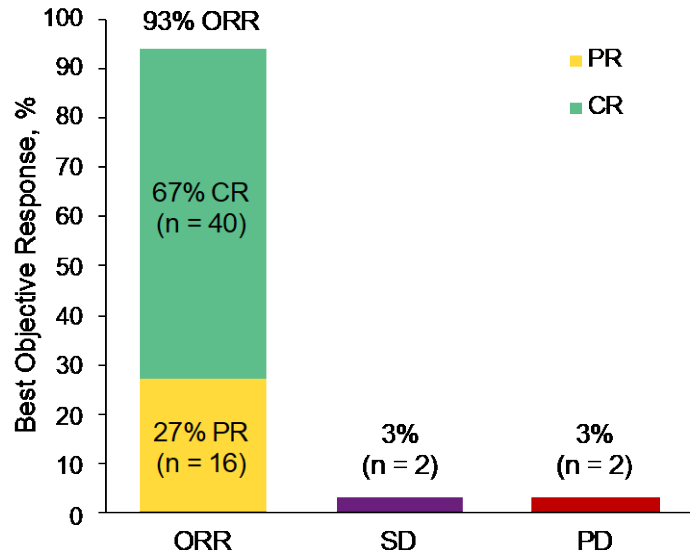
Treatment approach in rel/ref MCL

- 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!**
- } → Ven-len/ritux trial ongoing
(Philips ASCO 2021)

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

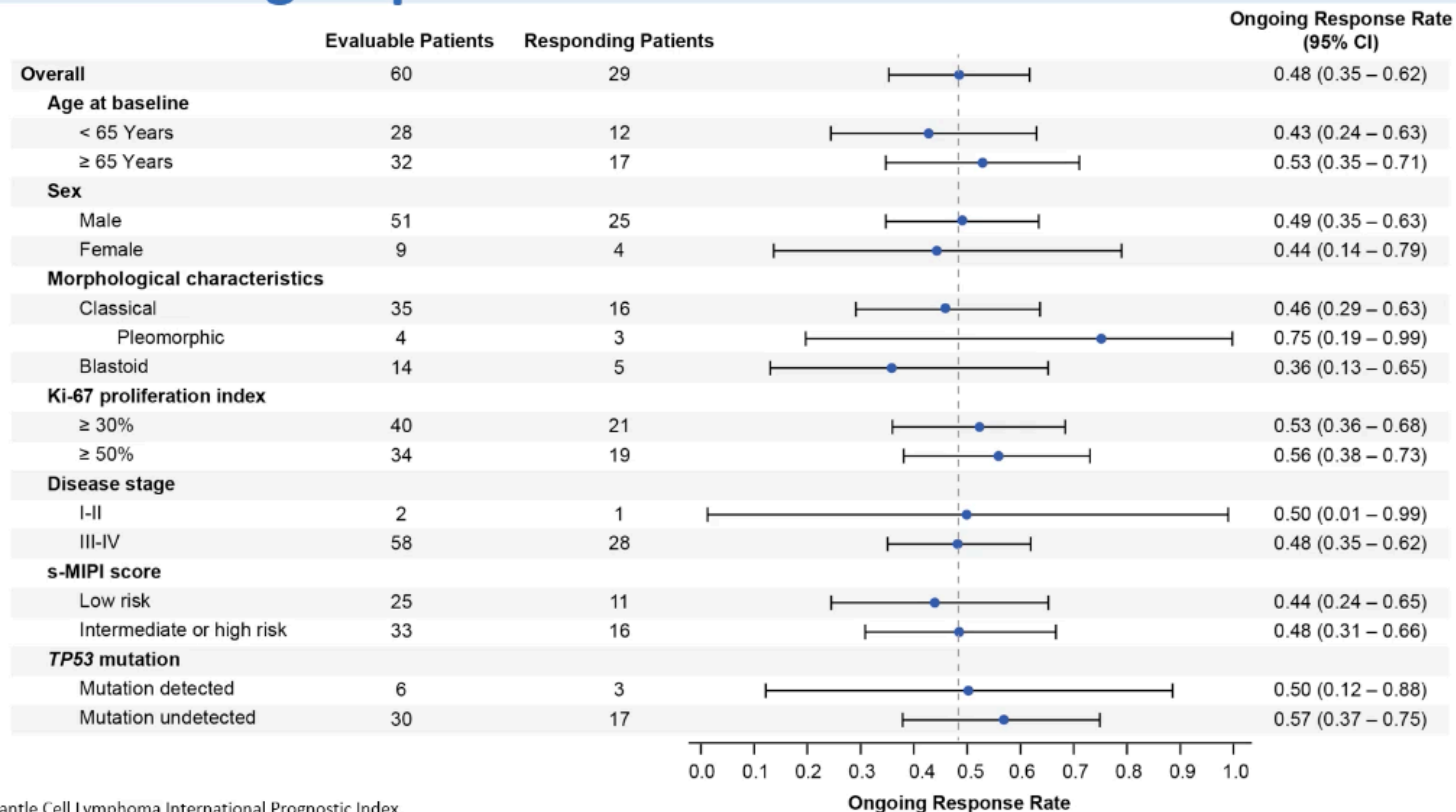
Key patient features:

- N=74 enrolled patients
- Med age 65 y
- Ki-67 \geq 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

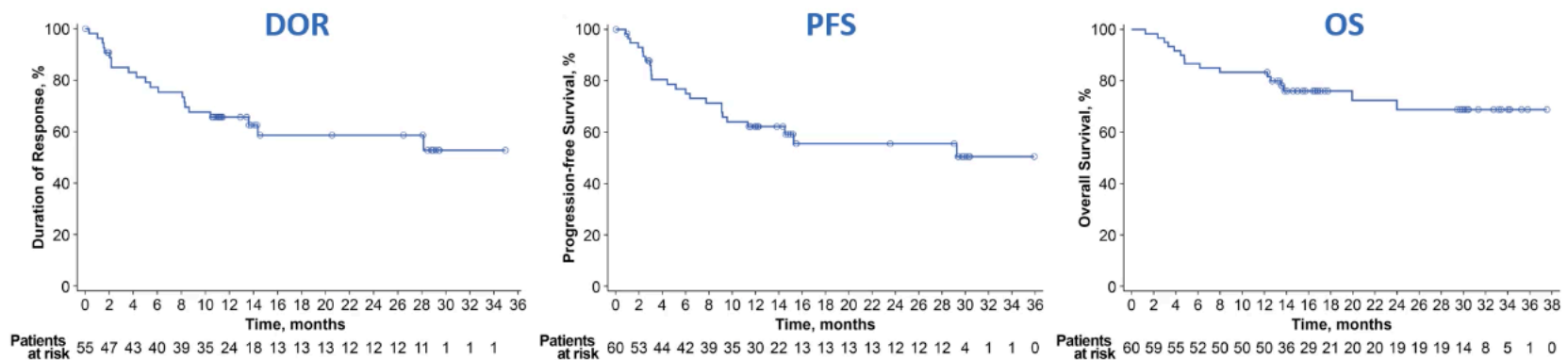


s-MIPI, simplified Mantle Cell Lymphoma International Prognostic Index.

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	
	Median (95% CI), mo	15-Mo Rate (95% CI), %	Median (95% CI), mo	15-Mo Rate (95% CI), %	Median (95% CI), mo	15-Mo Rate (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)

^aOf 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.

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



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