

# Dose Modification Dynamics of Ponatinib in Patients With Chronic-Phase Chronic Myeloid Leukemia (CP-CML) From the PACE and OPTIC Trials



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## Introduction

- Ponatinib is a potent, oral, third-generation tyrosine kinase inhibitor (TKI) that is US Food and Drug Administration (FDA) and European Medicines Agency approved for the treatment of patients with relapsed CML<sup>1,2</sup>
- In the Ponatinib Ph+ ALL and CML Evaluation (PACE) trial, patients with highly resistant CP-CML with substantial prior second-generation (2G) TKI treatment demonstrated deep, lasting responses to 45 mg once daily ponatinib<sup>2</sup>
  - PACE did not have a response-based dose-reduction strategy
- The phase 2 Optimizing Ponatinib Treatment In CP-CML (OPTIC) trial prospectively evaluated a response-based dose-reduction strategy in an attempt to optimize the dose schedule of ponatinib in patients with CP-CML whose disease was resistant to 2G BCR::ABL1 TKI therapy or with a T315I mutation<sup>3</sup>
  - OPTIC was designed with required dose reductions upon achievement of  $\leq 1\%$  BCR::ABL1<sup>3</sup>

## Methods

Trial	PACE	OPTIC
<b>Inclusion Criteria</b>	<ul style="list-style-type: none"> <li>≥18 years of age</li> <li>ECOG <math>\leq 2</math></li> <li>Normal QT interval corrected by Friderica calculation (QTcF) on screening ECG evaluation<sup>a</sup></li> </ul>	<ul style="list-style-type: none"> <li>≥18 years of age</li> <li>ECOG <math>\leq 2</math></li> <li>Normal QT interval corrected by Friderica calculation (QTcF) on screening ECG evaluation<sup>a</sup></li> </ul>
<b>Exclusion Criteria</b>	<ul style="list-style-type: none"> <li>Significant or active CV</li> <li>Uncontrolled hypertriglyceridemia<sup>b</sup></li> </ul>	<ul style="list-style-type: none"> <li>Clinically significant, uncontrolled, or active CV disease</li> <li>Uncontrolled hypertension<sup>c</sup></li> <li>Poorly controlled diabetes<sup>d</sup></li> </ul>

<sup>a</sup>Defined as QTcF of  $\leq 450$  ms in males or  $\leq 470$  ms in females. <sup>b</sup>Triglycerides  $>450$  mg/dL. <sup>c</sup>Defined as  $>150$  mm Hg and  $>90$  mm Hg for systolic and diastolic blood pressure, respectively. <sup>d</sup>Hemoglobin A1c values of  $>7.5\%$ .

- The propensity score analysis was conducted to reduce potential bias from differences in baseline characteristics when comparing the TE-AOE incidence rates in PACE and OPTIC
- The propensity score calculation was adjusted for 14 parameters including baseline characteristics, disease parameters, and exposure

## Results

Table 2. Demographics and Baseline Disease Characteristics

Characteristic	PACE CP-CML (N=270) <sup>a</sup>	OPTIC 45 mg → 15 mg (N=94)
Median age, y	60	46
Male gender, n (%)	144 (53)	50 (53)
Median time since diagnosis, y	7.03	5.5
Patients with CV risk factors, n (%)		
Hypercholesterolemia	65 (24)	3 (3.2)
Baseline BMI $\geq 30$ kg/m <sup>2</sup>	73 (27)	26 (28)
Diabetes mellitus	32 (12)	5 (5.3)
Vascular disorders	119 (44)	30 (32)
Arterial hypertension	102 (38)	29 (31)
Deep vein thrombosis	4 (1.5)	0
Concomitant medications, n (%)		
Statins	64 (24)	24 (26)
Acetylsalicylic acid	87 (32)	18 (19)
Best response to last prior TKI <sup>b</sup> , n (%)		
CHR or worse	171 (63)	66 (70)
Better than CHR	99 (37)	28 (30)
BCR::ABL1 mutation at baseline <sup>c,e</sup> , n (%)		
$\leq 1\%$ BCR::ABL1 <sup>3</sup>	6 (2.2)	3 (3.2)
$>1$ to $10\%$ BCR::ABL1 <sup>3</sup>	54 (20)	26 (28)
$>10\%$ BCR::ABL1 <sup>3</sup>	200 (74)	74 (79)
Mutations <sup>d</sup> , n (%)		
No mutation	138 (51)	51 (54)
T315I	64 (24)	25 (27)
Mutation other than T315I	68 (25)	26 (28)
$\geq 2$ mutations detected	29 (11)	10 (11)
Stopped prior therapy due to resistance, n (%)	260 (96)	92 (98)
Prior TKIs, n (%)		
1	18 (6.7)	1 (1.1)
$\geq 2$	252 (93)	93 (99)
$\geq 3$	162 (60)	50 (53)

<sup>a</sup>PACE: Includes 3 patients with CP-CML who were not T315I+ at study entry and not resistant to dasatinib or nilotinib. <sup>b</sup>OPTIC: Baseline response to last prior TKI is missing for 4 patients. <sup>c</sup>Sanger sequencing was used for mutation testing. <sup>d</sup>PACE: Four patients had an e1a2 variant. 4 had atypical transcripts, and 2 had missing data. <sup>e</sup>OPTIC: One patient in the 45-mg cohort does not have BCR::ABL1. <sup>f</sup>OPTIC: Two patients in the 45-mg cohort did not have any mutation testing performed at baseline.

## References

- Cortes JE, et al. *Engl J Med*. 2013;369:1783-96.
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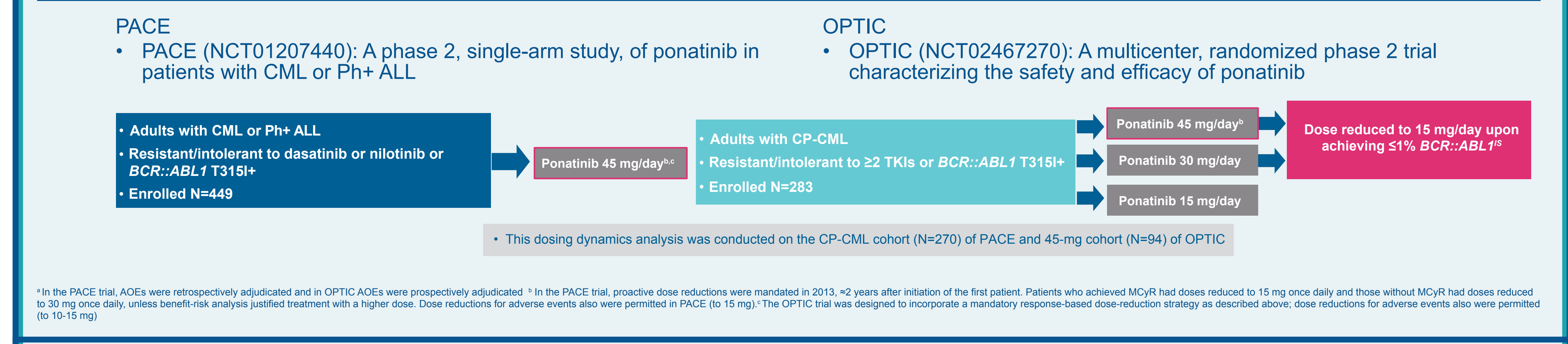
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## Objective

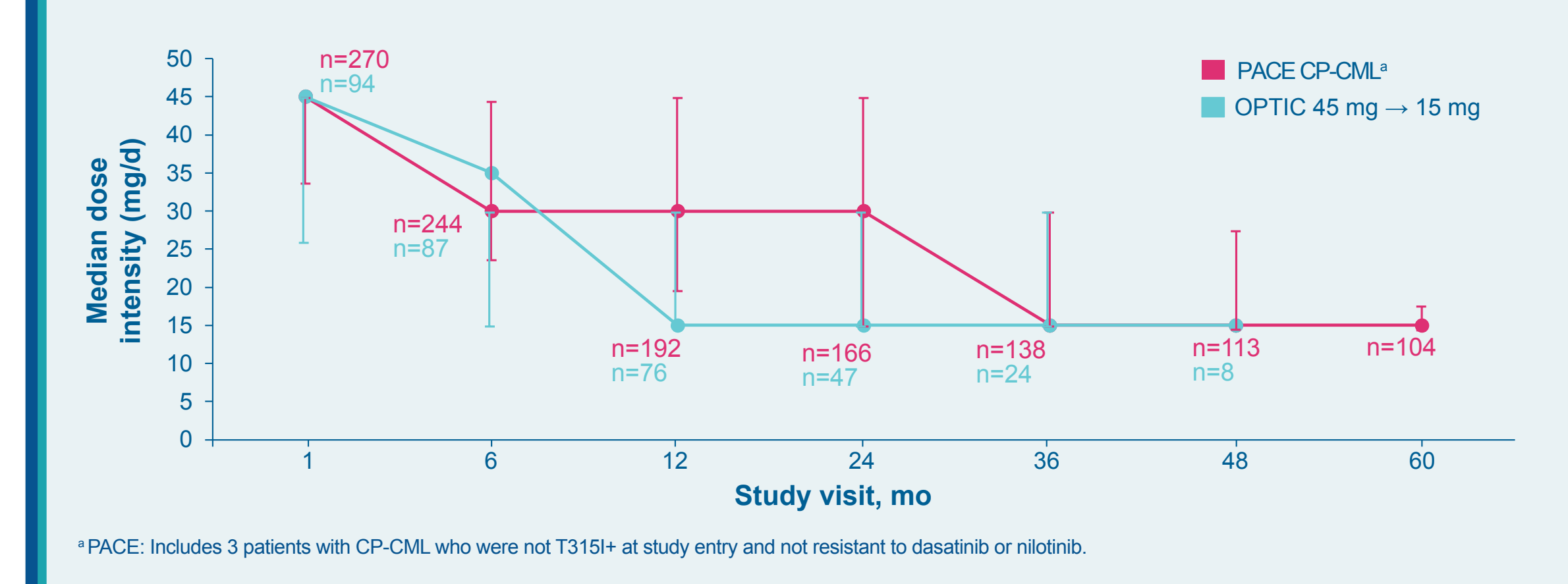
Here, we descriptively compared the potential impact of dosing differences between the 2 trials on efficacy and safety, as well as explored the impact of dosing differences on AOE

Figure 1. PACE and OPTIC Trial Designs<sup>a</sup>



## Results

Figure 2. Median Dose Intensity Over Time



<sup>a</sup>PACE: Includes 3 patients with CP-CML who were not T315I+ at study entry and not resistant to dasatinib or nilotinib.

## Key Takeaways

The response-based dose-reduction strategy in the OPTIC trial resulted in more rapid dose reductions, fewer dose reductions related to AEs, and longer median time on therapy in OPTIC compared with PACE, further demonstrating the benefit of the response-based dosing regimen used in OPTIC

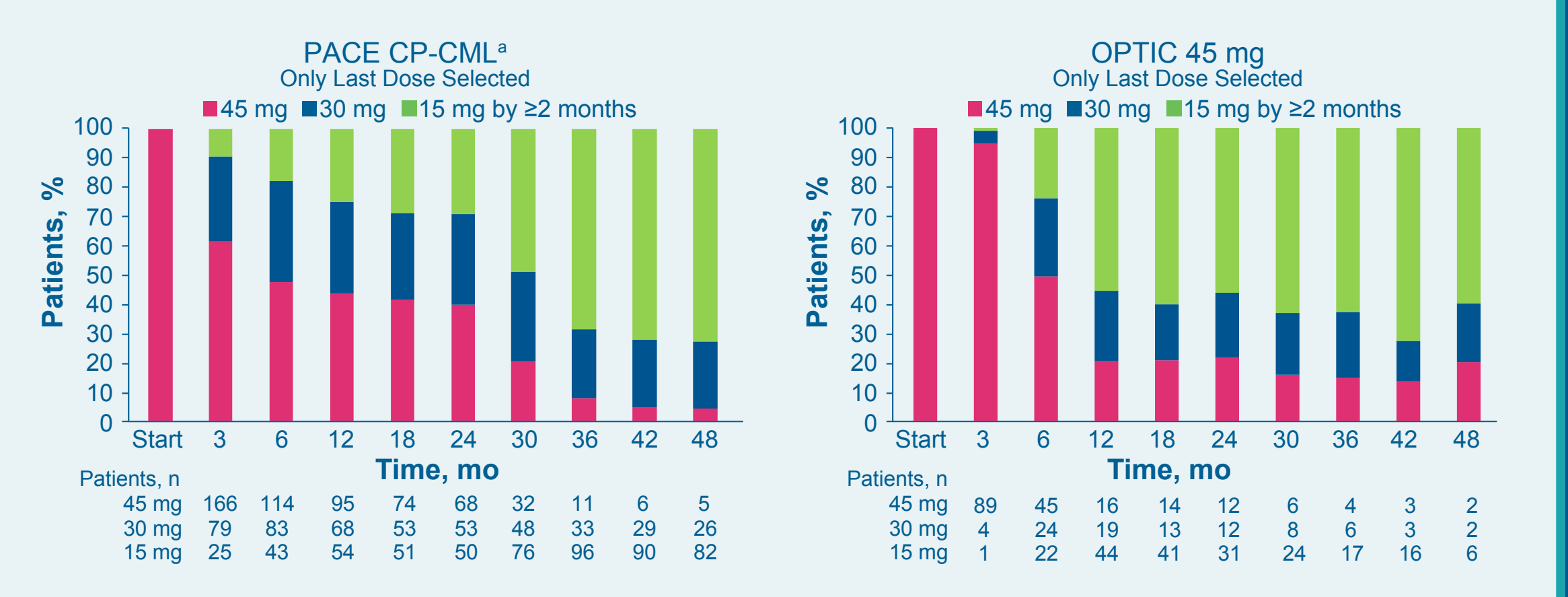
Table 3. Dose Reductions and Dose Intensity in PACE and OPTIC

	PACE CP-CML (N=270)	OPTIC 45 mg → 15 mg (N=94)
<b>Dose reduction</b>		
Dose reduction due to AEs, n (%)	175 (65)	42 (45)
Dose reductions per FDA mandate or per protocol upon response <sup>a</sup> , n (%)	46 (17)	33 (35)
No dose reductions, n (%)	49 (18)	19 (20)
Median time to dose reduction, mo	2.9	3.6
For safety	2.1	6.3
For efficacy	23.8	2.6

<sup>a</sup>Dose reductions in PACE were FDA mandated after 2013 for safety concerns; for OPTIC, dose reductions were due to efficacy according to study design

- Efficacy outcomes were generally comparable or better in OPTIC when compared with PACE (Figures 4–6, Table 4)

Figure 3. Change in Dose Over Time



<sup>a</sup>PACE: Includes 3 patients with CP-CML who were not T315I+ at study entry and not resistant to dasatinib or nilotinib.

Figure 4. Efficacy Outcomes in PACE and OPTIC by 12, 24, and 60 Months

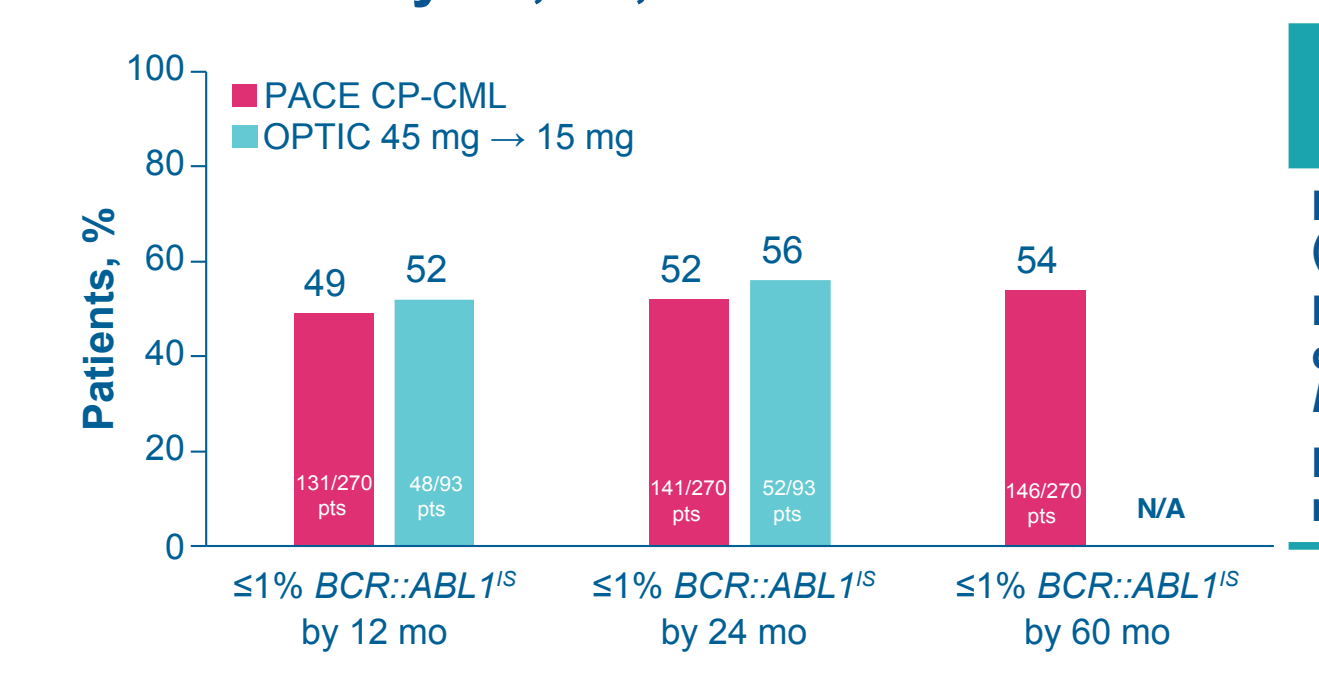


Table 4. Efficacy Outcomes in PACE and OPTIC

	PACE CP-CML (N=270)	OPTIC 45 mg → 15 mg (N=94)
Median time to response ( $\leq 1\%$ BCR::ABL1 <sup>3</sup> ), mo	5.6	6
Median duration of response ( $\leq 1\%$ BCR::ABL1 <sup>3</sup> ), mo	NR	NR
Median time on therapy, mo	12.6	19.5

- The preemptive dose adjustment in OPTIC led to longer time on therapy (19.5 mo vs 12.6 mo median time on therapy)

Figure 5. Progression-Free Survival<sup>a,b</sup>

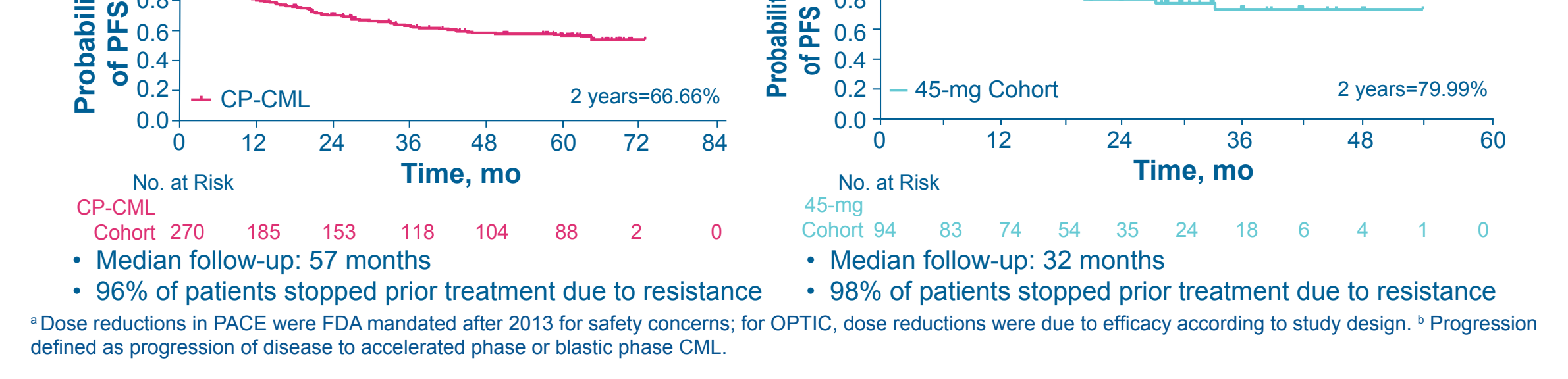
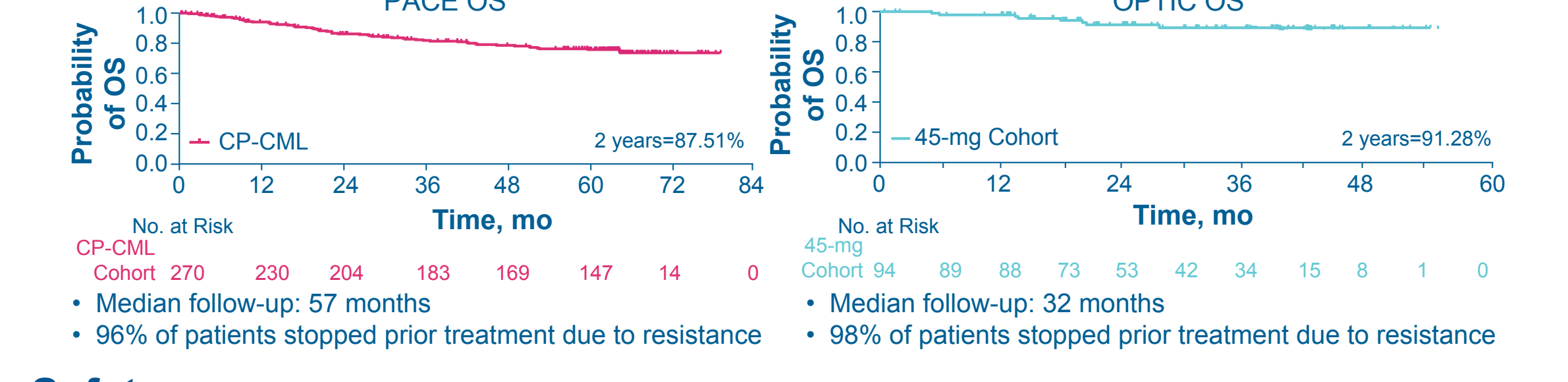


Figure 6. Overall Survival



## Safety

- The overall incidence of TEAEs was similar between the 2 trials, AOE were substantially lower in OPTIC when compared with PACE (Table 5, Figure 7)
- Propensity score analyses comparing AOE incidence showed a reduction in relative risk for AOE in OPTIC versus PACE (Table 6)

Table 5. Safety Summary for PACE and OPTIC

Safety Parameter	PACE CP-CML (N=270)	OPTIC 45 mg → 15 mg (N=94)
Any TEAE, n (%)	269 (100)	94 (100)
Grade 3/4	227 (84)	64 (68)

Figure 7. Exposure-Adjusted AOE in PACE and OPTIC

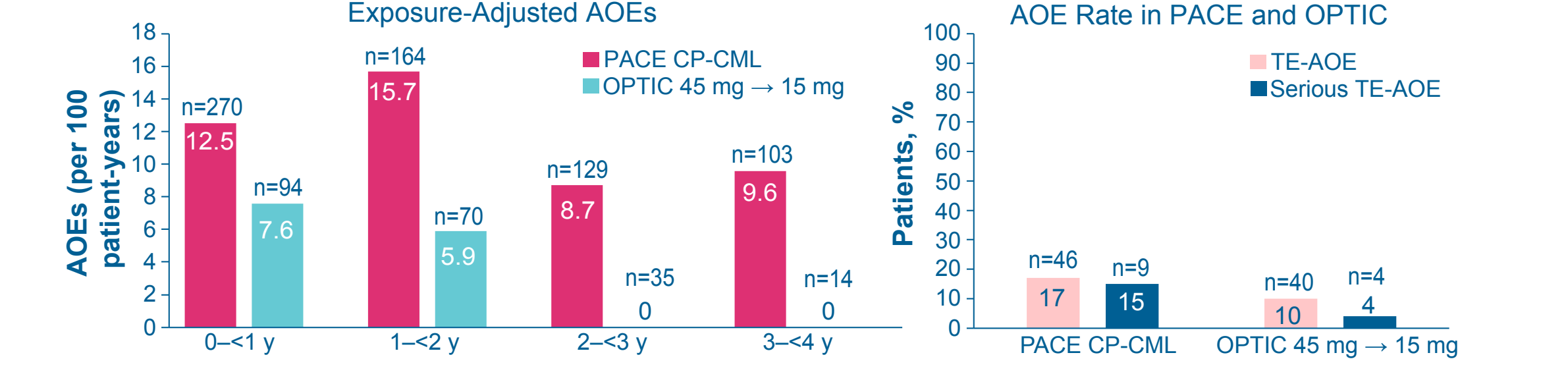


Table 6. Propensity Score Analysis Comparing AOE Incidence Between PACE and OPTIC

Safety Parameter	PACE CP-CML (N=270)	OPTIC 45 mg → 15 mg (N=94)
Patients with AOE	61	9
Unadjusted AOE rate (95% CI)	0.2230 (0.1733, 0.2728)	0.0879 (0.0297, 0.1461)
Adjusted AOE rate		
Odds ratio (95% CI)		0.3288 (0.1499, 0.7212)
Relative risk (95% CI)		0.4066 (0.2060, 0.8027)

- The propensity score analysis showed an overall risk reduction of 60% in OPTIC compared with PACE
- The parameters accounted for in the propensity score analysis were:
  - Baseline systolic blood pressure, time since diagnosis, baseline ECOG status, height at baseline, baseline weight, history of diabetes, mutation at baseline, geographic region, age, sex, race, ethnicity, total dose, total days on drug

## Conclusions

- The response-based dose-reduction strategy in the OPTIC trial resulted in more rapid dose reductions, fewer dose reductions related to AEs, and longer median time on therapy in OPTIC compared with PACE, further demonstrating the benefit of the response-based dosing regimen used in OPTIC
- These data from the PACE and OPTIC trials suggest that treatment with a response-based dose-reduction strategy may provide comparable or better efficacy while mitigating risk of AEs/AOEs with ponatinib
  - After adjusting for differences in baseline characteristics, OPTIC had lower AOE rates than PACE
- Furthermore, this analysis supports the rationale to explore response-based dose-modification strategies for other BCR::ABL1 TKIs