

B ENGELBERG CENTER for Health Care Reform at BROOKINGS

# 2013 Conference on Clinical Cancer Research

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



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### **Immune Checkpoint Modulators**

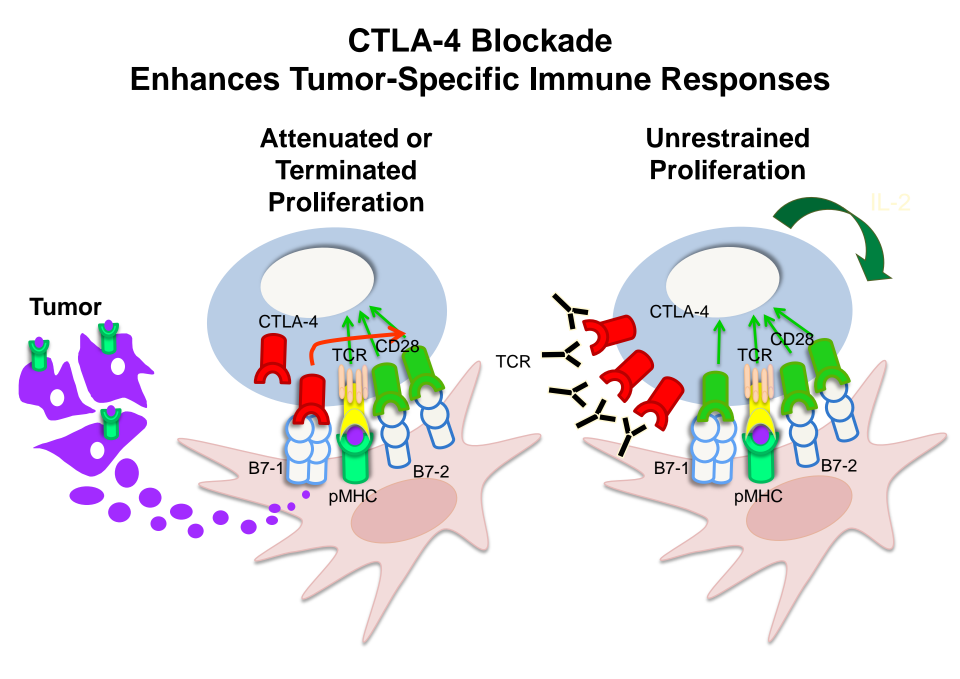
Jim Allison, Ph.D. The University of Texas MD Anderson Cancer Center

# Why immunotherapy?

Specificity

## Memory

## Adaptability



## **Evolution of Response: Patient Example**

#### Screening





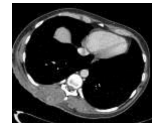
Week 12 Initial increase in total tumor burden (mWHO PD)



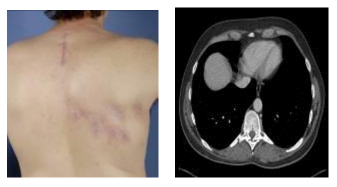


#### Week 16 Responding

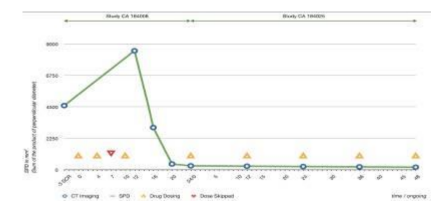




#### Week 72 Durable & ongoing response without signs of IRAEs

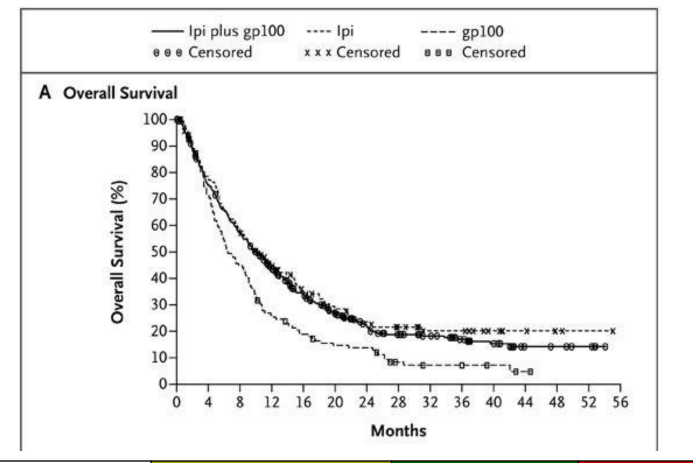


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#### Harmankaya

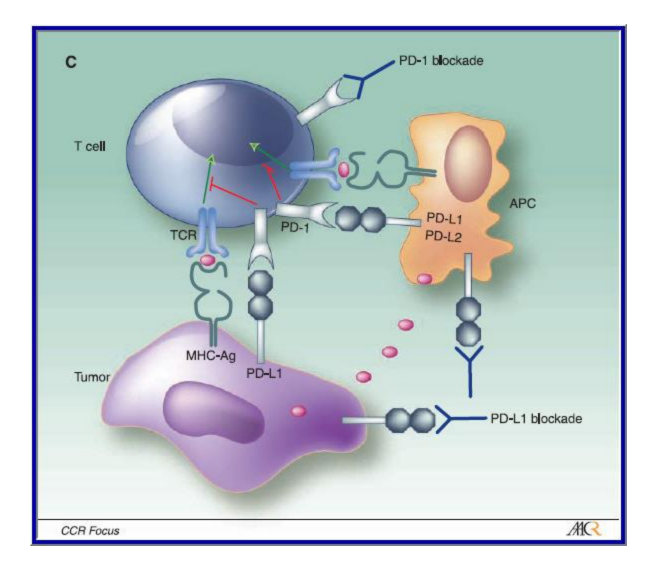
# **Kaplan-Meier Analysis of Survival**



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

#### Hodi et al. NEJM 2010

### **Programmed Death 1**



http://www.melanoma.org/community/mpip-melanoma-patients-information-page/video-how-anti-pd-1-therapy-works-imumne-system

## Anti – PD-1 (BMS-936558)

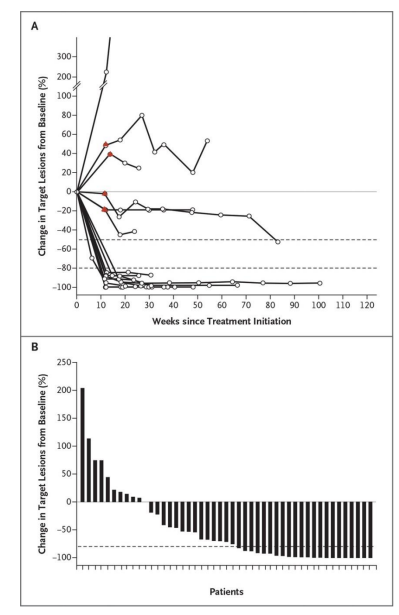
296 Patients with Metastatic Cancer 1, 3, 10 mg/kg, MTD not reached

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

Clinical Activity: Melamona (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* 

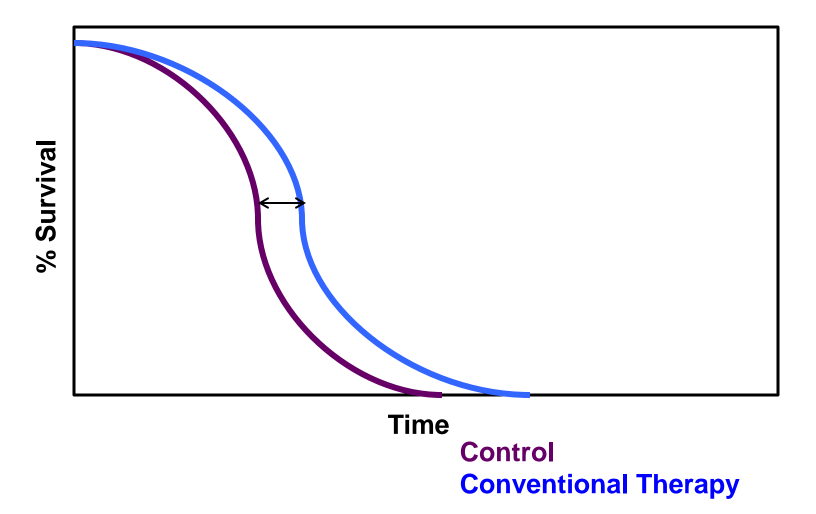
Topalian ASCO, NEJM 2012

### Clinical Activity in Melanoma Patients Receiving Ipilimumab (αCTLA-4) and Nivolumab (αPD-1)

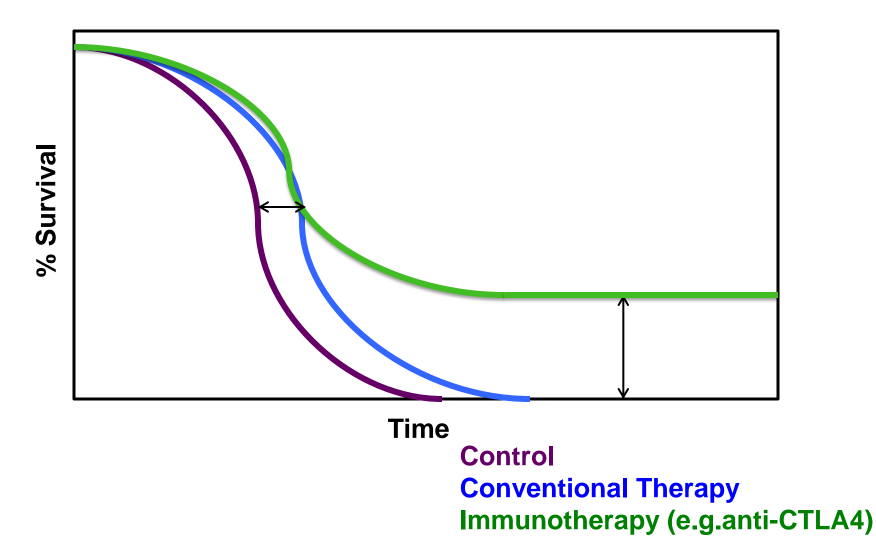


ASCO 2013 NEJM 6/2/2013

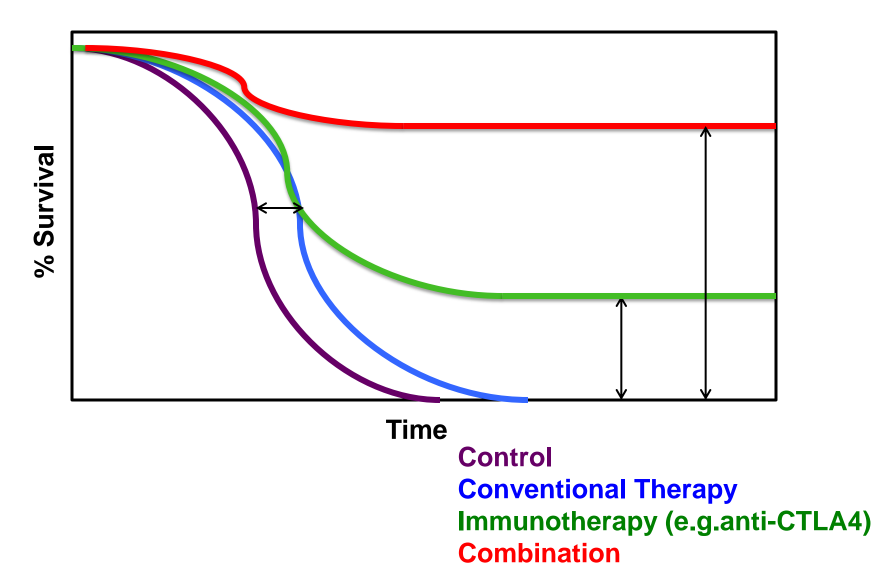
## **Improving Survival with Combination Therapy**



## **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 efficay
- 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

# <u>Challenge:</u> 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	Non-Commercial License (http://creativecommons.org/licenses/by-nc/2.5), which permits unrestricted non-commercial use, distribution, and reproduction in any medium, provided the original work is properly cited.	
the Cancer Vaccine Consortium (CVC)	Improved Endpoints for Cancer Immunotherapy Trials	
Lothar H. Finke <sup>a,g,*</sup> , Kerry Wentworth <sup>b,g</sup> , Brent Blumenstein <sup>c</sup> , Natalie S. Rudolph <sup>d</sup> , Hyam Levitsky <sup>e,g</sup> , Axel Hoos <sup>f,g</sup>	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	

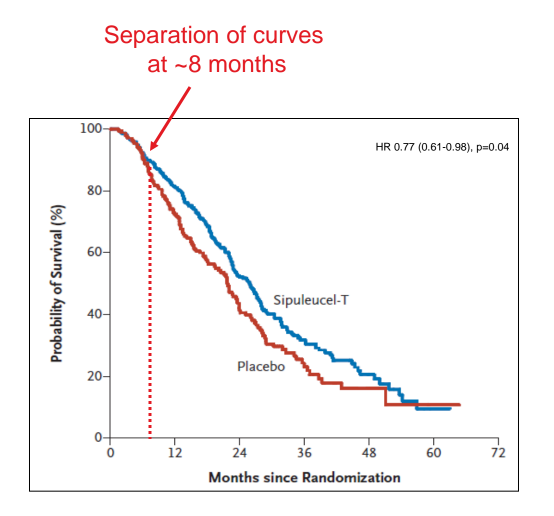


## Survival: Conventional Design Assumptions



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

# **Delayed Separation – Sipuleucel-T**



Sponsor: Dendreon

#### Agent:

autologous dendritic cell vaccine

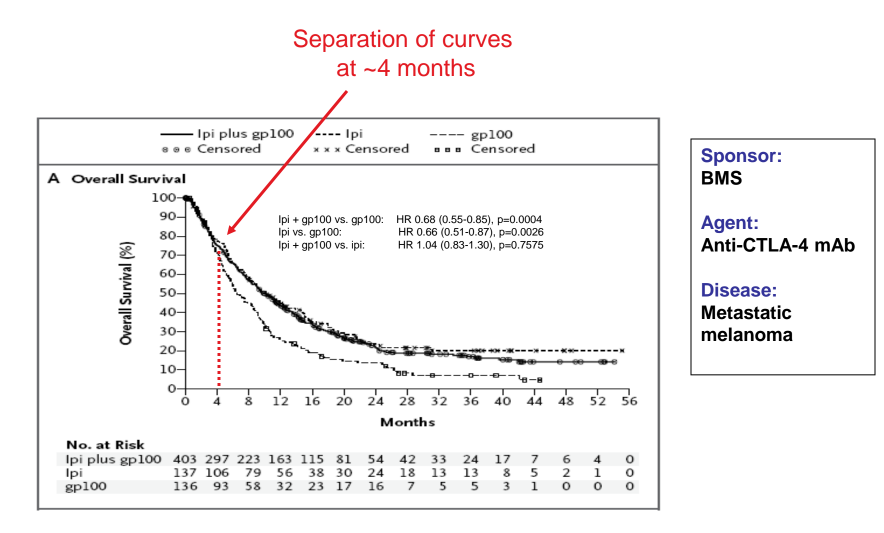
#### **Disease:**

hormone-refractory prostate cancer



#### Kantoff et al., New Engl. J. Med. 2010

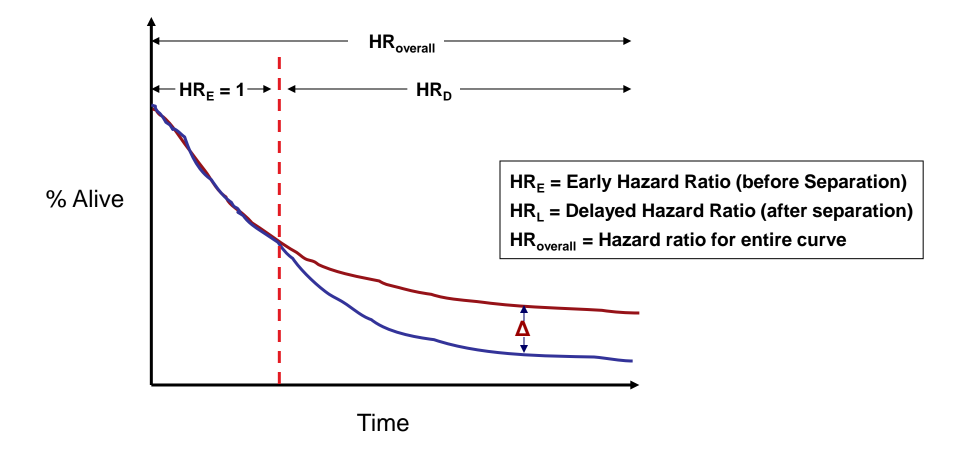
# **Delayed Separation – Ipilimumab**



#### Hodi et al., New Engl. J. Med. 2010

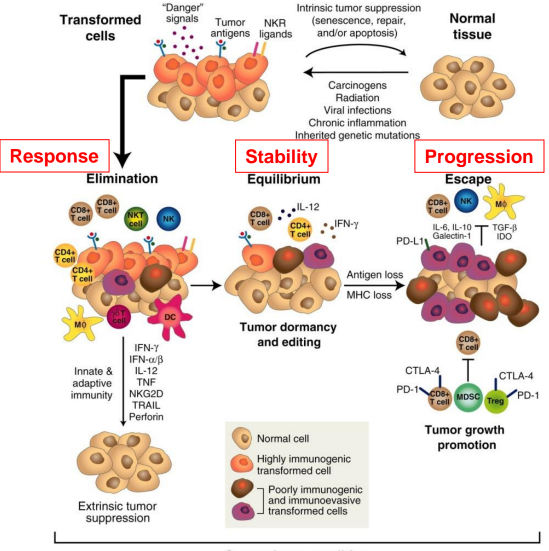


### Implications of Delayed Separation of Curves - Model Scenario -



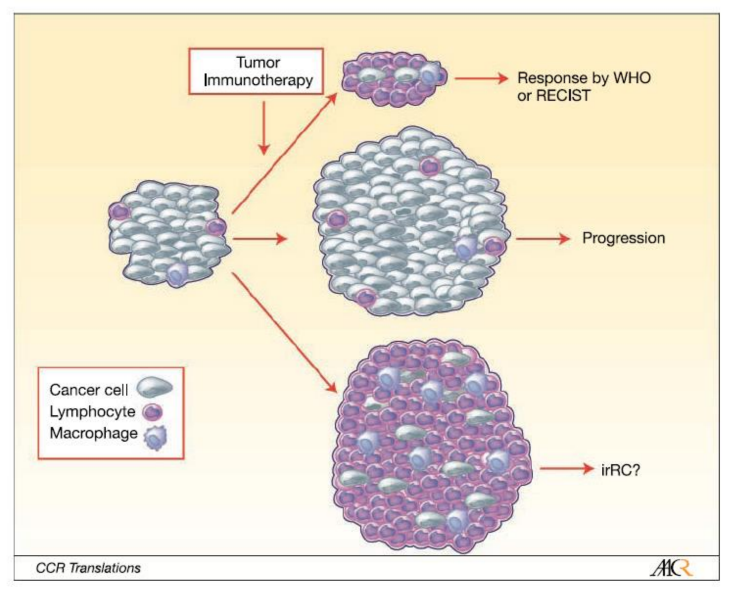
• Large  $\Delta$  after separation needed to compensate for no effect before separation

### Interactions between Immune System and Tumor



**Cancer Immunoediting** 

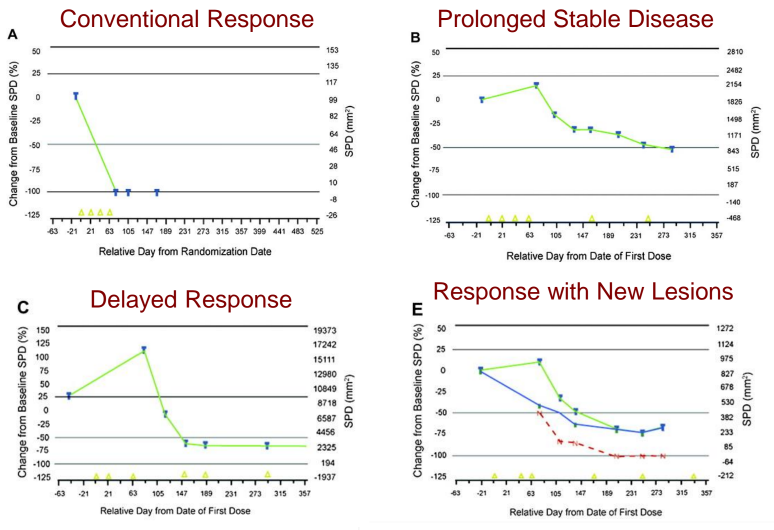
#### **Tumor Volume Increase Due to Lymphocyte Infiltration**





Ribas et al., Clin Cancer Res 2009; 15:7116–8

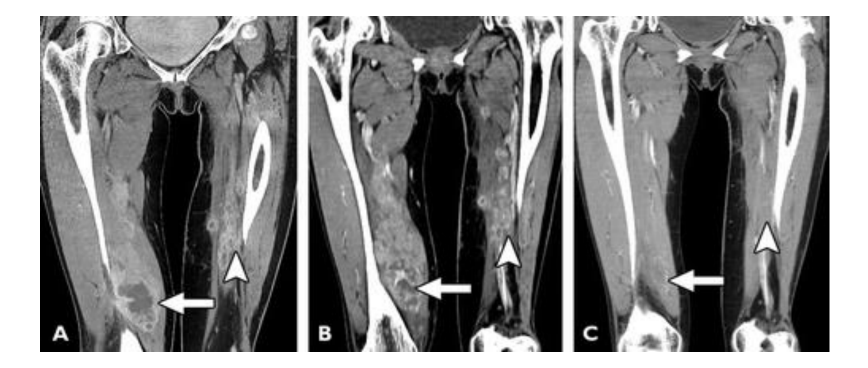
## **Immunotherapy Patterns of Response**



Wolchok JD, et al. Clin Cancer Res. 2009;15:7412-7420

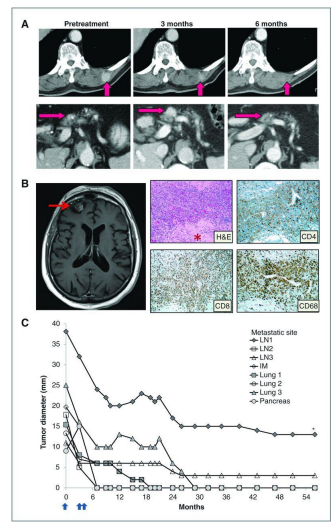
Hoos A et al. JNCI J Natl Cancer Inst 2010;102:1388-1397

### **Anti-CTLA-4 (Ipilimumab): Delayed Response**



O'Regan, KN, et al. AJR 2011

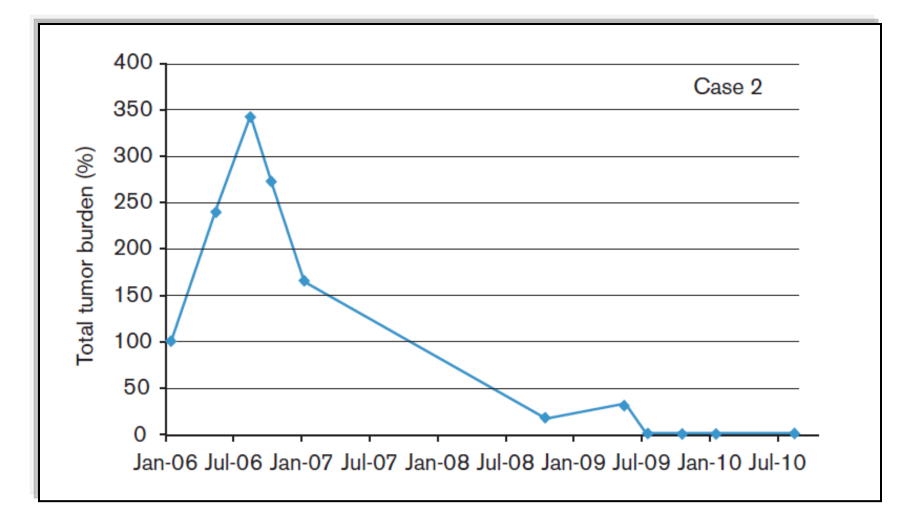
#### Regression of metastatic RCC following anti-PD-1 therapy, with "immune-related" response characteristics.



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

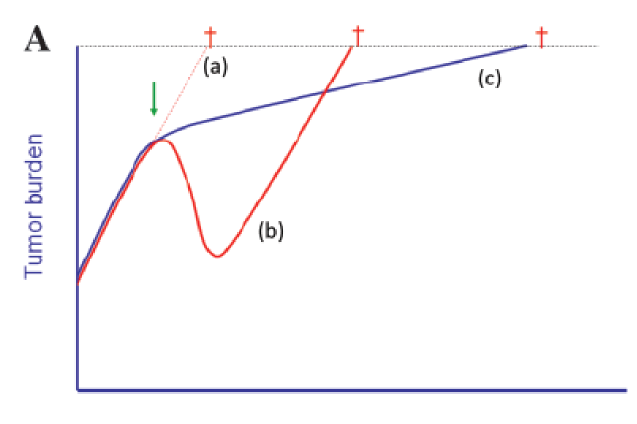


### Autologous DC + IFN α2b in Advanced Melanoma: Delayed Response



Wilgenhof S. et al., Melanoma Res. 2011

### Tumor Growth Rate: Potential Impact on Survival



#### Time

# **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

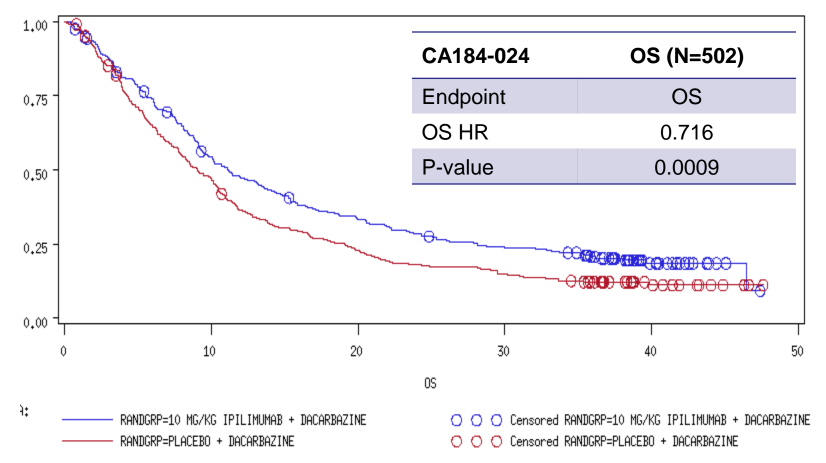
## Rationale

- 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**

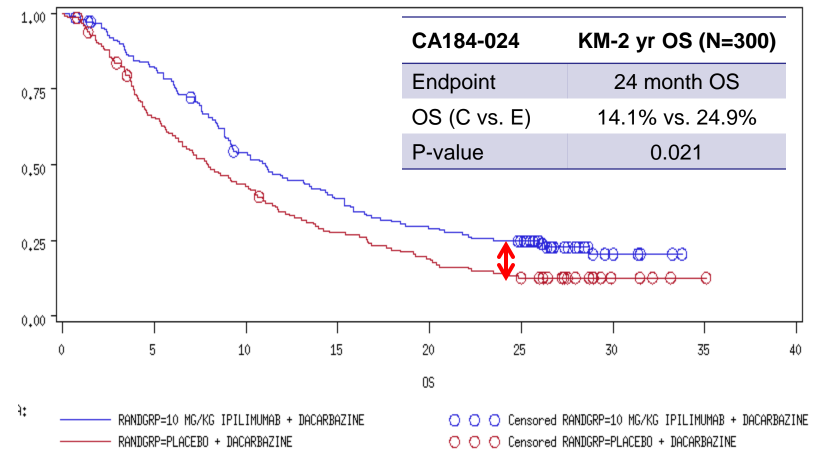
- Milestone survival is defined as the Kaplan-Meier survival probability at a pre-specified milestone, e.g., 2 years
- Study design and analysis consideration
  - Primary endpoint: overall survival
  - Intermediate endpoint: milestone survival probability
  - Population includes patients with a minimal follow-up duration,
    i.e., ≥ 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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# 2013 Conference on Clinical Cancer Research

### 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