Chronic Lymphocytic Leukemia and Hairy Cell Leukemia

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Disclosures

Research Funding:

 Beigene, Mustang Biopharma, Celgene, BMS, Genentech, Pharmacyclics, Acerta Pharma, Astra Zeneca, AbbVie, Gilead Sciences, Sunesis, TG Therapeutics, Merck

Advisory Board/Consultancy:

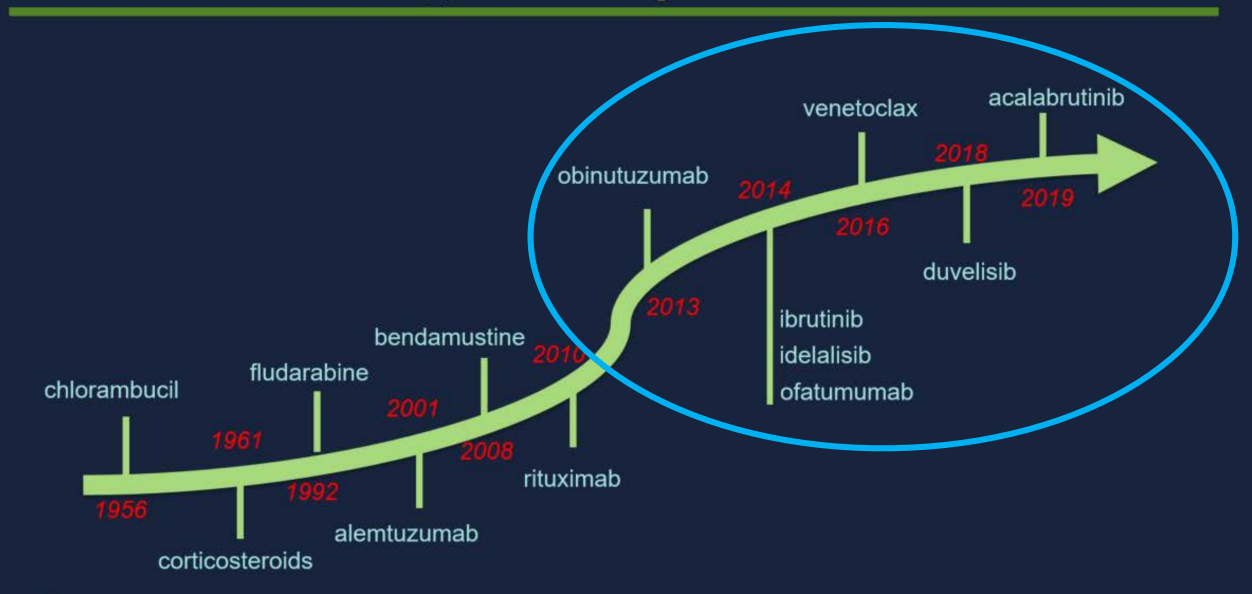
 AbbVie, AstraZeneca, Genentech, Beigene, Morphosys, Merck, Verastem, Pharmacyclics, ADC therapeutics, Atara Biotherapeutics, Sound Biologics, Gilead Sciences

Disclosures

Main purpose of this presentation is "Board Review"

- Will not discuss **experimental** treatments:
 - New biomarkers (prognostic and predictive)
 - New and unapproved BTKis or PI3Kis
 - Topic of MRD with venetoclax
 - Combination studies (ibrutinib + venetoclax, etc)
 - Details on CAR-T cell therapy (will have one slide)

Approved drugs for CLL





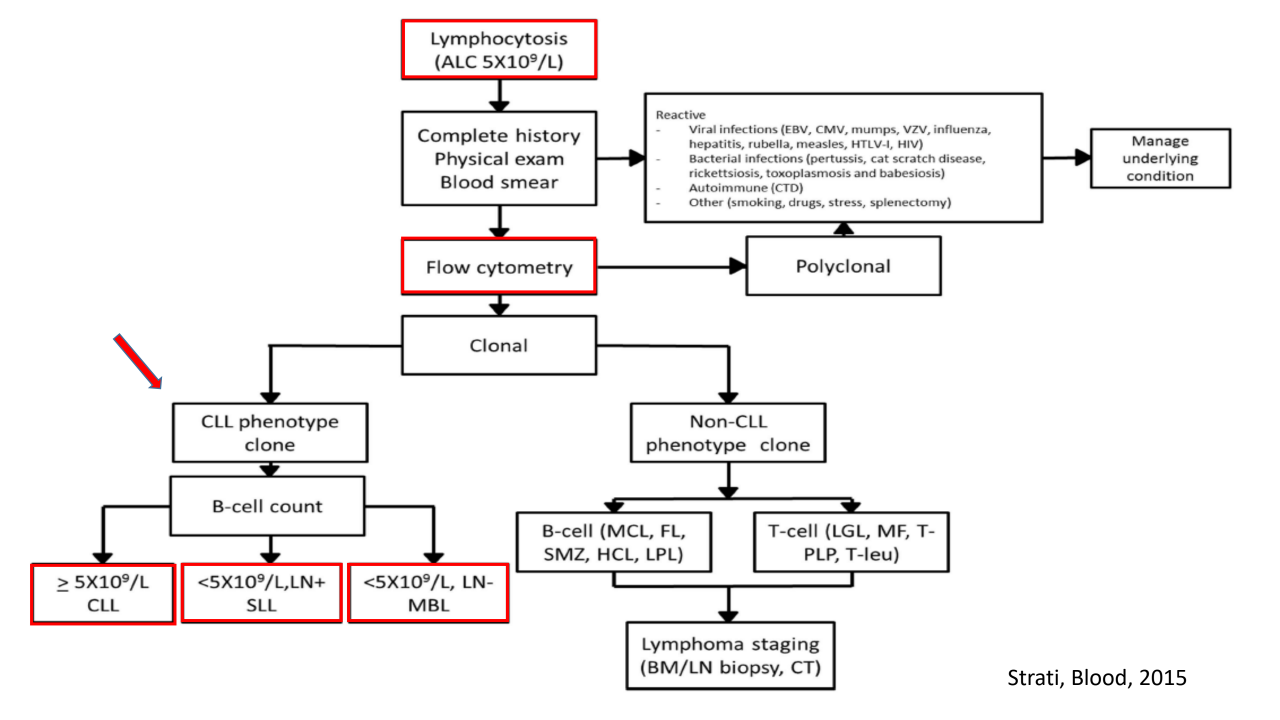
Epidemiology

- CLL/SLL is the most common leukemia in adults in western countries
 - 4.5 cases per 100,000
- Median age ~ 70 years
- Slight male predominance (1.7:1)
- Familial risk (7-8 fold)
- Caucasians > African Americans > Asian Pacific Islanders
- Genetic > Environmental

Top 10 topics in CLL

- 1. Initial diagnosis and appropriate work-up
- 2. Prognostic and predictive markers
- 3. Important therapeutic agents for CLL
- 4. Who should be treated?
- 5. Is there a role for early intervention in "high-risk" patients?
- 6. Treatment options for treatment-naïve patients (without del17p/P53 mutation)
- 7. Treatment options for previously treated patients (without del17p/P53 mutation)
- 8. Treatment options for patients with del17p/P53 mutation
- 9. Cellular therapies (CAR-T cell and Allogeneic Transplant)
- 10. Practical points about novel drugs

1. Initial diagnosis and appropriate work-up



Immunophenotypic Features

	CD5	CD10	CD23	CD103	BCL6	CD20	Cyclin D1
CLL/SLL	+	-	+	-	-	+	-
						(weak)	

Immunophenotypic Features

	CD5	CD10	CD23	CD103	BCL6	CD20	Cyclin D1
CLL/SLL	+	-	+	-	-	+ (weak)	-
MCL	+	-	-	-	_	+	+
LPL	-	-	-	-	_	+	-
sMZL	-	-	-	-	_	+	-
FL	-	+/-	-/+	-	+	+	-
HCL	-	-	-	+	-	+	+/-

	CD23	Cyclin D1	t(11,14)
CLL/SLL	+	-	-
MCL	-	+	+

MBL (monoclonal B cell lymphocytosis)

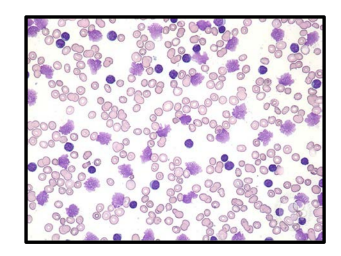
- < 5x 10⁹/L monoclonal B- cells in the PB <u>AND</u> no lymphadenopathy
- Almost all cases of CLL are preceded by MBL but only a small percentage of persons with MBL will ultimately develop CLL
- Low-count MBL (< 0.5x 10⁹/L) → rarely progresses to CLL
- High-count MBL (≥0.5x 10⁹/L) → progresses to CLL at a rate of 1-2% /year
- Up to 17 percent of first-degree family members of patients with CLL were found by flow cytometry to have MBL
- Screening of family members is NOT recommended

Diagnosis

- Flow cytometry of blood is essential and adequate to make the diagnosis
- ➤ Biopsy may be needed if PB flow cytometry is not conclusive
- > Cytogenetic and molecular studies are informative for prognostic and/or therapy determination .
- ➤ Baseline CT scan (or PET) is **NOT** required for asymptomatic patients (The ASH "Choosing Wisely" List)







2. Prognostic and predictive markers

Staging for CLL

	Rai Staging System for Chronic Lymphocytic Leukemia					
Stage	Risk	Clinical Features	Overall Survival, y			
Rai						
0	Low	Lymphocytosis in peripheral blood and bone marrow only	>10			
1/11	Intermediate	Lymphadenopathy ± hepatosplenomegaly	7			
III/IV	High	Anemia ± thrombocytopenia	<4			

Some use Ann Arbor staging for SLL

Molecular Biomarkers for CLL

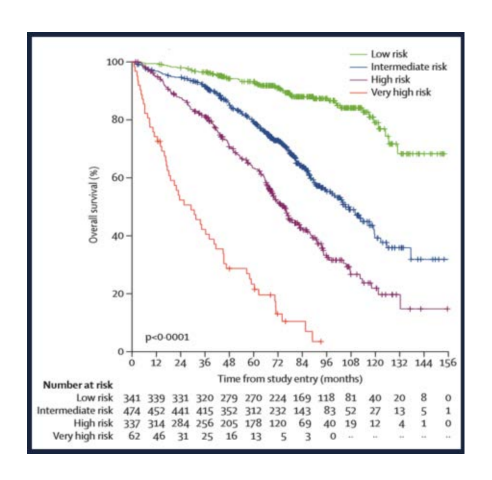
	FISH	Karyotype	Mutations
Unfavorable	del (17p) del (11q)	Complex (>3 abnormalities) (> 5?)	TP53 unmutated IGHV (≤ 2%) * NOTCH-1 SF3B1 BIRC3 ATM
Neutral	Normal +12		
Favorable	del (13q) (sole abnormality)		mutated IGVH (>2%)

Prognostic Models: CLL-IPI

Characteristic	Points
Del(17p) or TP53 mutation	4
Serum beta-2-macroglobulin ≥ 3.5mg/L	2
Un-mutated IgVH	2
Rai Stage I-IV	1
Age > 65 years	1

Points	Risk Group	5-y OS (%)	10-yr OS (%)
0-1	Low	93	79
2-3	Int	79	39
4-6	High	63	22
7-10	Very High	23	4

- ✓ Developed for chemoimmunotherapy
- ✓ Not validated for novel agents



3. Important therapeutic agents for CLL

Treatment options for CLL

Chemotherapy

anti-CD20 Abs

BCR inhibitors BCL-2 inhibitor

- fludarabine
- cyclophosphamide
- bendamustine
- chlorambucil

- rituximab
- ofatumumab
- obinutuzumab
- ublituximab *

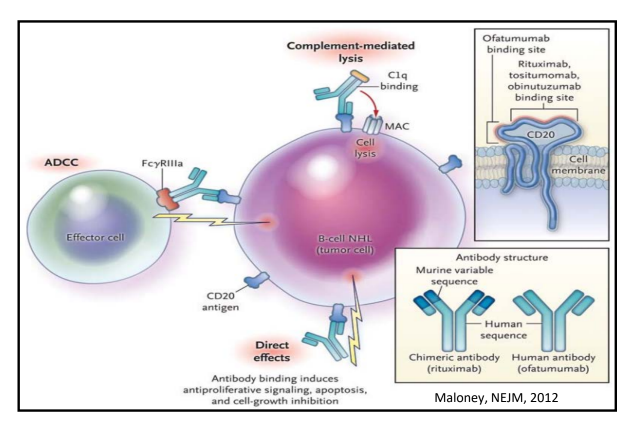
- BTK inhibitors
 - ibrutinib
 - acalabrutinib
 - zanubrutinib*†
- PI3K inhibitors
 - idelalisib
 - duvelisib
 - umbralisib *

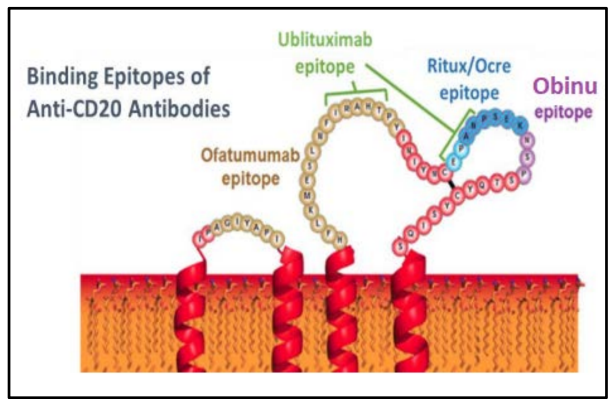
venetoclax

^{*} Not FDA approved for CLL as of August 2020

[†] Approved for MCL

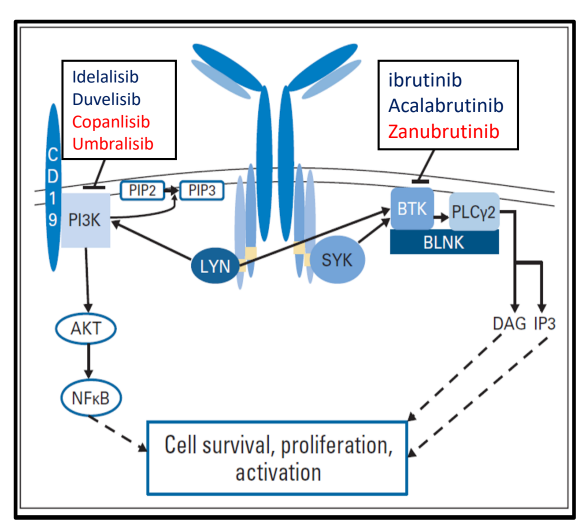
Anti-CD20 antibodies

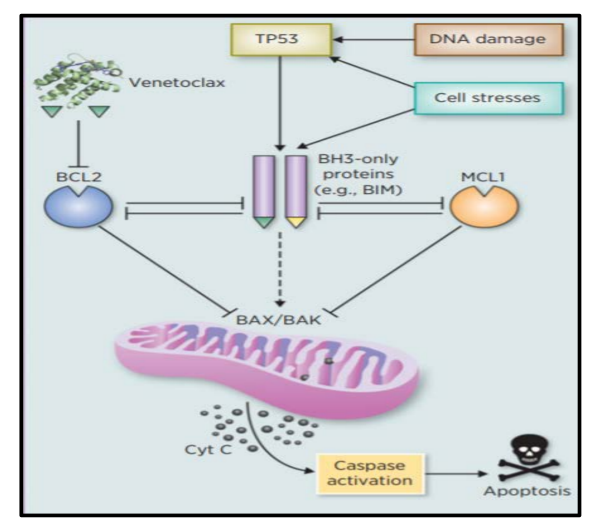




		Glycoengineered	Туре	Direct effect	CDCC	ADCC
Rituximab	chimeric	No	I	↑	$\uparrow\uparrow\uparrow\uparrow$	$\uparrow \uparrow$
ofatumumab	humanized	No	I	↑	$\uparrow\uparrow\uparrow\uparrow$	$\uparrow \uparrow$
obinutuzumab	humanized	Yes	II	$\uparrow\uparrow\uparrow\uparrow$	↑	个个个
ublituximab	chimeric	Yes	I	↑	<u> </u>	↑ ↑↑↑ 9

BCR Pathway inhibitors vs. BCL2 antagonist





Byrd, JCO, 2014

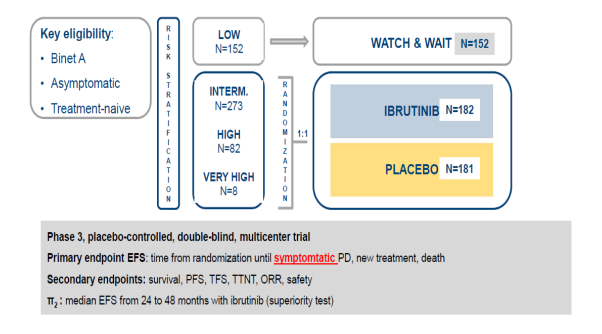
4. Who needs to be treated?

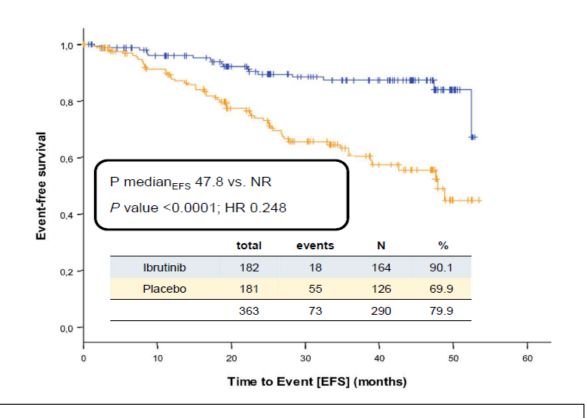
Indications for treatment

- Progressive marrow failure
- Massive , progressive or symptomatic lymphadenopathy or organomegaly
- Constitutional symptoms
- Autoimmune anemia and/or thrombocytopenia that is poorly responsive to corticosteroids or other standard therapy
- Lymphocyte doubling time

5. Is there a role for early intervention in "high-risk" patients?

CLL-12 Study — Early intervention with Ibrutinib





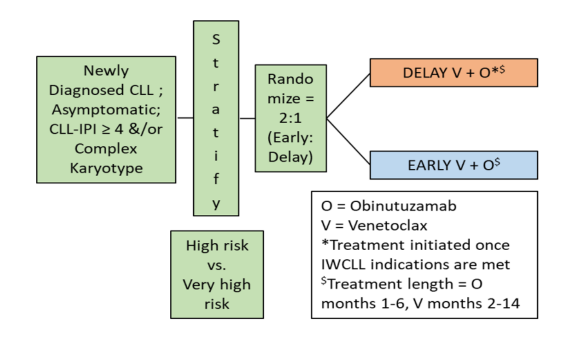
- No OS benefit
- Study is powered for OS so longer follow-up would be interesting
- Early intervention with ibrutinib is NOT recommended at this time

Upcoming US Intergroups Early Intervention Trial with Venetoclax

CLL-IPI

Characteristic	Points
Del(17p) or TP53 mutation	4
Serum beta-2-macroglobulin≥3.5mg/L	2
Un-mutated IgVH	2
Rai Stage I-IV	1
Age > 65 years	1

Points	Risk Group
0-1	Low
2-3	Int
4-6	High
7-10	Very High



Primary Endpoint: Overall Survival

Open 2020



6. Treatment options for treatment-naïve patients (without del17p/P53 mutation)

First line treatment for patients with normal TP53

For all pts:

Acalabrutinib ± G

OR

Ibrutinib

OR

Venetoclax + G

FCR is <u>not preferred</u> but can be a <u>reasonable option</u> for selected patients if:

- younger than 65 and fit
- mutated IGHV
- no evidence of del17p or TP53 mutation
- (no evidence of del 11q)

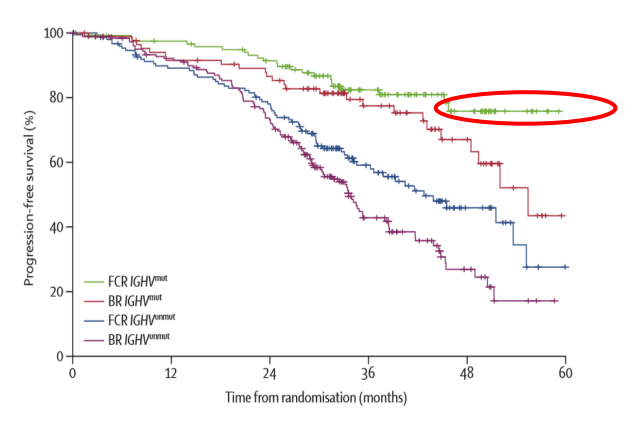
Frontline (normalTP53)

Historical studies from the "chemo era"

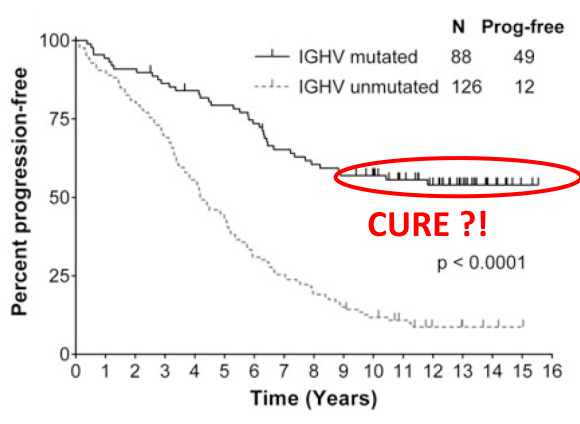
Study	Treatments	N	Result	Outcome	Notes
German CLL10	FCR vs. BR	564	FCR > BR	PFS but not OS	No benefit if > 65 AML/MDS: 5% with FCR
German CLL11	CHL-obino vs. CHL-ritux vs. CHL	780	CHL-obino > CHL-ritux > CHL	PFS and OS	
RESONATE-2	Ibrutinib vs. CHL	269	Ibrutinib > CHL	PFS and OS	

IGHV mutation as a predictive marker for FCR





MD Anderson



Eichhorst, Lancet Oncology, 2016

Thompson, Blood, 2106

First line – Summary of novel vs. chemo studies





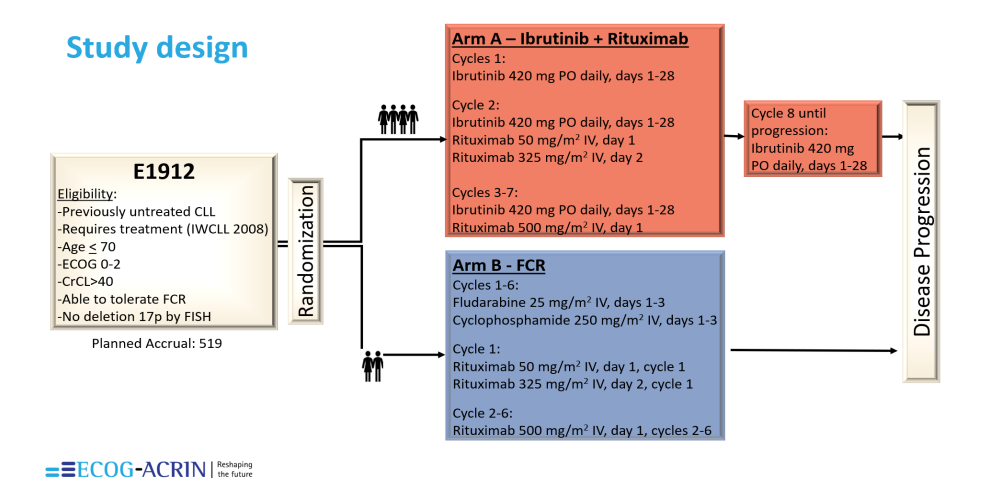






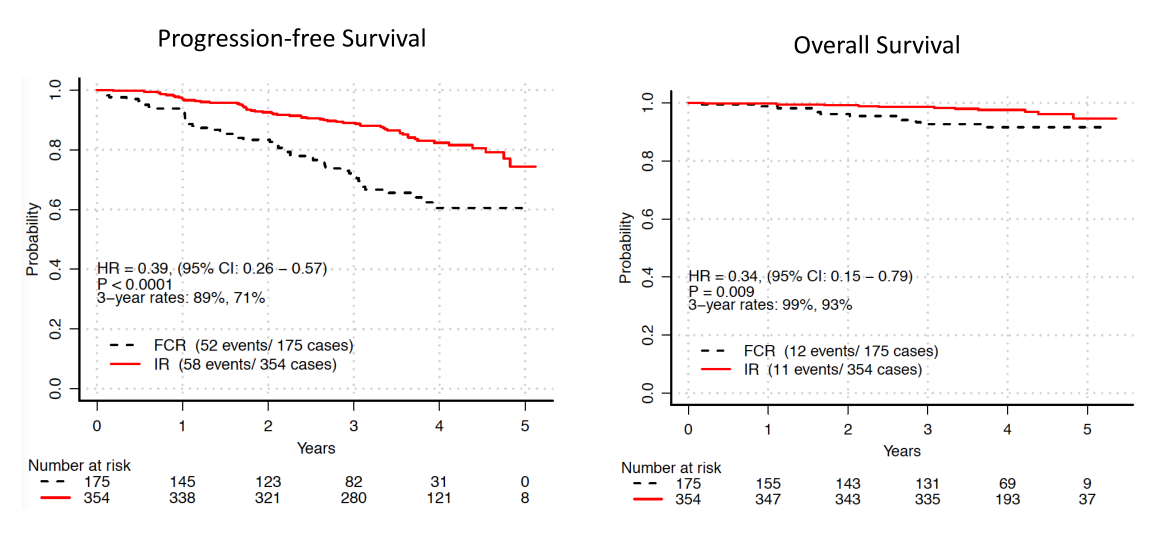
Fit and young	FCR	? (E1912)	Ibrutinib +R
Older	BR	(A041202)	Ibrutinib ± R
Older or with comorbid conditions	CHL+G	? (illuminate)	Ibrutinib +G
Older or with comorbid conditions	CHL+G	(ELEVATE)	acalabrutinib ± G
with comorbid conditions	CHL+G	? (CLL14)	Venetoclax+ G

FCR vs. IB+R (E1912 Study)



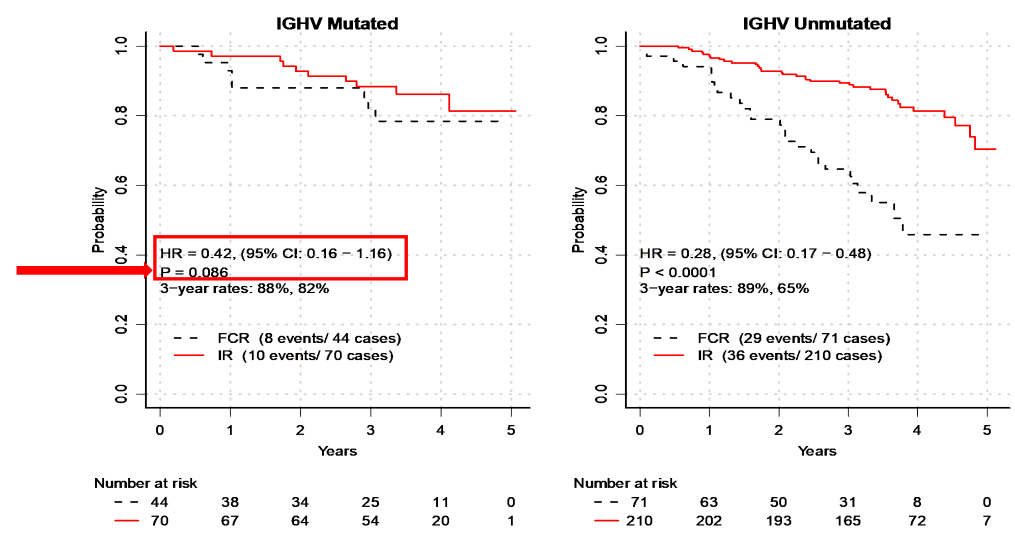
FCR vs. IB+R (E1912 Study)

(48 months follow-up)



FCR vs. IB+R (E1912 Study)

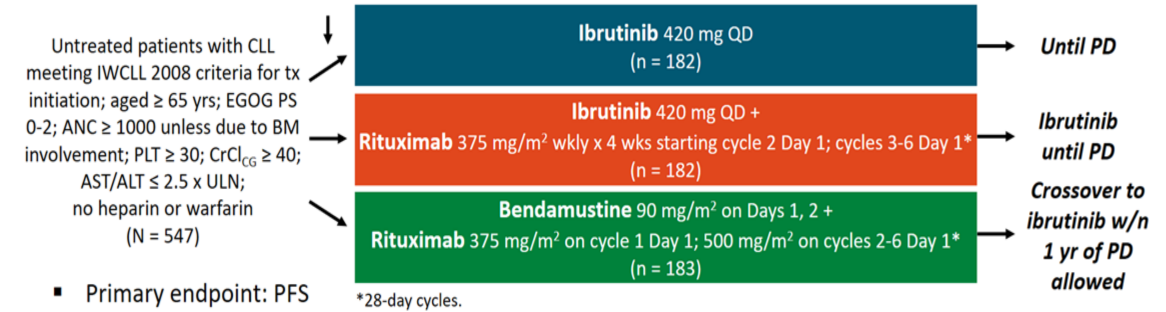
(48 months follow-up)



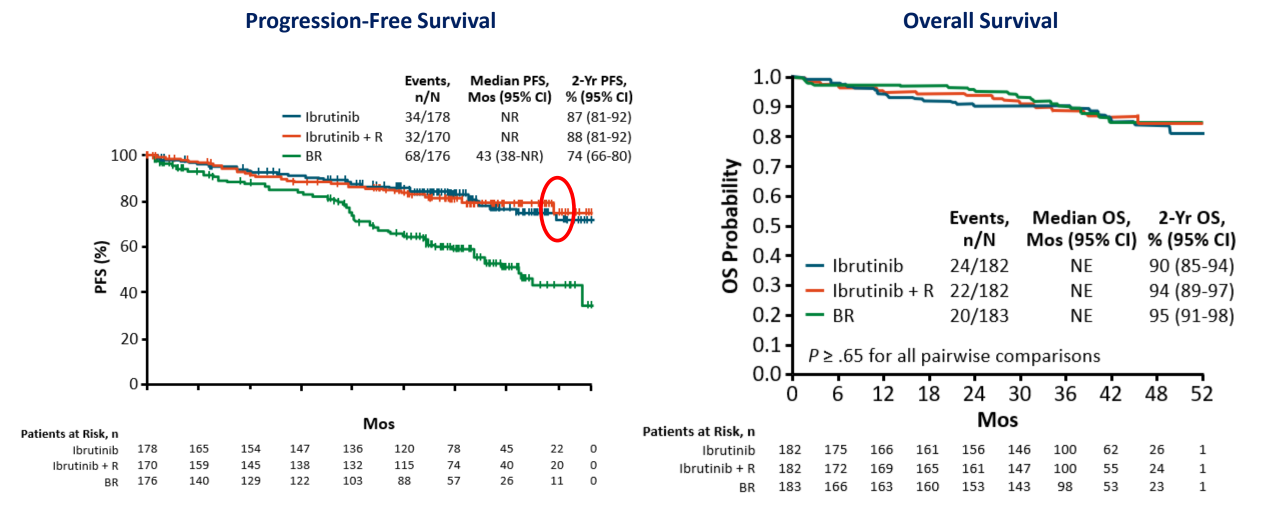
BR vs. IB+R vs. IB (A041202 Study)

Multicenter, randomized, double-blind phase III study (data cutoff: October 4, 2018)

Stratified by Rai stage (high vs intermediate risk), del(11q22.3) or del(17p13.1) (presence vs absence), ZAP-70 methylation (< vs ≥ 20%)



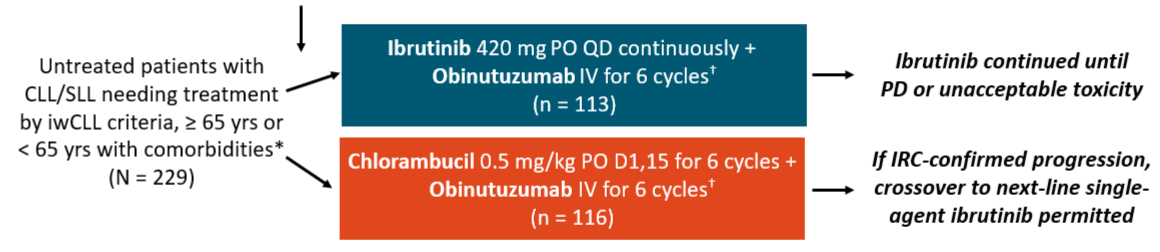
A041202: Results



IB+G vs. CHL+G (iLLUMINATE)

Randomized, open-label, multicenter phase III trial

Stratified by ECOG PS (0-1 vs 2), del(17p)/del(11q) (+/+ vs +/- vs -/+ vs -/-)



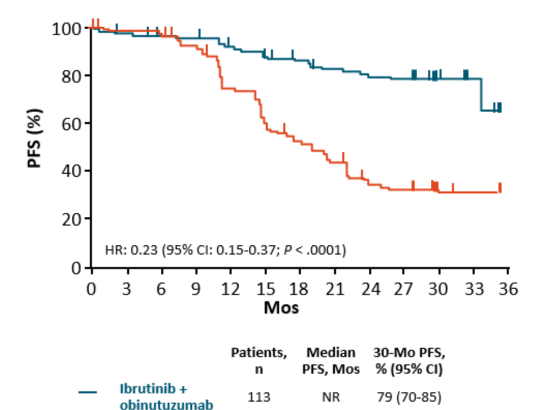
^{*}Cumulative Illness Rating Score > 6, creatinine clearance < 70 mL/min, and/or del(17p)/TP53 mutation.

Primary endpoint: PFS

[†]Cycle 1: 100 mg, Days 1; 900 mg, Day 2; 1000 mg, Days 8, 15. Cycle 2-6: 1000 mg, Day 1.

IB+G vs. CHL+G (iLLUMINATE) Results

Progression-free Survival



116

19.0

31 (23-40)

Chlorambucil +

obinutuzumab

No Overall Survival Benefit

Acalabrutinib ± G vs. CHL+G (ELEVATE)

Treatment-naive CLL (N=535)

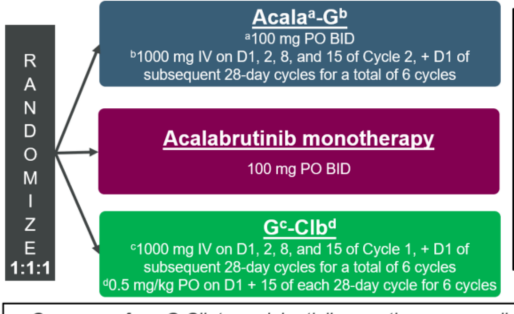
Age ≥65 or <65 years with

coexisting conditions:

- CIRS score >6, or
- creatinine clearance
 <70 mL/min

Stratification

- del(17p), y vs n
- ECOG PS 0-1 vs 2
- Geographic region (N America, W Europe, or other)



Primary endpoint

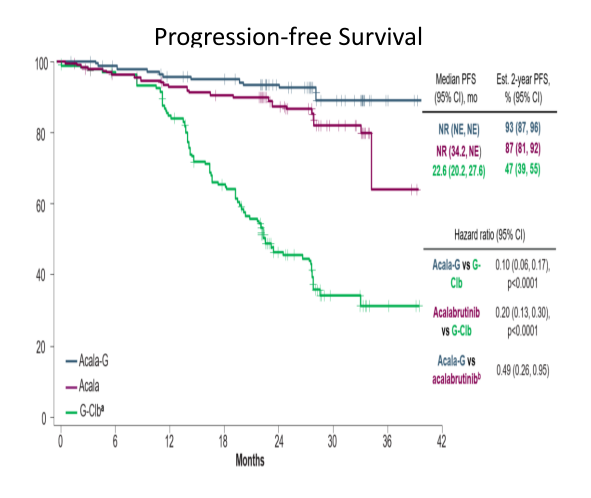
 PFS (assessed by IRC) Acala-G vs G-Clb

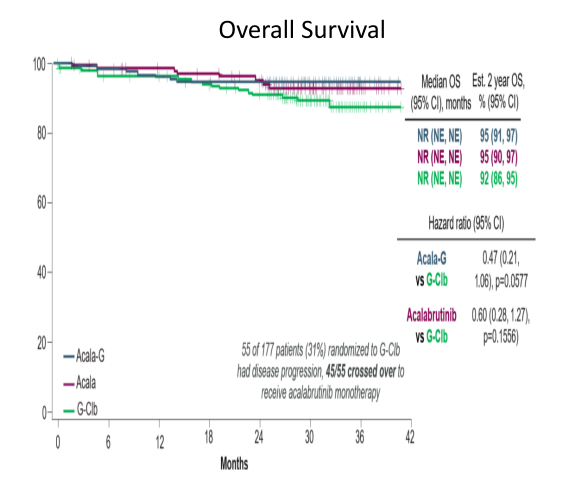
Key secondary endpoints

- PFS acalabrutinib vs G-Clb
- ORR (assessed by IRC and investigator)
- Time to next treatment
- OS
- Safety

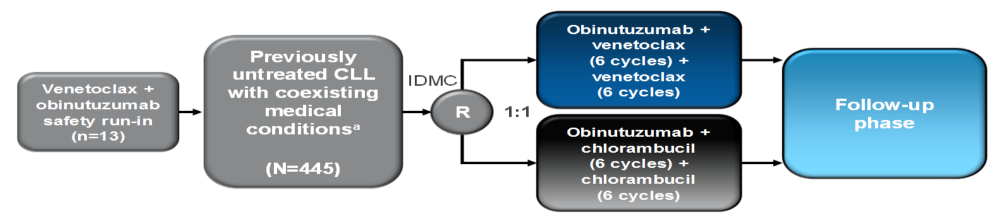
Crossover from G-Clb to acalabrutinib monotherapy was allowed after IRC-confirmed progression

Acalabrutinib ± G vs. CHL+G (ELEVATE)





Venetoclax + G vs CHL + G (CLL-14)



Primary endpoint:

 PFS as assessed by investigator³

Secondary endpoints³:

- PFS as assessed by IRC
- MRD
- ORR
- CR rate
- DOR

- EFS
- OS
- TTNT
- Safety

aCIRS >6 and/or CrCl <70 mL/min

Venetoclax and TLS

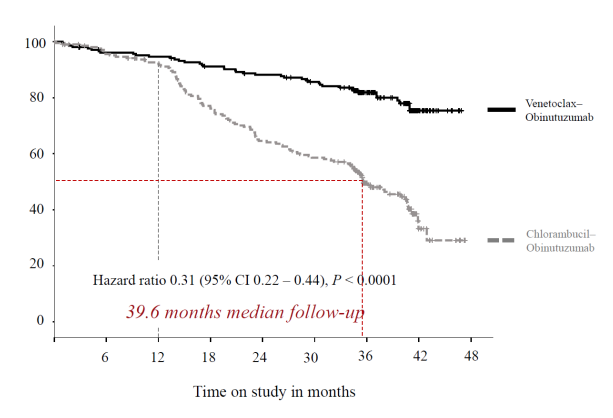


Tumor Burden		Prophylaxis		Blood Chemistry Monitoring ^c	
		Hydration ^a	Anti- hyperuricemics	Setting and Frequency of Assessments	
Low	All LN <5 cm AND ALC <25 × 10 ⁹ /L	Oral (1.5-2 L)	Allopurinol ^b	Outpatient Pre-dose, 6-8 hours, 24 hours at first dose of 20 mg and 50 mg Pre-dose at subsequent ramp up doses	
Medium	Any LN 5 cm to <10 cm OR ALC ≥25 × 10 ⁹ /L	Oral (1.5-2 L) and consider additional intravenous	Allopurinol	Outpatient Pre-dose, 6-8 hours, 24 hours at first dose of 20 mg and 50 mg Pre-dose at subsequent ramp up doses Consider hospitalization for patients with CrCl <80 mL/min at first dose of 20 mg and 50 mg; see below for monitoring in hospital	
High	Any LN ≥10 cm OR ALC ≥25 × 10 ⁹ /L AND any LN ≥5 cm	Oral (1.5-2 L) and intravenous (150-200 mL/hr as tolerated)	Allopurinol;consider rasburicase ifbaseline uric acid is elevated	In hospital at first dose of 20 mg and 50 mg • Pre-dose, 4, 8,12 and 24 hours Outpatient at subsequent ramp-up doses • Pre-dose, 6-8 hours, 24 hours	

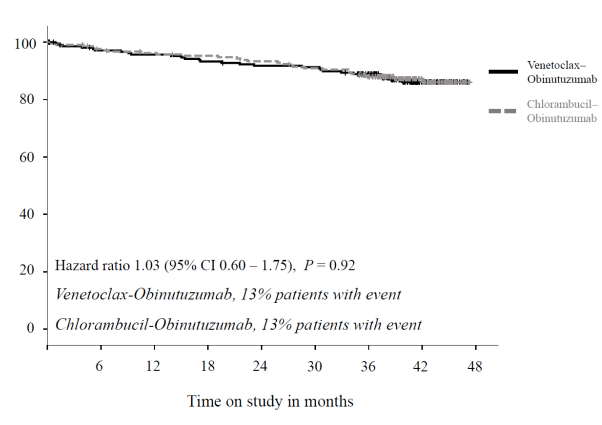
For patients at risk of TLS, monitor blood chemistries at 6-8 hours and at 24 hours after each subsequent ramp-up dose

Venetoclax + G vs CHL + G (CLL-14)

Progression-free Survival

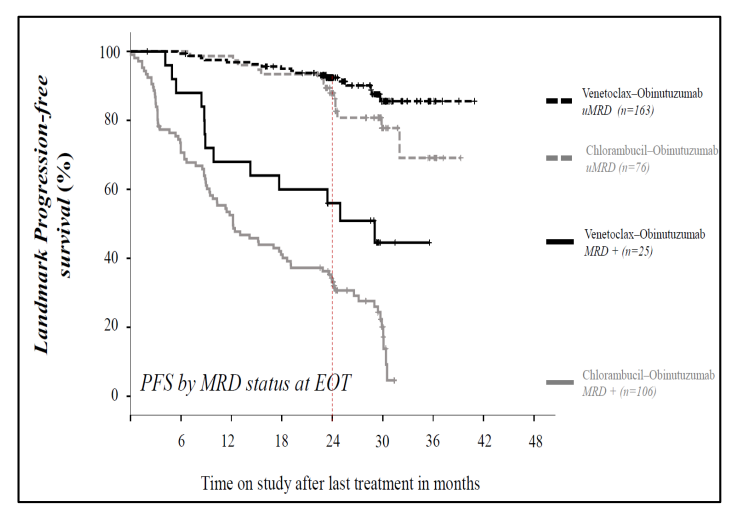


Overall Survival



uMRD4 at the end of treatment (12 months) and PFS

	Venetoclax- Obinutuzumab	Chlorambucil- Obinutuzumab	P value
Number of patients, N	216	216	
Peripheral blood			
Negative (<10 ⁻⁴)	76 %	35 %	< 0.001
Negative (<10-4) in complete response	42 %	14 %	< 0.001
Bone marrow			
Negative (<10 ⁻⁴)	57 %	17 %	< 0.001
Negative (<10 ⁻⁴) in complete response	34 %	11 %	< 0.001
By ASO-PCR 3 months after completion of treatm	nent		



First line – Summary of novel vs. chemo studies











Fit and young	FCR	(E1912)	Ibrutinib +R
Older	BR	(A041202)	Ibrutinib ± R
Older or with comorbid conditions	CHL+G	(iLLUMINATE)	Ibrutinib +G
Older or with comorbid conditions	CHL+G	(ELEVATE)	acalabrutinib ± G
with comorbid conditions	CHL+G	(CLL14)	Venetoclax+ G

First line treatment for patients normal TP53

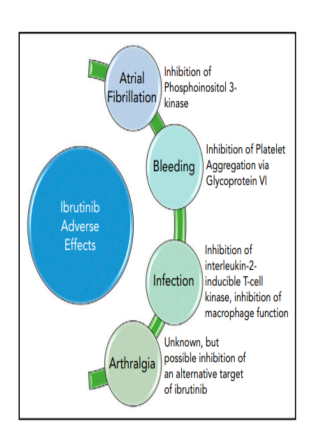
For all pts: Acalabrutinib ± G OR Ibrutinib OR Venetoclax + G

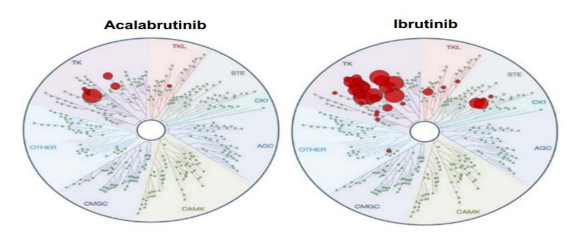
FCR is not preferred but can be a reasonable option for selected patients if:

- younger than 65 and fit
- mutated IGHV
- no evidence of del17p or TP53 mutation
- (no evidence of del 11q)

Acalabrutinib or Ibrutinib

Head-to-head trial is done in the relapsed setting and will be reported "soon"

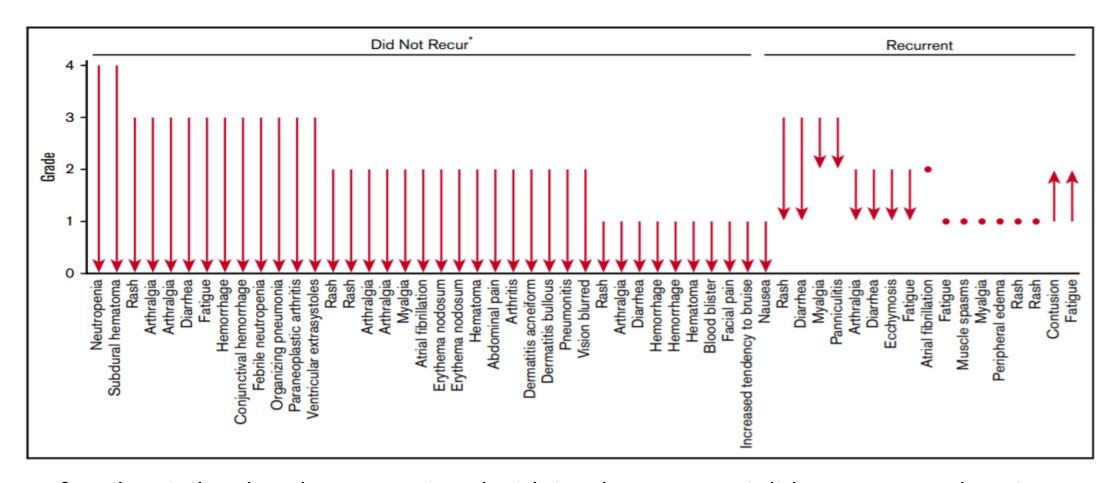




Treatment discontinuation rates due to toxicity

Ibrutinib	Frontline: 15% Relapsed: 22%
Acalabrutinib	Frontline: no data Relapsed: 12%

Acalabrutinib in Ibrutinib intolerant patients



of 61 ibrutinib-related AEs associated with intolerance, 72% did not recur and 13% recurred at a lower grade with acalabrutinib

BTKis vs. Ven-G

BTKi (Acalabrutinib/Ibrutinib)	Ven-G		
Indefinite treatment (responses mostly PR)	Fixed-duration; High CR and uMRD rate		
Long-term efficacy data available	<u>Time-limited treatment</u>		
Easier to start	Better tolerated and easier to continue		
 Preferred in patients who: Can't follow the ramp-up schedule for venetoclax Significant/unstable renal issues 	Preferred in patients with:Cardiac (arrythmia, HTN)Bleeding issues		
IB is studied against stronger regimens: (FCR and BR)	Deep remissions (@ MRD level) – would expect the same in younger pts		
Can use after Ven and is effective	Can use after BTKi and is effective		

- No head-to-head comparison
- Both are reasonable options
- Consider patient and disease factors
- Look at pros and cons for each

7. Treatment options for previously treated patients (without del17p/P53 mutation)

Previously Treated CLL Summary

Acalabrutinib Ibrutinib For all pts: Venetoclax + R OR OR **Duvelisib Idelalisib** OR R = rituximab

Previously Treated CLL Summary

1. First

- Venetoclax + Rituximab or
- BTKi: Ibrutinib or acalabrutinib

Second

- Ibrutinib/acalabrutinib if previously treatment with Ven
- Ven-R if previously treated with BTKi (ibrutinib or acalabrutinib)

3. Third

Idelalisib+ rituximab OR duvelisib

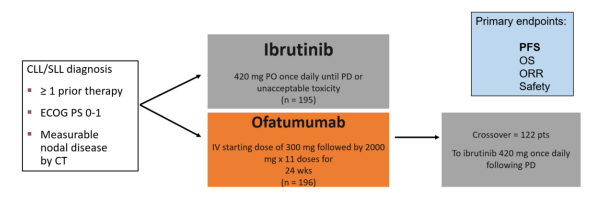
Previously treated CLL: Principles

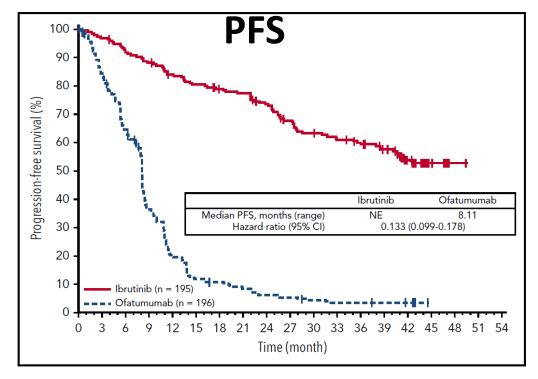
1. Repeat FISH panel - look for del (17p) or TP53 mutation

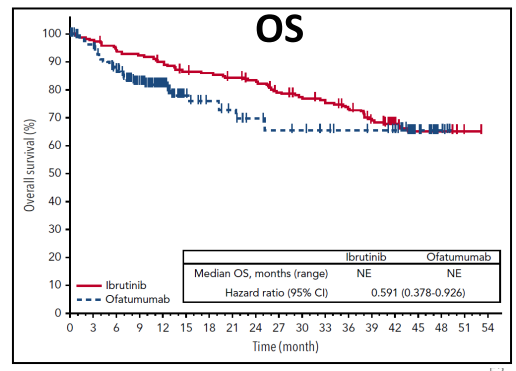
2. Bone marrow needs to be repeated to assess for MDS if prior FCR

3. Very limited role for chemoimmuntherapy (almost never)

Ibrutinib vs Ofatumumab in R/R CLL (RESONATE: Phase III)







Byrd, NEJM, 2014; Byrd, Blood, 2019

Acalabrutinib vs. Investigator choice for relapsed CLL (ASCEND Study)

Relapsed/Refractory CLL (N= 310)

Stratification:

del(17p), y vs n

ECOG PS 0-1 vs 2

1-3 vs ≥4 prior therapies

Acalabrutinib

100 mg PO BID

Idelalisib plus Rituximab (IdR)

Idelalisib 150 mg PO BID + rituximaba

- or
Bendamustine plus Rituximab (BR)

Bendamustine 70 mg/m² IVb + rituximabc

Primary endpoint:

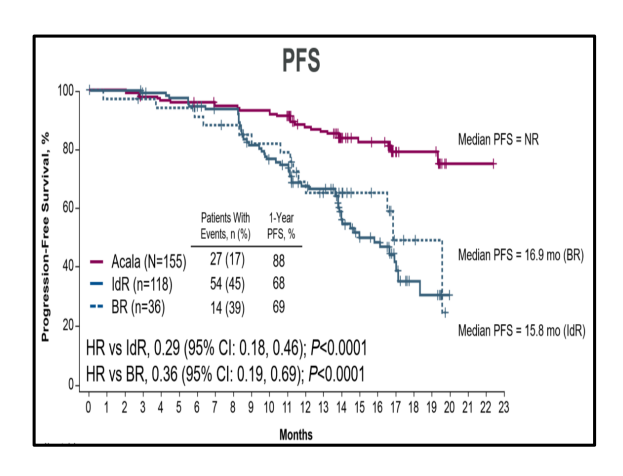
PFS (assessed by IRC)

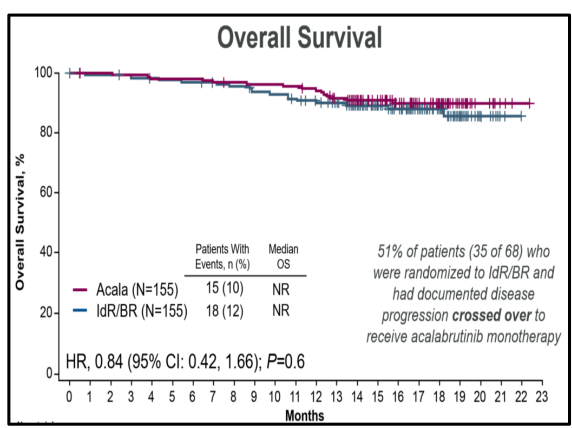
Key secondary endpoints:

- ORR (assessed by IRC and investigator)
- Duration of response
- PFS (assessed by investigator)
- OS

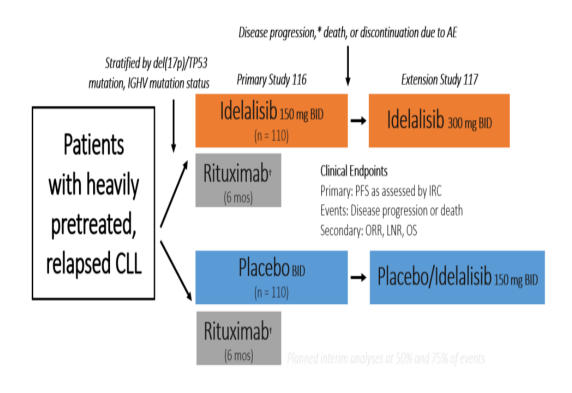
Crossover from IdR/BR arm allowed after confirmed disease progression

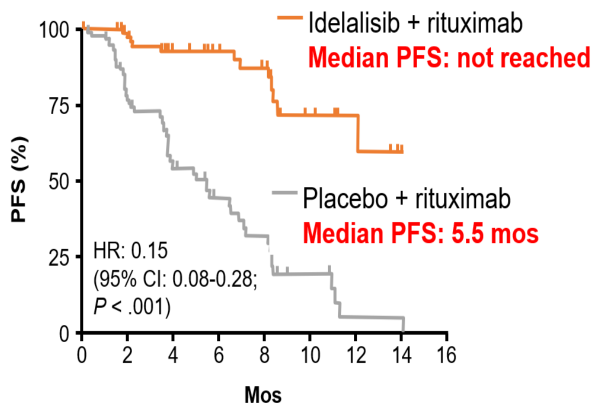
Acalabrutinib vs. Investigator choice for relapsed CLL (ASCEND Study)



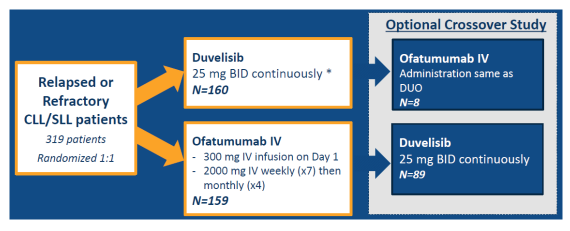


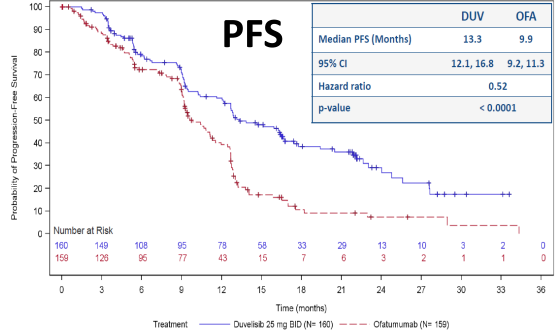
Idelalisib and Rituximab for Previously Treated Patients

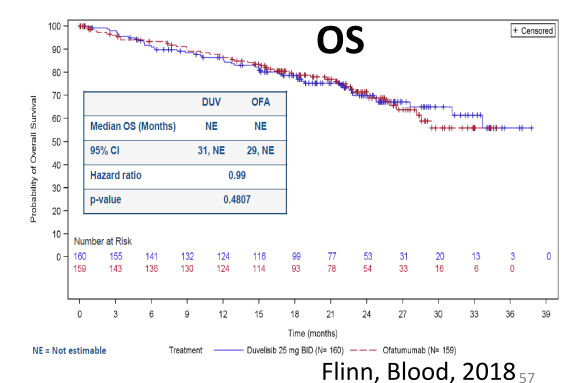




Duvelisib vs Ofatumumab (DUO trial) - Relapsed/Refractory







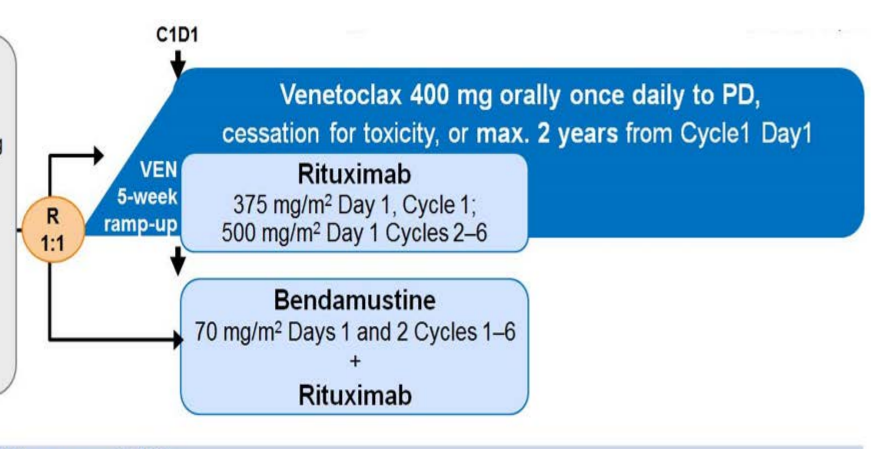
Ven-R vs. BR in R/R CLL (MURANO Study)

Relapsed/refractory CLL (N=389)

- ≥18 years of age
- Prior 1–3 lines of therapy, including ≥1 chemo-containing regimen
- Prior bendamustine only if DoR ≥24 months

Stratified by:

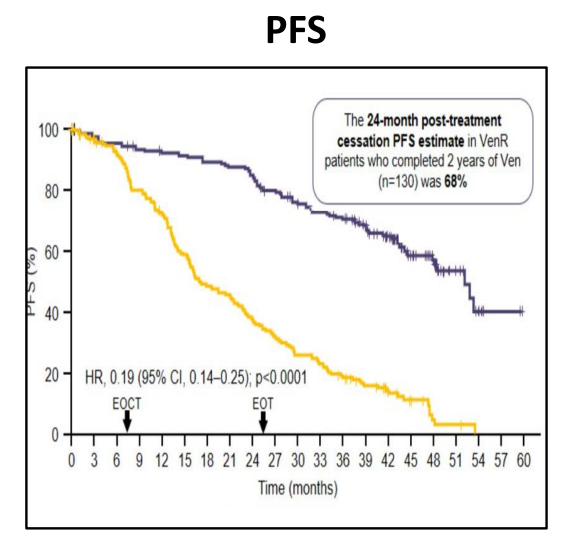
- Del(17p) by local labs
- Responsiveness to prior therapy*
- Geographic region

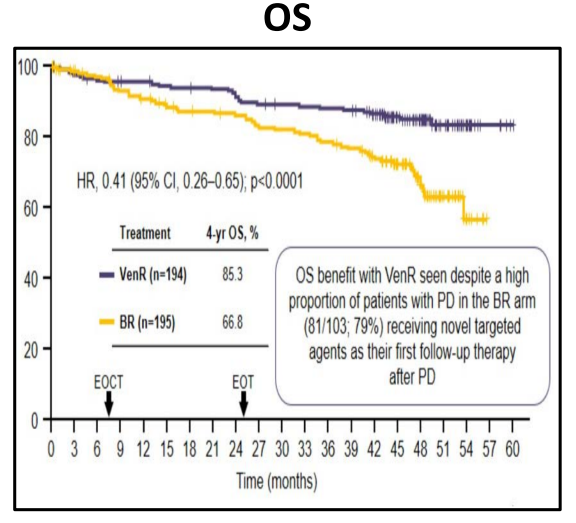


Primary Endpoint

INV-assessed PFS

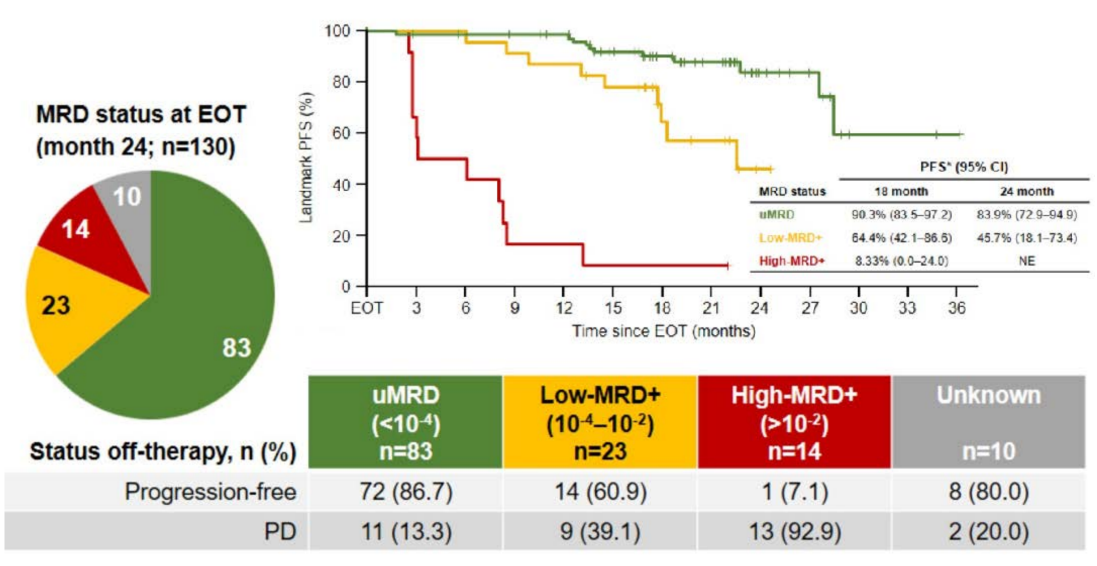
Ven-R vs. BR in R/R CLL (MURANO Study)





Seymour, ASH,2019

Ven-R outcomes (MRD and PFS)



Novel Agents for R/R setting

	Acalabrutinib/ Ibrutinib	Venetoclax	Duvelisib/ Idelalisib
Target	ВТК	BCL-2	PI3K delta+gamma / delta
Duration	Indefinite	2-years	Indefinite
Addition of Anti CD20 Ab	No major benefit Faster "response"	Recommended	Idelalisib + R Duvelisib monotherapy
Major side effect (concern)	Bleeding (anticoagulation)	TLS (initially)	Colitis (diarrhea) Infections (FDA alert)
Other side effects	Body painFatigueHypertensionA fib	Neutropenia	PneumonitisTransaminitis (mainly idela)PJPCMV
FDA label for CLL	All settings	All settings	Relapsed

Previously Treated CLL Summary

1. First

- Venetoclax + Rituximab or
- Ibrutinib or acalabrutinib

Second

- Ibrutinib/acalabrutinib if previously treatment with Ven
- Ven-R if previously treated with BTKi (ibrutinib or acalabrutinib)

3. Third

Idelalisib+ rituximab OR duvelisib

8. Treatment options for patients with del17p/P53 mutation

CLL with del17p or TP53

Acalabrutinib OR **Ibrutinib** Frist line Second line Venetoclax + R Third line **Duvelisib Idelalisib** OR

R = rituximab

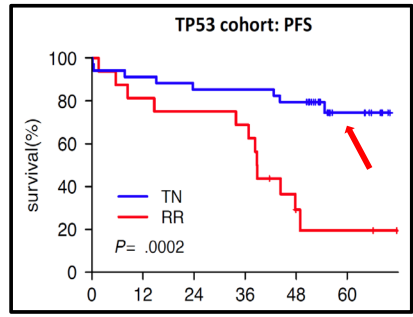
There is no role for chemotherapy in abnormal TP53 (deletion or mutation)

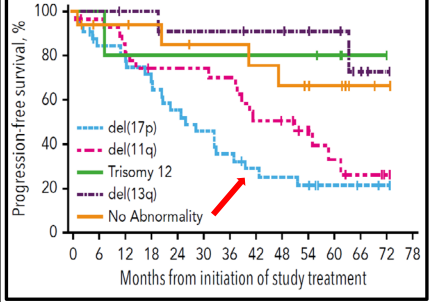
Regimen	PFS
FCR (frontline)	11.3 m
Alemtuzumab (frontline)	11 m
BR (frontline)	7.9 m
HDMP + R	7.5m
BR (relapsed)	7 m
FC (frontline)	6.5 m
FCR (relapsed)	5 m

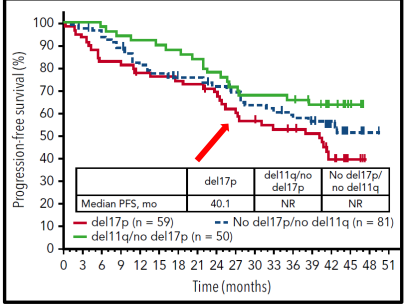
5 – 11 months

Ibrutinib for abnormal TP53

Study	Setting	Outcome
NIH study	TN	5-year PFS 74%
	R/R	5-year PFS 19%
PCYC-1102/1103 5-year f/u	R/R	median PFS 26 m
RESONATE f/u	R/R	Median PFS 40 m

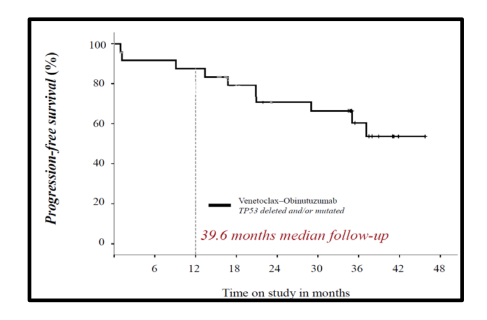


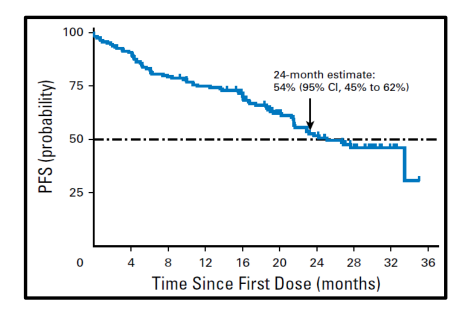




Venetoclax for abnormal TP53

Study	Setting	Outcome
CLL14	TN	24 m PFS 74%
M13-982 study	R/R	24 m PFS 54% m PFS 27 m
MURANO	R/R	m PFS 48 m

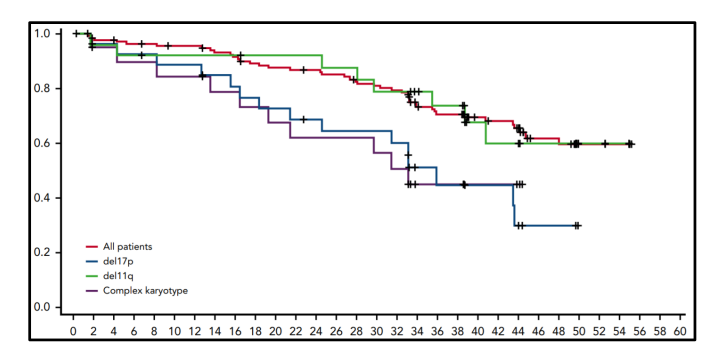




Fischer, ASH, 2019; Stilgenbauer, JCO, 2018; Seymour, ASH, 2019

Acalabrutinib for abnormal TP53

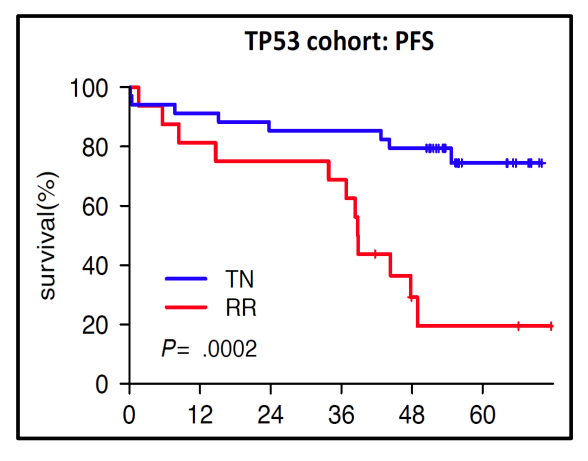
Study	Setting	Outcome
ELEVATE TN	TN	Not reported
ACE-CL-001	R/R	PFS 36m (21 – NR)
ASCEND	R/R	Not reported



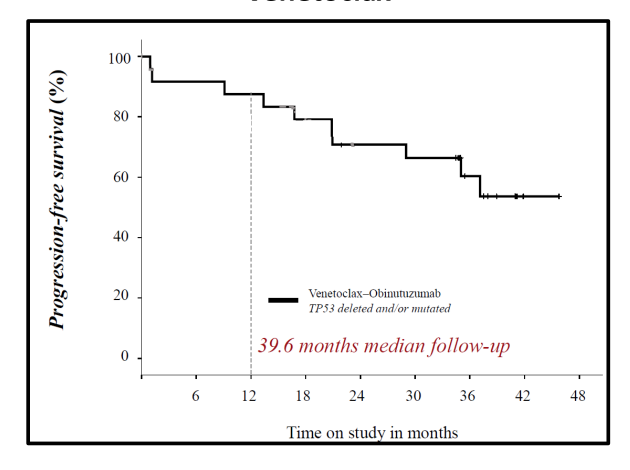
Sharman, ASH, 2019; Byrd, Blood, 2020; Ghia, 15-ICML, 2019

Ibrutinib vs. Ven-G for first-line treatment in CLL patients with abnormal TP53

Ibrutinib



Venetoclax



CLL with del17p or TP53

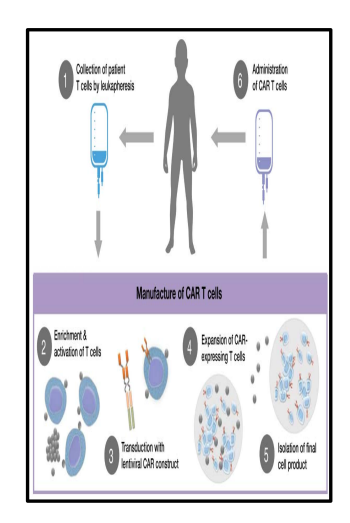
Acalabrutinib OR **Ibrutinib** Frist line Second line Venetoclax + R Third line **Duvelisib Idelalisib** OR

R = rituximab

9. Cellular therapies for CLL

CAR-T for CLL

- Experimental, not FDA approved
- Registration studies are currently ongoing
- Long-term remissions ~ 30-35%
- Best predictor od response: MRD neg after treatment
- Recommend before alloSCT, if available



Allogeneic SCT for High Risk CLL

• Reduced intensity/ Nonmyeloablative allogeneic transplant

Author	Shadman	Kramer	Sorror	Dreger	Brown	Khouri	Khouri	Michallet
Year	2019	2017	2008	2013	2013	2011	2017	2013
N	55	90	82	90	76	86	26	40
Conditioning	Flu-TBI-R	variable	Flu-TBI	FC± ATG	Flu-Bu	FCR	BFR	FCR
Follow-up (yr)	3	10	5	6	5	5	3	3
os	54	51	50	58	63	51	82	55
PFS	45	34	39	38	43	36	63	46
NRM	38 (<12)*	20	23	23	16	17	8	27
aGVHD	20	?	16-23	14	17	7	4	23
Extensive cGVHD	66	?	49-53	55	48	56	45	29

50 40 20-25

* in pts without comorbidities

10. Practical points about novel drugs

New Agents: Practical Considerations

- •BTKi: ibrutinib and acalabrutinib
- •PI3Ki: idelalisib and duvelisib
- Venetoclax

BTKis (ibrutinib/acalabrutinib)

Common side effects:

- Muscle/bone pain
- Cytopenia
- Hypertension
- Diarrhea (early, reversible)

Serious side effect:

- Bleeding: (peri-procedural management)
- Atrial fibrillation
- Opportunistic infections: PJP, aspergillosis (?) (case reports)
- Second generation BTKi, acalabrutinib has a better toxicity profile
 - Acalabrutinib

PI3Kis (idelalisib/duvelisib)

- Important side effects
 - LFT abnormalities (idelalisib)
 - Pneumonitis
 - CMV reactivation and PJP (FDA alert 2016)
- Colitis/Diarrhea
 - Median time to grade III/IV: 7 months
 - Not responsive to anti-motility agents
 - Corticosteroids ; treat as aGVHD



Management of adverse events associated with idelalisib treatment: expert panel opinion

Steven E. Coutré, Jacqueline C. Barrientos, Jennifer R. Brown, Sven de Vos, Richard R. Furman, Michael J. Keating, Daniel Li, Susan M. O'Brien, John M. Pagel, Martin H. Poleski, Jeff P. Sharman, Nai-Shun Yao & Andrew D. Zelenetz

Don't use Idelalisib in treatment naïve patients!

Toxicity Frequency

	Phase I	Overall relapsed	Upfront Pts ≥ 65 yo	Upfront younger Pts	
Number of patients	54	760	64	24	
Median prior treatments	5 (2-14)	≥ 1	0	0	
Median age	63 (37-82)	66 (21-91)	71 (65-90)	67 (58-85)	
Median time to therapy (months)	15 (0.2-49)	-	22 (0.8 – 46)	8 (0.7-16)	
Grade≥ 3 transaminitis	1.9%	14%	23%	52%	
Grade ≥3 Colitis/diarrhea	5.6%	14%	42%	13%	
Any grade pneumonitis	5.6%	3%	3%	13%	
Reference	Brown Blood 2014	Coutre EHA 2015	O'Brein Blood 2015	Lampson ASH 2015	

Venetoclax



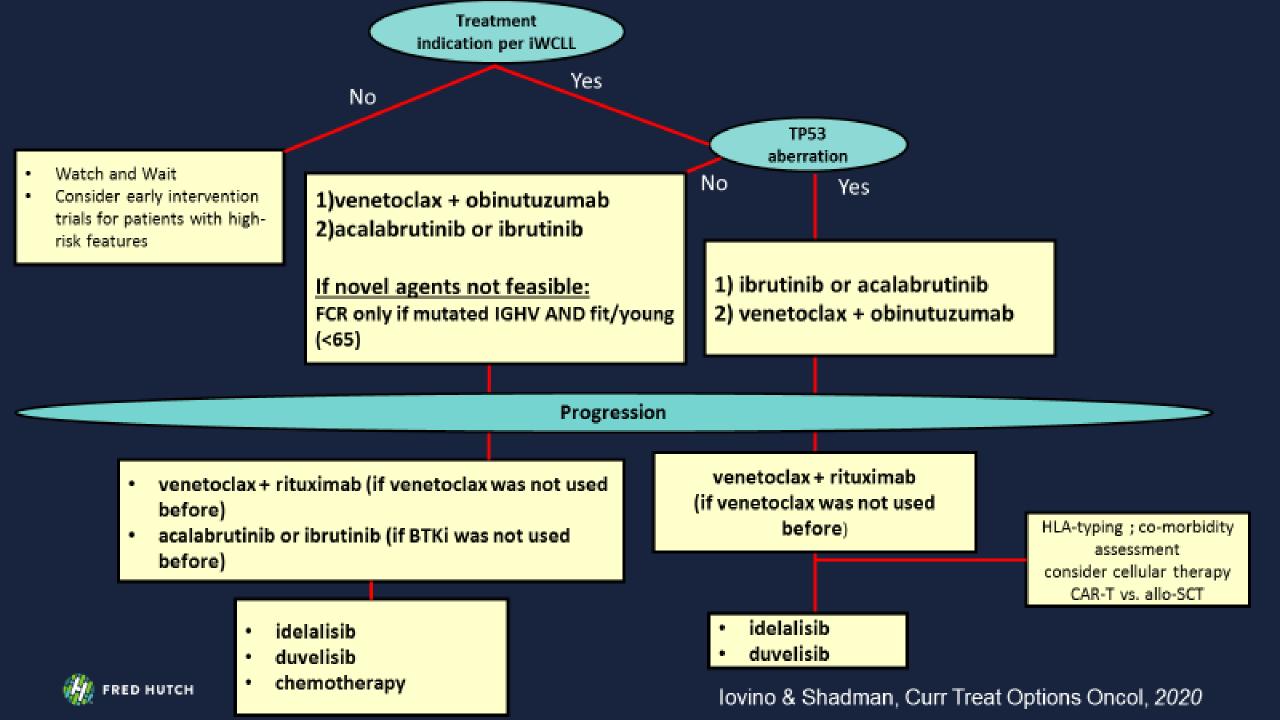
Tumor Burden		Pro	phylaxis	Blood Chemistry Monitoring ^c			
		Hydration ^a	Antí- hyperuricemics	Setting and Frequency of Assessments			
Low	All LN <5 cm AND ALC <25 × 10 ⁹ /L	Oral (1.5-2 L)	Allopurinol ^b	Outpatient Pre-dose, 6-8 hours, 24 hours at first dose of 20 mg and 50 mg Pre-dose at subsequent ramp up doses			
Medium	Any LN 5 cm to <10 cm OR ALC ≥25 × 10 ⁹ /L	Oral (1.5-2 L) and consider additional intravenous	Allopurinol	Outpatient Pre-dose, 6-8 hours, 24 hours at first dose of 20 mg and 50 mg Pre-dose at subsequent ramp up doses Consider hospitalization for patients with CrCl <80 mL/min at first dose of 20 mg and 50 mg; see below for monitoring in hospital			
High	Any LN ≥10 cm OR ALC ≥25 × 10 ⁹ /L AND any LN ≥5 cm	Oral (1.5-2 L) and intravenous (150-200 mL/hr as tolerated)	Allopurinol;consider rasburicase ifbaseline uric acid is elevated	In hospital at first dose of 20 mg and 50 mg • Pre-dose, 4, 8,12 and 24 hours Outpatient at subsequent ramp-up doses • Pre-dose, 6-8 hours, 24 hours			

 For patients at risk of TLS, monitor blood chemistries at 6-8 hours and at 24 hours after each subsequent ramp-up dose

Venetoclax

- Consider debulking strategies
- Follow the standard ramp-up schedule
- Coordinate with the inpatient team
- Selected patients can be treated using the "escalated inpatient ramp-up" *
- Follow ALL TLS labs (not just uric acid!)

Will take some effort to start

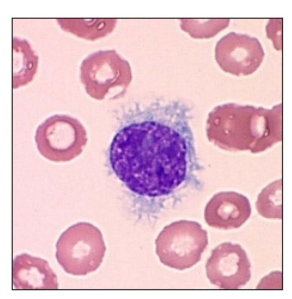


CLL (Night before the test)

- 1. Flow cytometry is critical (and adequate) to make the diagnosis
- 2. Remember CLL immunophenotype (and differences with MCL and other lymphomas)
- 3. Review Indications for treatment. This hasn't change even with new agents.
- 4. Check FISH before each line of treatment (r/o del 17p/P53 mutation)
- 5. Frontline (no del17p or P53mutation): Ven-G or BTKi (acalabrutinib is better tolerated). FCR reasonable for: fit,<65 and mutated IGHV.
- 6. Relapsed setting: Ven-R or BTKi (acalabrutinib better tolerated), idelalisib/duvelisib.
- 7. For del 17p patients: BTKi (more data with ibrutinib), Ven-R, cellular therapy, idelalisib/duvelisib.
- 8. BTKi: initial lymphocytosis (is OK), bleeding, Afib, HTN, body pain (acalabrutinib is better tolerated)
- 9. Idelalisib/duvelisib: lymphocytosis (is OK), colitis, pneumonitis, hepatitis (more with idela), PJP, CMV Don't use in frontline setting
- 10. Venetoclax: watch for TLS at the beginning. Ramp-up HAS to be done!

Hairy Cell Leukemia

- ➤ Uncommon chronic B cell lymphoid neoplasm
- ➤ Small mature B cell lymphoid cells with abundant cytoplasm and "hairy" projections within the peripheral blood, bone marrow, and splenic red pulp
- > Splenomegaly and cytopenias



Hairy cell Leukemia (Diagnosis)

	CD11c	CD25	CD103	CD123	CD10	CD21	CD23	CD5	CD20	CD19	CD22	Annexin A1
HCL	+	+	+	+	-	•	•	-	+	+	+	+

BRAF V600E mutation is a disease-defining event

HCL variant:

CD25 (-), CD123 (-), annexin A1 (-) and BRAF V600E (-)

Hairy cell Leukemia

Clinical presentation

- Splenomegaly
- Cytopenias (infections, bleeding)
- Constitutional symptoms

• Treatment Indications:

- Systemic symptoms
- Splenic discomfort
- Recurrent infections
- Cytopenias (Hb <11, ANC < 1000, bleeding due to plt <100,000)

Hairy Cell Leukemia

Treatment

- First Line
 - Purine analogs
 - Cladrabine (2-CdA) Up to 80% CR with a CR duration of 57 months (7 – 246) after a single cycle
 - Pentostatin
- Refractory (failure in less than a year) or Relapsed disease
 - Purine analogs ± Rituximab
 - INF-alfa
 - Rituximab
 - BRAF targeting agents (Vemurafenib)
 - Moxetumomab Pasudotox (anti CD22 immunotoxin conjugate)

Moxetumomab Pasudotox for R/R HCL

- Anti CD22 immunotoxin conjugate
- IV; D1,3,5 of 28D cycle (up to 6 cycles)
- At least 2 prior systemic therapies, including a purine analog

• Efficacy:

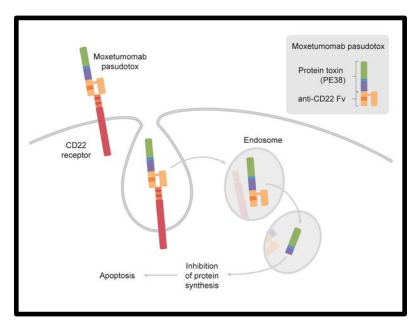
• ORR: 75%

durable CR: 30%

• MRD eradication 34% of all CRs

• Unique side effects

- 1. Hemolytic-uremic syndrome
- 2. Capillary leak syndrome
- supportive care and discontinuation were effective
- could occur at any cycle



Please Consider Clinical Trials!

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