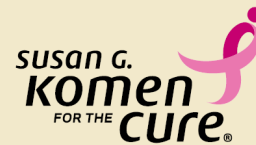




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Facilitating the Development of Immunotherapies:
Intermediate Endpoints for Immune
Checkpoint Modulators

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Immune Checkpoint Modulators

Jim Allison, Ph.D.

The University of Texas

MD Anderson Cancer Center

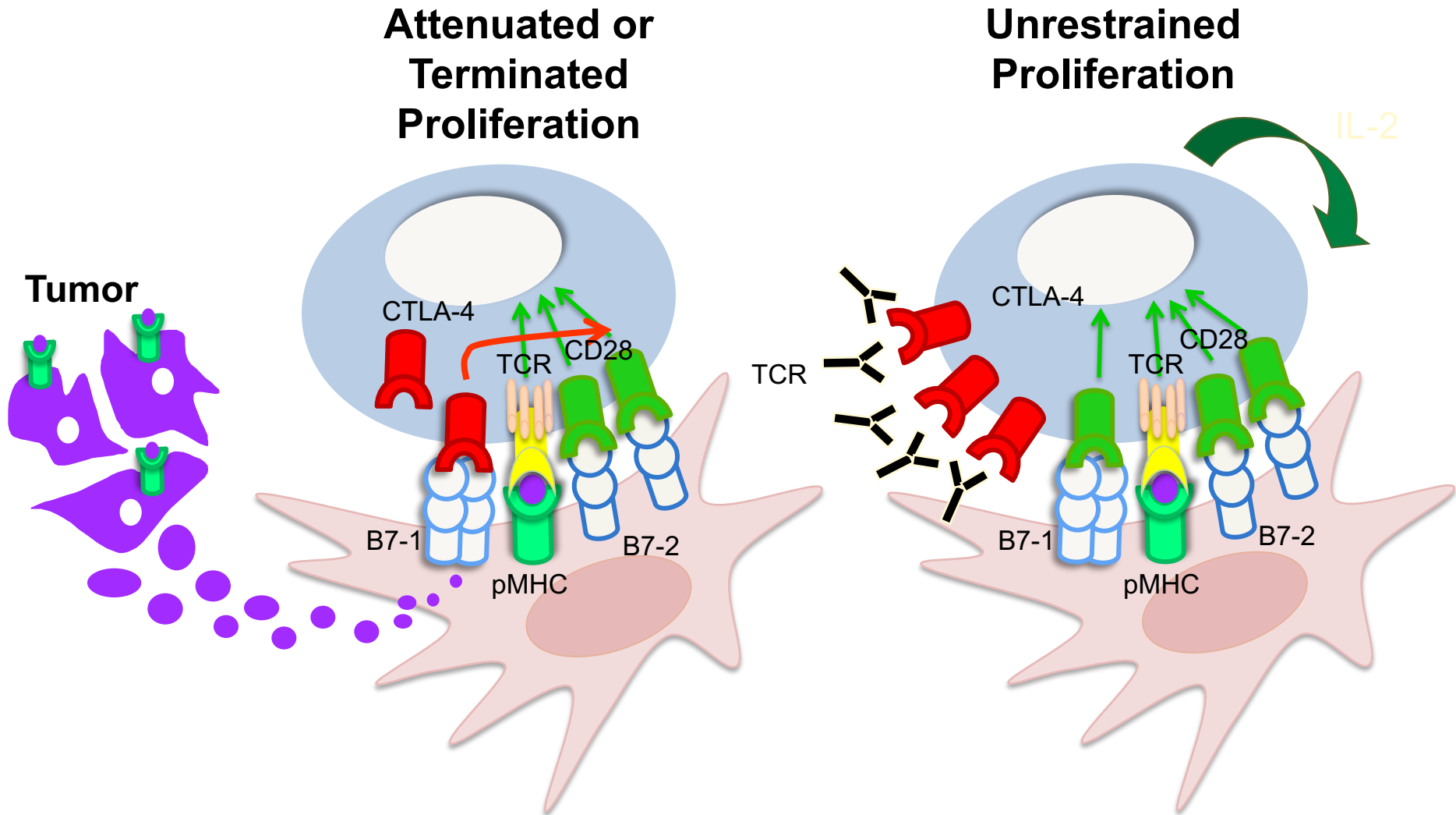
Why immunotherapy?

Specificity

Memory

Adaptability

CTLA-4 Blockade Enhances Tumor-Specific Immune Responses



Evolution of Response: Patient Example

Screening



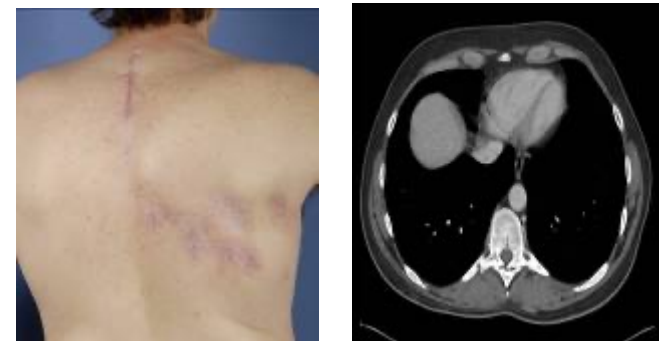
Week 12
Initial increase in
total tumor burden (mWHO PD)



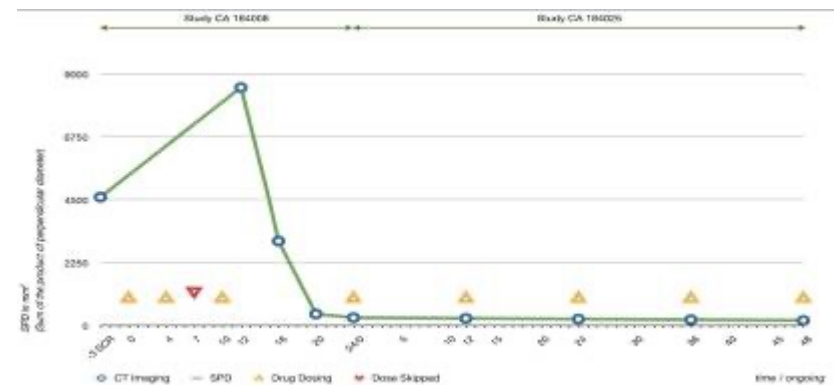
Week 16
Responding



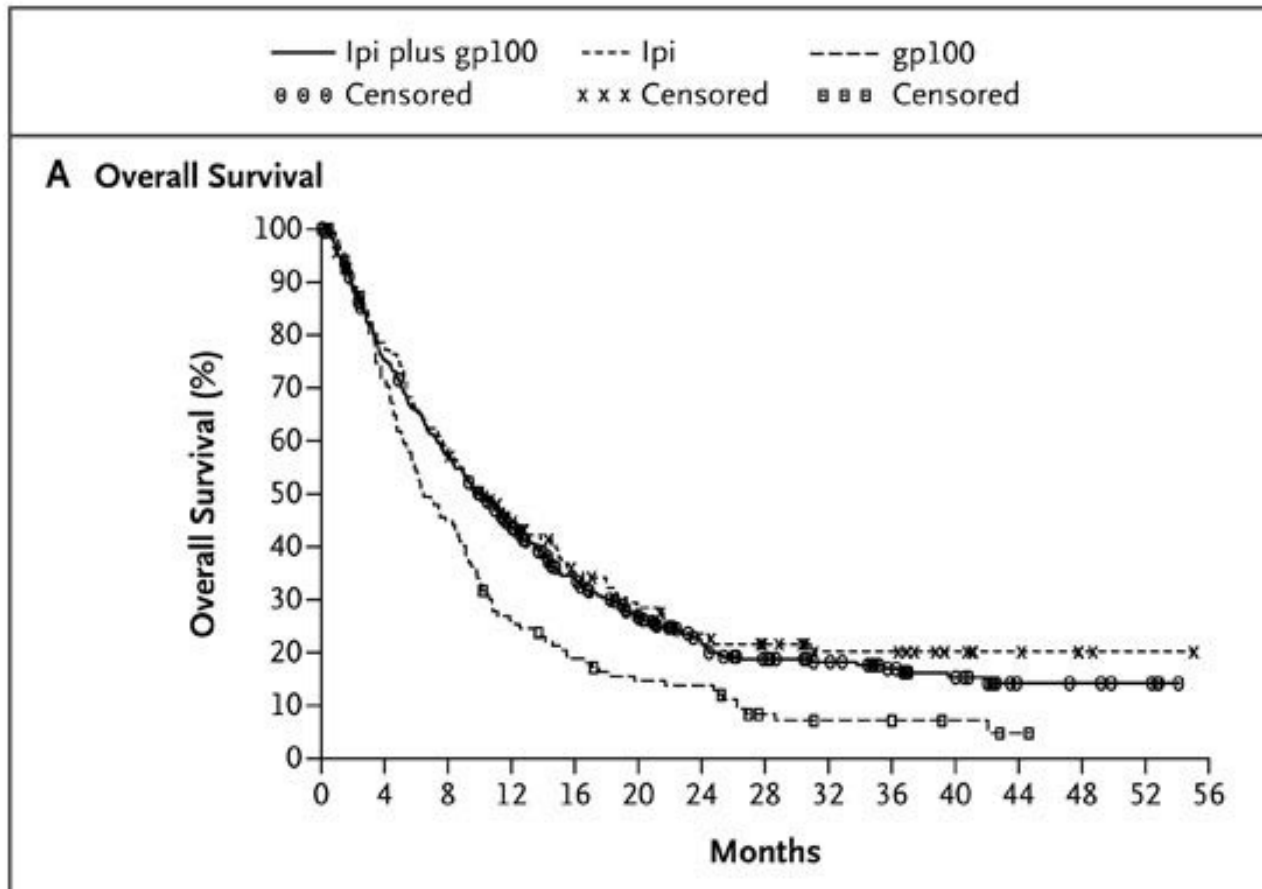
Week 72
Durable & ongoing response
without signs of IRAEs



20006

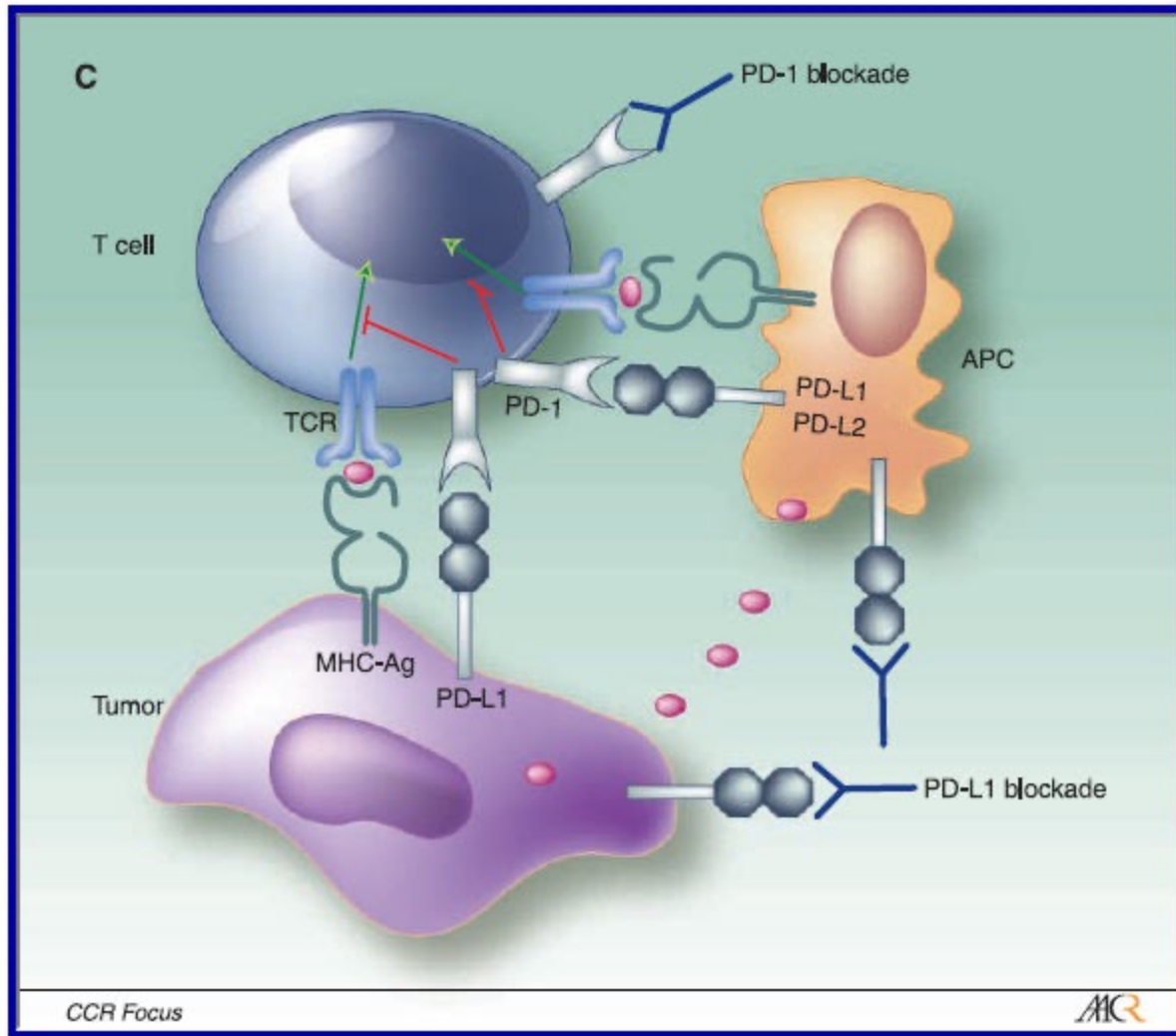


Kaplan-Meier Analysis of Survival



Survival Rate	Ipi + gp100 N=403	Ipi + pbo N=137	gp100 + pbo N=136
1 year	44%	46%	25%
2 year	22%	24%	14%

Programmed Death 1



Anti – PD-1 (BMS-936558)

**296 Patients with Metastatic Cancer
1, 3, 10 mg/kg, MTD not reached**

**Safety: Adverse events similar to Ipilimumab, but
4% pneumonitis (3 deaths)**

Clinical Activity:

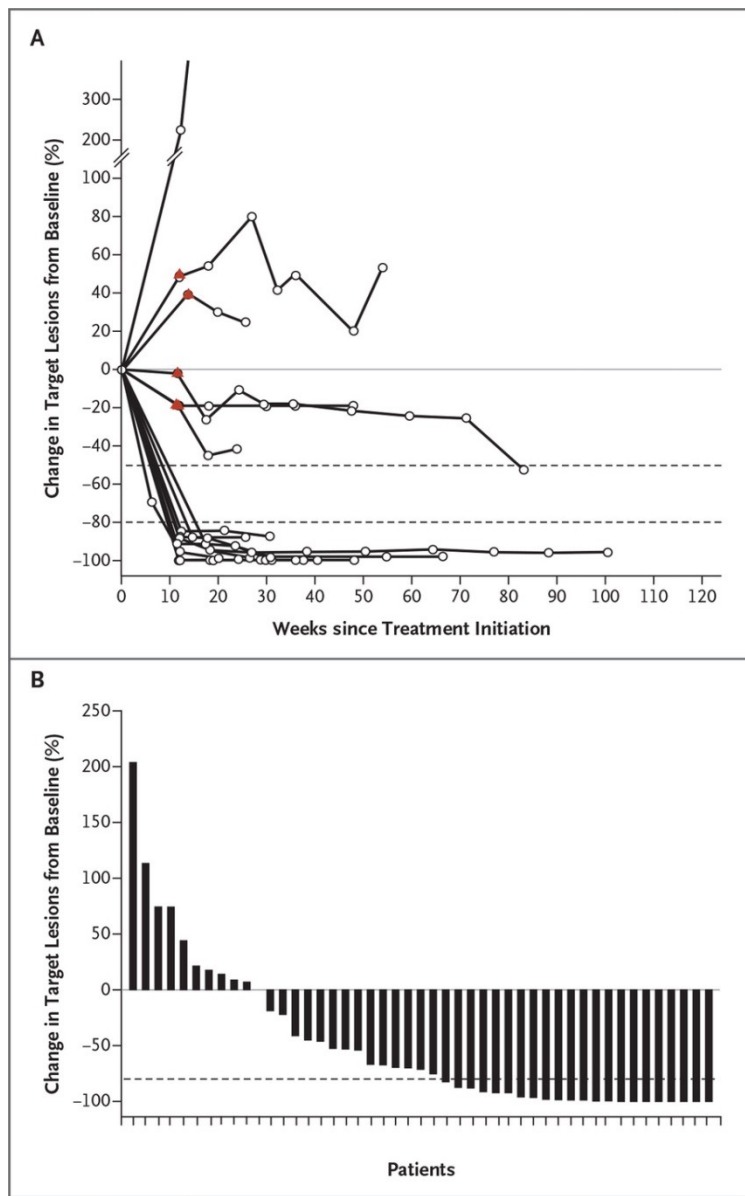
Melanoma (n= 94): 28% CR/PR, 6% SD

NSCLC (n=76): 18% CR/PR, 7% SD

RCC (n= 33): 27% CR/PR, 27% SD

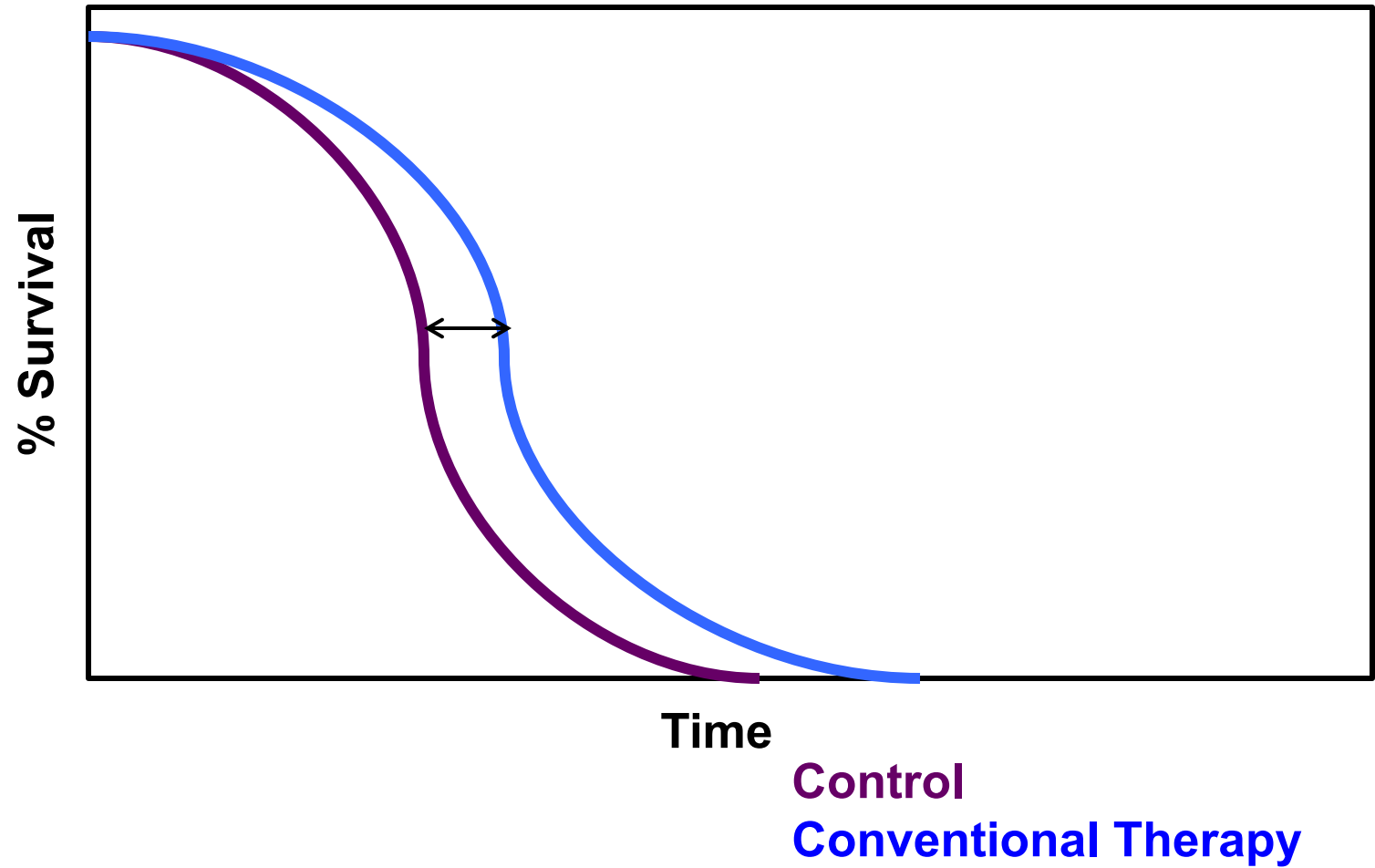
CRC (n=19), CRPC (n=13): No responses

Clinical Activity in Melanoma Patients Receiving Ipilimumab (α CTLA-4) and Nivolumab (α PD-1)

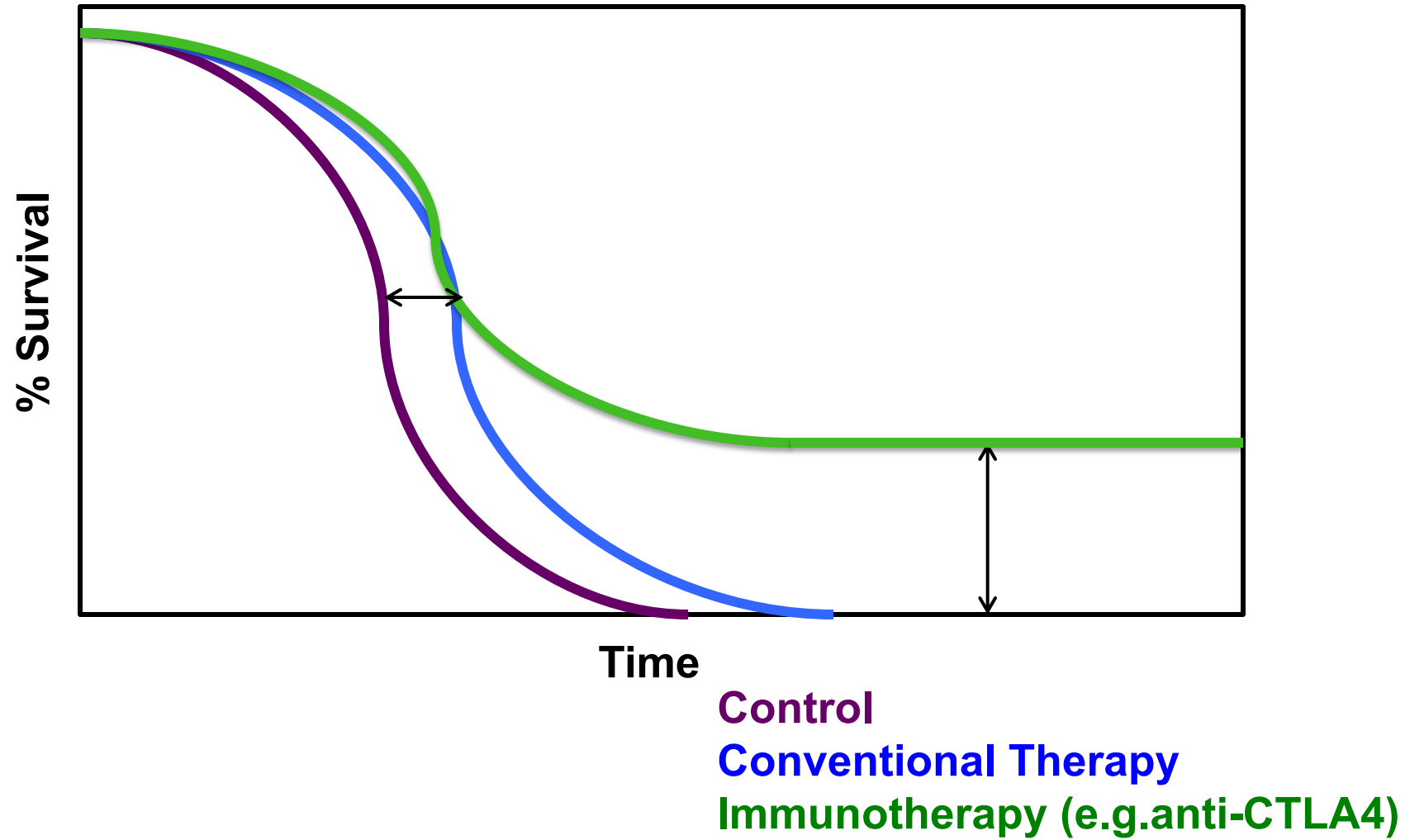


ASCO 2013
NEJM 6/2/2013

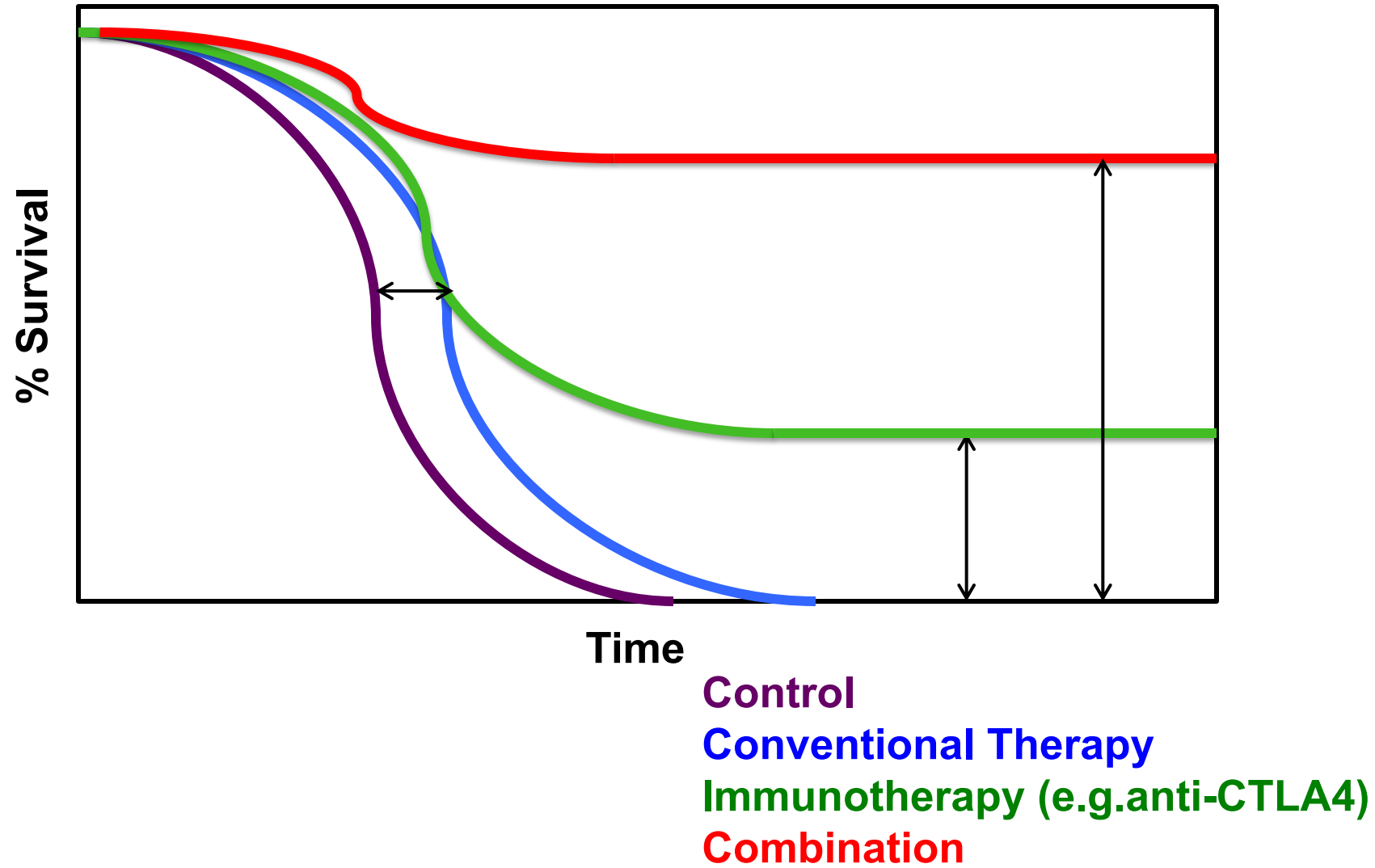
Improving Survival with Combination Therapy



Improving Survival with Combination Therapy



Improving Survival with Combination Therapy



Speakers

- **Jim Allison, Ph.D.**, U.Texas MD Anderson Cancer Center
- **Mark Gorman**, Survivor and Advocate
- **Ramy Ibrahim, M.D.** MedImmune
- **Axel Hoos, M.D., Ph.D.**, Glaxo-Smith Kline
- **Tai-Tsang Chen, Ph.D.**, Bristol-Myers Squibb
- **Steve Rosenberg, M.D., Ph.D.**, National Cancer Institute
- **Amy McKee, M.D.**, FDA-CDER
- **Celia Witten, M.D., Ph.D.**, FDA-CBER
- *Contributors:* **Renzo Canetta, M.D.**, **Suzanne Topalian, M.D.**

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Mark Gorman

**Long-term Survivor of Metastatic Melanoma
And Patient Advocate**

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Immunotherapies: Dosing Challenges

Ramy Ibrahim, M.D.
MedImmune

Conventional dose/schedule selection and anti-cancer development

- Preclinical data from efficacy studies to identify target exposures in human
- Escalate doses in FTIH studies to assess safety and achieve target exposures (or higher) to increase likelihood of early signal
- Determine the MTD after DLTs are observed
- Select MTD for further development in randomized studies to assess efficacy
- Initiate registrational studies

Novelties with immune-modulators and implications on dose/schedule selection

- Animal data might not inform dose selection
 - Cross reactivity and finding surrogate has limitations
- The “target” is the immune system and not the cancer
 - Complexity of the interaction between the immune system and cancer
 - Patients might have different threshold or sensitivity to immune priming
 - We need to identify a dose that achieves appropriate exposure while accommodating inter-patient variability
- Immune targets are dynamic
 - Variability in target level, site of expression, tumor type and tumor burden
- Animal data and PK modeling might only inform the starting dose and identify a target exposure range

Novelties with immune-modulators and implications on dose/schedule selection (cont)

- Dose escalation till “toxicity” is not a viable approach
 - None of the PD1/PDL1 targeting antibodies reached an MTD
 - Activity observed at multiple dose levels
 - Early phase clinical PK, target related biomarkers, markers of immune response and clinical activity should be leveraged
 - Need for novel phase I designs to inform dose selection

Novelties with immune-modulators and implications on dose/schedule selection (cont)

- ◆ Dose-ranging comparative studies may not necessarily better inform dose selection
 - Tremelimumab development
 - Randomized phase II suggested 15 mg/kg q3mo to be associated with more favorable risk: benefit
 - Phase III study suggested 15mg/kg Q 3 months not to maintain desired AUC
 - Currently exploring monthly dosing
- ◆ Beside dose/schedule, what about duration of treatment?
- ◆ Due to the early and sometimes dramatic signal of activity, programs progress quickly from large phase 1 to phase 3
 - How to design better studies to inform registrational studies

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**Delayed Treatment Effects of Cancer
Immunotherapies**

Axel Hoos, M.D., Ph.D.
Glaxo-Smith Kline

A Methodological Framework for Immuno-Oncology

Challenge: Clinical trial endpoints are not immunotherapy-focused

Solution: Adjustment of endpoints to immunotherapy biology

Review

Lessons from randomized phase III studies with active cancer immunotherapies—Outcomes from the 2006 Meeting of the Cancer Vaccine Consortium (CVC)

Lothar H. Finke^{a,g,*}, Kerry Wentworth^{b,g}, Brent Blumenstein^c, Natalie S. Rudolph^d, Hyam Levitsky^{e,g}, Axel Hoos^{f,g}

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REVIEW

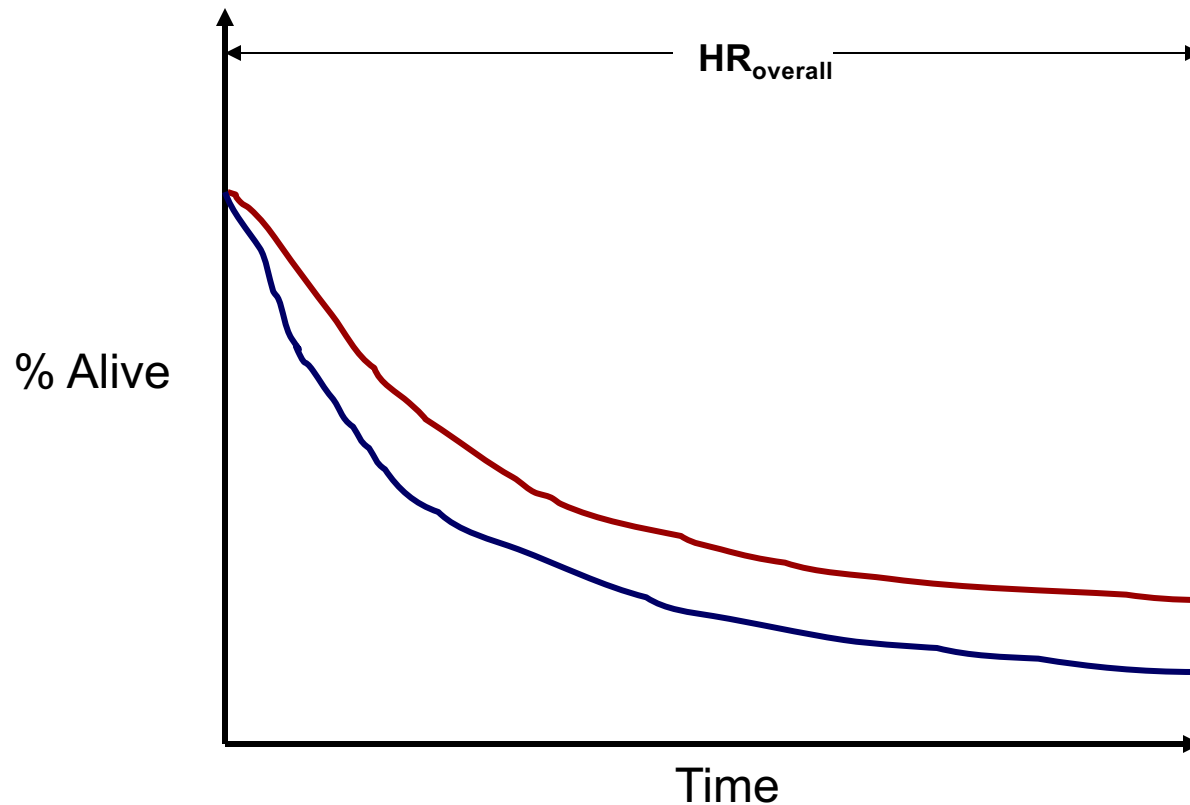
Improved Endpoints for Cancer Immunotherapy Trials

Axel Hoos, Alexander M. M. Eggermont, Sylvia Janetzki, F. Stephen Hodi, Ramy Ibrahim, Aparna Anderson, Rachel Humphrey, Brent Blumenstein, Lloyd Old, Jedd Wolchok

Vaccine 2007

J Natl Cancer Inst 2010

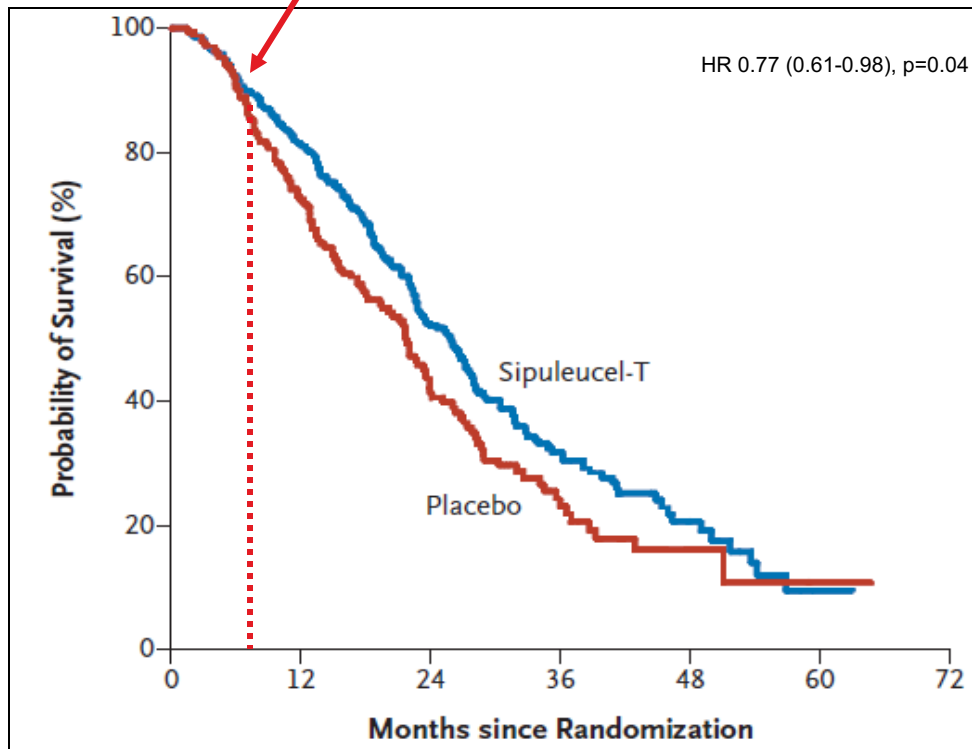
Survival: Conventional Design Assumptions



- No events occur before separation of curve
- Proportional hazard applies

Delayed Separation – Sipuleucel-T

Separation of curves
at ~8 months



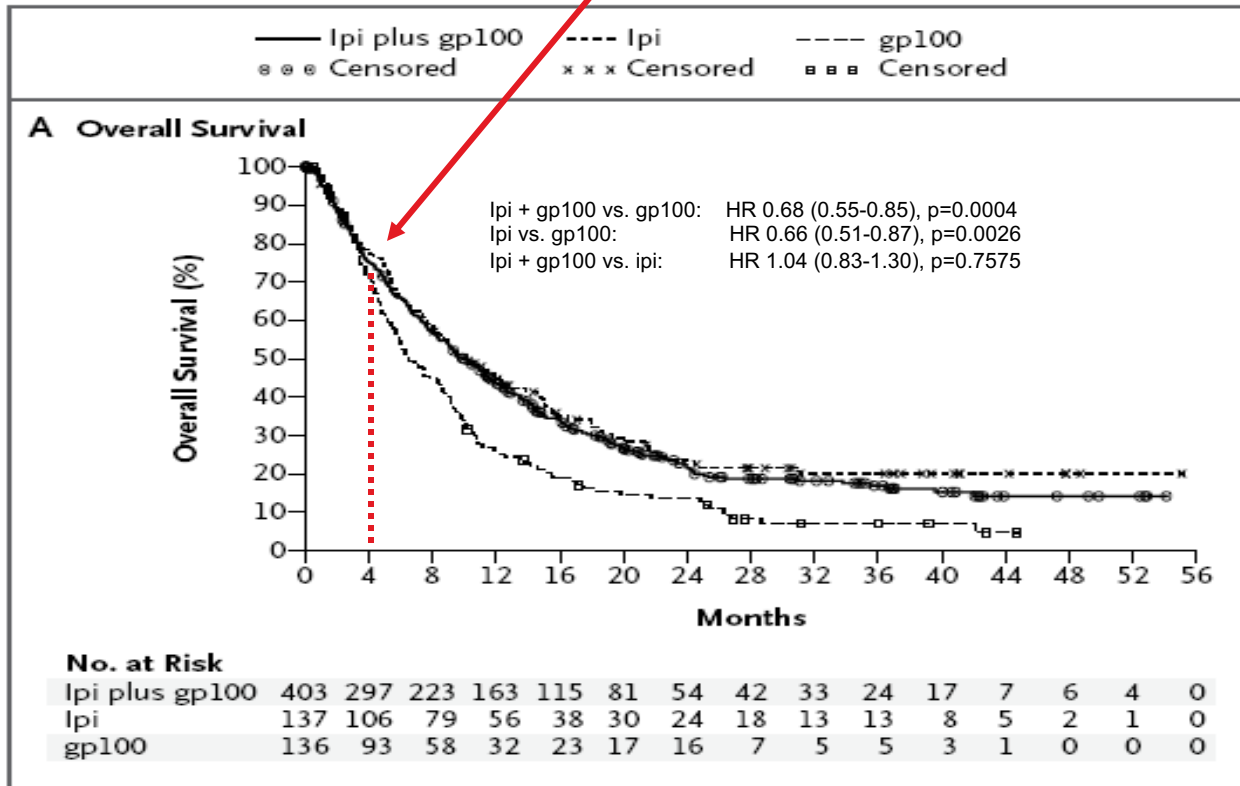
Sponsor:
Dendreon

Agent:
autologous dendritic
cell vaccine

Disease:
hormone-refractory
prostate cancer

Delayed Separation – Ipilimumab

Separation of curves
at ~4 months



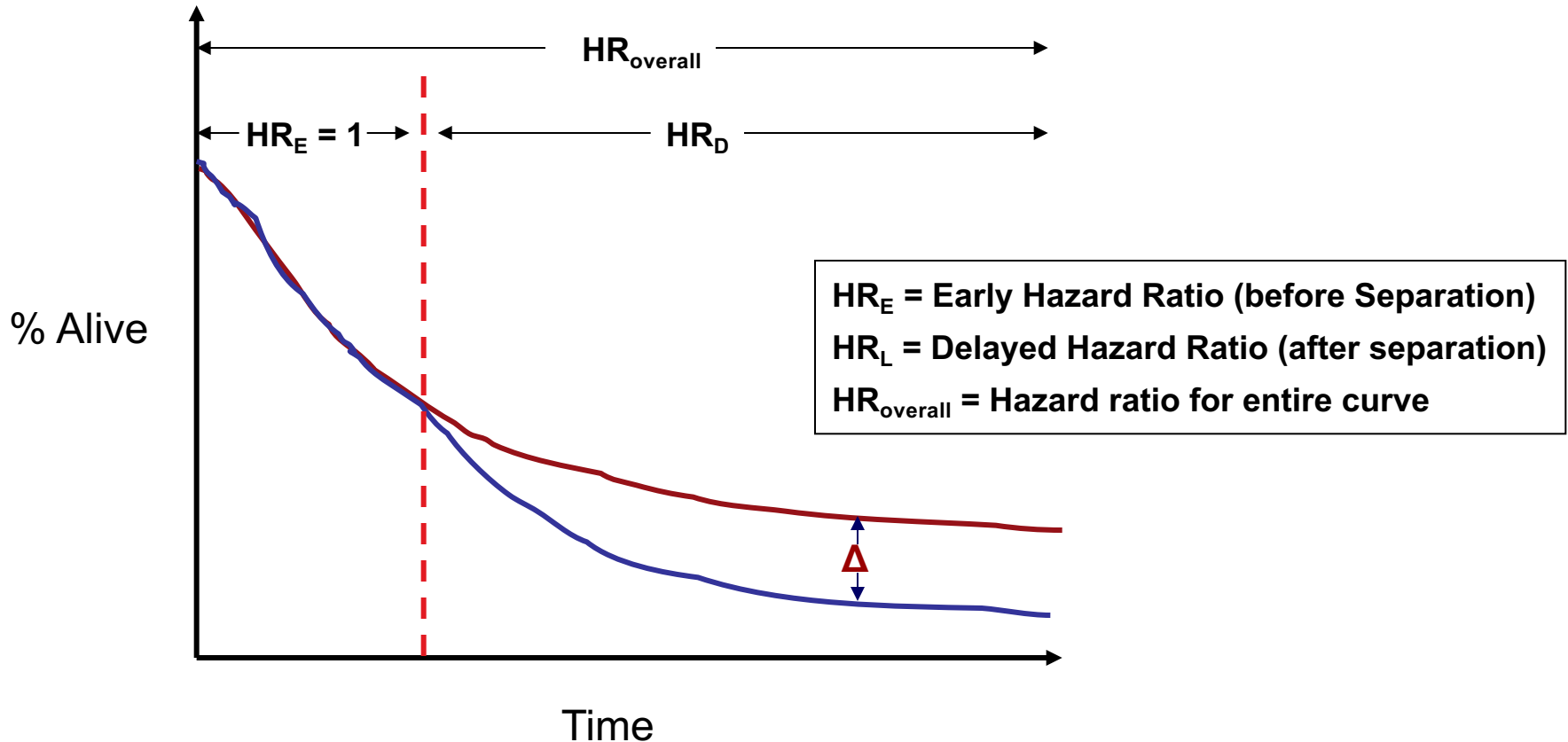
Sponsor:
BMS

Agent:
Anti-CTLA-4 mAb

Disease:
Metastatic
melanoma

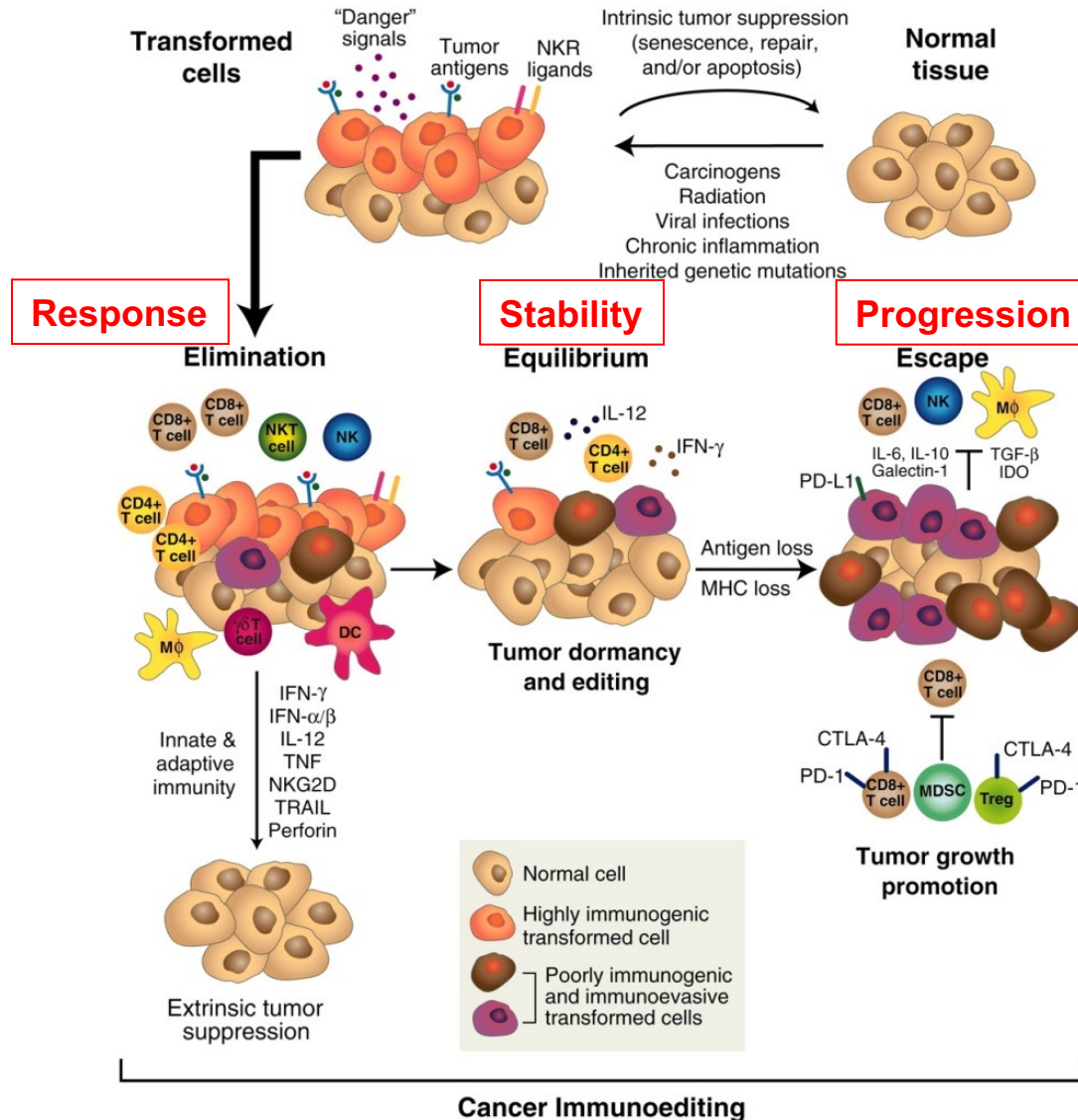
Implications of Delayed Separation of Curves

- Model Scenario -

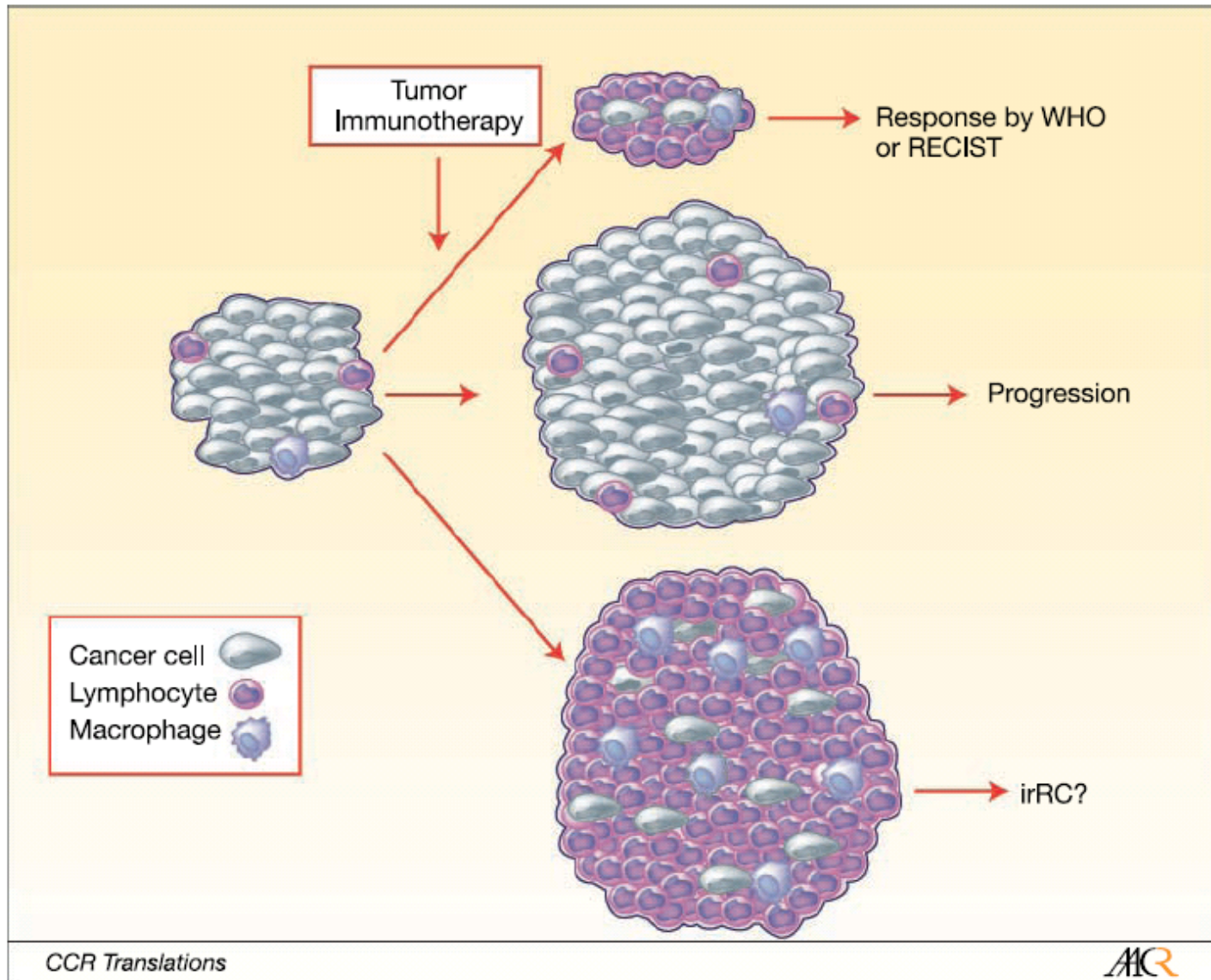


- Large Δ after separation needed to compensate for no effect before separation

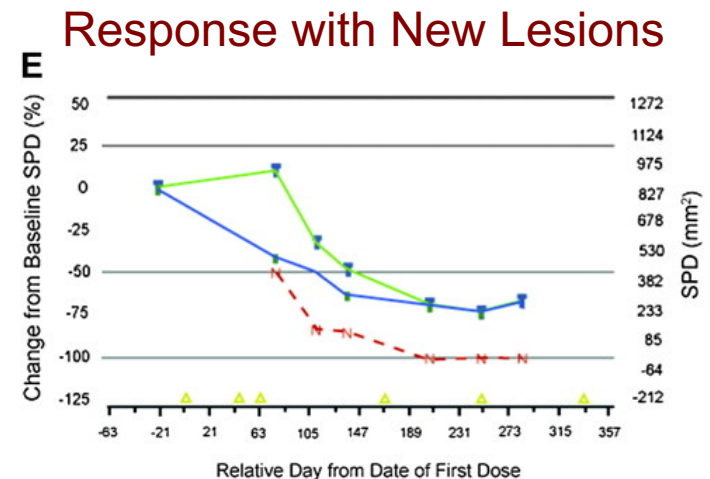
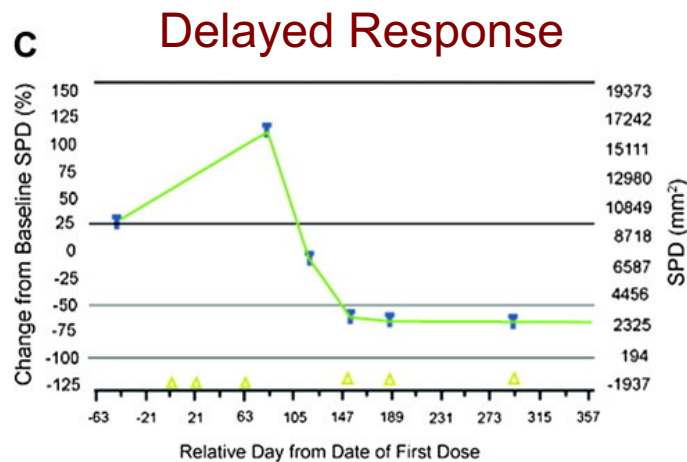
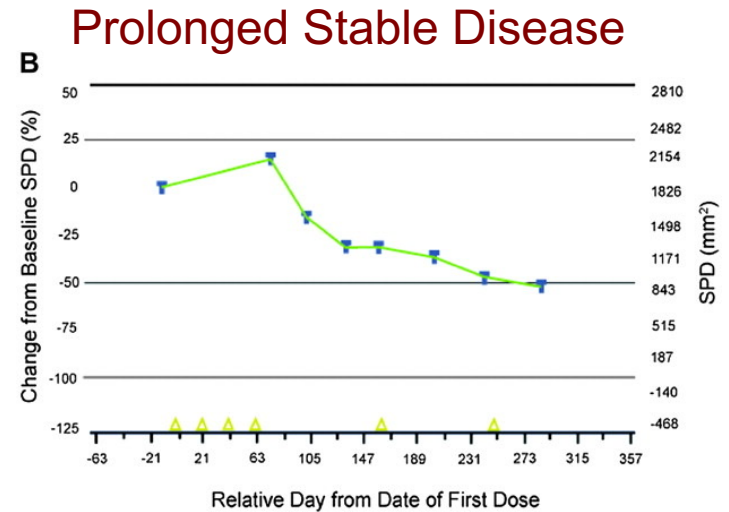
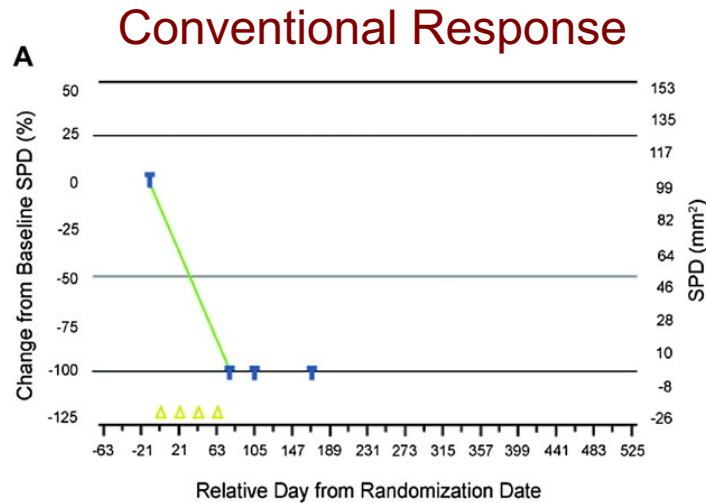
Interactions between Immune System and Tumor



Tumor Volume Increase Due to Lymphocyte Infiltration

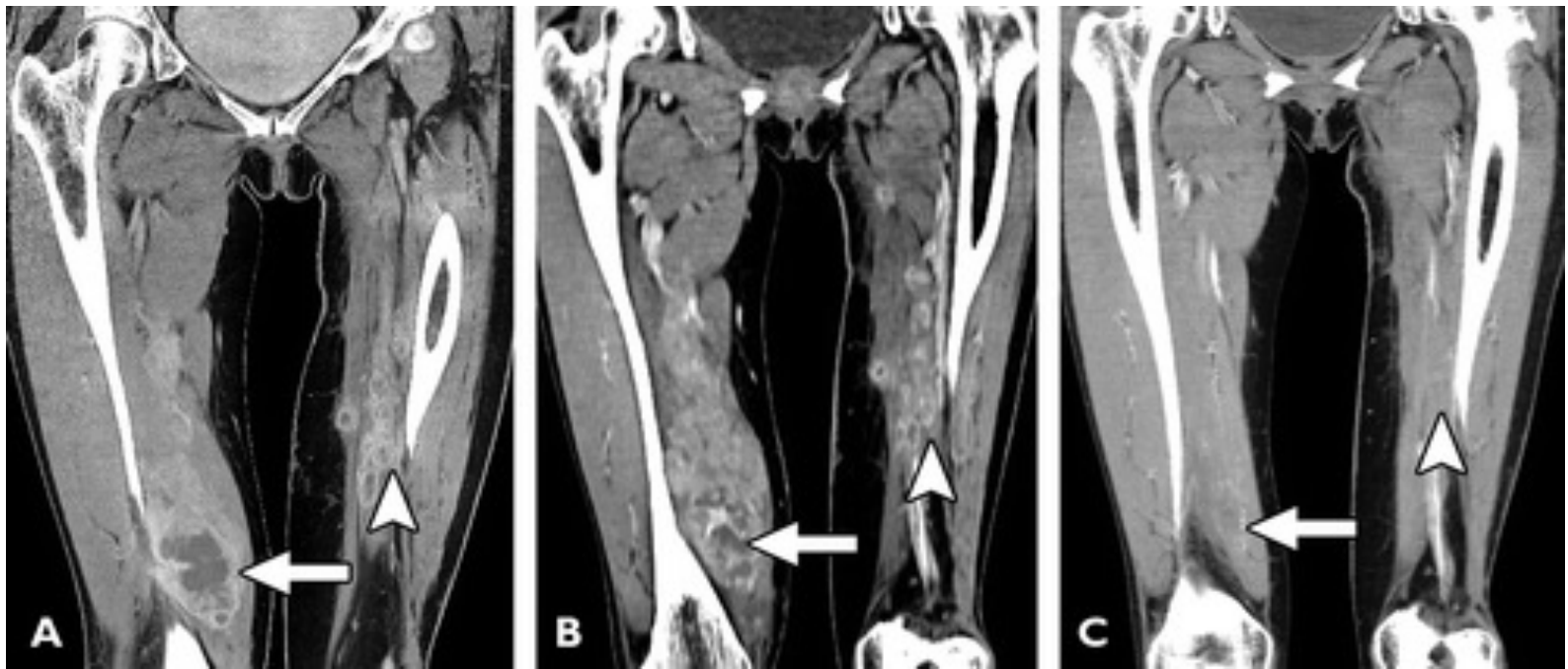


Immunotherapy Patterns of Response

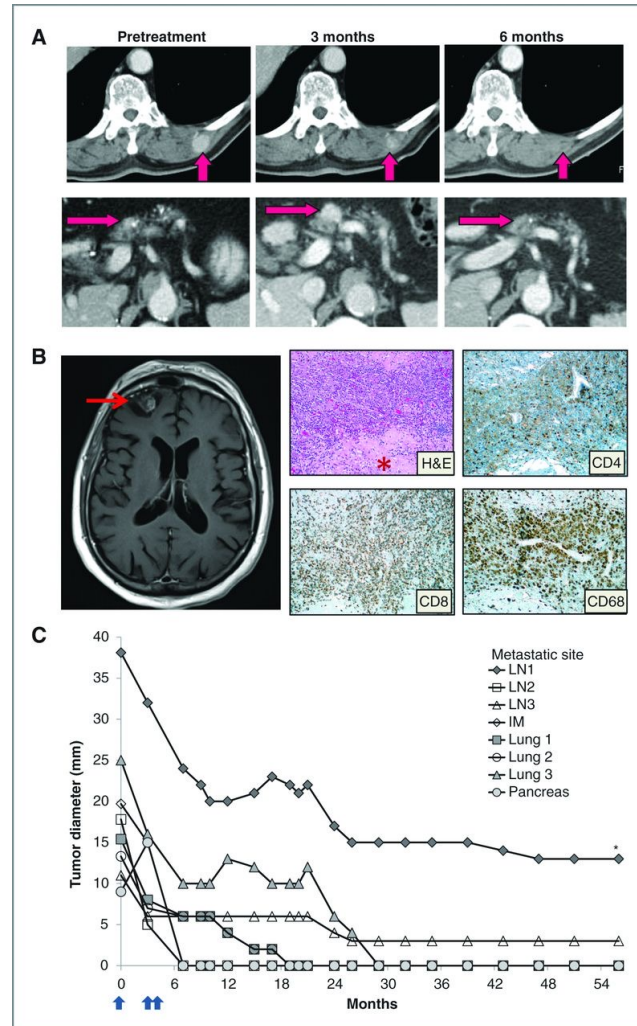


Wolchok JD, et al. *Clin Cancer Res.* 2009;15:7412-7420
 Hoos A et al. *JNCI J Natl Cancer Inst* 2010;102:1388-1397

Anti-CTLA-4 (Ipilimumab): Delayed Response

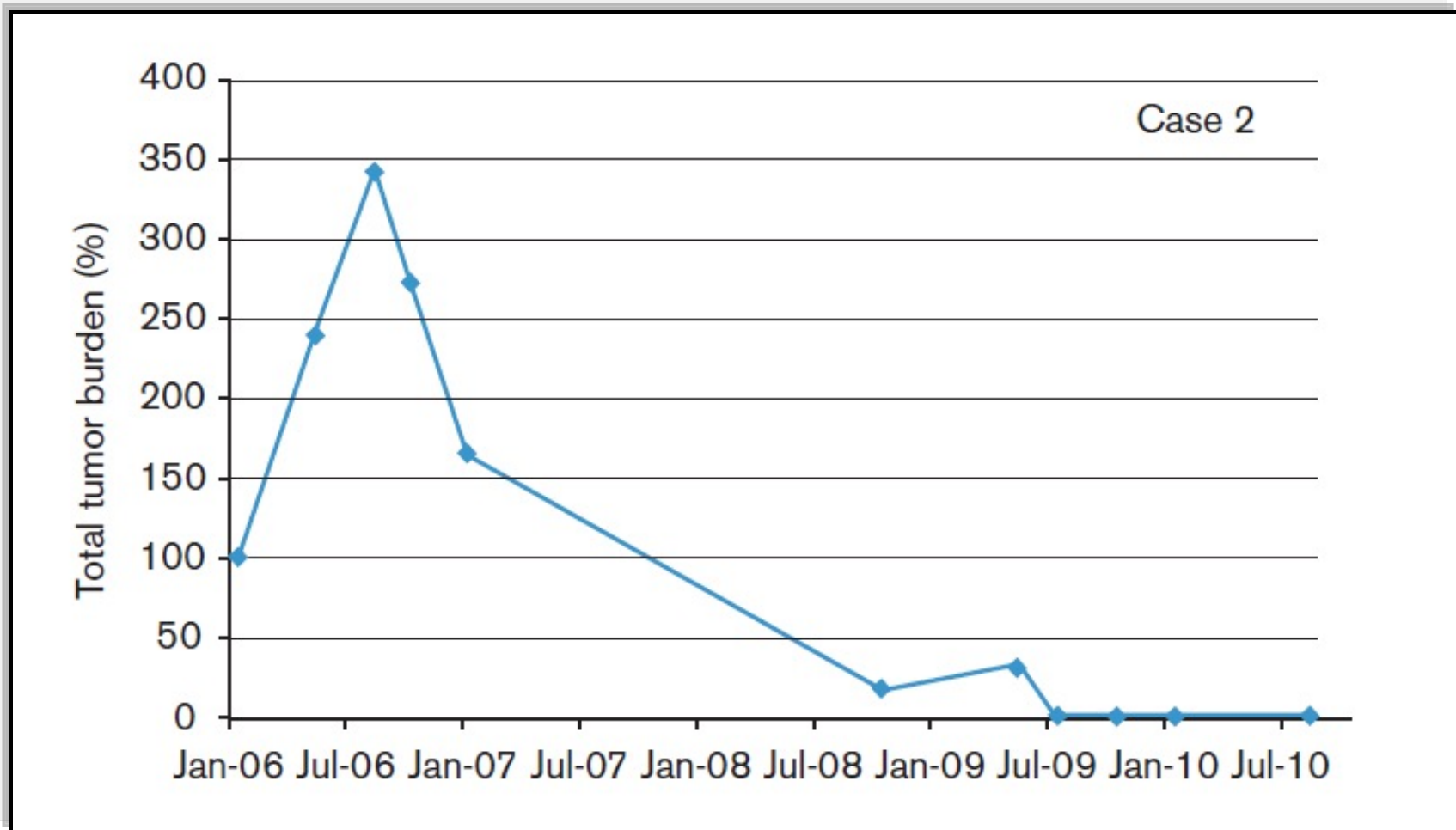


Regression of metastatic RCC following anti-PD-1 therapy, with “immune-related” response characteristics.

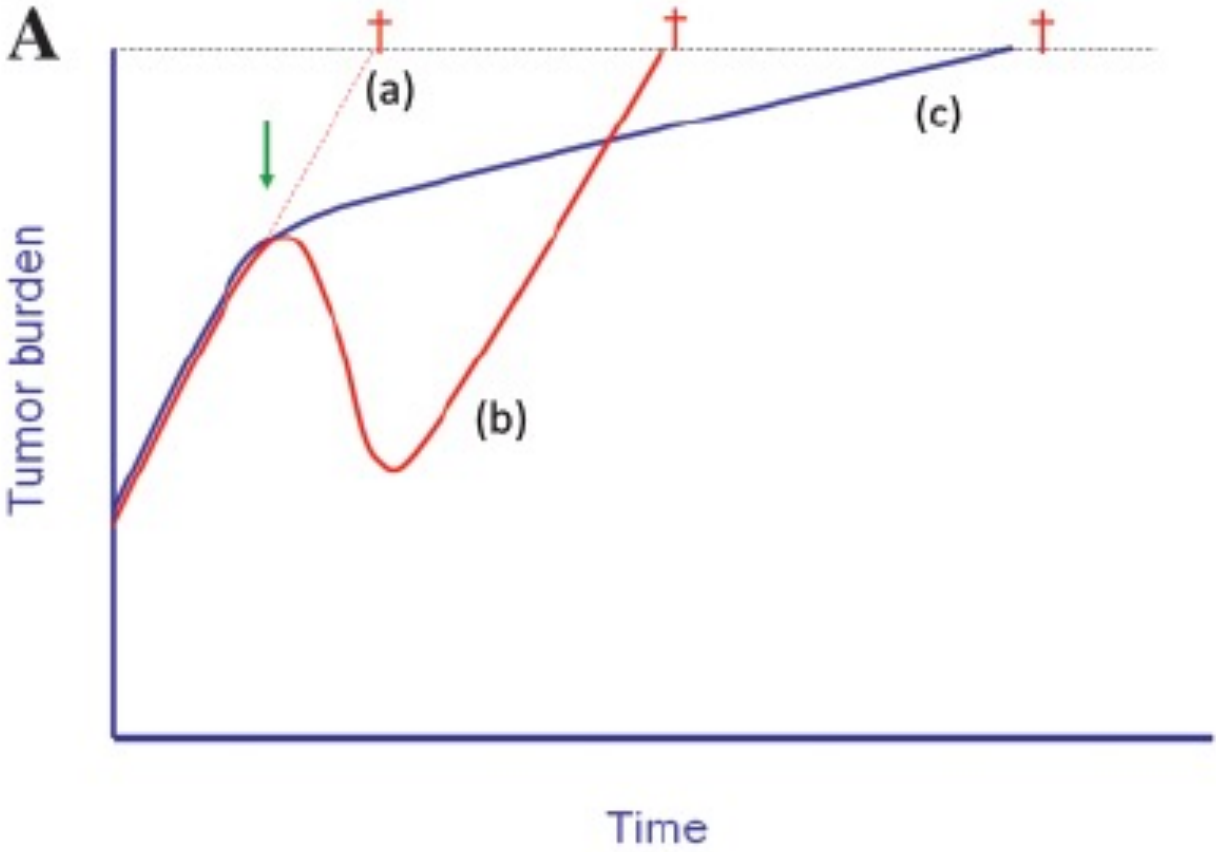


Lipson E J et al. Clin Cancer Res 2013;19:462-468

Autologous DC + IFN α 2b in Advanced Melanoma: Delayed Response



Tumor Growth Rate: Potential Impact on Survival



Available Tools

- Statistical methods for analyzing survival
- Immune-related Response Criteria
- Tumor growth kinetics

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**Intermediate Endpoints for Immune
Checkpoint Modulators: Milestone OS Analysis**

Tai-Tsang Chen, Ph.D.
Bristol-Myers Squibb

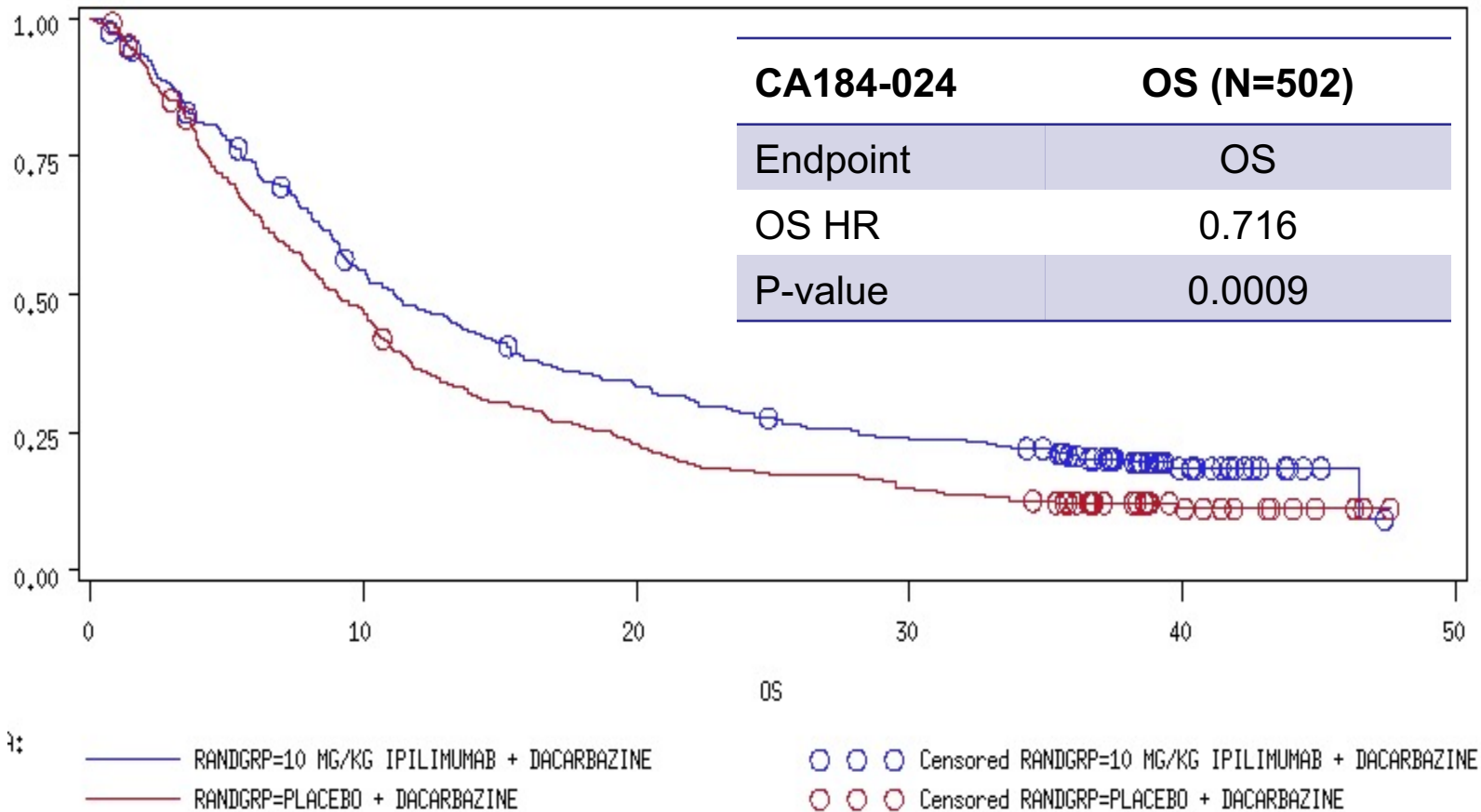
Rationale

- Unique characteristics of immune checkpoint modulators
 - Survival probability (long term survival)
 - Delayed clinical effect
- Key challenges of log-rank analysis as sole characterization of overall survival
 - Does not capture key attribute of survival probability (or long term survival)
 - Time to final analysis may continue to lengthen based on kinetics of survival effect

Milestone OS Analysis

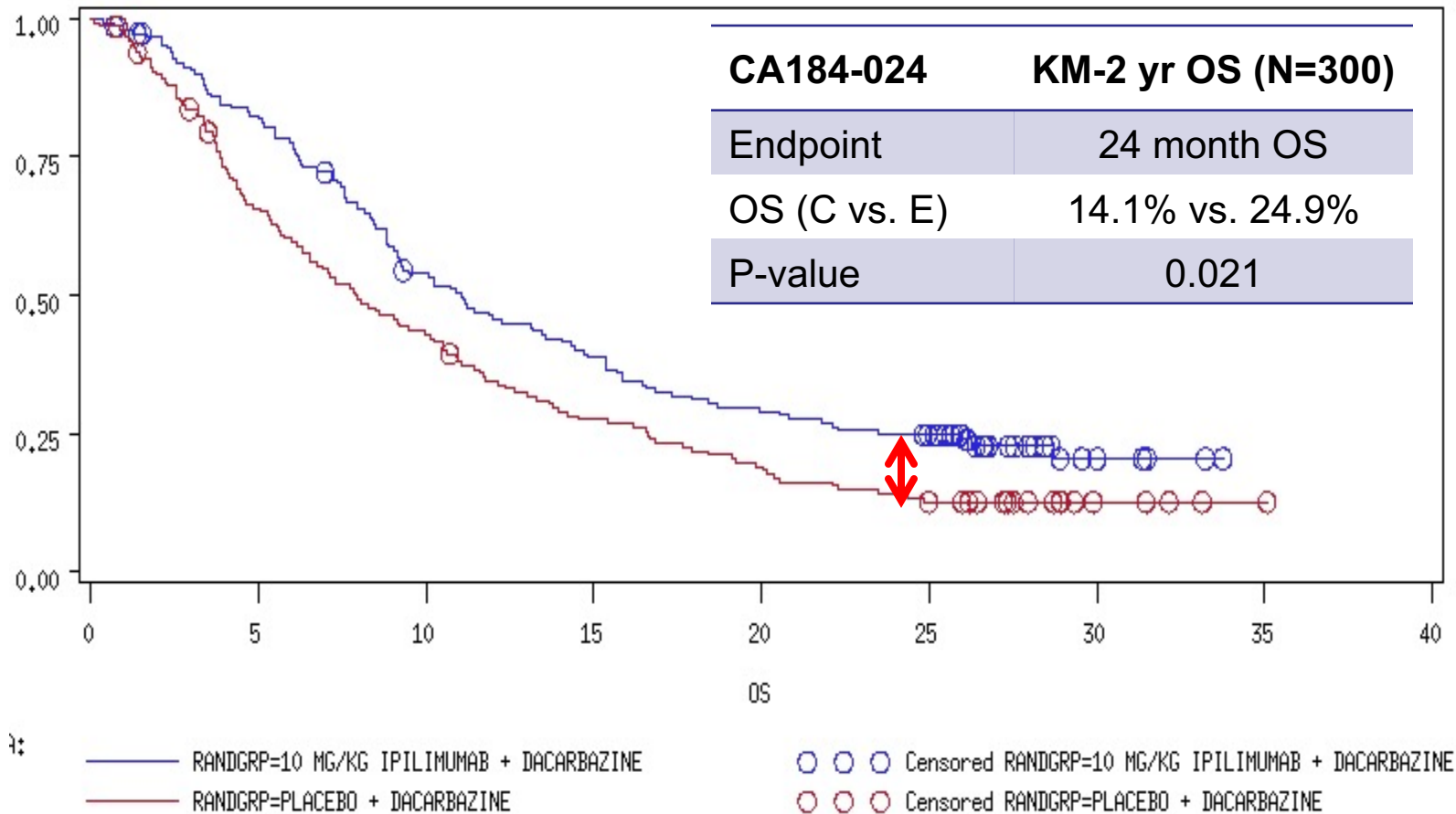
- Milestone survival is defined as the Kaplan-Meier survival probability at a pre-specified milestone, e.g., 2 years
- Study design and analysis consideration
 - Primary endpoint: overall survival
 - Intermediate endpoint: milestone survival probability
 - Population includes patients with a minimal follow-up duration, i.e., \geq milestone duration
 - Hierarchical testing procedure

Example*: Ipilimumab+DTIC vs. DTIC Final OS Analysis



* Roberts, C. et al. NEJM, 2011, 364: 2517-2526.

Example*: Ipilimumab+DTIC vs. DTIC Intermediate 2-year Milestone OS Analysis



* Roberts, C. et al. NEJM, 2011, 364: 2517-2526.

Pros and Cons

- Pros
 - Potential earlier assessment of benefit/risk
 - Greater statistical power when delayed treatment effect is present
 - Direct characterization of survival probability (long term survival effect)
 - Predictable timing of analysis
 - Both intermediate and final endpoints are overall survival
- Cons
 - Challenge in maintaining study integrity post milestone analysis, i.e., unblinding prior to final OS analysis
 - Does not account for the totality of OS data
 - Only appropriate for a registration trial when prior data enable an understanding of appropriate milestone time point selection

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Steven Rosenberg, M.D., Ph.D.

National Cancer Institute

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Amy McKee, M.D.

FDA

Center for Drug Evaluation and Research



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Celia Witten, M.D., Ph.D.

FDA

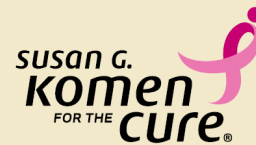
Center for Biologics Evaluation and Research



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