

LEARNING OBJECTIVES

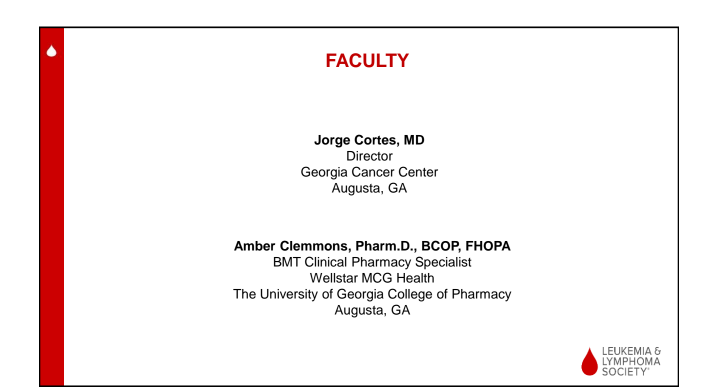
- Describe the various types and subtypes of chronic myeloid leukemia (CML)
- · Identify tests used to diagnose disease and monitor treatment of CML
- · Explain the overarching goals of treatment for the types of CML
- Explain approved and emerging treatment options for CML, including stem cell transplantation, and the role of clinical trials.
- Describe strategies to manage treatment side effects as well as potential long-term and late effects of treatments for CML
- Describe the healthcare professional's role in managing patients with CML



1

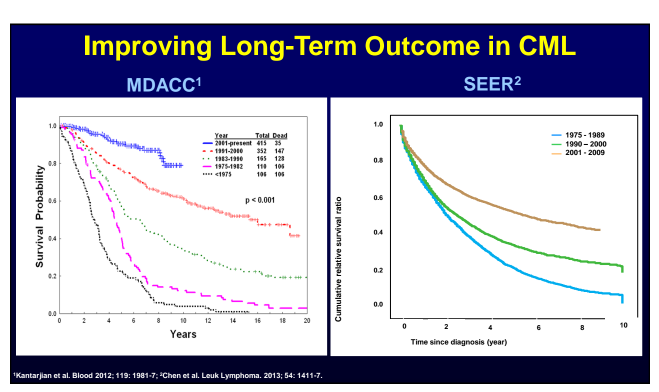
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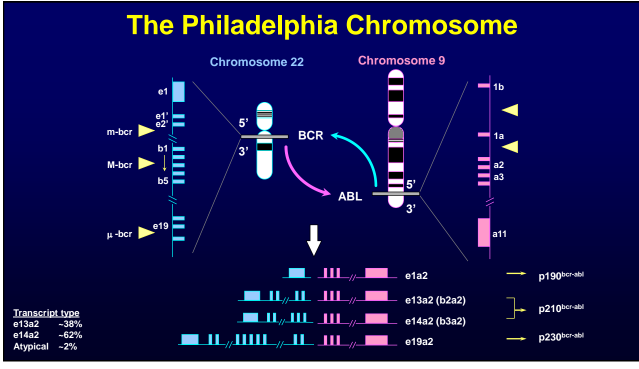
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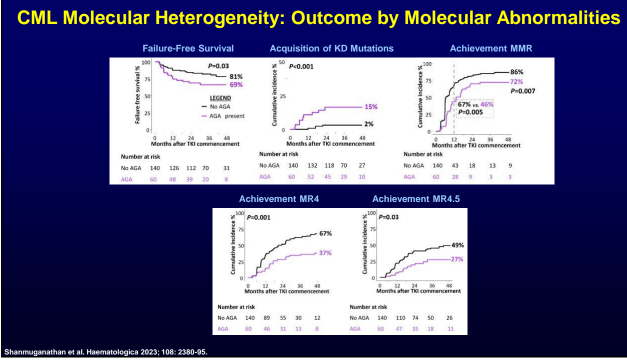
Disclosure Information Jorge Cortes

- I have the following financial relationships to disclose:
 - Grant or research support (to my institution) from: Ascentage, *Novartis, Sun Pharma*
 - Paid Consultant for: Novartis, Pfizer, Sun Pharma, Terns
 - Clinical Investigator for: Ascentage, Novartis, Sun Pharma
- AND
 - I will NOT include discussion of investigational or off-label use of a product in my presentation.



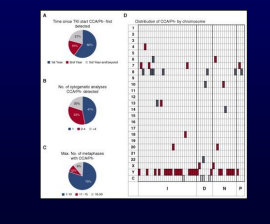


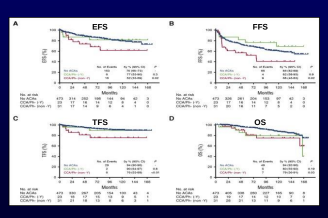
The Molecular Heterogeneity of CML • 200 patients from TIDEL-II sequenced with RNA capture panel (126 genes) • At diagnosis: 40 SNV and indels in cancer genes in 33 pts (16%) (across 10 genes); Ph-associated rearrangements in 36 patients (18%). Ph-associated rearrangements Additional genetic abnormalities at diagnosis ale ID Variant subtype Missense Frameshift Nonsense Splice variant Structural variant Ph-associated rearrangement BC na 5 Ech me 6 Chro ma 7 🔲 Cł nosome 16 Chromosome 17 Chromosome 19 Chromosome 20 Chromosome 22 Shanmuganathan et al. Haematologica 2023; 108: 2380-95.



Additional Chromosomal Abnormalities (ACA) in CML

- 598 pts treated frontline with TKI
- Clonal chromosomal abnormalities in Ph- metaphases occurred in 58 (10%) pts
- Median time to appearance: 6 mo (3-78 mo) (1st occurrence after 12 mo in 39%)
- Transient (≤2 times) in 36 (62%)





Issa et al. Blood 2017; 130: 2084-91.

Chronic	Accelerated	Blast*
Past 3-5 years	12-18 months	3-9 months
Present 25+ years	4-5 years	6-12 months
 Asymptomatic (if treated) None of criteria for accelerated or blast phase 	 Blasts ≥ 15% BI + pros ≥ 30% Basophils ≥ 20% Plts < 100,000/mcl Clonal evolution 	 Blasts ≥ 30% Extramedullary disease with localized immature blasts Increased lymphoblasts in PB or BM

* Blast phase is not AML or ALL

2022 CML Classifications

• WHO

- AP is omitted in favors of an emphasis on high-risk features associated with CP progression and resistance to TKI.
- BP criteria: (1) ≥20% myeloid blasts in PB or BM; or (2) extramedullary proliferation of blasts; or (3) increased lymphoblasts in PB or BM. (The optimal cutoff for lymphoblasts and the significance of low-level B-lymphoblasts remain unclear and require additional studies.)
- ICC
 - AP criteria: (1) 10%-19% blasts in PB or BM; or (2) PB basophils ≥20%; or (3) additional clonal cytogenetic abnormality in Ph+ cells (ACA)^{1.}
 - BP criteria: (1) ≥20% blasts in PB or BM; or (2) myeloid sarcoma²; or (3) morphologically apparent lymphoblasts (>5%)³ ("warrants consideration of lymphoblastic crisis")

¹Second Ph, trisomy 8, isochromosome 17q, trisomy 19, complex karyotype, or 3q26.2 abnormalities. ²Extramedullary blast proliferation. ³Immunophenotypic analysis is required to confirm lymphoid lineage. Khoury et al. Leukemia 2022; 36: 1703-19; Arber et al. Blood 2022; 140: 120-8.

11

ELN 2020 - Diagnostic Work-up at Baseline

- Physical examination (spleen and liver size)
- Complete blood cell count with microscopic differential
- Bone marrow aspirate for cytologic examination and cytogenetics
 - Core biopsy on dry tap
- Chromosome banding analysis
- FISH (only in case of Ph-negativity)
- Qualitative PCR (for detection of BCR-ABL1 transcripts and identification of transcript type)
- ECG
- Standard biochemistry with hepatitis B-serology
- Consider mutation analysis for AP/BP (NCCN)

Prognostic Scores in CML

- Sokal: age, spleen, platelets, blasts
- Hasford (EURO): age, spleen, platelets, blasts, eosinophils, basophils
- EUTOS: spleen, basophils

https://www.leukemia-net.org/content/leukemias/cml/euro__and_sokal_score/index_eng.html https://www.leukemia-net.org/content/leukemias/cml/eutos_score/index_eng.html

• ELTS: age, spleen, platelets, blasts

Response Criteria for CML

Cytogenetic response	% Ph*	Molecular response	% BCR-ABL/ ABL (IS)
Minor (mCyR)	36-95		>10
Partial (PCyR)	1-35		1 to <10
			>0.01 to <1
		Major (MMR; MR3)	≤0.1
Complete		MR4	≤0.01
(CCyR)		MR4.5 Better	
		Deep mo	
		Questionable	

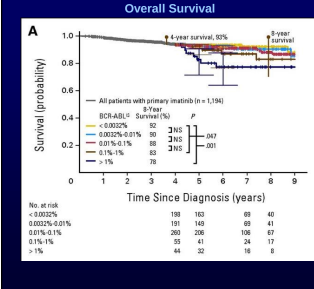
7-Year Outcome by Molecular Response Among Patients with CCyR

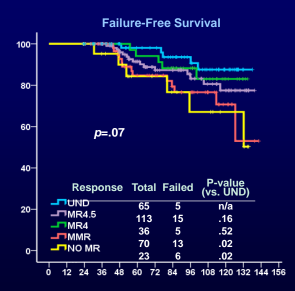
Londmork		Percentage			
Landmark		MMR	No MMR		
	EFS	85	93		
6 mo	TFS	96	98		
	OS	90	93		
	EFS	91	92		
12 mo	TFS	99	96		
	OS	93	97		
	EFS	95	86		
18 mo	TFS	99	96		
	OS	95	96		

Hughes T, et al. Blood 2010; 116: 3758-65

15

Outcome by Depth of Molecular Response





Hehlmann et al. JCO 2014; 32: 415-23. Falchi et al. Am J Hem 2013; 88: 1024-9.

Relative Survival with TKI by Response to Therapy • 483 pts with CML treated with imatinib 400mg (n=71), imatinib 800 mg (n=201), dasatinib (n=111) or nilotinib (n=101) • 5-yr relative survival 94.8% [92.1 - 97.4] 100 Relative Survival 80 60 40 20 0 120 0 12 36 48 60 72 96 108 All 100.0 99.6 99.5 96.7 94.8 95.0 92.7 92.5 90.6 88.3 98.2 97.2 -CCyR 100.0 99.7 100.1 99.7 98.4 96.7 95.1 95.3 94.0 91.7 - MMR 94.5 100.0 99.9 100.3 100.4 100.0 98.9 99.5 97.2 96.4 ~* 101.4 100.4 101.0 96.7 100.0 100.4 100.8 99.2 99.4 98.3 CMR 100.0 100.4 100.8 101.3 101.8 101.4 102.0 100.2 100.1 98.5 98.2 Month Sasaki et al. Lancet Hematology 2015

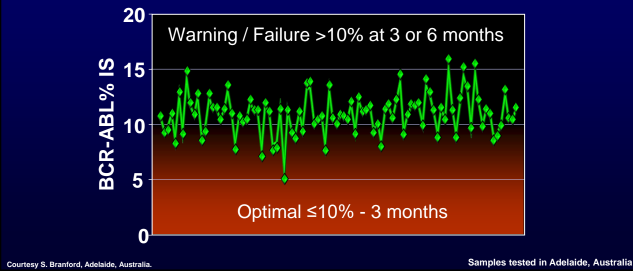
17

When	ELN	NCCN
At diagnosis	•CG (BM aspiration) •FISH (in case of Ph-) •PCR	 CG (BM aspiration) FISH (in case of Ph-) PCR
During treatment	 PCR (IS) every 3 months In patients with atypical translocations, rare or atypical BCR-ABL1 transcripts that cannot be measured by qPCR, treatment failure/resistance to exclude ACA, and with progression to AP or BP FISH may be needed in patients with atypical transcripts 	years after BCR-ABL1 ≤1% IS
Failure, progression	 PCR (IS), mutation analysis, cytogenetics Immunophenotype for BP 	 PCR (IS), mutation analysis, cytogenetics

Hochhaus et al. Leukemia 2020; 34: 966-984.

Variability of a BCR-ABL1 Positive Sample Measured 96 Times in a Single Centre Over Several Months

Mean BCR-ABL1 value = 11% (range 5-17%); CV 18%

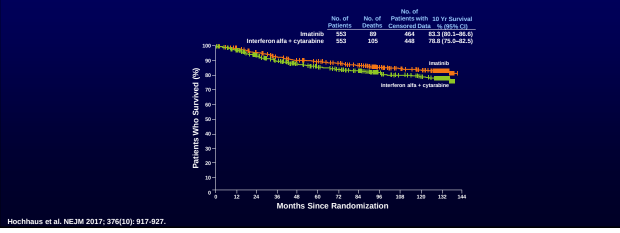


ELN 2020 Treatment Recommendations

	Imatinib
•	Dasatinib
	Nilotinib
	Generic imatinib
	Bosutinib
S	econd line and subsequent lines
•	The criteria for the choice of 2 nd -line TKI almost entirely patient related and depend on factors such as age, comorbidities and toxicity of 1 st TKI
,	Imatinib
	Dasatinib
	Nilotinib
	Bosutinib: 2 nd -line in patients with prior TKI failure/resistance or intolerance
•	
	Ponatinib: after failure of 2GTKI or T315I

Results With Imatinib in Early CP CML the IRIS Trial at 10 Years

- 49% discontinued therapy
- 10 yr CCyR 92%, MMR 93%, MR4.5 63% (ITT 22%, 34%, 23%, respectively)
- 38 pts (7%) transformed to AP/BP (34 during 1st 4 yrs)
- 10-yr freedom from transformation 92%, EFS 80%



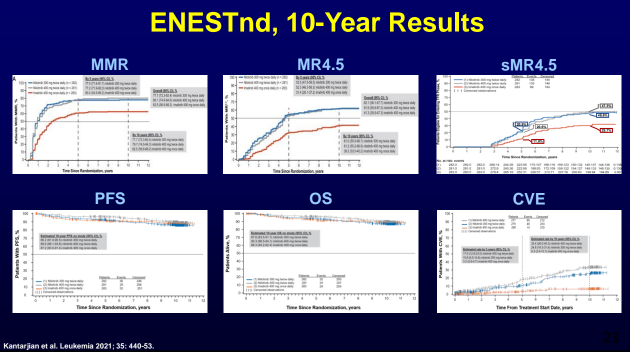
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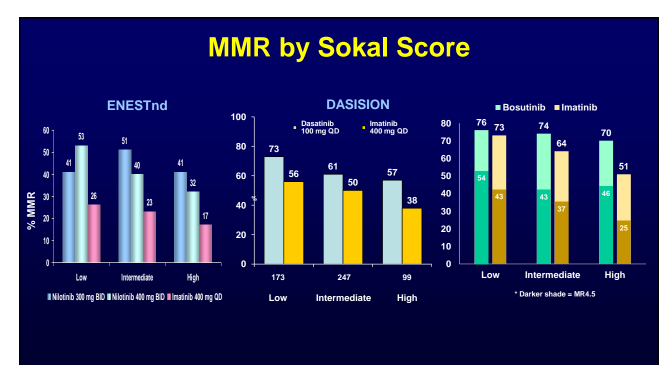
Outcome Across 1st Line CML Studies

Response	DASI	SION	ENE	STnd	BFC	RE	ТО	PS
at, %	DAS	IMA	NIL ^b	IMA	BOS	IMA	IMA	IMA
CCyR 12m	77	66	80	65	77	66	70	66
MMR 3m	8	0.4	9	1	4.1	1.7	12	3
MMR 12m	46 ^a	28 ª	44	22	47	37	47	40
MMR 5 yr	76	64	77	60	74	66		
MR4.5 5 yr	42	33	54	31	46	36		
AP/BP	2.3	5.0	1	6	2.2	2.6	1.9	3.2
PFS	94	92	96	94	NR	NR	97	94
OS	95.3	95.2	97	96	99	97	99	98

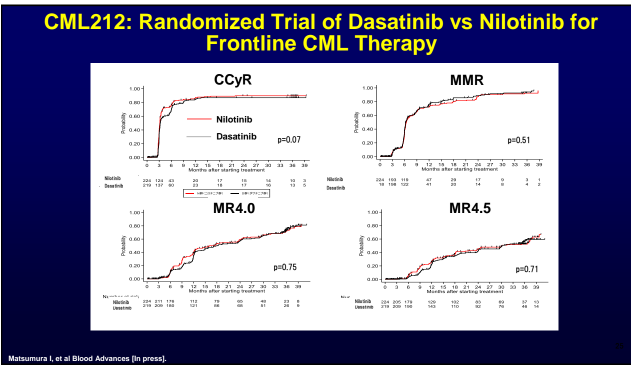
^a MMR by 12 mo; ^b Nilotinib 300 mg BID

Cortes et al. JCO 2016; 34: 2333-40; Hochhaus et al. Leukemia 2016; 30: 1044-54; Brümmendorf TH, et al. Leukemia 2022; 36: 1825-33; Cortes et al. JCO 2010; 28: 424-30.



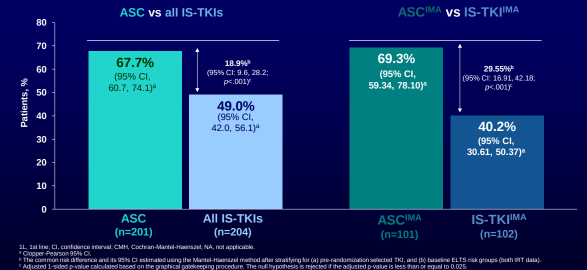


Saglio et al. Blood 2009; 114: abst# LBA-1; Kantarjian et al. ASCO 2011; abstract #6510; Brümendorf et al. ASH 2020; abstract #46



25

MMR Rate at Week 48 (Primary endpoint)



Recommendations for Management According to Response – ELN 2020

- **Optimal: Continue** ۲
- Failure/resistance: Change ۲
- ۲ Warning:
 - Carefully consider continuation or change, depending on • patients' characteristics, comorbidities and tolerance.
 - Additional qPCR testing may be indicated if the kinetics of the response are not clear, or if toxicity or intolerance cause dose interruptions or reductions.

27

CML Recommendations ELN & NCCN*

	ELN 2020							
	Optimal	Warnings	Failure					
Baseline	NA	High-risk ACA, high-risk ELTS score	NA					
3 months	BCR-ABL1 ≤10%	BCR-ABL1 >10%	BCR-ABL1 >10%, if confirmed within 1-3 months					
6 months	BCR-ABL1 ≤1%	BCR-ABL1 >1%-10%	BCR-ABL1 >10%					
12 months	BCR-ABL1 ≤0.1%	BCR-ABL1 >0.1%- 1%	BCR-ABL1 >1%					
Then, at any time	BCR-ABL1 ≤0.1%	BCR-ABL1 >0.1%- 1% Loss of ≤0.1% (MMR)ª	BCR-ABL1 >1%, resistance mutations, high-risk ACA					

			N	CCN		
BCR:	CR::ABL1 (IS) 3 months 6 month			hs	12 months ^m	
>10% ⁿ YELLOW			R	ED		
>1%-10% GREEN				YELLOW		
>0.1%-1%			GREE	N		LIGHT GREEN
≤0.1% GREEN						
COLOR	CONCERN	CL	INICAL CONSIDERATIONS ^P	1	RECOMMENDATIONSP	
RED	TKI-resistant disease ⁰	• <u>c</u>	valuate patient adherence and drug inter onsider <u>BCR::ABL1 kinase domain muta</u> onsider bone marrow cytogenetic analys dditional chromosomal abnormalities (Al	itional analysis ^q sis to assess	Switch to alternate TKI (<u>CML_5</u>) (other than imatinib) and evaluate for allogeneic HCT	
YELLOW	Possible TKI resistance ⁰	0.0	valuate patient adherence and drug inter onsider <u>BCR::ABL1 kinase domain muta</u> onsider bone marrow cytogenetic analys ICyR at 3 mo or CCyR at 12 mo	tional analysis ^q	Continue same	mate TKI (<u>CML-5)</u> or e TKI (<u>CML-G)</u> ' evaluation for allogeneic HCT
LIGHT Green	TKI-sensitive disease	۰lf	valuate patient adherence and drug inter treatment goal is long-term survival: ≤1 treatment goal is treatment-free remission	% optimal	 If not optimal 	ontinue same TKI (<u>CML-G</u>) I: shared decision-making with
GREEN	TKI-sensitive disease		lonitor response (<u>CML-E)</u> valuate patient adherence and drug inter	ractions	Continue same	e TKI (<u>CML-G</u>) ^t

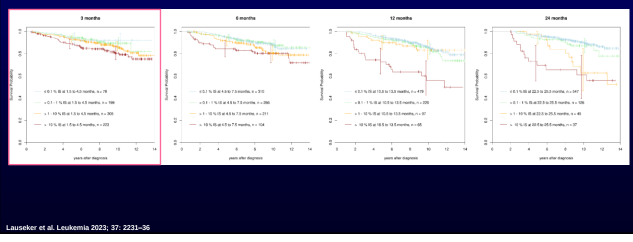
*For the latest information, access NCCN guidelines at www.NCCN.org.

Major differences:

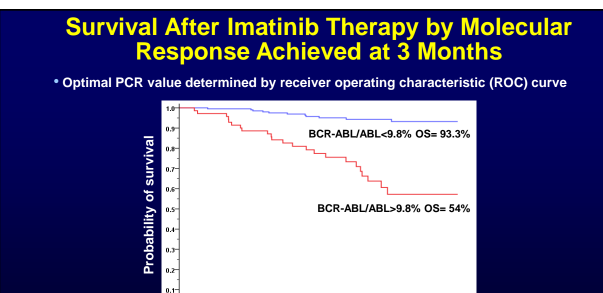
- No baseline features in NCCN (warning only in ELN) 6-month >1-10% optimal by NCCN, warning by ELN 12-month >1-10% "yellow" by NCCN, failure by ELN Losses not included in NCCN ("Early treatment milestones")

Hochhaus et al. Leukemia 2020; 34: 966-984; NCCN Version 2.2024 – December 5, 2023 (<u>https://www.nccn.org/professionals/physician_gls/pdf/cml.pdf</u>)

Benefit of TKI Treatment After Failing Milestones





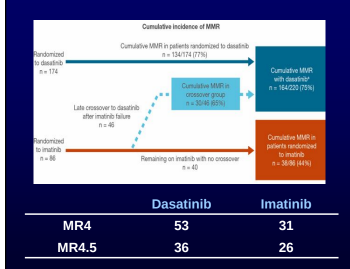


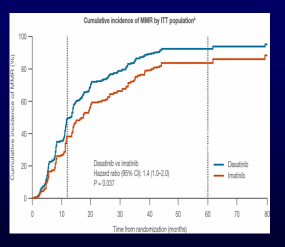
Time from onset of imatinib therapy (years)

p<0.0001

0.0

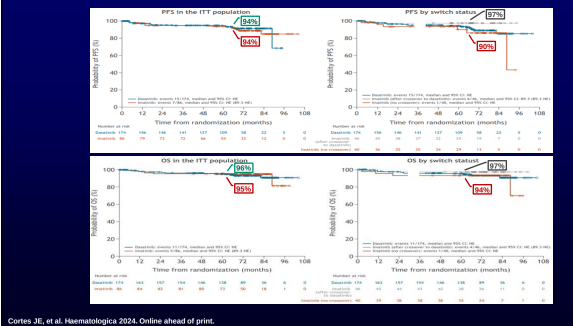
DASCERN - Cumulative Incidence of Response





^aFour patients achieved then lost MMR and subsequently crossed over to dasatinib; ^bThe Kaplan–Meier curve accounts for competing risk and censored patients. Cortes JE, et al. Haematologica 2024. Online ahead of print.





Treatment Discontinuation by TKI

	DASISION		ENESTnd		BFORE		
	Dasatinib	Imatinib	Nilotinib	Imatinib	Bosutinib	Imatinib	
2 yrs	23	25	26	33	29	31	
Efficacy	9	11	9	17	5	15	
Safety	9	5	9	10	19	11	
5 yrs	39	37	39	50	40	42	
Efficacy	11	14	13	25	6	18	
Safety	21	9	12	14	25	13	
10 yrs		-	53ª	48 ^b	-	-	
Efficacy			5 ^a	6 ^b			
Safety			22	35			

^a 62% including those who switched to imatinib or increased to nilotinib 400 mg BID (14% for efficacy)
 ^b 65% including those who switched to nilotinib or increased imatinib dose (24% for efficacy)

Cortes et al. JCO 2016; 34: 2333-40; Hochhaus et al. Leukemia 2016; 30: 1044-54; Kantarjian et al. Blood 2012; 119: 1123-9; Kantarjian et al. Lancet Oncology 2011; 12: 841-51; Cortes et al. J Clin Oncol 36, 2018 (suppl; abstr 7002); Brümmendorf et al. ASH 2020; abstract #46.

Mechanisms of Resistance to Imatinib

- BCR-ABL-Dependent
 - Mutations in ABL
 - Amplification/overexpression
 - Remigration of BCR-ABL to cytoplasm
- BCR-ABL-Independent
 - Decreased hOCT1 expression
 - Increased MDR expression
 - Increased alpha-1 acid glycoprotein
 - Overexpression of Src-related kinases
- Quiescent stem cells (Persistence)

LeCoutre Blood 95: 1758, 2000.Weisberg Blood 95: 3498, 2000. Mahon Blood 96: 1070, 2000. JNCI 92:1641, 2000. Vigneri Nature Medicine 7: 228, 2001.

Recommended TKIs in Case of BCR-ABL1 Resistance Mutations – ELN 2020

Mutation	Recommended TKI
T315I	Ponatinib
F317L/V/I/C, T315A	Nilotinib, bosutinib* or ponatinib
V299L	Nilotinib or ponatinib
Y253H, E255V/K, F359V/I/C	Dasatinib, bosutinib* or ponatinib

* There are limited data available regarding mutations associated with clinical resistance to bosutinib in vivo. Some in vitro data suggest that the E255K and, to a lesser extent, the E255V mutation, might be poorly sensitive to bosutinib.

 Asciminib not in 2020 ELN because it was not available at the time, but it can be recommended in all instances

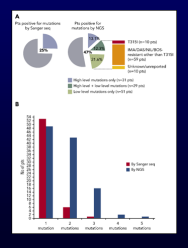
Hochhaus et al. Leukemia 2020; 34: 966-984.

35

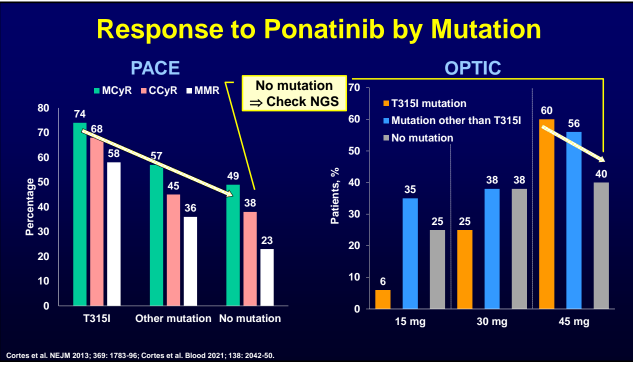
BCR-ABL1 KD Mutations After TKI Failure/Warning

 236 consecutive CML patients with a non-optimal response to TKI therapy, 124 (53%) failure, 112 (47%) warning

	Mutations by SS	Mutations by NGS
1 st -line failure	13/57 (23)	27/57 (47)
1 st -line warning	7/68 (10)	23/68 (34)
2 nd -line failure	15/39 (38)	20/39 (51)
2 nd -line warning	6/37 (18)	17/37 (49)
3 rd -line failure	14/21 (67)	17/21 (80)
3 rd -line warning	1/7	3/7
4 th or 5 th -line failure	4/7	4/7
Total	60/236 (25)	111/236 (47)



Soverini et al. Blood 2020; 135: 534-41.



2nd Generation TKI in CML CP Post-Imatinib Resistance

Paspapa	Percentage				
Response	Dasatinib [†]	Nilotinib [‡]	Bosutinib		
FU (mo)	>24	>24	>24		
CHR	89	77	85		
MCyR	59	56	57		
CCyR	44	41	41		
24 mo PFS*	80%	64%	79%		
24 mo OS*	91%	87%	92%		

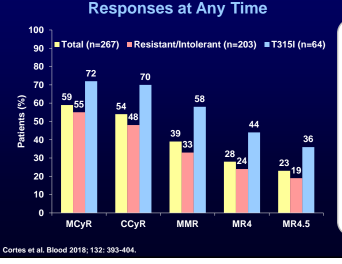
7-yr MMR 43%, PFS 42%, OS 65%; discontinued 78% : 4-yr PFS 57%, OS 78%; discontinued 70% All patients (resistant + intolerant)

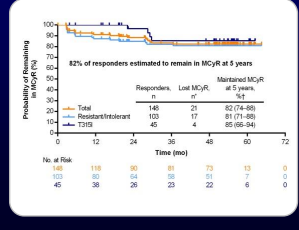
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Shah et al. Haematologica 2010; 95: 232-40; Shah et al. Am J Hematol 2016; 91: 869-74. Kantarjian et al. Blood 2011; 117: 1141-45; Giles et al. Leukemia 2013; 27: 107-112 Cortes et al. Blood 2011; 118; 456-776; Gambacorti-Passerini et al. Am J Hematol 2014; 89: 732-42.

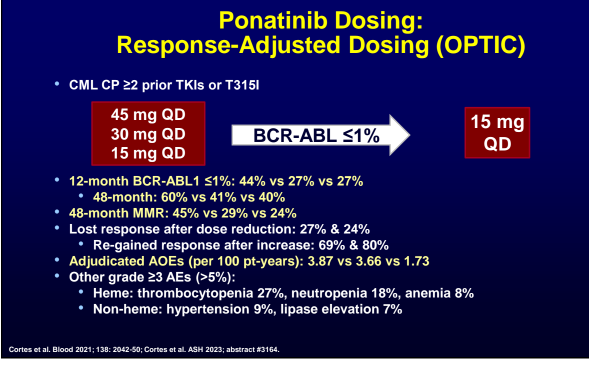
Efficacy of Ponatinib in CP-CML

 Median times to MCyR 2.8 (1.6–24.5) mo, CCyR 2.8 (1.6–35.7) mo, and MMR 5.5 (1.8–32.9) mo



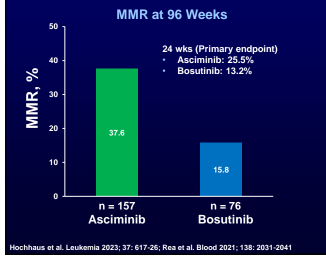


Duration of MCyR



ASCEMBL – Asciminib vs Bosutinib in R/R CML CP

- 233 pts previously treated with ≥2 TKIs randomized 2:1 to asciminib 40 mg BID or bosutinib 500 mg QD
- T315I and V299L excluded



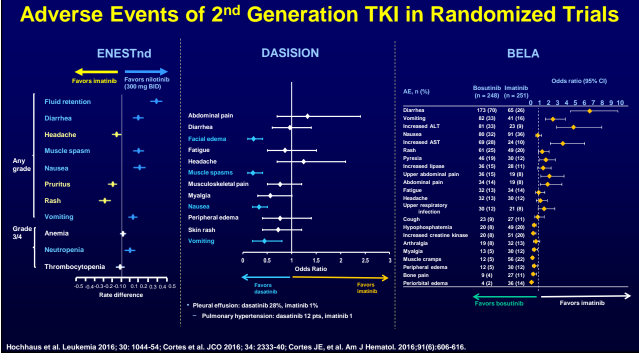
- Median wks to MMR: asciminib 12.7 vs bosutinib 14.3
- Median wks exposure: asciminib 43.4 (0.1-129.9), bosutinib 29.2 (1.0-117.0)
- Other efficacy endpoints:
 - CCyR: 40.8% v 24.2% (96 w: 45.1% v 19.4%)
 - MR4: 10.8% v 5.3%
 - MR4.5: 8.9% v 1.3%
- TEAEs ≥G3 >2%: thrombocytopenia 22%, neutropenia 19%, hypertension 6.4%, ↑ lipase 3.8%
- AOEs (per 100 pts-years): asciminib 3.0, bosutinib 1.4

Asciminib for T315I CML - Response

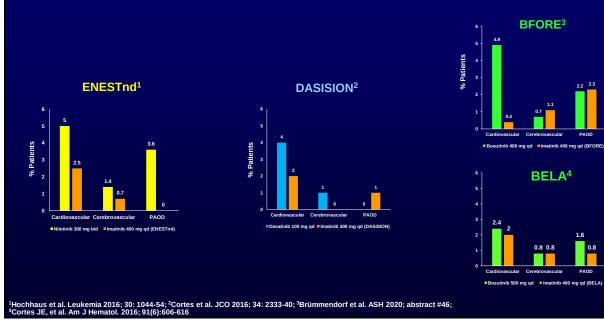
•	Ascimini	b 200 mc	g twice daily	V

Patients, n (%)	MMR	MR4	MR4.5
All patients (n = 49)	23 (46.9)	13 (26.5)	10 (20.4)
Ponatinib naive (n = 21)	12 (57.8)	8 (38.1)	7 (33.3)
Ponatinib pretreated (n = 28)	8 (28.6)	5 (17.9)	3 (10.7)

- Median time to MMR: 12.1 weeks (range, 4 to 48 weeks)
 - Kaplan-Meier-estimated MMR duration at 144 weeks (2.8 years): 87% (95% CI: 68.4%, 100%)
- Median time to MR4: 20 weeks (range, 8 to 33 weeks)
- Median time to MR4.5: 20 weeks (range, 8 to 49 weeks)



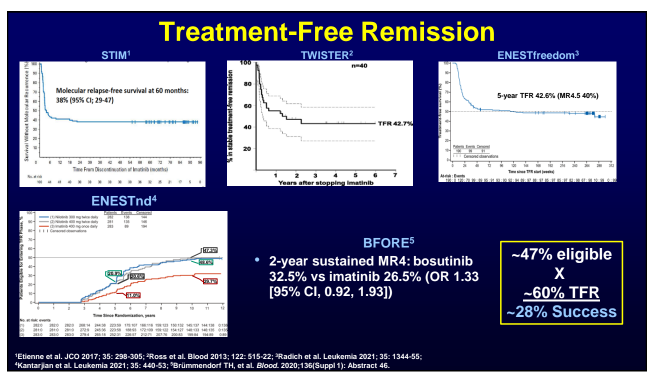
Ischemic Events by TKI From Randomized Trials at 5 years

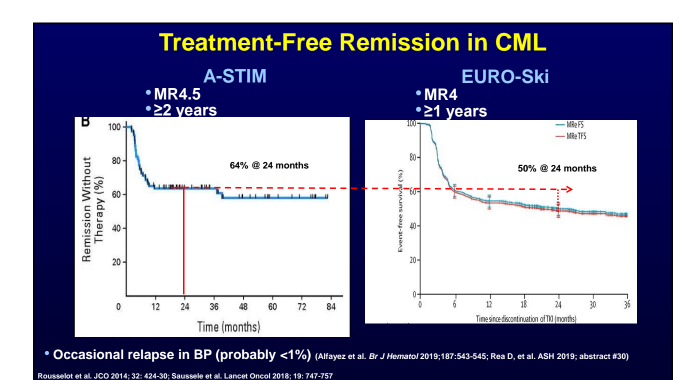


Requirements for TKI Discontinuation – ELN & NCCN* 2020

ELN	NCCN
CML 1 st CP only (Mand)	CP CML. No history of AP or BP
TKI therapy >5 y (>4 y for 2GTKI) (Min)	On approved TKI ≥3 y
e13a2- or e14a2-BCR–ABL1 transcripts (Min)	Prior evidence of quantifiable BCR-ABL1 transcript.
Duration DMR (MR ⁴ or better) >2 years (Min)	MR ⁴ for ≥2 years (≥4 tests, performed ≥3 mo apart)
Access to high quality quantitative PCR using IS with rapid turn- around for results (Mand)	Access to a reliable qPCR test with sensitivity of at least MR4.5 IS and that provides results within 2 wks.
Patient's agreement to more frequent monitoring after stopping. Monthly for the 1 st 6 mo, every 2 mo for mo 6-12, and every 3 mo thereafter. (Mand)	
Motivated patient with structured communication (Mand)	Age ≥18 years
1 st -line therapy or 2 nd -line if intolerance was the only reason for changing TKI (Min)	Prompt resumption of TKI within 4 wks of loss of MMR with monthly monitoring until MMR. If fail to achieve MMR after 3 mo of resumption, mutation testing continue monthly molecular monitoring for another 6 mo.
No prior treatment failure (Min)	

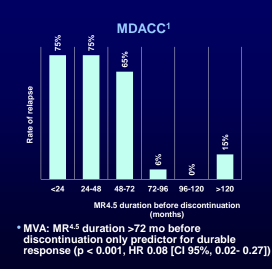
Hochhaus et al. Leukemia 2020; 34: 966-984; NCCN Version 3.2021 – January 13, 2021 (https://www.nccn.org/professionals/physician_gls/pdf/cml.pdf).





47

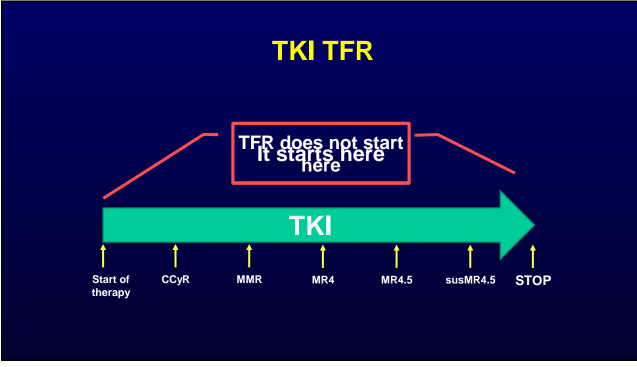
How to Improve TFR Success? - Wait Longer



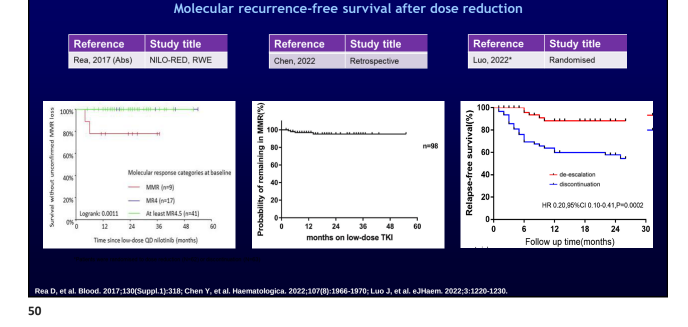
	EURO-SKI ²	
Years DMR	n/N	% (95% CI) without relapse
1–2	13/28	46% (28–66)
>2 to ≤3	17/31	55% (36–73)
>3 to ≤4	22/37	59% (42–75)
>4 to ≤5	11/17	65% (38–86)
>5 to ≤6	14/20	70% (46–88)
>6 to ≤7	9/15	60% (32–84)
>7	15/23	65% (43–84)

Yearly increase of \sim 3% in the probability of staying in MMR at 6 months over the observed range of DMR durations.

Chamoun et al. J Hematol Oncol 2019; 12: 1-10; Saussele et al. Lancet Oncology 2018; 19: 747-57.



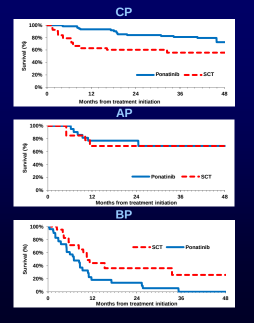
Dose Reduction after MMR/DMR achievement



Ponatinib or SCT for T315I CML

- Pts ≥18 yrs with CML T315I in any stage enrolled in PACE (n=449) or EBMT (1999-2010; n=222)
- Median age (yr): CP 53 vs 48; AP 55 vs 46; BP 47 vs 44; Ph+ ALL 55 vs 36

Disease group –	Median survival (mo)		
	PACE	EBMT	
СР	NR	103	
AP	NR	56	
BP	7	11	
Ph+ ALL	7	32	



Nicolini et al et al. Cancer 2017; 123: 2875-80.

51

Summary - CML 2024

- Excellent therapy available
- CCyR: gold standard for response (improves OS)
 - Deeper molecular responses: improve EFS (MMR) and option of treatment discontinuation (MR4.5)
- Early response (3-6 mos) predictive
 - Benefit of early switch uncertain
- Adequate management and monitoring mandatory for optimal outcome
- Change of therapy indicated for failure (not warning)
- Advanced phase:
 - AP: TKI as good as SCT
 - BP: TKI (± chemo, depending on goals) + SCT in CHR

Questions?

jorge.cortes@augusta.edu 706-721-0570

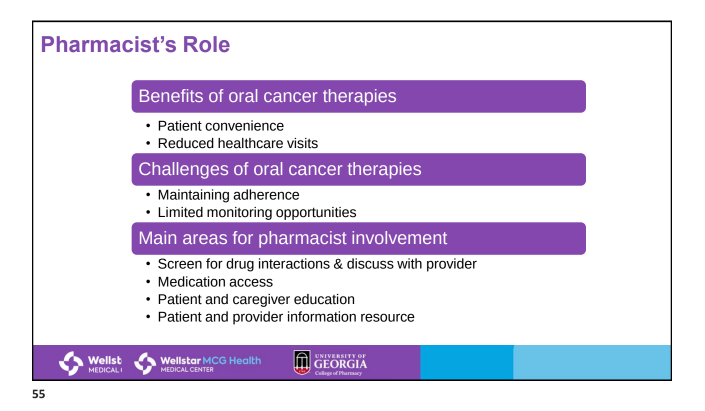


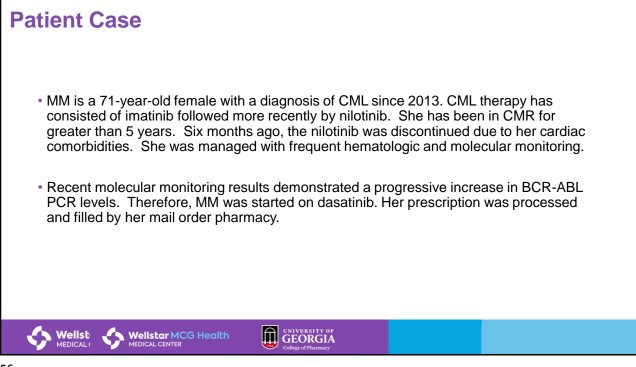
Pharmacist Role in Managing Patients with Chronic Myeloid Leukemia

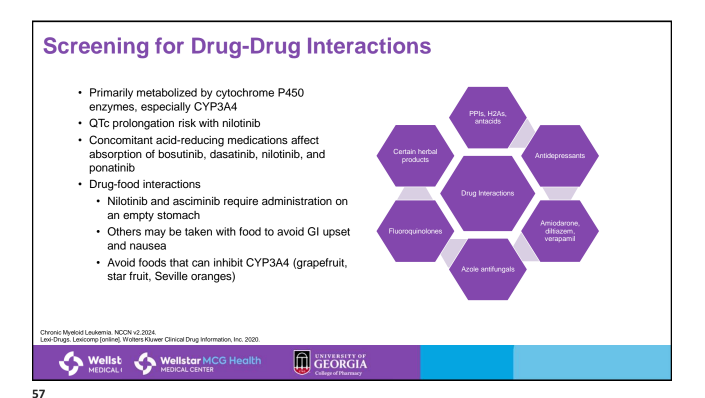
Amber B. Clemmons, PharmD, BCOP, FHOPA Heme/BMT Clinical Pharmacy Specialist Wellstar MCG Health The University of Georgia College of Pharmacy Augusta, GA

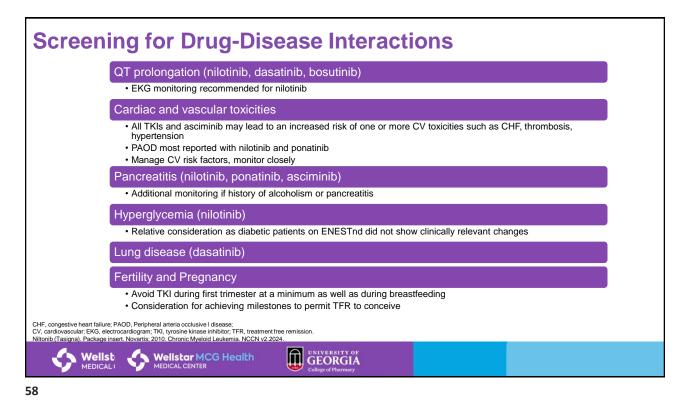


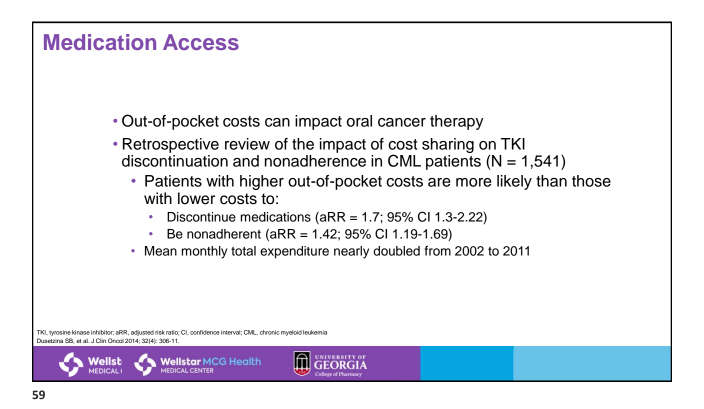
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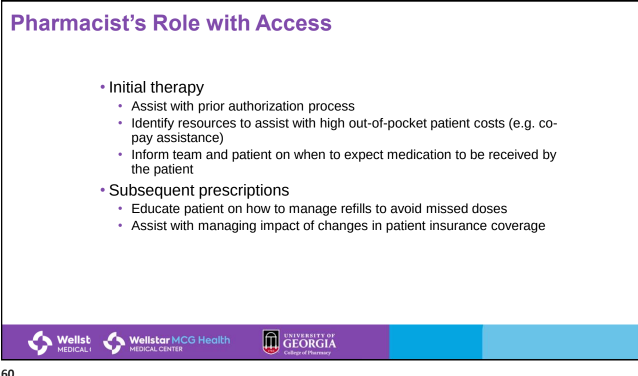


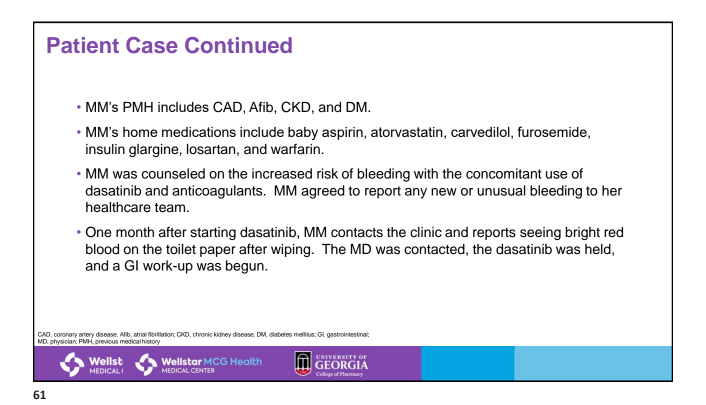


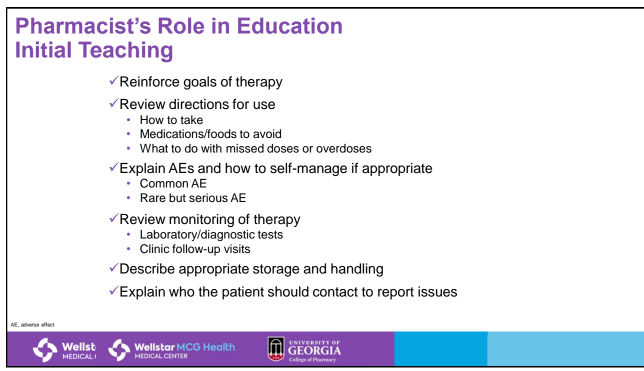












Pharmacist's Role in Education Follow-up

- Best practice phone follow-up shortly after the patient receives the first prescription, regularly for a time after initiating therapy, then periodically thereafter depending on need
- Reinforce

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- Goals of therapy
- Directions for use
- Ask open-ended questions regarding missed doses and barriers to taking the oral therapy

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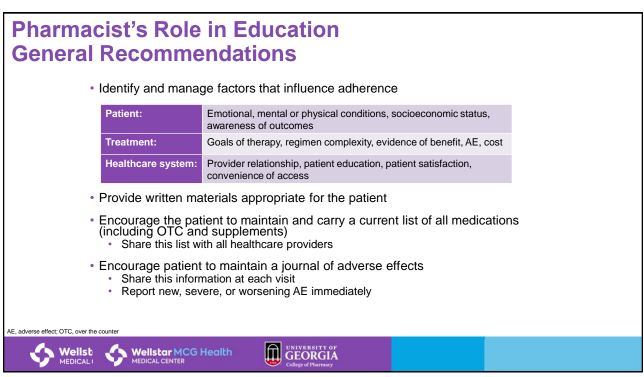
Review the AE profile and patient reported AEs

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· Ask about any changes to other medications and medical conditions

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Medication Adherence
 Adherence: extent to which patients comply with prescribed therapy Barriers: access, toxicity / tolerability, health literacy, etc.
 Adherence impacts outcomes Chronic phase CML patients in CCR on imatinib for at least 2 years had adherence electronically monitored during a 3-month period (N = 87) Adherence rate (≤90% vs >90%) was the <i>only</i> independent predictor of CMR on the multivariate analysis (RR = 19.35; p = 0.004)
 Pharmacist-managed oral anticancer therapy program can improve outcomes Pharmacist-managed oral anticancer therapies in CML patients (N = 56) Higher adherence rate than usual care (88.6% vs. 65.8%, p =0.0046)
CCR, complete cytogenetic response; RR, relative risk; PFS, progression-free survival; NR, not reached; CMR, complete molecular response; CML, chronic myeloid leukemia Marin D, et al. J Clin Oncol 2010; 28(14): 2381-8. Lam MSH, forburg N. J Oncol Pharm Practice 2016; 22(6): 741-8. Chronic Myeloid Leukemia. NCCN v2:2024.
Wellster MCG Health MEDICAL CENTER WEISCAL CENTER UNIVERSITY OF GEORGIA College of Pharmacy

Patient Case Continued
 MM's work-up revealed a lower GI bleed. Her INR was therapeutic at the time of the bleeding event. Given the concern of increased bleeding risk with resuming dasatinib and previous concerns of nilotinib affecting MM's cardiac comorbidities, it was decided to switch MM's CML therapy to bosutinib.
 MM was educated on bosutinib therapy, received her medication from her mail order pharmacy and continues on bosutinib with no issues to date.
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Pharmacist's Role as an Information Resource

Patients and Caregivers

- Follow-up phone calls
- Adherence aids
 - Diaries, pillboxes, electronic reminders
- Financial assistance resources
- Local and national support groups

Healthcare Providers

- Assess and manage adherence barriers
- Medication access
- Drug-drug and drug-disease interaction screening
- AE management recommendations

67

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Selected Resources

Medication Access

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- · Manufacturer's patient assistance programs
- Leukemia & Lymphoma Society co-pay
- co-pay assistance (www.lls.org/copay)
- NeedyMeds (<u>www.needymeds.org</u>)
- Disease Information for Patients
 - The Leukemia & Lymphoma Society (<u>www.lls.org</u>)
 - American Cancer Society (<u>www.cancer.org</u>)
 - ASCO (www.cancer.net)
- Disease Information for Healthcare Providers

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NCCN (<u>www.nccn.org</u>)

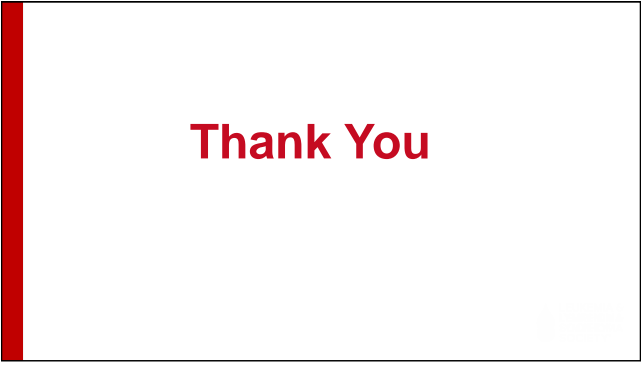
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- Standards for Safe Administration & Management of Oral Cancer Therapies
 - ASCO/ONS (http://ascopubs.org/doi/pdf/10.1200/JOP.2016.017905)
- Patient Education Tools for Oral Cancer Therapies
 - MASCC (https://mascc.org/resources/assessment-tools/mascc-oral-agent-teaching-tool-moatt/)

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ONS Oral Anticancer Medication Toolkit (<u>https://www.ons.org/clinical-practice-resources/oral-adherence-toolkit</u>)





CML: TKI Side Effects and Management

Sarah Jimenez DNP, AGACNP-BC, AOCNP Blood and Marrow Transplantation/Immune Effector Cell Therapy Program Wellstar MCG Health/Georgia Cancer Center Augusta, GA



The What, Where, and When of TKI's

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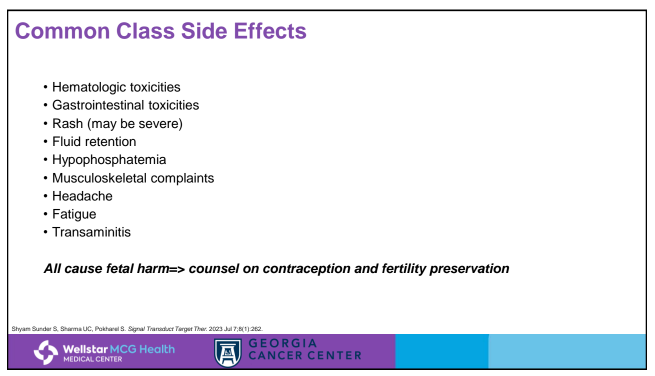
Drug Name	Place In Therapy	Dosing Schedule
Imatinib mesylate (Gleevec®)	First line	Daily or twice daily dosing
Dasatinib (Sprycel®)	First line or subsequent	Daily dosing
Bosutinib (Bosulif®)	First line or subsequent	Daily dosing
Nilotinib (Tasigna®)	First line or subsequent	Twice daily dosing
Ponatinib (Iclusig®)	Subsequent and/or T315i mutation	Daily dosing
Asciminib (Scemblix®)	Subsequent and/or T315i mutation	Daily or twice daily dosing

Imatinib package insert, 3/2022; Dasatinib package insert, 2023; Bosutinib package insert, 2023; Nilotinib package insert, 2021; Ponatinib package insert, 2024; Asciminib package insert, 2023

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Agent Specific Side Effect					
• Imatinib: fluid retention/edema, hepatotoxicity, congestive heart failure, renal impairment, hypothyroidism					
• Dasatinib: pleural effusion, QT prolongation, pulmonary arterial hypertension, cardiac dysfunction, bleeding					
 Nilotinib: hyperglycemia, elevated amylase/lipase, dyslipidemia, QT prolongation/sudden death, pancreatitis, hepatotoxicity, pleural effusion, arterial thrombotic events 					
• Bosutinib: diarrhea, hepatotoxicity, pleural effusion, pancreatitis, hypersensitivity					
 Ponatinib: hypertension, elevated amylase/lipase, pancreatitis, arterial thrombotic events, events, hepatotoxicity, cardiac arrhythmias, congestive heart failure, bleeding 	venous thrombotic				
 Asciminib: pancreatitis, hypertension, hypersensitivity, cardiovascular toxicity (ischemic car thrombotic, and embolic), heart failure, prolonged QT 	diac, CNS, arterial				
Institute and reacting a 19000. Description and a 1000. Description and and a 2000.	Italicized = rare/serious adverse effect				
Imatinib package insert, 3/2022; Dasatinib package insert, 2023; Bosutinib package insert, 2023; Nilotinib package insert, 2021; Ponatinib package insert, 2024; Asciminib package insert, 2023	Bold = Prescribing Information Boxed Warning				
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Edema/Fluid Retention

Drug	Manifestation	Diagnostic Testing	Supportive Care		
Imatinib Dasatinib Bosutinib Nilotinib Ponatinib	 Periorbital edema Pleural effusions Pericardial effusions Pulmonary edema Peripheral edema 	Monitor weight Chest X-ray ECHO	Use of diuretics as needed Low Sodium diet Hold/dose adjustment/discontinuation Thoracentesis Oxygen if needed Cold compress to eyes Topical Hydrocortisone cream Use of corticosteroids		
Imatinib Nilotinib	Ascites	Abd ultrasound Monitor weight	Use of diuretics as needed Low Sodium diet Hold/dose adjustment/discontinue Paracentesis		
	natinib package insert, 3/2022; Dasatinib package insert, 2023; Bosutinib package insert, 2023; ilotinib package insert, 2021; Ponatinib package insert, 2024; NCCN, 2024				
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75

Myelosuppression Drug **Manifestation Diagnostic Testing Supportive Care** ALL TKIs Anemia Monitor CBC regularly Hold/dose adjustment • ٠ Neutropenia Check iron and nutritional Growth factor support Thrombocytopenia labs Blood product transfusion support • Hemorrhage Correct any nutritional deficiencies **Bleeding events Review medications** • - Concomitant use of antiplatelet or anticoagulants Imatinib package insert, 3/2022; Dasatinib package insert, 2023; Bosutinib package insert, 2023; Nilotinib package insert, 2021; Ponatinib package insert, 2024; Asciminib package insert, 2023 NCCN, 2024 GEORGIA CANCER CENTER

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Drug	Manifestation	Diagnostic Testing	Supportive Care
Imatinib Bosutinib Ponatinib Asciminib	 Congestive Heart Failure Left Ventricular Dysfunction 	ECHO ECG Monitor Electrolytes Monitor heart rate Check Pro-BNP	Treat CV event per standard of care Correct electrolytes abnormalities Referral to Cardiology or Cardio- Oncology Review medications
Imatinib Dasatinib Bosutinib Ponatinib Asciminib	Prolonged QTArrhythmias		Hold/adjust dose/Discontinue

Cardiovascular Toxicities

Drug	Manifestation	Diagnostic Testing	Supportive Care
Nilotinib **Assess Cardiac risk prior to start	Prolonged QT/Sudden death	Monitor Electrolytes (K+ and Mg) -prior to start then periodically -correct deficiencies before starting Monitor ECG -Baseline, 7 days after start, then periodically and 7 days after dose adjustments	 DO NOT ADMINISTER TO PATIENTS WITH HYPOKALEMIA, HYPOMAGNESIA, OR LONG QT SYNDROME. Correct electrolytes Medication review-> avoid concomitant drugs known to prolong QT Consult Cardiology/Cardio- Oncology Hold/dose adjustment/discontinuation
RED= Black Box W	arning		
tinib package insert, 2021; NCCN, 2024		PGIA	
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Cardiovascular Toxicities

Drug	Manifestation	Diagnostic Testing	Supportive Care
Nilotinib	 Arterial vascular occlusive events Ischemia heart disease related events Peripheral arterial occlusive disease 	ECHO ECG Cardiac enzymes Duplex u/s	If confirmed = Discontinue treatment Treat any cardiac events per standard of care
Nilotinib Asciminib	Hyperlipidemia	Monitor lipid profile prior to start and periodically during first year then annually	May need to start lipid lowering agent - Review for drug-drug interactions
inib package insert, 2021; Asciminib pac	xkage insert, 2023 NCCN, 2024		
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79

Cardiovascular Toxicities

Manifestation	Diagnostic Testing	Supportive Care
Arterial Occlusive Events - MI, stroke, etc	ECHO ECG Cardiac enzymes Duplex u/s CT head Revascularization/heart cath	Per standard of care Hold/discontinue based on severity
Venous thromboembolic events (VTE's)	Monitor for evidence of VTE's	Per standards of care Hold/discontinue based on severity
arning		
	Arterial Occlusive Events - MI, stroke, etc Venous thromboembolic events (VTE's)	Arterial Occlusive Events ECHO - MI, stroke, etc ECG Cardiac enzymes Duplex u/s Duplex u/s CT head Revascularization/heart cath Venous thromboembolic Monitor for evidence of events (VTE's) VTE's

rug	Manifestation	Diagnostic Testing	Supportive Care
onatinib	Heart failure	Monitor for s/sx of heart failure ECHO Pro-BNP	Per standards of care Hold/discontinue for new or worsening heart failure
onatinib sciminib	Hypertension Hypertensive Crisis	Monitor blood pressure	Manage blood pressure with antihypertensives Hold/adjust dose/discontinue
D= Black Box	Warning		

Cardiovascular Toxicities Drug Manifestation **Diagnostic Testing Supportive Care** Dasatinib **Pulmonary Arterial** ECHO If PAH is confirmed = Hypertension (PAH) CT Chest discontinuation PFT's *Evaluate for s/sx of underlying cardiopulmonary disease prior to starting Dasatinib package insert, 2023; NCCN, 2024 GEORGIA CANCER CENTER Wellstar MCG Health MEDICAL CENTER A

Drug	Manifestation	Diagnostic Testing	Supportive Care
ALL TKI's	 Nausea Vomiting Diarrhea Abd pain 	Evaluate CBC and CMP Stool studies Abd imaging (i.e. ultrasound or CT) Check Amylase & Lipase	Antiemetics prior to dose and as needed Ginger hard candy Imodium [®] (loperamide)/Lomotil [®] (diphenoxylate and atropine) as needed for diarrhea Adequate Hydration Diet modification (BRAT diet) Avoid spicy, fatty foods, caffeine
Bosutinib Nilotinib Ponatinib Asciminib	Pancreatitis Elevated serum lipase	Evaluate CBC and CMP Check Amylase & Lipase Abd imaging (i.e. ultrasound or CT)	Per standard of care Hold medication until resolution/ decrease dose May require discontinuation

Gastr	ointe	stinal	Toxicities	S
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Drug	Manifestation	Diagnostic Testing	Supportive Care
Bosutinib Nilotinib Ponatinib	Constipation		Stool softeners/laxatives for constipation
Ponatinib	GI perforation	Evaluate CBC and CMP Abd imaging (i.e. ultrasound or CT)	GI Perforation= Discontinue
tinih nackane insert 2023: Nilo	tinib package insert, 2021; Ponatinib package insert,	2024: NGCN, 2024	

Dietary Considerations/Restrictions

Drug	Food?	Restrictions	Antacids?
Imatinib	Take with food and water	No Grapefruit juice	No restrictions
Dasatinib	Take with food and water	Avoid in lactose intolerant patients No Grapefruit juice	No Antacids 2 hours before or after administration. Avoid H2 Antagonists & PPI
Bosutinib	Take with food and water	No Grapefruit juice	No Antacids 2 hours before or after administration. Avoid H2 Antagonists & PPI
Nilotinib	Take on empty stomach - Avoid food 2 hours before and 1 hour after dose	Avoid in lactose intolerant patients No Grapefruit juice	No Antacids 2 hours before or after administration. Avoid H2 Blockers for 10 hours before and 2 hours after
Ponatinib	Take with or without food	Avoid in lactose intolerant patients No Grapefruit juice	No restrictions
Asciminib	Take on empty stomach - Avoid food 2 hours before and 1 hour after dose	No restrictions	No restrictions
nib package insert, 3/2022; Dasatinib pac tinib package insert, 2024; Asciminib pac	ckage insert, 2023; Bosutinib package insert, 2023; Nilo ckage insert, 2023 NCCN, 2024	tinib package insert, 2021;	
	Health GEO	RGIA CER CENTER	

85

Hepatotoxicity

Drug	Manifestation	Diagnostic Testing	Supportive Care	
Imatinib Dasatinib Bosutinib Nilotinib	Elevated liver functions Abd pain Jaundice	Check liver function testPrior to start of therapyAs clinically indicated (q2-3 months)	Hold/dose adjustment May require discontinuation Medication review	
Ponatinib	Liver Failure	Check liver function testBaseline, then monthly or as clinically indicated	Hold/discontinue based on severity	
RED= Black Box Warning				
matinib package insert, 3/2022; Dasat vilotinib package insert, 2021; Ponatin	inib package insert, 2023; Bosutinib package insert, 20 ib package insert, 2024; NCCN, 2024	023;		
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Drug	Manifestation	Diagnostic Testing	Supportive Care
matinib Bosutinib	Renal dysfunction	Evaluate renal function prior to start of therapy then as clinically indicated	Adequate hydration Dose adjustment
matinib Dasatinib Nilotinib Ponatinib	Electrolyte Abnormalities - Tumor Lysis Syndrome	Monitor electrolytes Monitor uric acid	Correct uric acid prior to start Adequate hydration Allopurinol prophylaxis Correct electrolyte abnormalities Dialysis
Ponatinib			Correct electrolyte abnormali

Dermatological Toxicity

Drug	Manifestation	Diagnostic Testing	Supportive Care
ALL TKIS	Rash - bullous dermatologic reactions (erythema multiforme and Stevens-Johnson Syndrome) Itching	Skin assessment Skin biopsy	Moisturizing skin cream Avoid sun exposure Hold/adjust dose/discontinue Antihistamines Corticosteroids Dermatology referral
Ponatinib	Impaired wound healing		Hold for 1 week prior to surgery and 2 weeks after surgery or until adequate wound healing
	. Dasatinib package insert, 2023; Bosutinib package insert, 2023; onatinib package insert, 2024, Asciminib package insert, 2023 N	CCN, 2024	
Nilotinib package insert, 2021; P	Ponatinib package insert, 2024; Asciminib package insert, 2023 N	CCN, 2024 ORGIA NCERCENTER	

Drug	Manifestation	Diagnostic Testing	Supportive Care
Ponatinib	Blurred vision Dry eyes	Comprehensive Eye Exam - Baseline then periodically	Lubricating eye drops for dry eyes

Endocrine Complications

Drug	Manifestation	Diagnostic Testing	Supportive Care
Nilotinib	Elevated Blood Glucose - Avoid in patient with diabetes	Monitor Blood Glucose - Prior to start of therapy then as clinically indicated	May need to start medications for glucose management (per standards of care
Imatinib	 Hypothyroidism Patients with history of thyroidectomy and on levothyroxine 	Monitor TSH	Adjust levothyroxine
		RGIA CER CENTER	

ug	Manifestation	Diagnostic Testing	Supportive Care
ALL TKI's	Fatigue No energy	Fatigue Assessment Score	Adequate Hydration Exercise Adjust time of administration

Musculoskeletal Complications

Drug	Manifestation	Diagnostic Testing	Supportive Care
matinib Dasatinib Ponatinib Asciminib	 Joint pain Musculoskeletal pain Myalgia Muscle cramps Arthralgias 	Monitor electrolytes (BMP, Mg, Phos)	Adequate Hydration Tonic water, tomato juice Potassium supplements Magnesium supplements Calcium supplements Muscle relaxers
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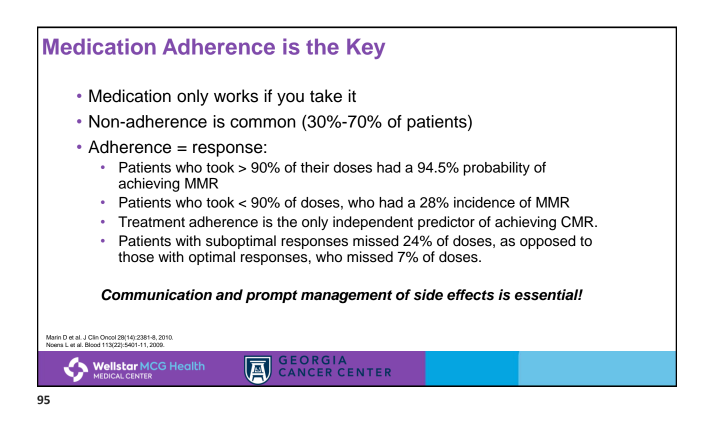


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Drug	Manifestation	Diagnostic Testing	Supportive Care		
Bosutinib Asciminib	Respiratory Tract Infections	Respiratory Viral Panel Chest imaging (Chest XR, CT)	Cough suppressants/expectorants Albuterol inhalers Antibiotics/antivirals if needed		
Bosutinib	Fevers	Evaluate for infections - Blood cultures - CBC - UA and urine culture	Acetaminophen or NSAIDS in moderation Antibiotics		
osutinib package insert, 2023;	Asciminib package insert, 2023 NCCN, 2024				
Bosutinib package insert, 2023; Asciminib package insert, 2023; NCCN, 2024					

g	Manifestation	Diagnostic Testing	Supportive Care
ALL TKI's	Headaches	Consider CT Head	Acetaminophen or NSAIDs in moderation
Ponatinib	Neuropathy	Monitor for symptoms of peripheral or cranial neuropathy	Hold/dose reduce/discontinue
Ponatinib	Reversible Posterior Leukoencephalopathy syndrome (RPLS)	MRI brain	Hold dose until resolution/discontinue

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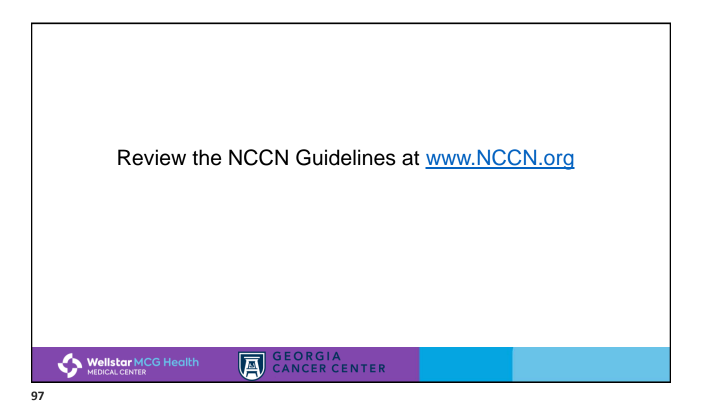


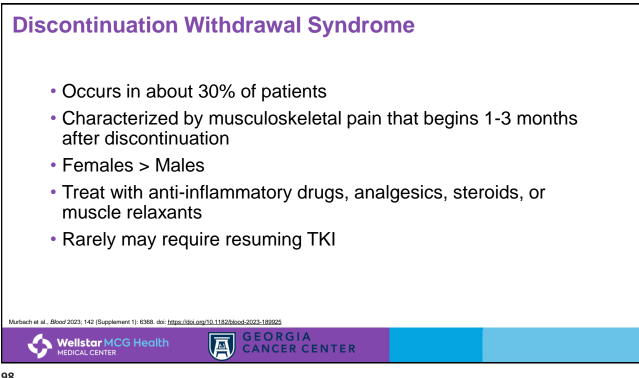
Discontinuation of TKI therapy should only be performed in consenting patients after a thorough discussion of the potential risks and benefits

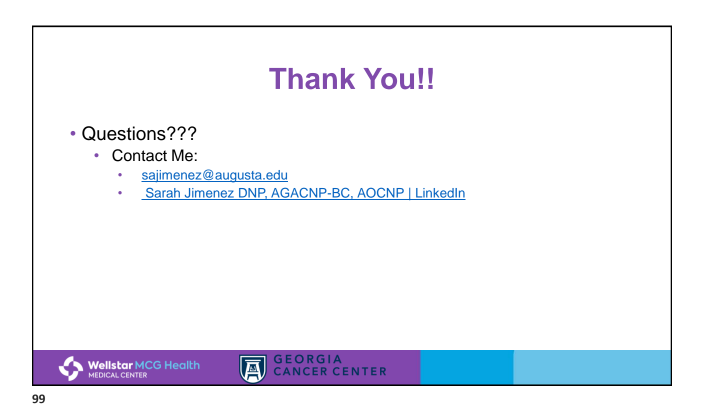
NCCN Clinical Practice Guidelines in Oncology (NCCN Guidelines®). Chronic Myeloid Leukemia Version 1.2025 — August 8, 2024. https://www.nccn.org/professionals/physician_gls/pdf/cml.pdf.



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۵	FREE LLS RESOURCES FOR PATIENTS	
	 Information Specialists – Personalized assistance for managing treatment decisions, side effects, and dealing with financial and psychosocial challenges (IRC). www.LLS.org/IRC Nutrition Education Services Center – one-on-one consultation with a registered dietician for patients/caregivers of all cancer types (NESC). www.LLS.org/Nutrition Clinical Trial Nurse Navigators – RNs and NPs provide a personalized service for patients seeking treatment in a clinical trial, sift through the information and provide information to bring back to their HC team (CTSC). www.LLS.org/CTSC Reach out Monday–Friday, 9 am to 9 pm ET Phone: (800) 955-4572 	
	 Live chat: <u>www.LLS.org/IRC</u> Email: infocenter@LLS.org HCP Patient Referral Form: <u>www.LLS.org/HCPreferral</u> 	LEUKEMIA & LYMPHOMA SOCIETY



