

FDA ADVISORY PANEL

Zilver[®] PTX[®] Drug-Eluting Peripheral Stent

Circulatory Systems
Device Panel Meeting



Introduction

Aaron Lottes, PhD

Lead Scientist for Zilver PTX

Director Regulatory Science
PAD Therapies

Cook Medical

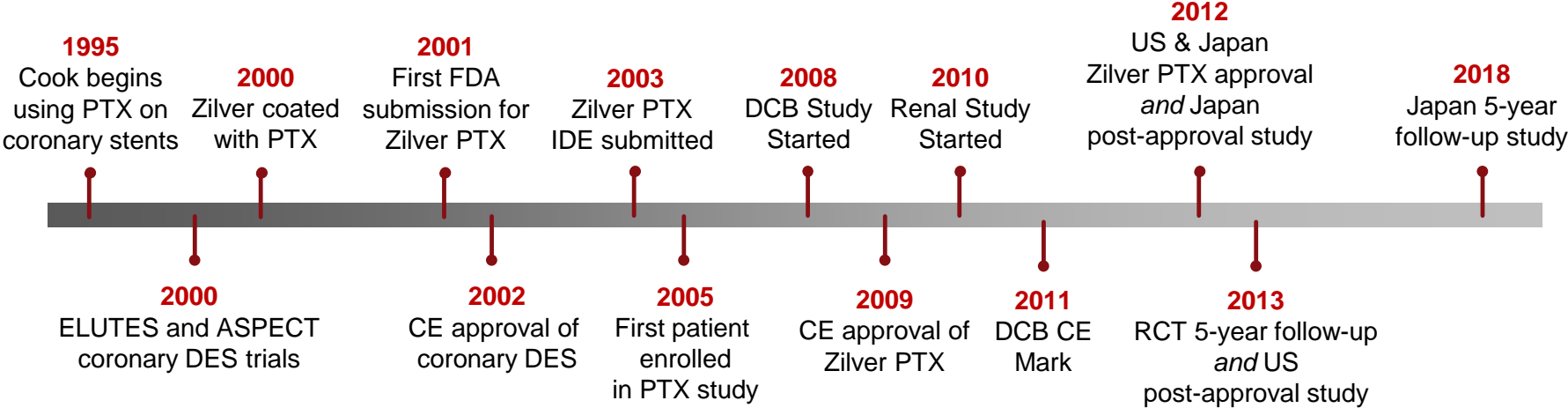
Michael Dake, MD

Global Principal Investigator for Zilver PTX

Senior Vice President of Health Sciences,
Professor of Medical Imaging, Medicine,
and Surgery, University of Arizona,
Tucson/Phoenix

Paid consultant of Cook Medical

Cook Medical's 25 Year History with Paclitaxel



▶ **No mortality signal in 25 years, across multiple studies and devices**



Overview

▶ Purest Data on Paclitaxel

Because other paclitaxel devices were not yet approved, the Zilver PTX RCT and Japan PMS provide the best data available to look at paclitaxel treatment

▶ Actual Treatment

Any analysis that does not consider known paclitaxel treatment is inappropriate for analyzing mortality and simply does not make sense

▶ Patient Impact

There is no mortality signal with Zilver PTX and the current situation is limiting patient access to the proven benefits of paclitaxel devices

ZILVER PTX

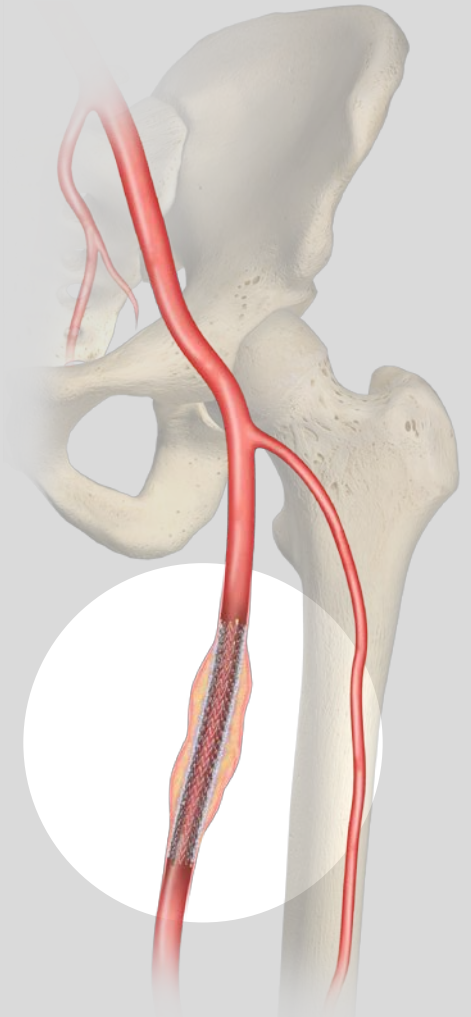
Patient Benefit

Durable results through 5 years

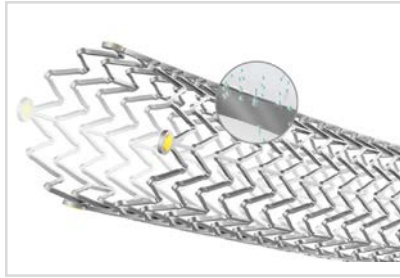
- Greater than 40% reduction in restenosis
- Greater than 40% reduction in reinterventions
- Proven clinical benefit in real-world patients



Dake MD, et al. *Circulation*. 2016;133:1472-1483
Yokoi H, et al. *J Am Coll Cardiol Interv*. 2016;9:271-277

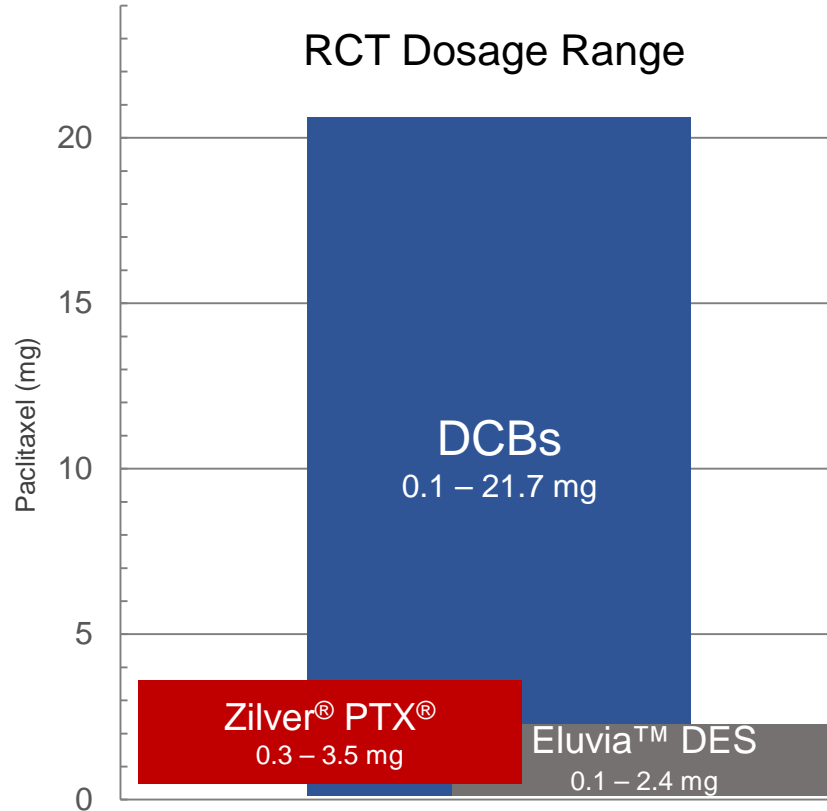


Device Overview

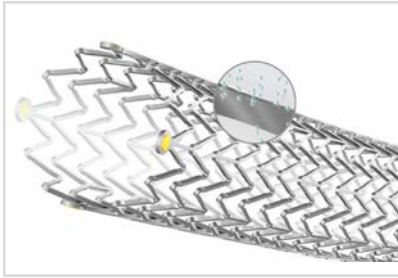


► Coating

Low dose, amorphous coating with no polymer or excipient

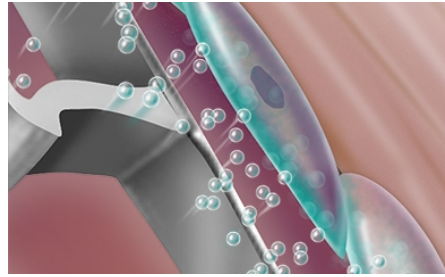


Device Overview



Coating

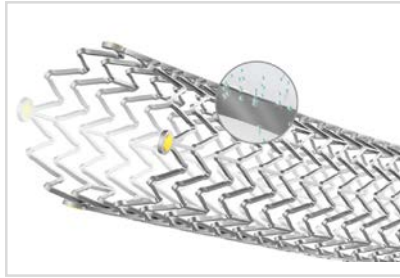
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▶ Local Drug Delivery

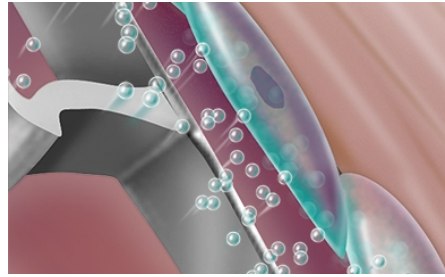
Short-term drug delivery, no long-term paclitaxel exposure, only BMS remains

Device Overview



Coating

Low dose, amorphous coating with no polymer or excipient



Local Drug Delivery

Short-term drug delivery, no long-term paclitaxel exposure, only BMS remains



▶ Long-term data

Only peripheral DES with long-term safety data

Zilver PTX Clinical Program

Study	Device	Follow-up	# of Patients
RCT	Zilver PTX	5 years	336
	PTA/BMS		143
Japan PMS	Zilver PTX	5 years	904
	BMS	3 years	190
EU BMS	BMS	5 years	110
US PAS	Zilver PTX	5 years ¹	200
Single-arm Study	Zilver PTX	2 years	787
French Reimbursement	Zilver PTX	2 years	119
China	Zilver PTX	1 year	178
REAL PTX	Zilver PTX	3 years	75
	DCB ²	3 years	75

- ▶ >1,000 patients to support US approval
- ▶ >2,500 patients in global pre- and post-market studies
- ▶ >300,000 stents to treat patients globally

¹ Ongoing ² 77.3% INPact, 21.3% Lutonix, 1.4% Other.

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- ▶ Large studies
- ▶ Long-term follow-up
- ▶ Concurrent comparator groups

¹ Ongoing ² 77.3% INPact, 21.3% Lutonix, 1.4% Other.

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- ▶ No exclusion criteria
- ▶ All treated patients enrolled
- ▶ Pure treatment comparison

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Zilver PTX Clinical Program

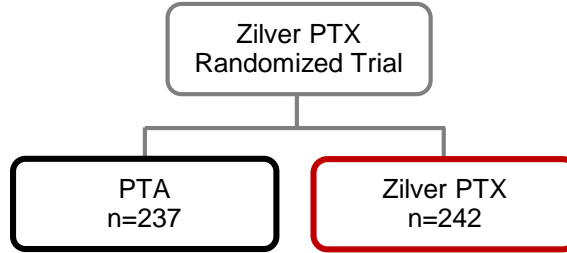
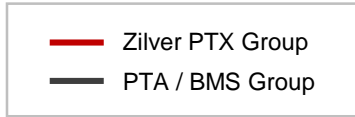
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- ▶ Trial designed with multidisciplinary physician input and approval from FDA, PMDA, and BfArM

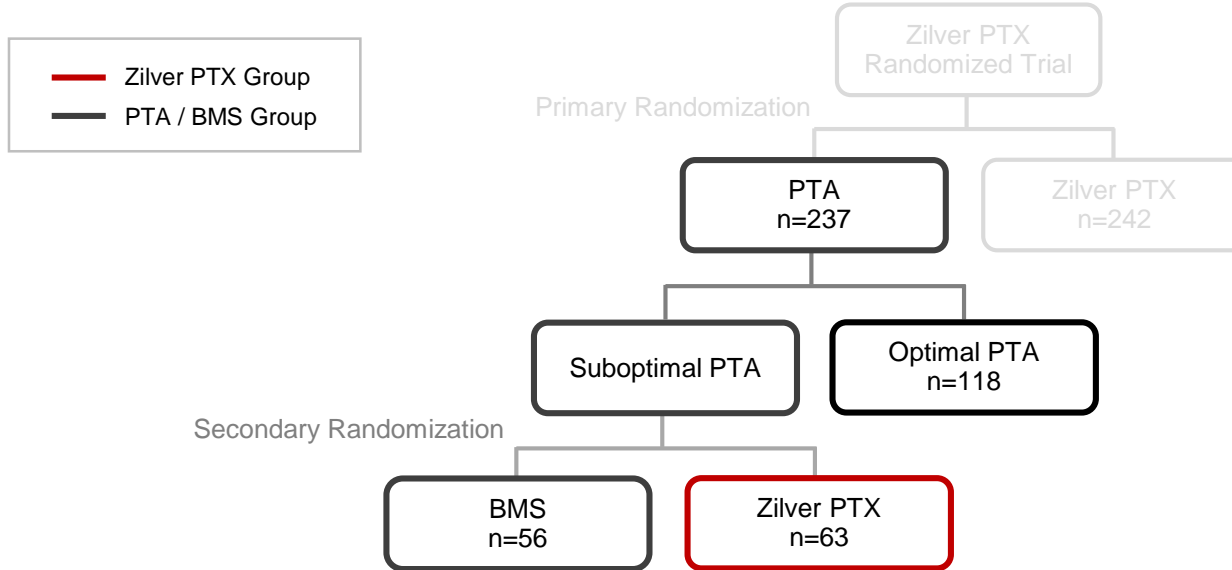
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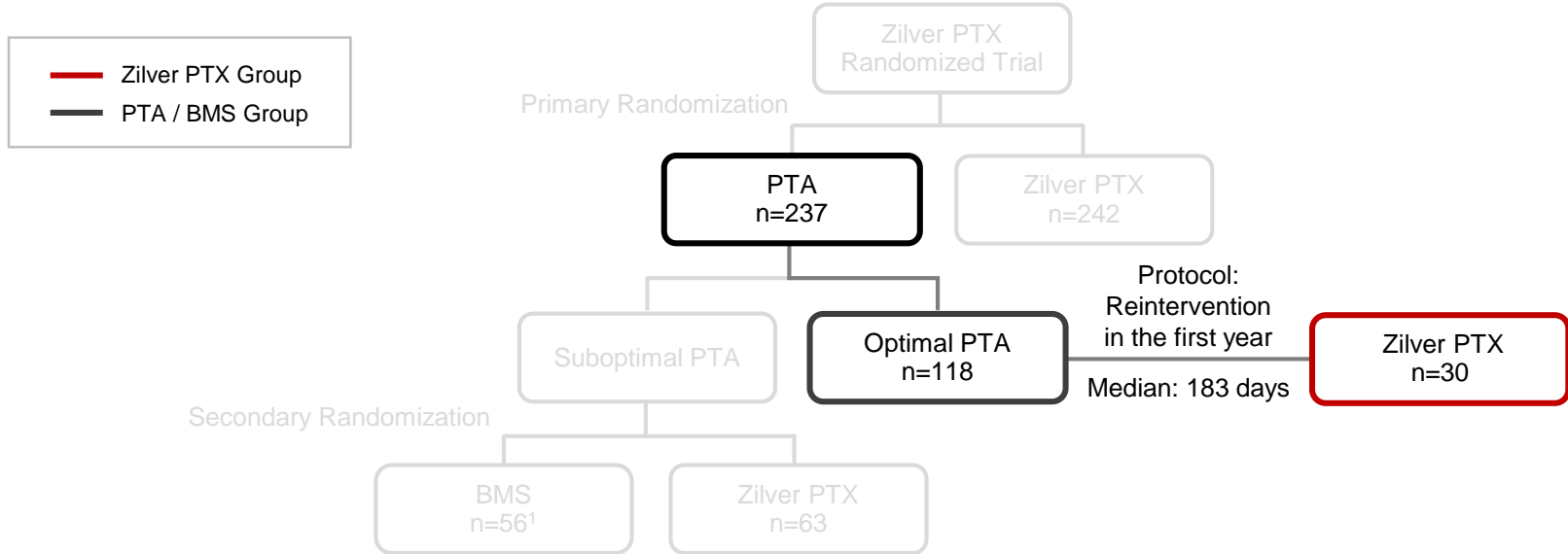
Primary Randomization



Secondary Randomization

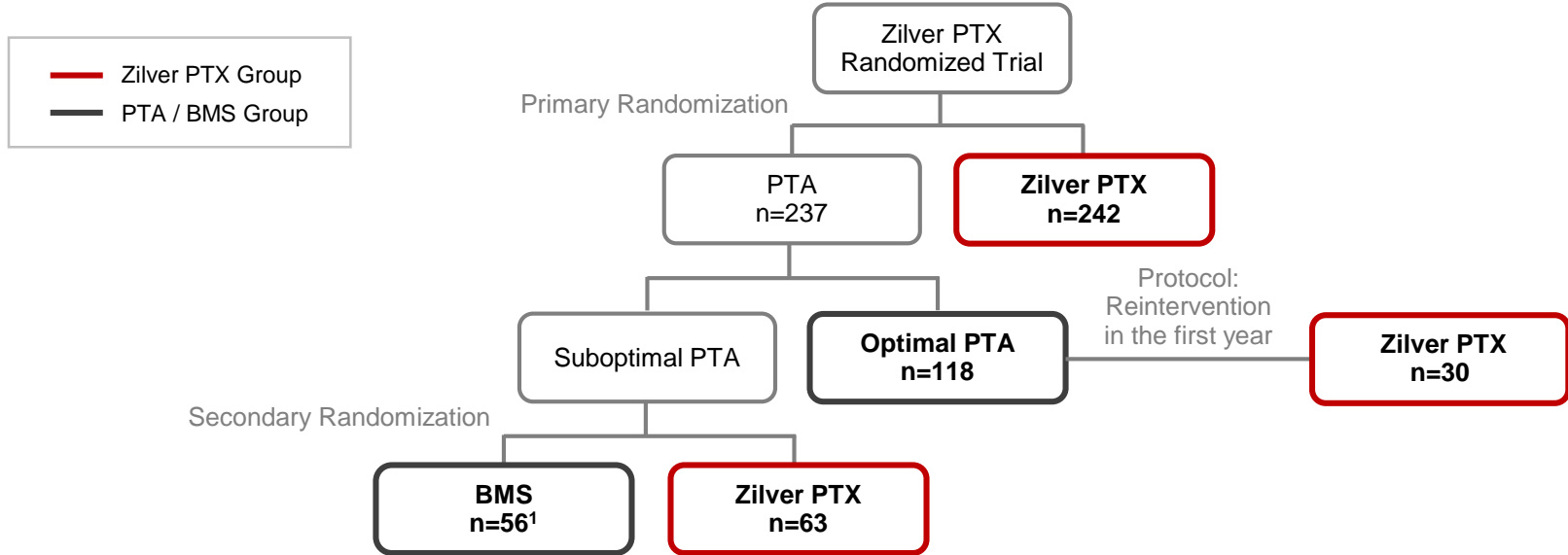


Early Crossover



¹ One BMS patient received Zilver PTX during reintervention within the first year.

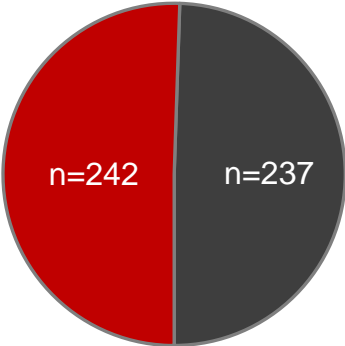
Actual Treatment



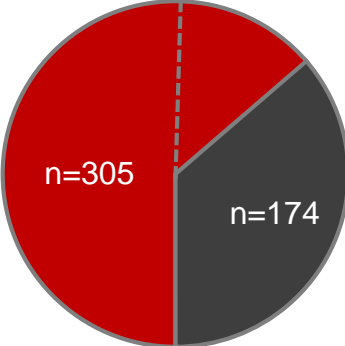
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Treatment Results

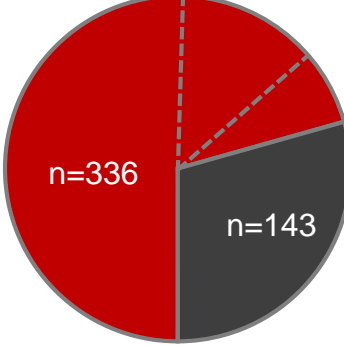
Primary Randomization



Primary + Secondary Randomization

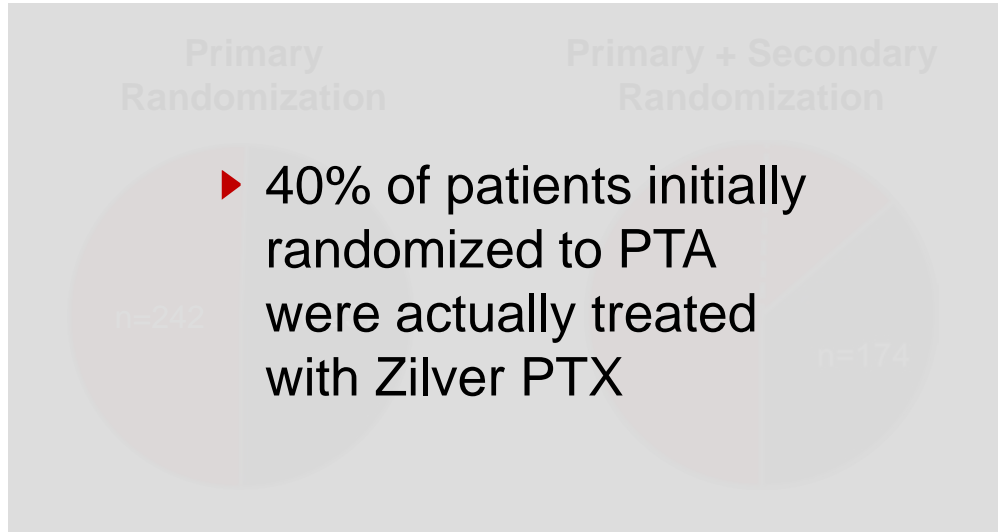


Actual Treatment = Primary + Secondary + Crossover

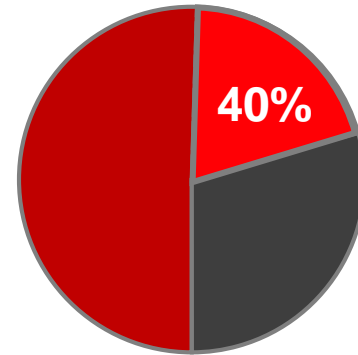


Treatment Results

- Zilver PTX
- PTA / BMS



**Actual Treatment =
Primary + Secondary + Crossover**

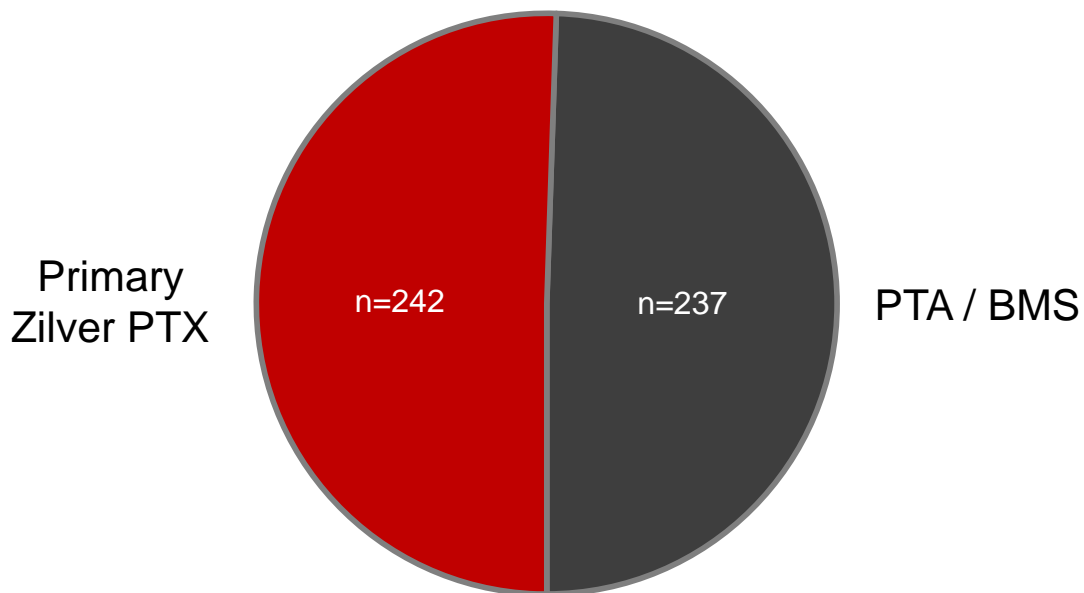


Results

Michael Dake, MD
Global Principal Investigator for Zilver PTX

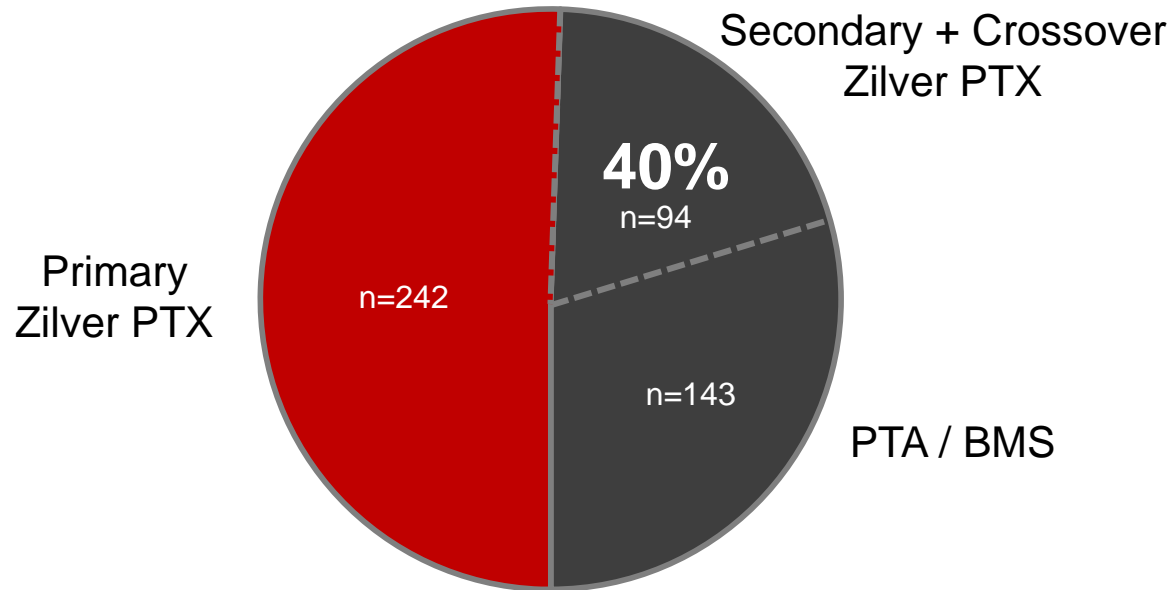


Primary Randomization

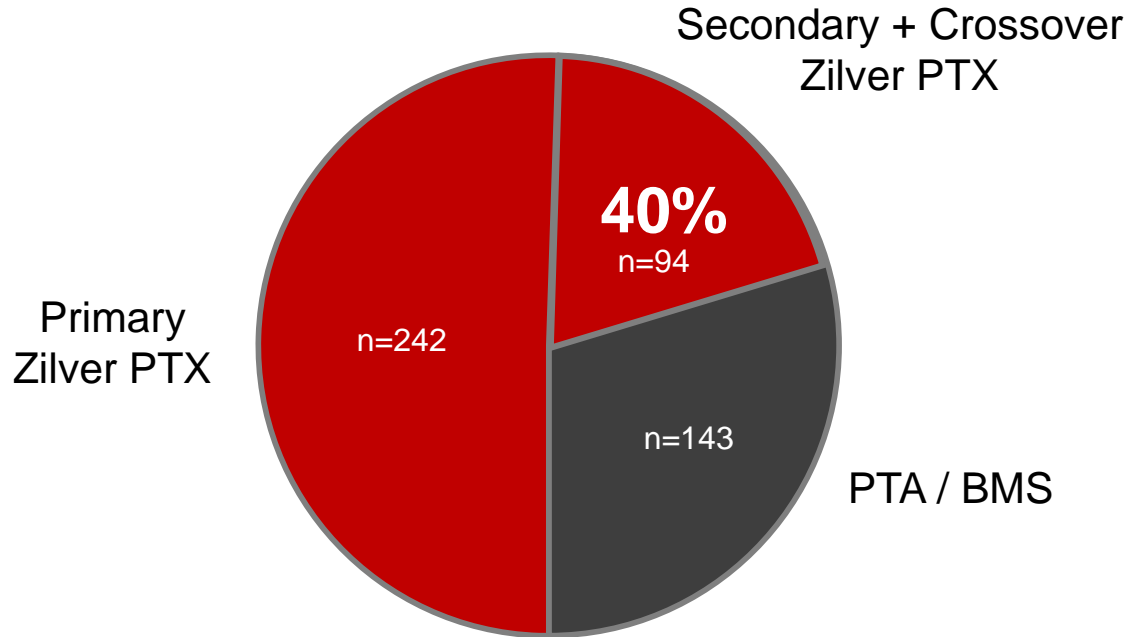


- ▶ Intent to treat is considered the standard for effectiveness
- ▶ Based on international standards, to evaluate safety we must analyze how patients were treated¹, not how they were randomized

Zilver PTX Patients in PTA/BMS Group

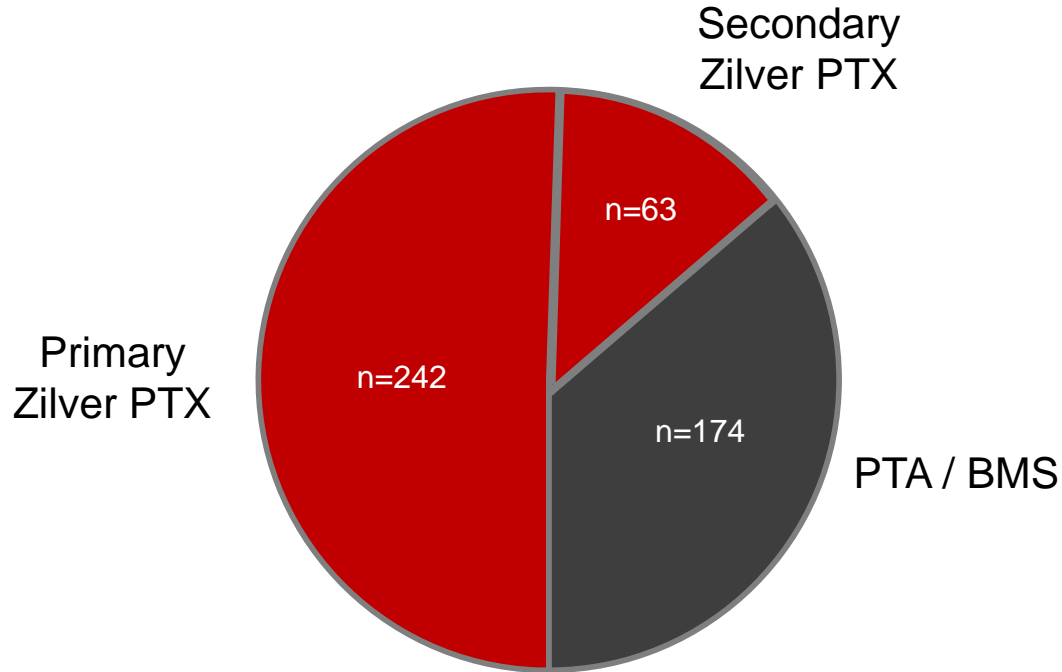


Zilver PTX Results Attributed to PTA/BMS

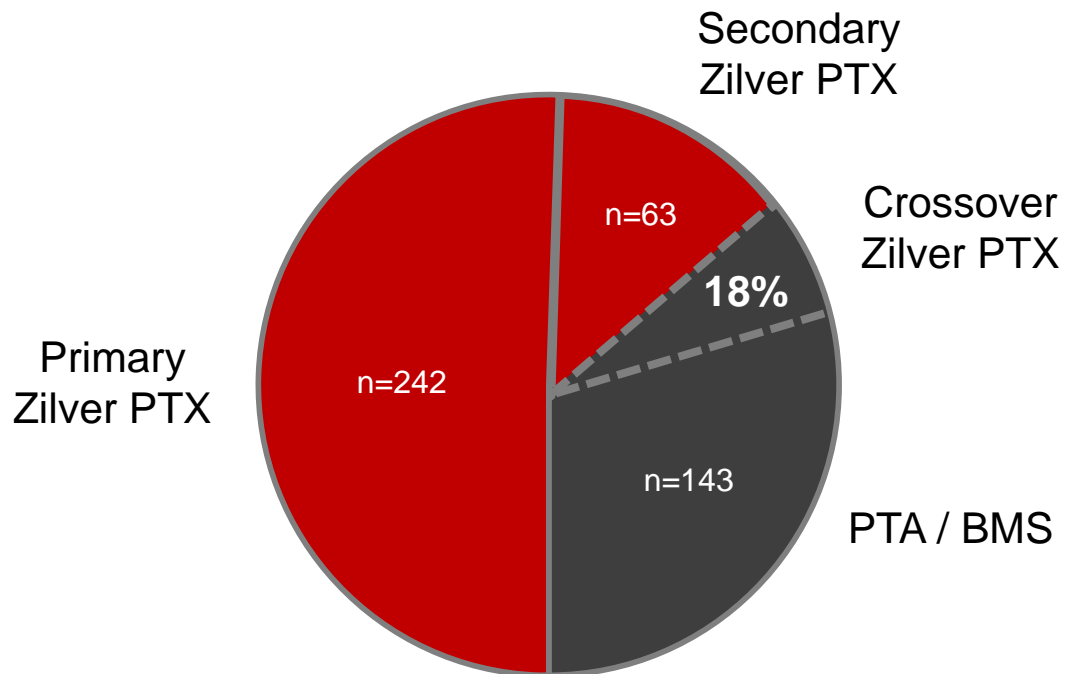


- ▶ 40% of the PTA/BMS group was treated with Zilver PTX
- ▶ Any analysis based on intent to treat is inappropriate for assessing paclitaxel mortality

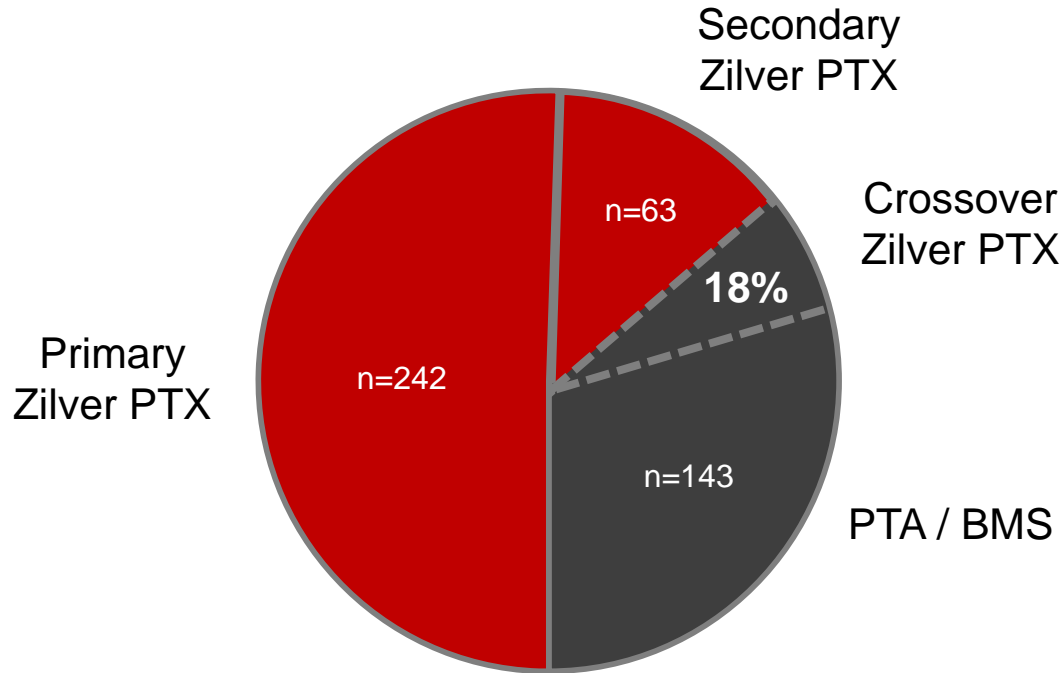
Secondary Randomization



Early Crossover

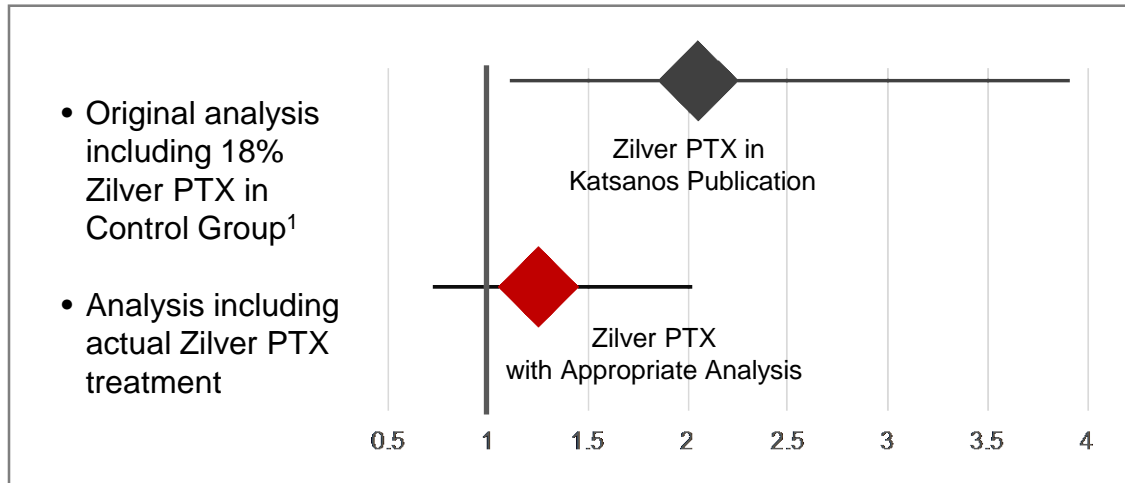


Zilver PTX Results Attributed to PTA/BMS



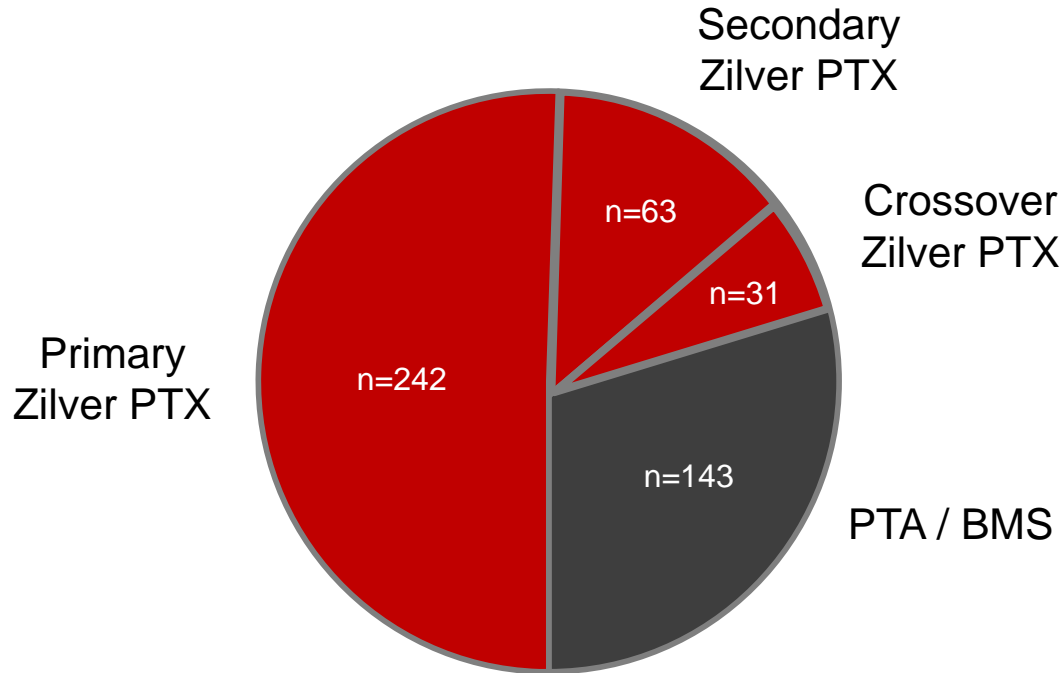
- ▶ Analyses by Katsanos, et al and FDA do not account for 18% of patients treated with Zilver PTX
- ▶ Zilver PTX mortality results were attributed to PTA/BMS group

Paclitaxel Mortality Meta-Analysis



- ▶ Evaluating all patients treated with Zilver PTX changes the conclusion
- ▶ In addition, the result of the meta-analysis becomes non-significant

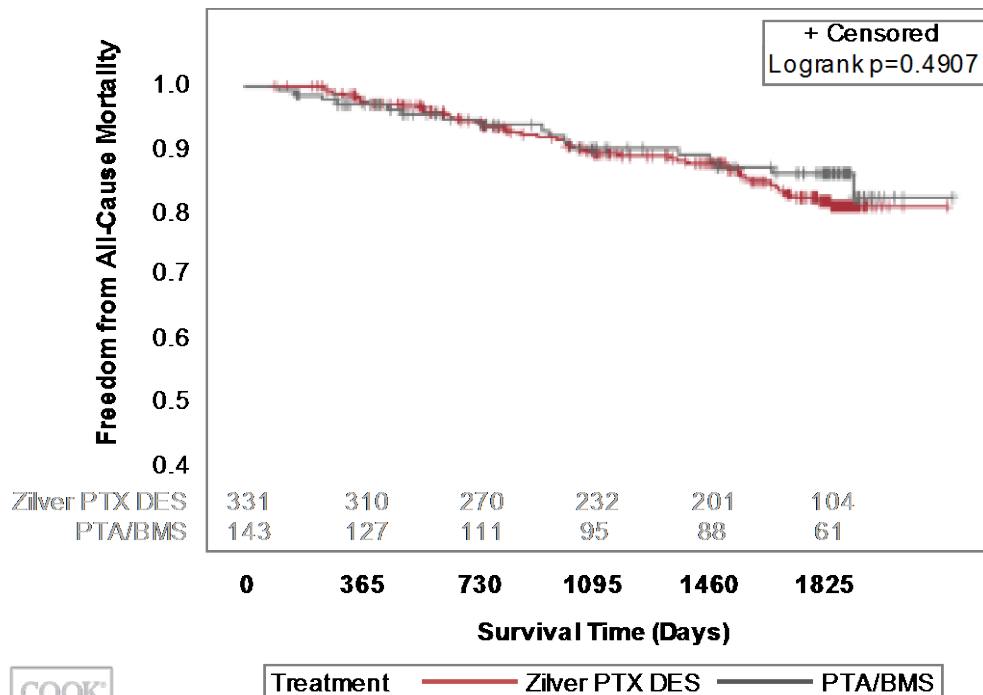
Actual Treatment



- ▶ Actual treatment is an appropriate assessment of paclitaxel-related mortality
- ▶ FDA modified as-treated analysis and Cook analysis include actual treatment

FDA Analysis of Actual Treatment

FDA Panel Pack (Appendix E)

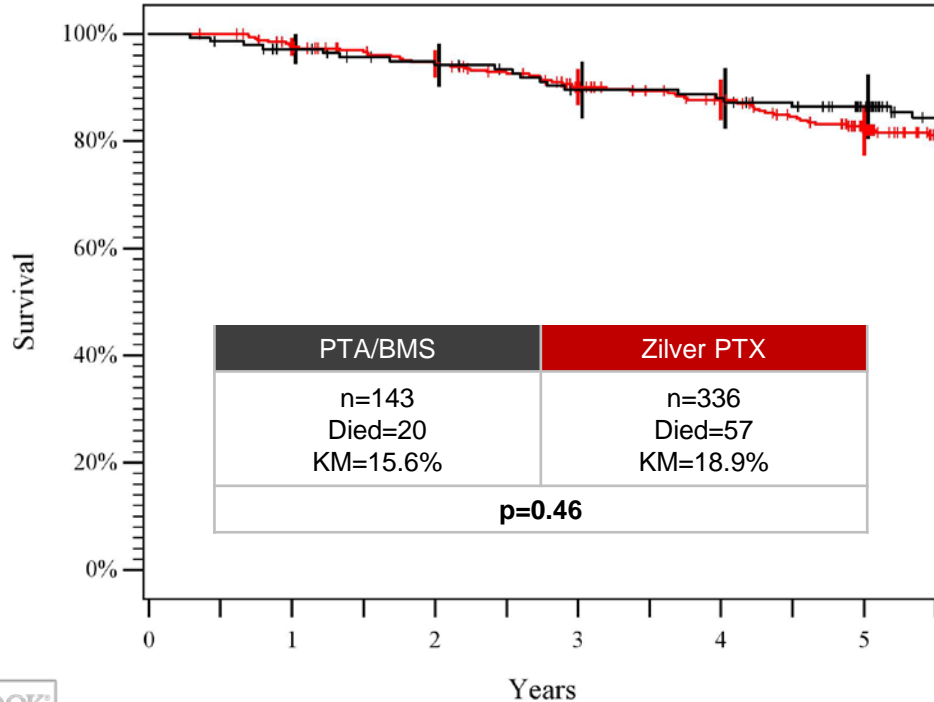


- ▶ All patients analyzed by actual treatment
- ▶ No mortality signal



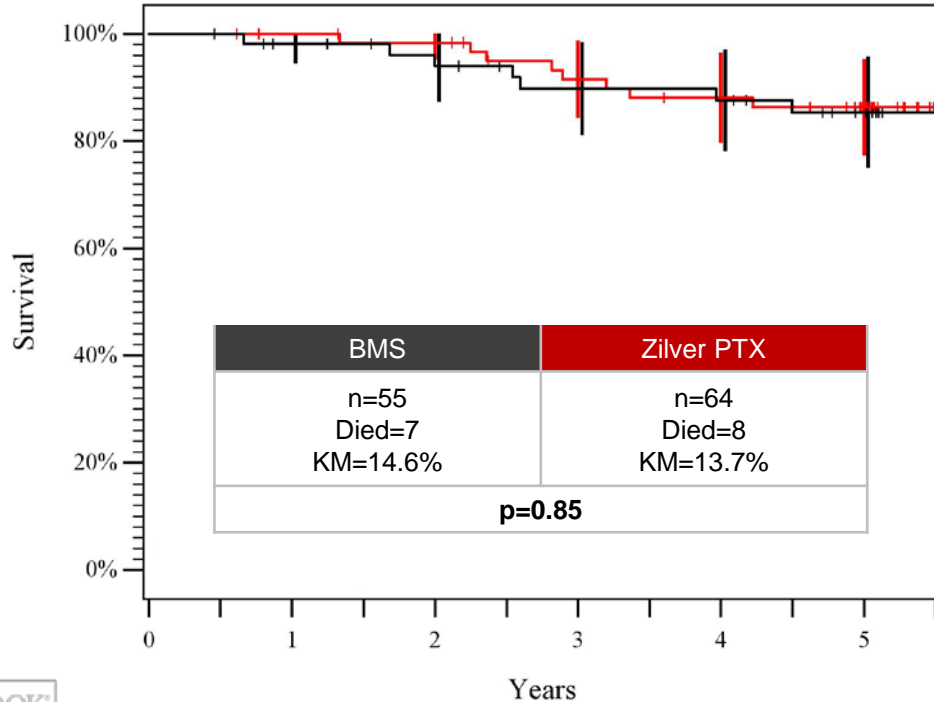
Note: The colors of the above graph were adjusted to be consistent with this presentation. No other changes were applied.

Cook Analysis of Actual Treatment



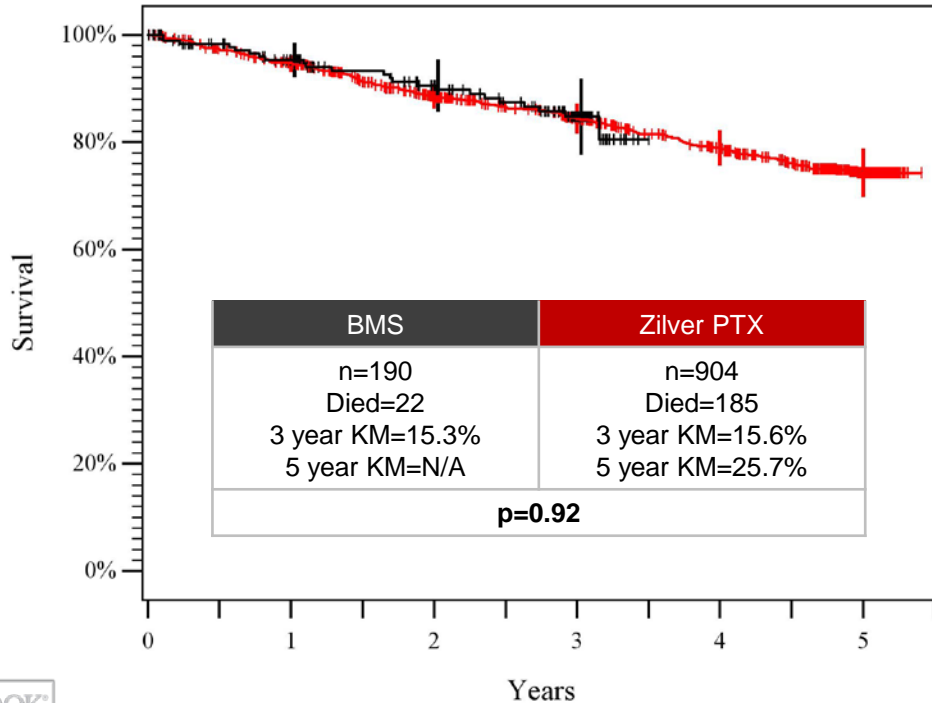
- ▶ Includes new patient status for 92% of patients previously lost-to follow-up
- ▶ Added data confirmed no mortality signal

Randomized Comparison to BMS



- ▶ Head-to-head comparison of Zilver PTX to BMS
- ▶ No mortality signal

Japan PMS: No Mortality Signal

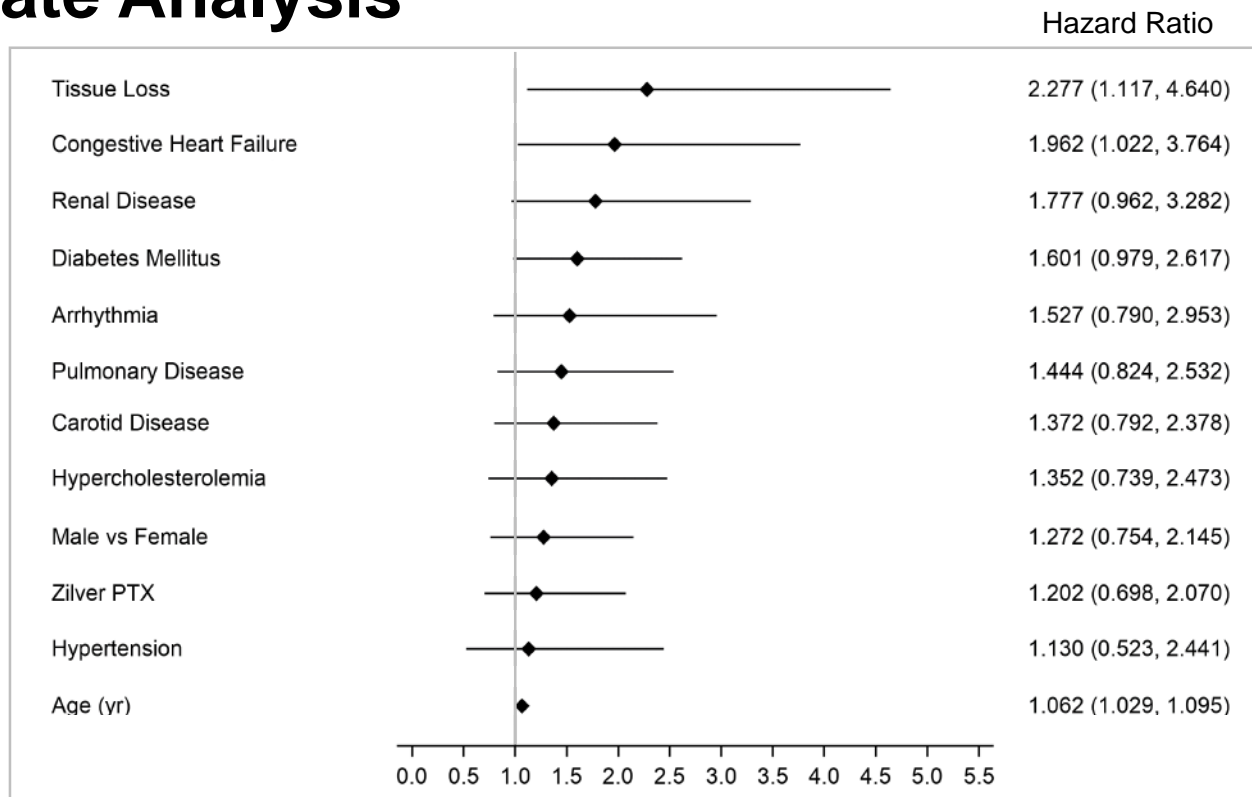


- ▶ Large, real-world, post-market studies
- ▶ No increase in rate of mortality after 3 years
- ▶ No mortality signal

Covariate Analysis

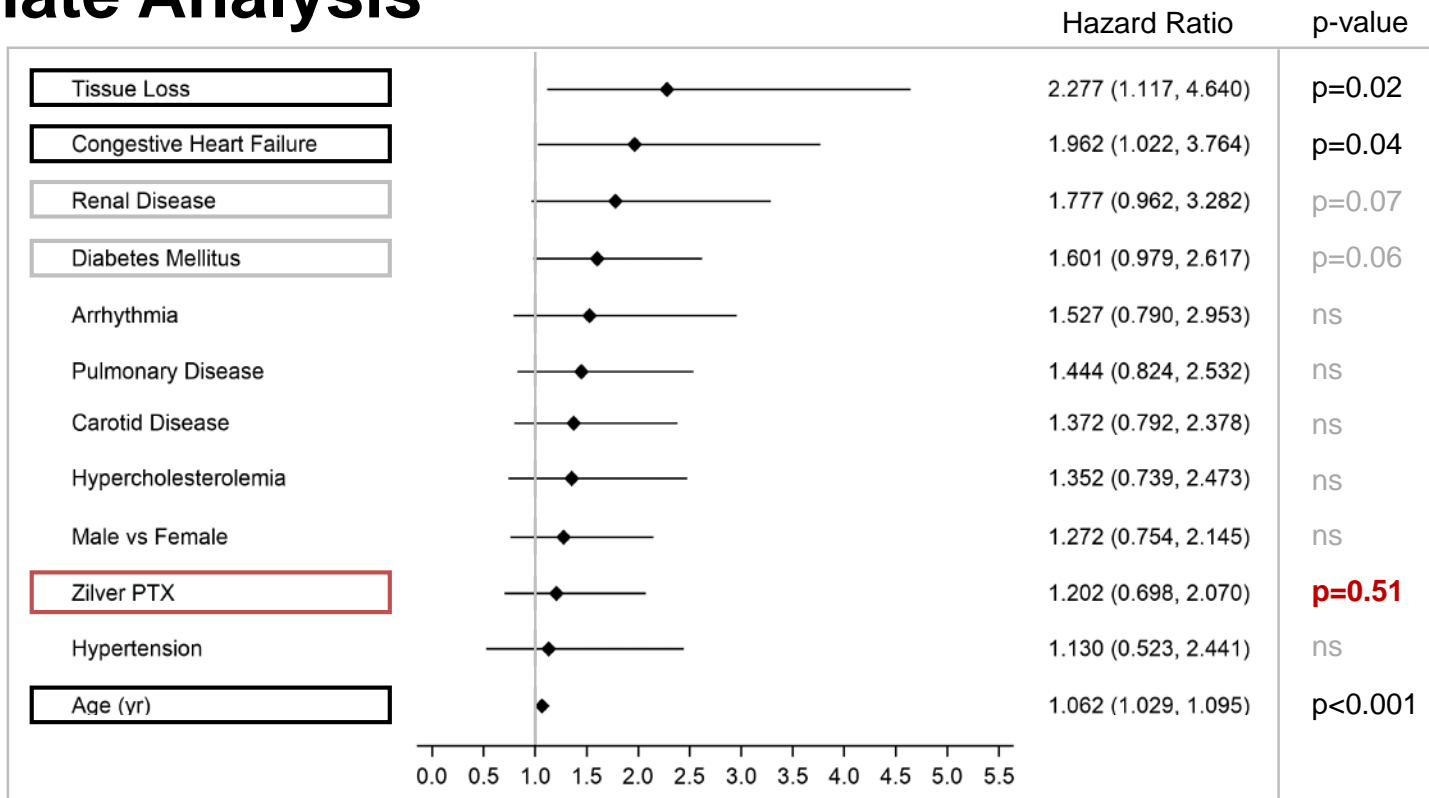
- No mortality signal for Zilver PTX when evaluating actual treatment
- What factors were associated with mortality?

Covariate Analysis



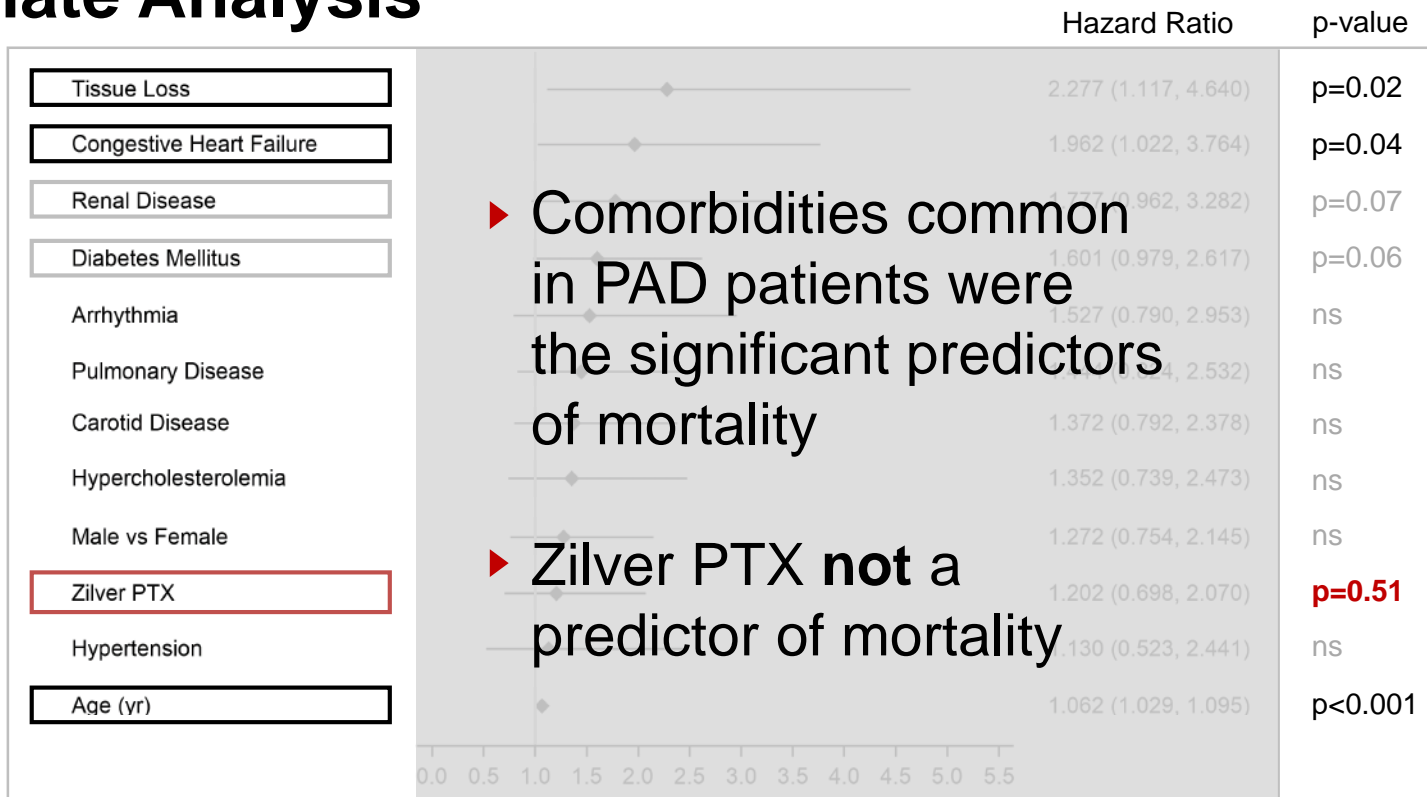
Additional non-significant factors included: smoking status, country, CLI/ Claudication, lesion length, previous MI, BMI

Covariate Analysis



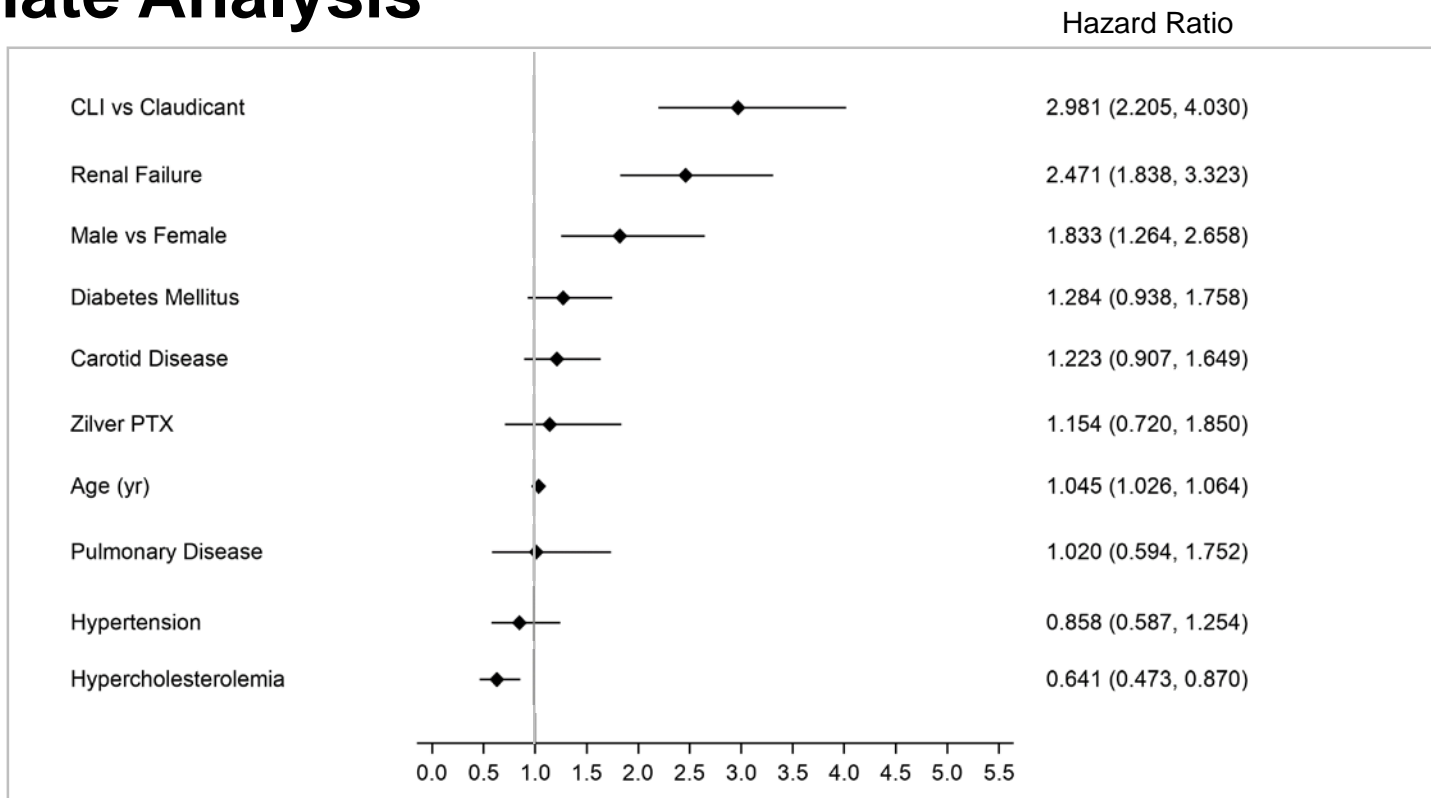
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Covariate Analysis



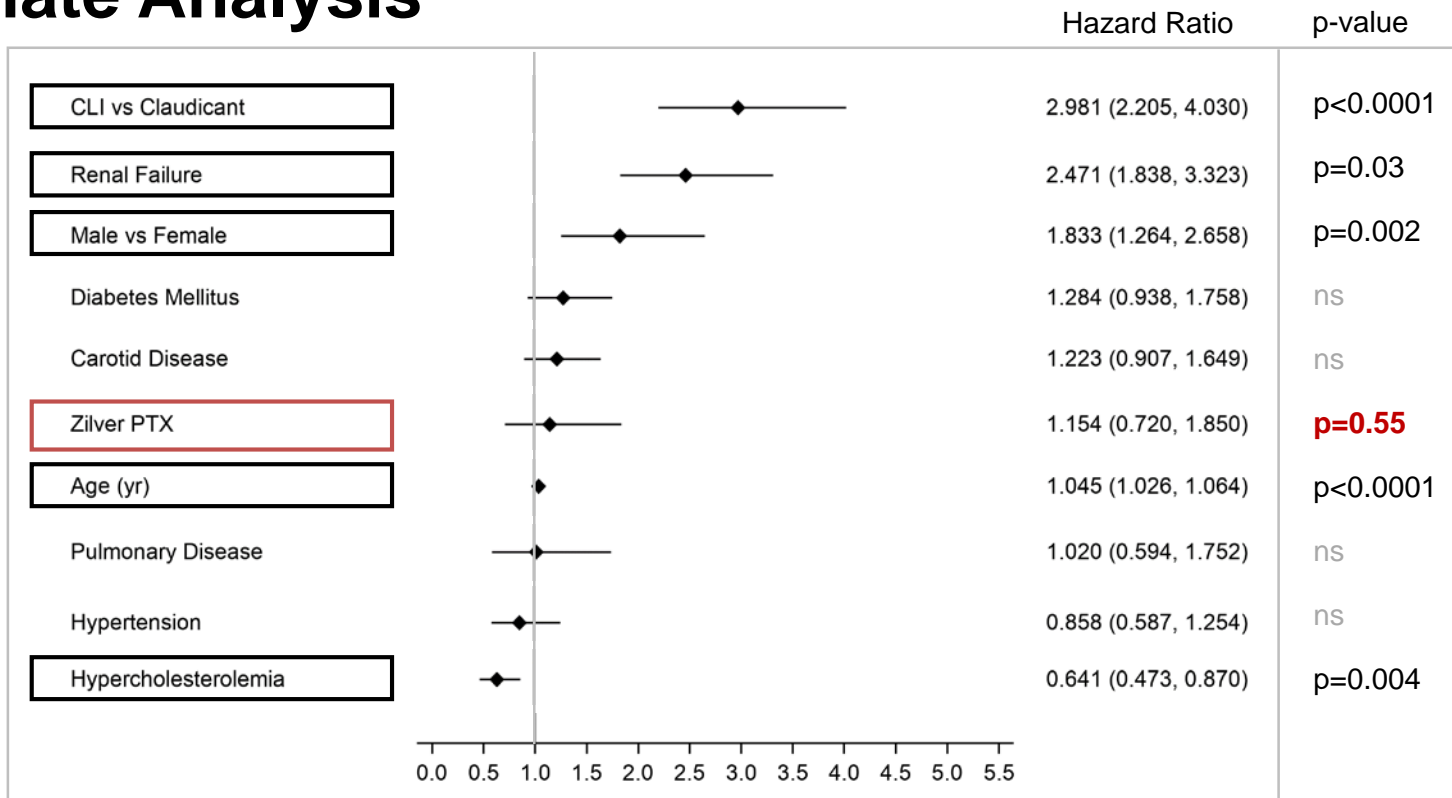
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Covariate Analysis



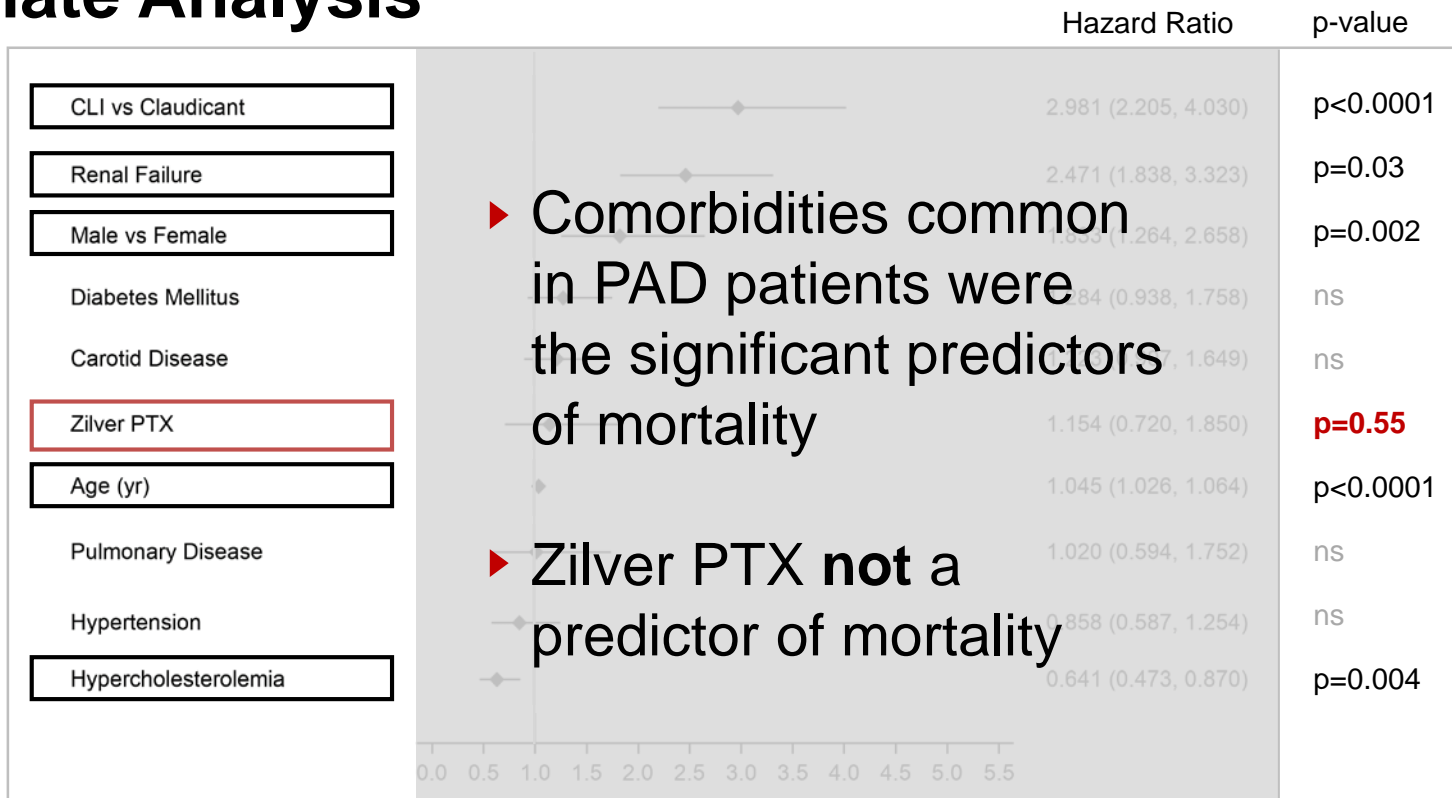
Additional non-significant factors included: smoking status, lesion length

Covariate Analysis



Additional non-significant factors included: smoking status, lesion length

Covariate Analysis



▶ Comorbidities common in PAD patients were the significant predictors of mortality

▶ Zilver PTX not a predictor of mortality

Additional non-significant factors included: smoking status, lesion length

Covariate Analysis: Dose

- Paclitaxel analyzed by dose (mg) per patient
- Significant predictors same as treatment arm analysis
 - RCT: Age, tissue loss, CHF
 - Japan: CLI, age, gender, renal, hypercholesterolemia

Study	Hazard Ratio	p-value
RCT	0.968 (0.659, 1.422)	0.87
Japan	1.162 (0.961, 1.404)	0.13

Covariate Analysis: Dose

- Paclitaxel analyzed by dose (mg) per patient
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Study	Hazard Ratio	p-value
▶ Paclitaxel dose	not a predictor of mortality	0.87
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Conclusion

- ▶ **Analysis must be based on actual treatment**

Protocol defined secondary randomization and crossover must not be ignored

- ▶ **No mortality signal with Zilver PTX**

When data are appropriately analyzed

- ▶ **Patient care is being negatively impacted**