

# **Interim Results of an Ongoing Phase 1, Dose Escalation Study of MGA271 (Enoblituzumab), an Fc-optimized Humanized Anti-B7-H3 Monoclonal Antibody, in Patients with Advanced Solid Cancer**

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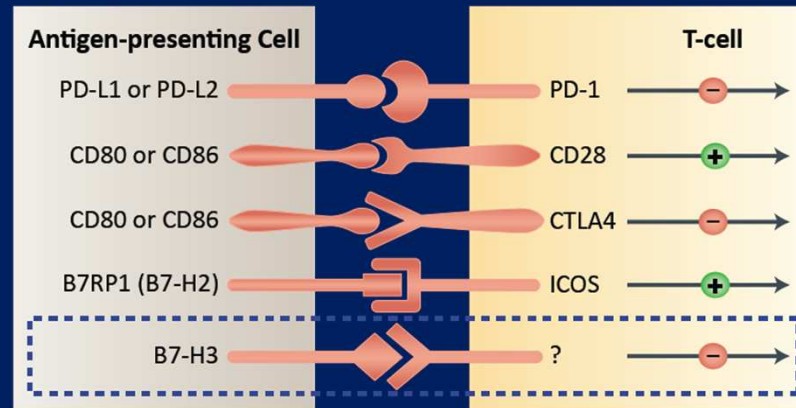
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  - Merck
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## B7-H3 (CD276): Member of B7 Family of Immune Regulators



Adapted from Pardoll, et al., Nature, April 2012.

### Immunosuppressive Role

- Expression on lung cancer cells and macrophages suppresses T-cell mediated anti-tumor immune response (*Chen 2013*)
- B7-H3-positive myeloid-derived suppressor cells found in tumor microenvironment (*Zhang 2015*)
- Crystal structure resolved: T-cell inhibitory domain mapped (*Vigdorovich 2013*)

### Tumor Invasion and Metastatic Role

- Silencing reduces migration and invasion of melanoma and breast cancer cell lines (*Chen 2008*)
- Enhances metastatic potential of melanoma cells (*Tekle 2012*)

# B7-H3: Tissue Expression and Prognosis

		IHC Summary of Samples Screened		
		B7-H3 Positive		2+ or Above
Fixed Tumor MicroArray				
Lead Potential Indications:				
Head and Neck	19/19	100%	19/19	100%
Kidney Cancer	77 / 78	99%	75 / 78	96%
Lung Cancer	226/272	83%	211/272	78%
Breast Cancer	119/164	73%	115/164	70%
Prostate Cancer	88/99	89%	51/99	52%
Melanoma	66/70	94%	32/70	46%
Bladder	14/20	70%	9/20	45%
Other Potential Indications:				
Glioblastoma	65/66	98%	63/66	95%
Thyroid Cancer	34/35	97%	33/35	94%
Mesothelioma	41/44	93%	39/44	89%
Pancreas Cancer	69/78	88%	45/78	58%
Ovarian Cancer	59/79	75%	36/79	46%

## B7-H3 Tissue Expression

- High level expression in a broad range of tumors
- Minimal expression on normal tissue
- Expressed on tumor neo-vasculature
- Correlation of high expression with advanced disease, presence of metastases and poor outcome

## Timeline of selected B7-H3 articles in peer-reviewed publications

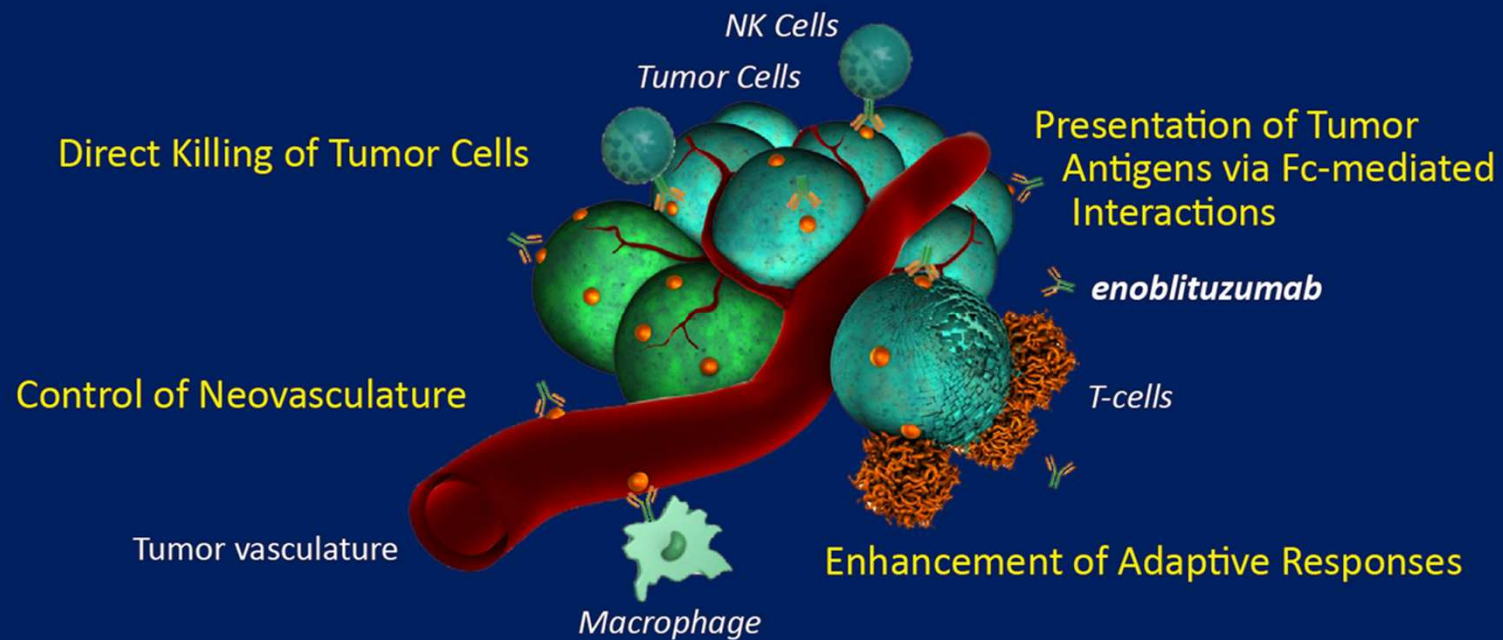


## Enoblituzumab (MGA271, Anti-B7-H3 Antibody)

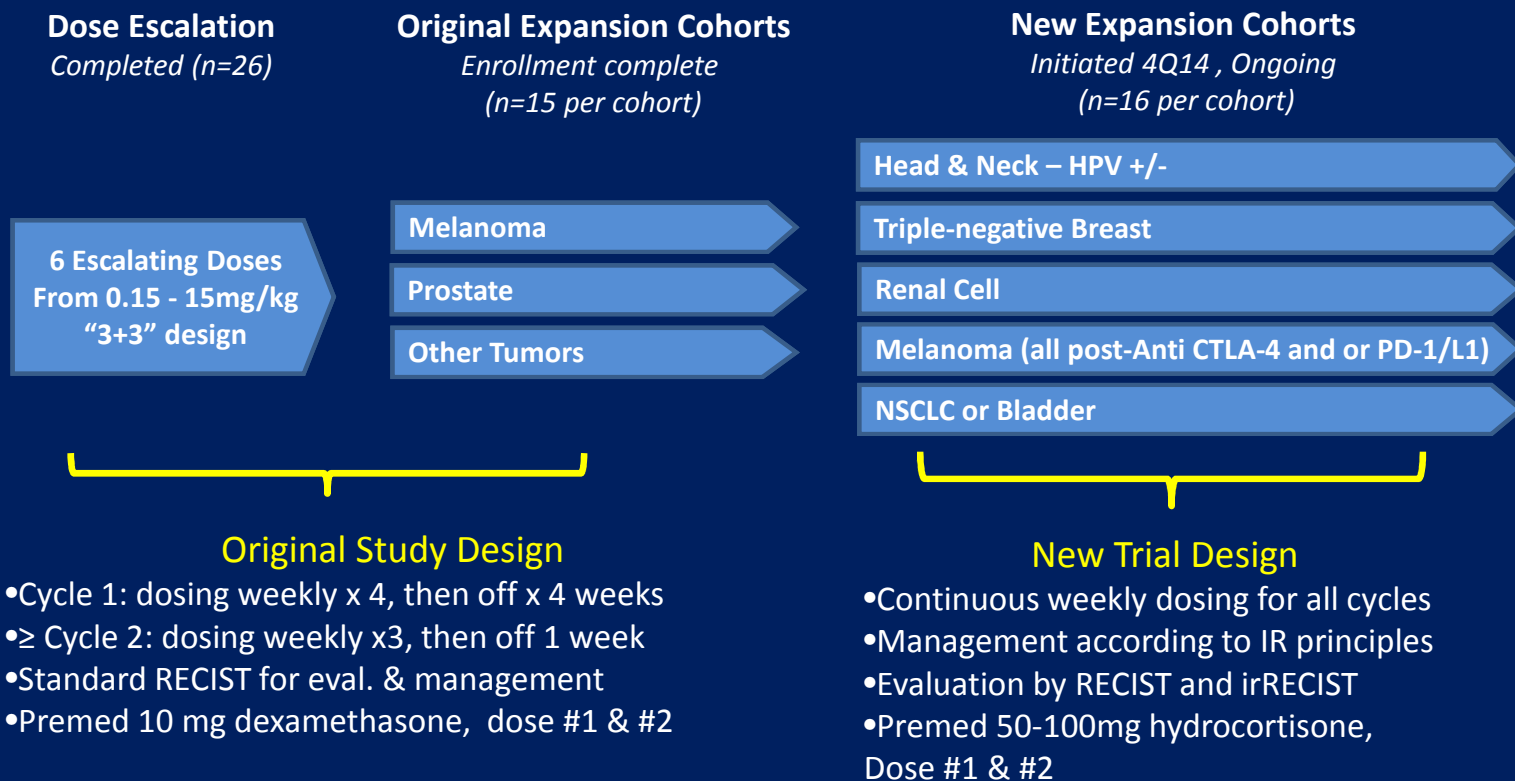
- Humanized IgG1 monoclonal antibody recognizing human B7-H3 with high affinity ( $KD \approx 7$  nM)
- Terminal Half Life  $\approx 3$  weeks
- Fc-optimized via mutation to enhance effector function (e.g., ADCC)
  - Increased affinity for activating Fc $\gamma$  receptor (Fc $\gamma$ RII, CD16A)
  - Decreased affinity for the inhibitory Fc $\gamma$  receptor (Fc $\gamma$ RIIB, CD32B)
- Once-weekly intravenous dosing
- Currently in clinical trials as monotherapy (described today) and in combination with checkpoint inhibitors including pembrolizumab and ipilimumab (see SITC Trials-In-Progress Poster Session)

# Enoblituzumab

## Potential Mechanisms of Action



# Study Design: Ongoing Phase 1 Dose Escalation and Cohort Expansion



# Study Objectives

- **Primary Objective**

- Describe safety profile of enoblituzumab in patients with advanced cancer that expresses B7-H3 in tumor and/or tumor-associated vasculature

- **Secondary Objectives**

- Determine Maximum Tolerated Dose or Maximum Administered Dose of enoblituzumab
- Evaluate preliminary anti-tumor activity of enoblituzumab
- Determine enoblituzumab pharmacokinetics/pharmacodynamics

- **Exploratory Objectives**

- Evaluate and assess IHC diagnostic test for B7-H3 expression on tumor cells and tumor vasculature



# Key Inclusion/Exclusion Criteria

## Inclusion

- B7-H3 expression on tumor cells or tumor vasculature
  - $\geq 10\%$  of tumor cells with 2 or 3+ IHC\* staining or  $\geq 25\%$  of tumor vasculature having 2 or 3+ IHC staining
- Progressive disease during or following last treatment regimen
  - Up to 4 to 5 prior treatments allowed depending on tumor type
- Prior checkpoint inhibitor therapy allowed (mandated for melanoma)
- ECOG Performance Status  $\leq 1$
- Measurable disease by RECIST 1.1
  - Prostate cancer required measurable disease in new trial design
- Completed systemic anticancer therapy  $\geq 28$  days prior to enrollment

## Exclusion

- $\geq$  Grade 3 autoimmune toxicity with prior immune checkpoint inhibitor
- Concurrent systemic steroids  $>10$  mg/day of oral prednisone/equivalent
- Active brain metastases

\*IHC: Immunohistochemistry with B7-H3 cell surface staining

## Baseline Characteristics

Baseline Characteristics	Escalation n=26	Original Expansion n= 48	Additional Expansion n= 42	Total n= 116
Median age, (range), years	62 (42-77)	64 (26-88)	67 (24-83)	63 (24-88)
Male, no. (%)	17 (65)	33 (69)	28 (67)	78 (67)
Prior Cancer Therapy				
Median no. (range): Chemo and Immunotherapy	2 (1-5)	3 (0-8)	3 (0-5)	3 (0-8)
Prior Chemotherapy, no. (%)	21 (81)	34 (71)	37 (88)	92 (79)
Prior Immunotherapy, no. (%)	6 (23)	18 (38)	7 (17)	31 (27)
ECOG Performance Status, no. (%)				
0	16 (62)	20 (42)	10 (24)	46 (40)
1	10 (38)	28 (58)	32 (76)	70 (60)

Data as of September 21, 2015

# Enoblituzumab-Related Adverse Events

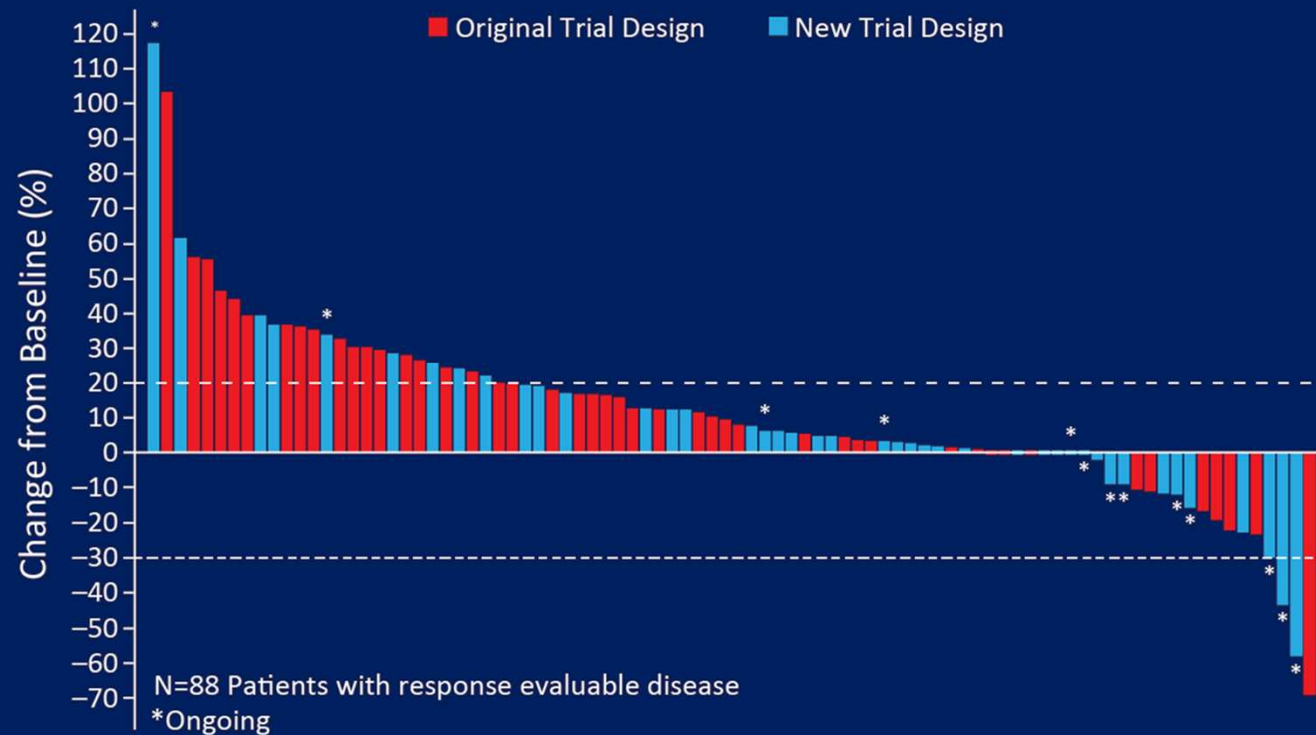
- Acceptable safety profile
- No drug-related treatment discontinuation
- Mild-moderate infusion reactions readily managed with conventional supportive care including corticosteroids, decreased infusion rate

Drug-Related Adverse Event ≥10% of Patients	No. (%) of Patients			
	All Grades		Grades 3-4	
	Total Population (N=116)	New Study Design* (N=55)	Total Population (N=116)	New Study Design* (N=55)
<b>Any adverse event</b>	<b>86 (74)</b>	<b>42 (76)</b>	<b>5 (4)</b>	<b>3(5)</b>
<b>Infusion related reaction/ cytokine release syndrome</b>	<b>39(34)</b>	<b>24 (44)</b>	<b>1(1)</b>	<b>1(2)</b>
<b>Fatigue</b>	<b>37 (32)</b>	<b>15 (27)</b>	<b>0</b>	<b>0</b>
<b>Nausea</b>	<b>22 (19)</b>	<b>14 (25)</b>	<b>0</b>	<b>0</b>
<b>Vomiting</b>	<b>15 (13)</b>	<b>10(18)</b>	<b>0</b>	<b>0</b>

\*New study design is continuous, uninterrupted weekly infusion of enoblituzumab with reduced steroid pre-med

# Best Change in Target Lesion Size

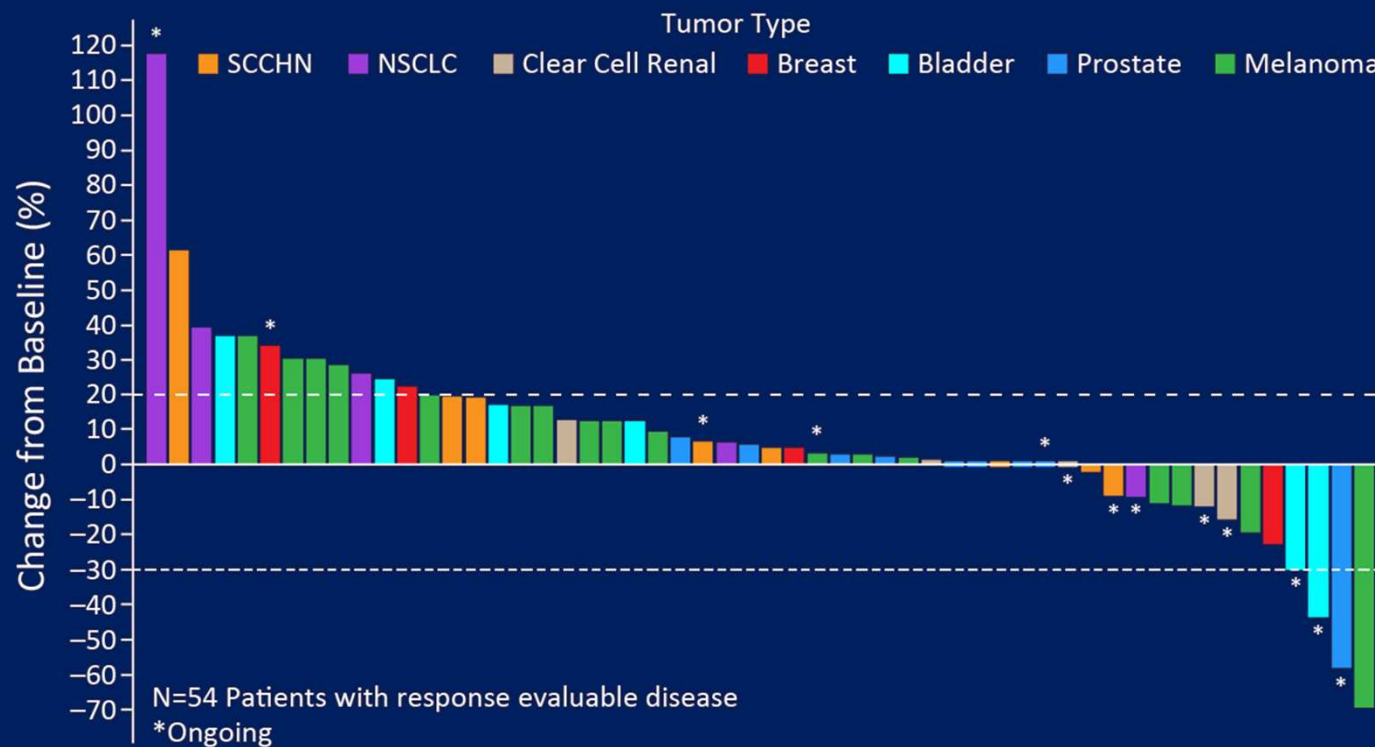
All Response Evaluable Patients: Escalation and Expansion



- Tumor regression at multiple dose levels (0.15mg/kg – 15mg/kg)
- Enrollment continues under new trial design:  $\approx$  half of planned patients enrolled

# Best Change in Target Lesion Size

Response Evaluable, Tumor-Specific Expansion Cohorts: 15 mg/kg  
Cohorts: Melanoma, Prostate, TNBC, SCCHN, NSCLC, Bladder, RCC

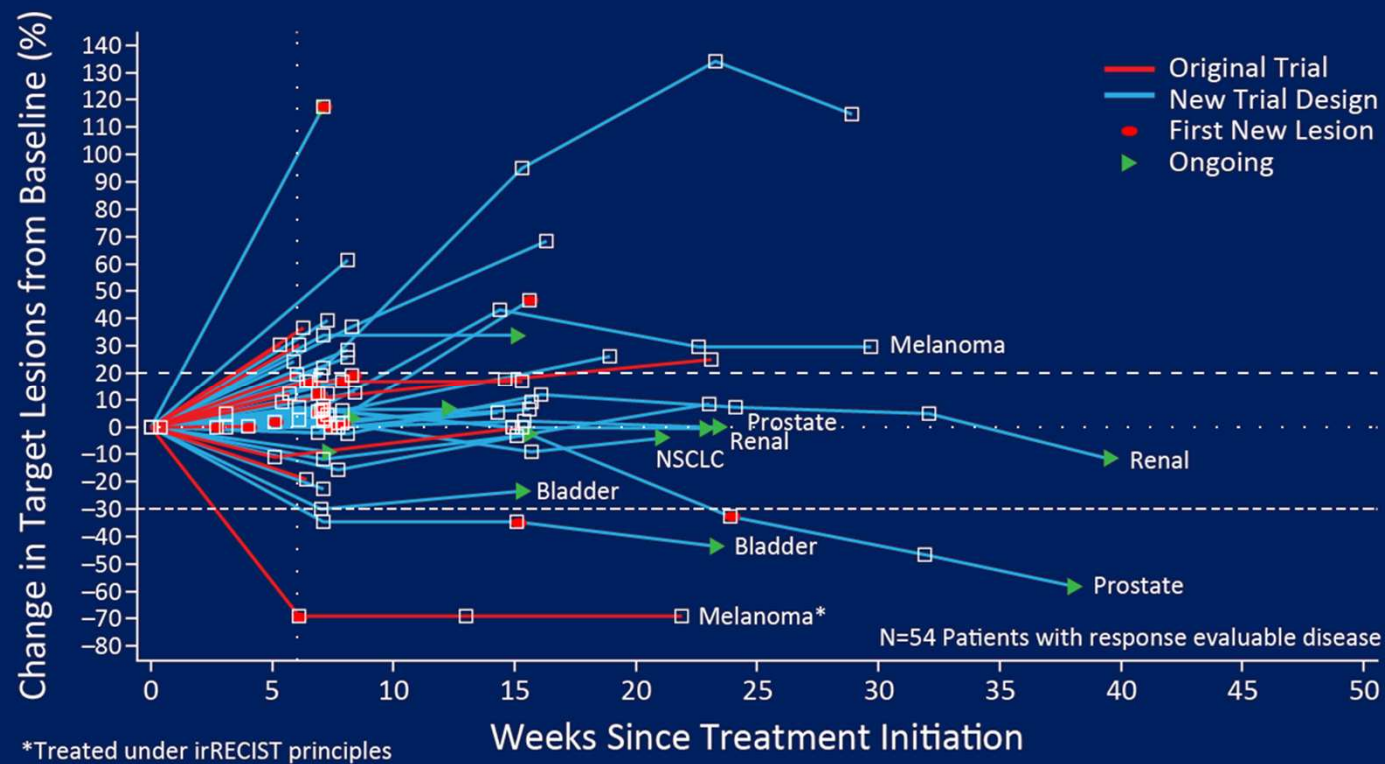


- Tumor regression observed in each disease cohort

Data as of September 21, 2015

# Change in Target Lesion Size Over Time

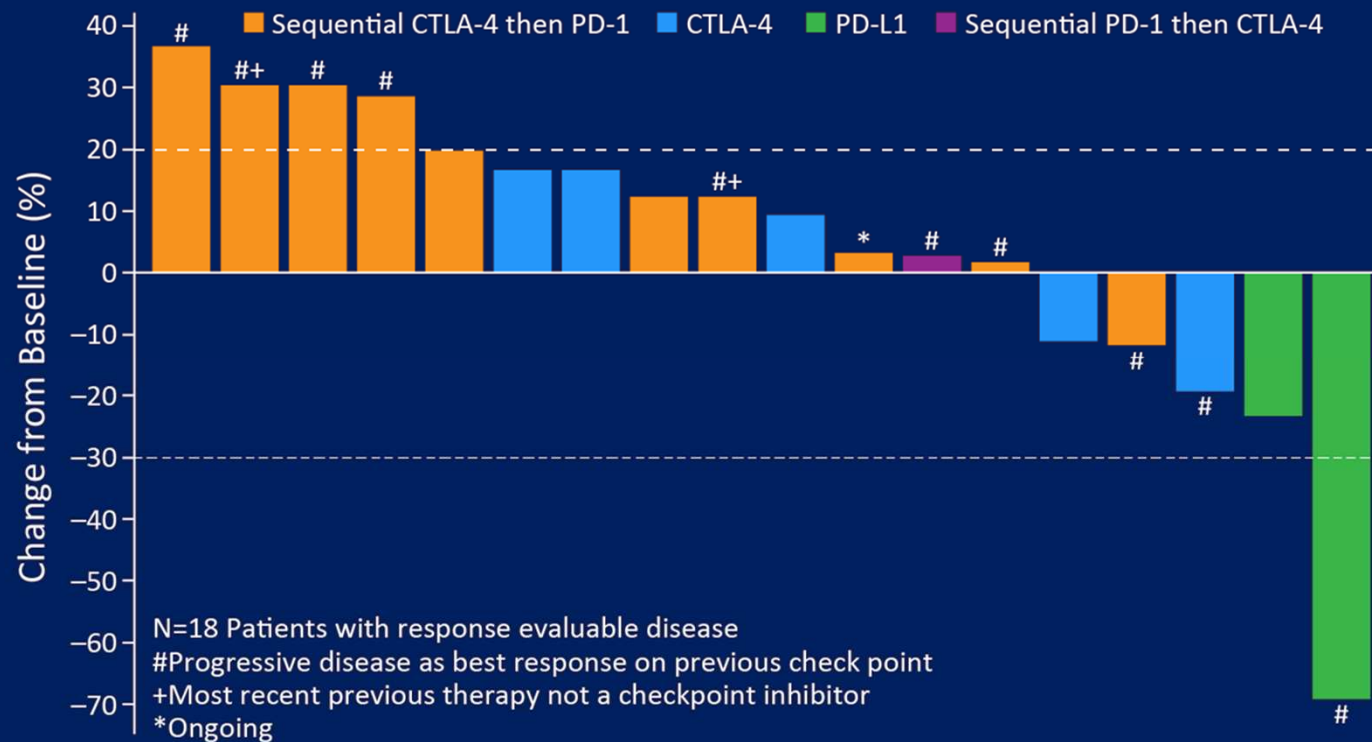
Response-Evaluable Tumor-Specific Expansion Cohorts: 15 mg/kg  
Cohorts: Melanoma, Prostate, TNBC, SCCHN, NSCLC, Bladder, RCC



Data as of September 21, 2015

# Best Change in Target Lesion Size: Melanoma

All patients are post-checkpoint inhibitor



All but one patient treated 15mg/kg enoblituzumab

Data as of September 21, 2015

# Metastatic Melanoma

73-year-old man previously progressed on Anti-PD-L1 And Trametinib



- Near complete regression of ulcerated 4 cm tumor in groin
- Regression of small pulmonary nodules on CT

Courtesy of Dr. Chmielowski at UCLA Jonsson  
Comprehensive Cancer Center

Data as of September 21, 2015

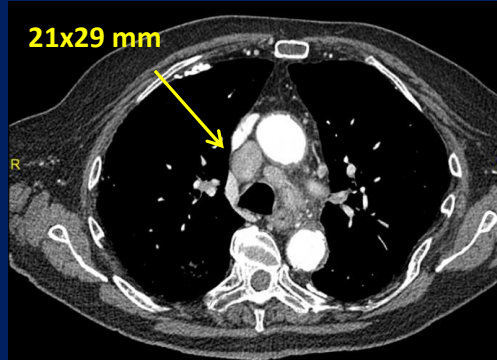


# Metastatic Prostate Cancer

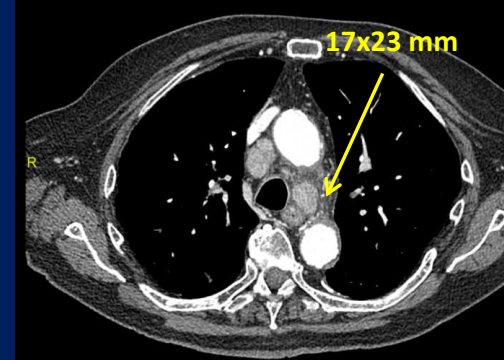
87-year-old man

Pre-Treatment  
Baseline

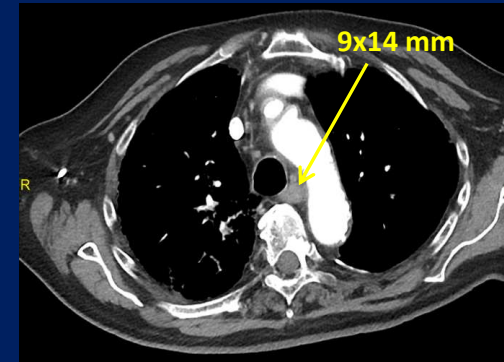
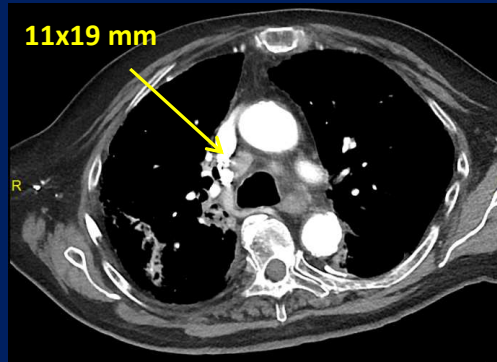
Right Paratracheal Lymph Node



Left Paratracheal Lymph Node



Day 287  
34 Doses  
enoblituzumab  
(15mg/kg)



Patient remains on therapy after 11 months of treatment

Courtesy of Dr. Chmielowski at UCLA Jonsson  
Comprehensive Cancer Center

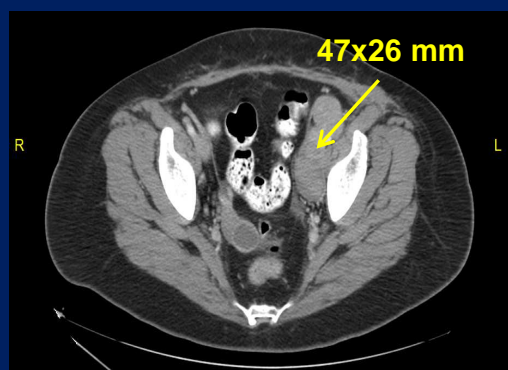
Data as of September 21, 2015

# Vitiligo in Melanoma Patient with Progression on Prior Therapy with Checkpoint Inhibitors

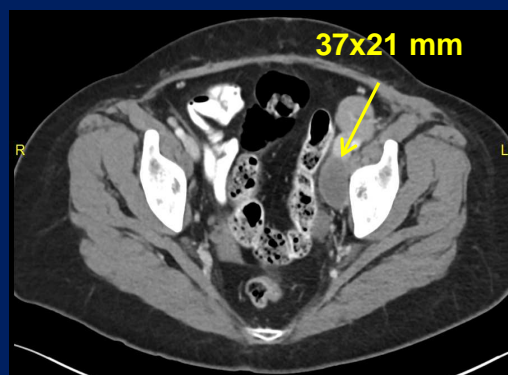
52-year-old woman previously progressed on anti-CTLA-4 and anti PD-1

Pre-Treatment  
Baseline

Left Ext Iliac Lymph Node #1



Day 58  
8 Doses  
enoblituzumab  
(15mg/kg)



Development of Vitiligo (Post-enoblituzumab)



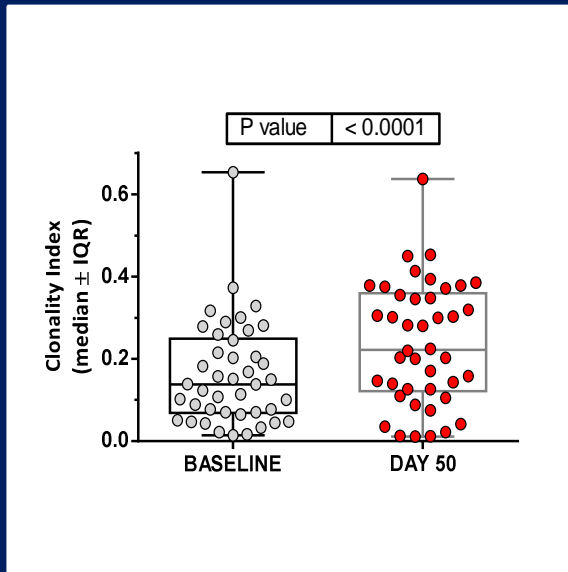
Data as of September 21, 2015

Courtesy of Dr. Chmielowski's patient at UCLA  
Jonsson Comprehensive Cancer Center

# Increase in T-Cell Receptor Repertoire Clonality Following Enoblituzumab

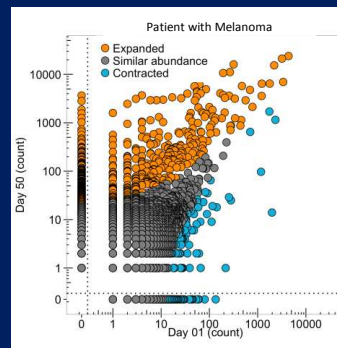
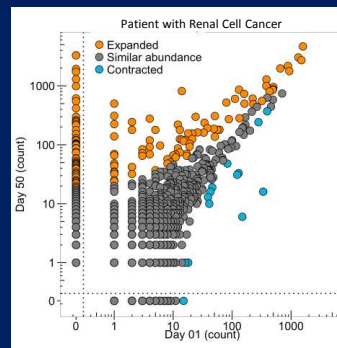
## Evaluation of T-Cell Clonality in the Peripheral Blood

### Population Clonality

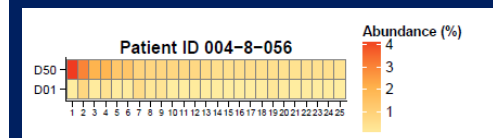
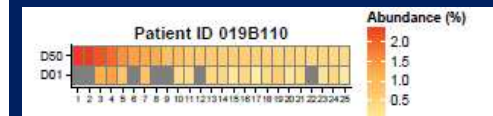


Baseline (Day 1) v D50 Post-treatment  
(42 patients)

### Clonality: 2 Patients with Tumor Shrinkage



### Top 25 Clones at Day 50 Comparison to Baseline



## Conclusions from Ongoing Enoblituzumab CP-MGA271-01 Study

- Manageable and tolerable safety profile
  - No treatment related discontinuation
  - No severe immune mediated toxicity
- Preliminary anti-tumor activity in broad range of tumors
  - Post check-point inhibitor failure melanoma
  - New study design: management principles used in immune oncology
- Initial demonstration of T-cell modulation with enoblituzumab
- Interim results:
  - **Support continued evaluation of enoblituzumab monotherapy**
  - **Support evaluation of enoblituzumab in combination with check-point inhibitors: anti PD-1 and anti CTLA-4**

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