

Immunotherapy for the Treatment of Genitourinary Malignancies

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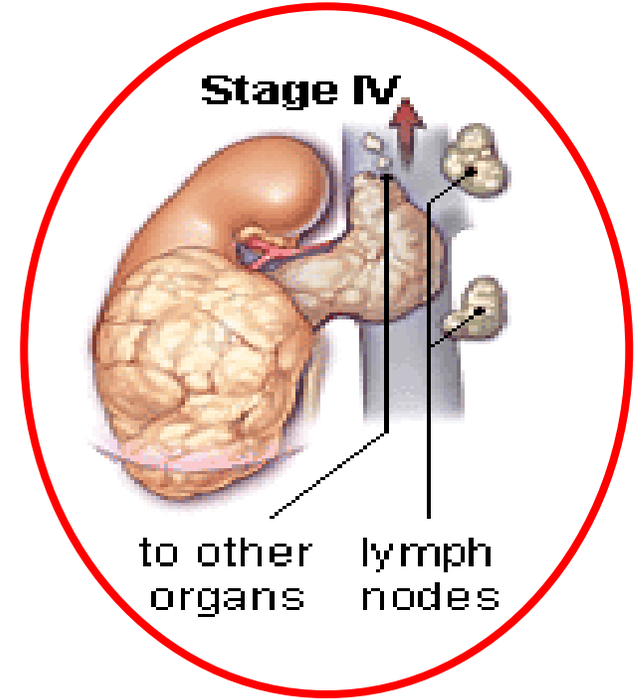
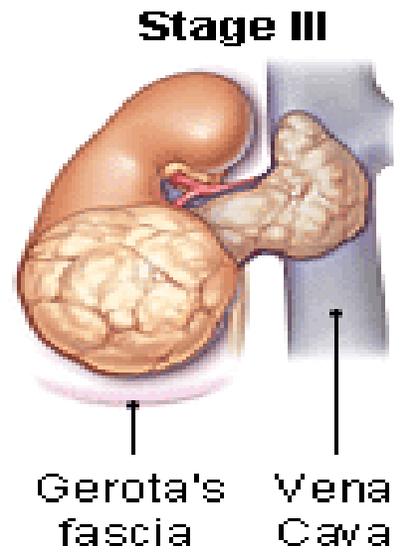
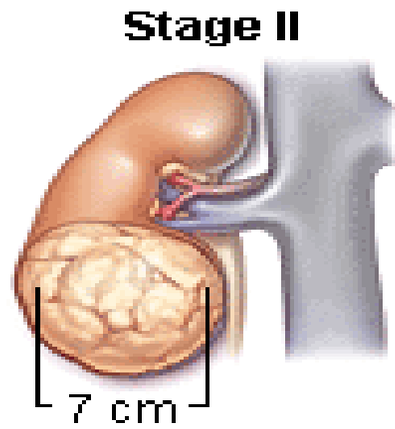
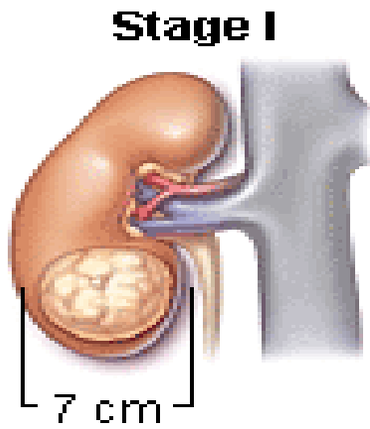
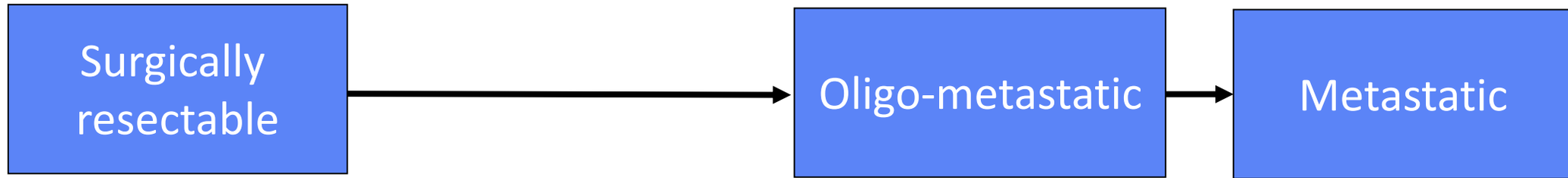
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Disclosures

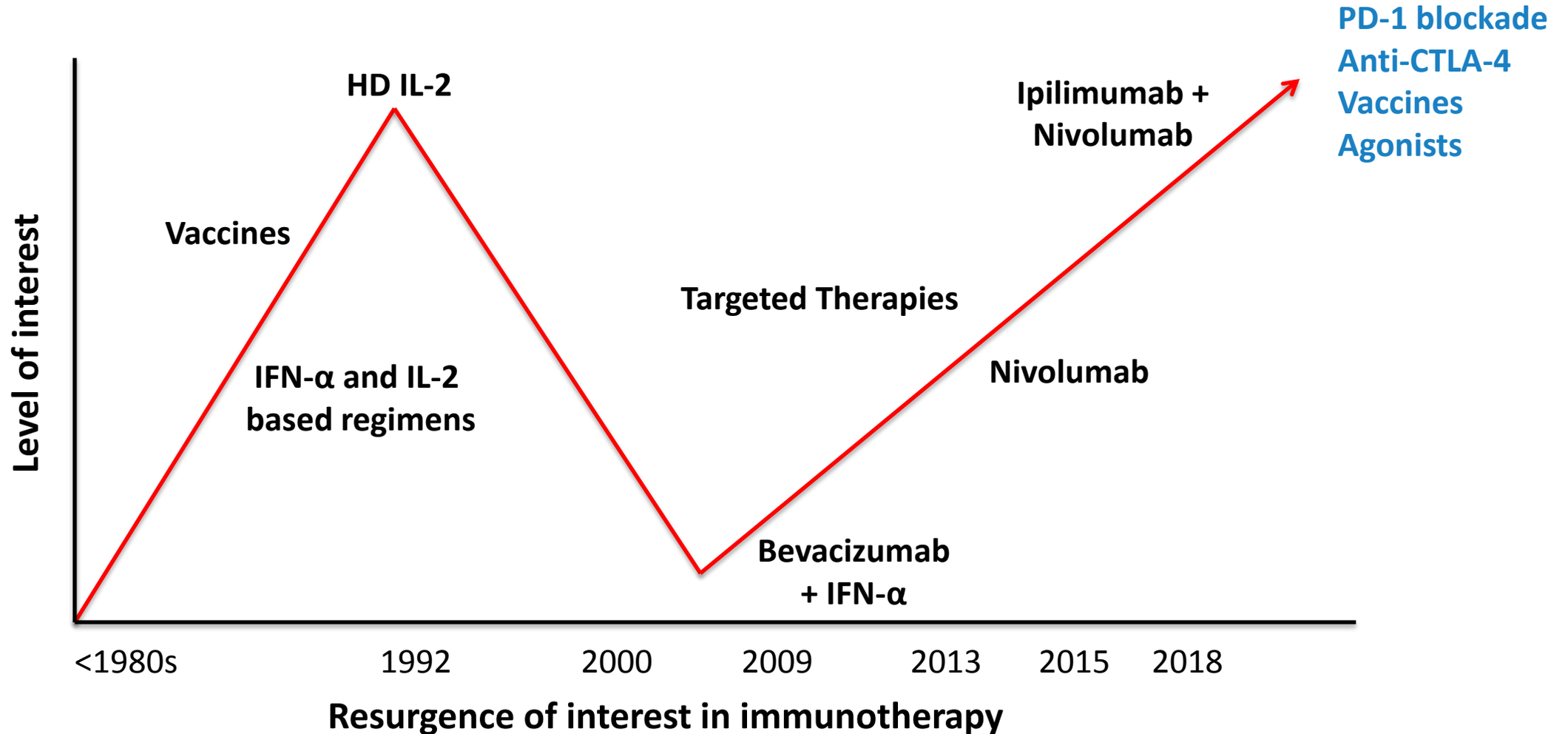
- Astellas pharmaceuticals, investigator funding
- I will not be discussing non-FDA approved indications during my presentation.

Immunotherapy for Metastatic Kidney Cancer (Renal Cell Carcinoma; RCC)



reemakeup.blogspot.com

History of Immunotherapy in mRCC



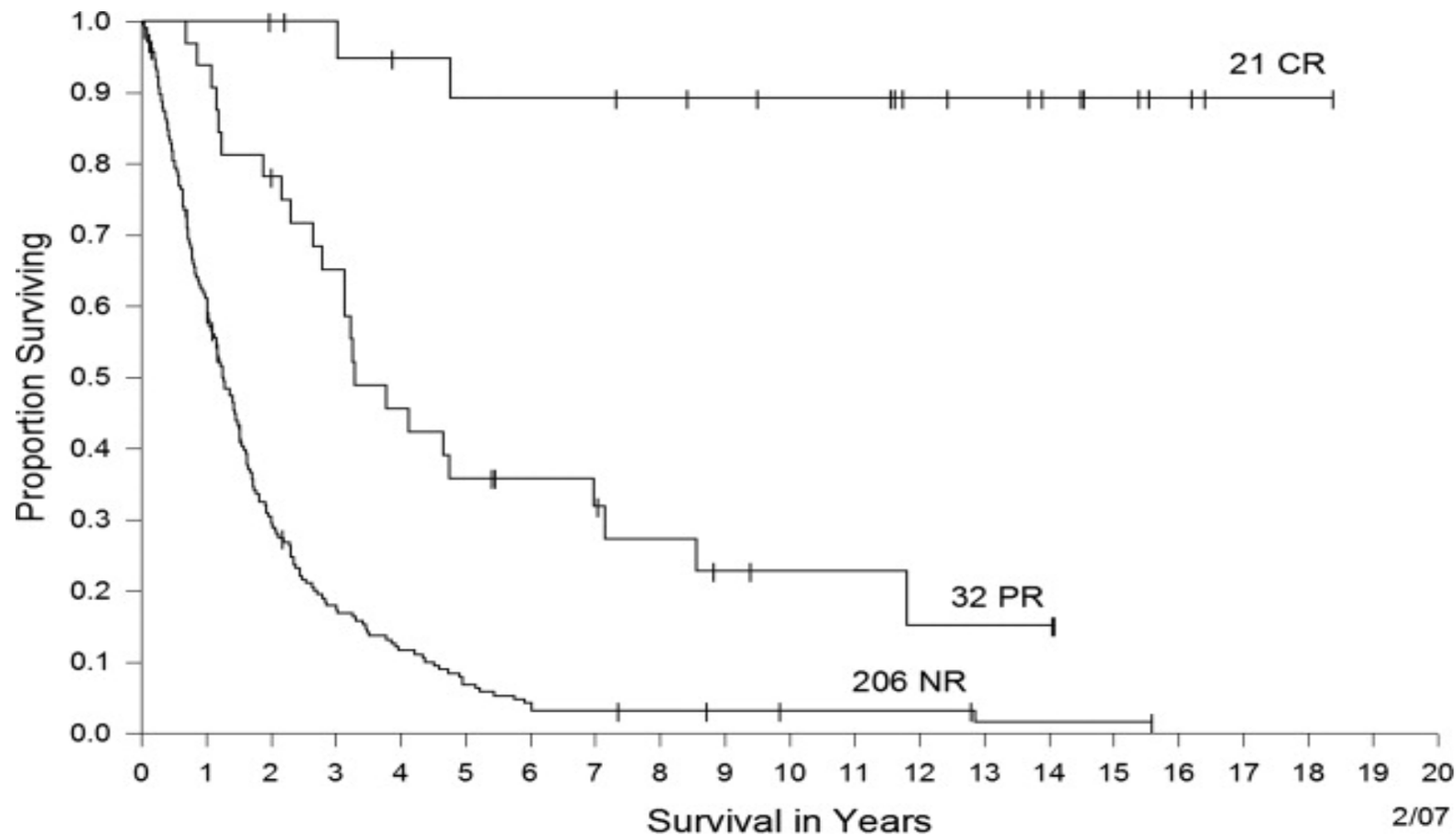
FDA-approved Immunotherapies for mRCC

Drug	Approved	Indication	Dose
High dose Interleukin-2	1992	Metastatic RCC	600,000 International Units/kg (0.037 mg/kg) IV q8hr infused over 15 minutes for a maximum 14 doses, THEN 9 days of rest, followed by a maximum of 14 more doses (1 course)*
Interferon-a (with bevacizumab)	2009	Clear cell RCC***	9 MIU s.c. three times a week
Nivolumab	2015	Clear cell RCC Refractory to prior VEGF Targeted therapy	3mg/kg 240mg IV q 2 week or 480mg IV q 4 wks
Nivolumab +ipilimumab	2018	Clear cell RCC, treatment naïve	3mg/kg nivo plus 1mg/kg ipi q3 wks x 4 doses then nivo maintenance at flat dosing

*Retreatment: Evaluate after 4 weeks, advisable only if tumor shrinkage and no retreatment contraindications (see package insert for details)

High Dose IL-2 in mRCC

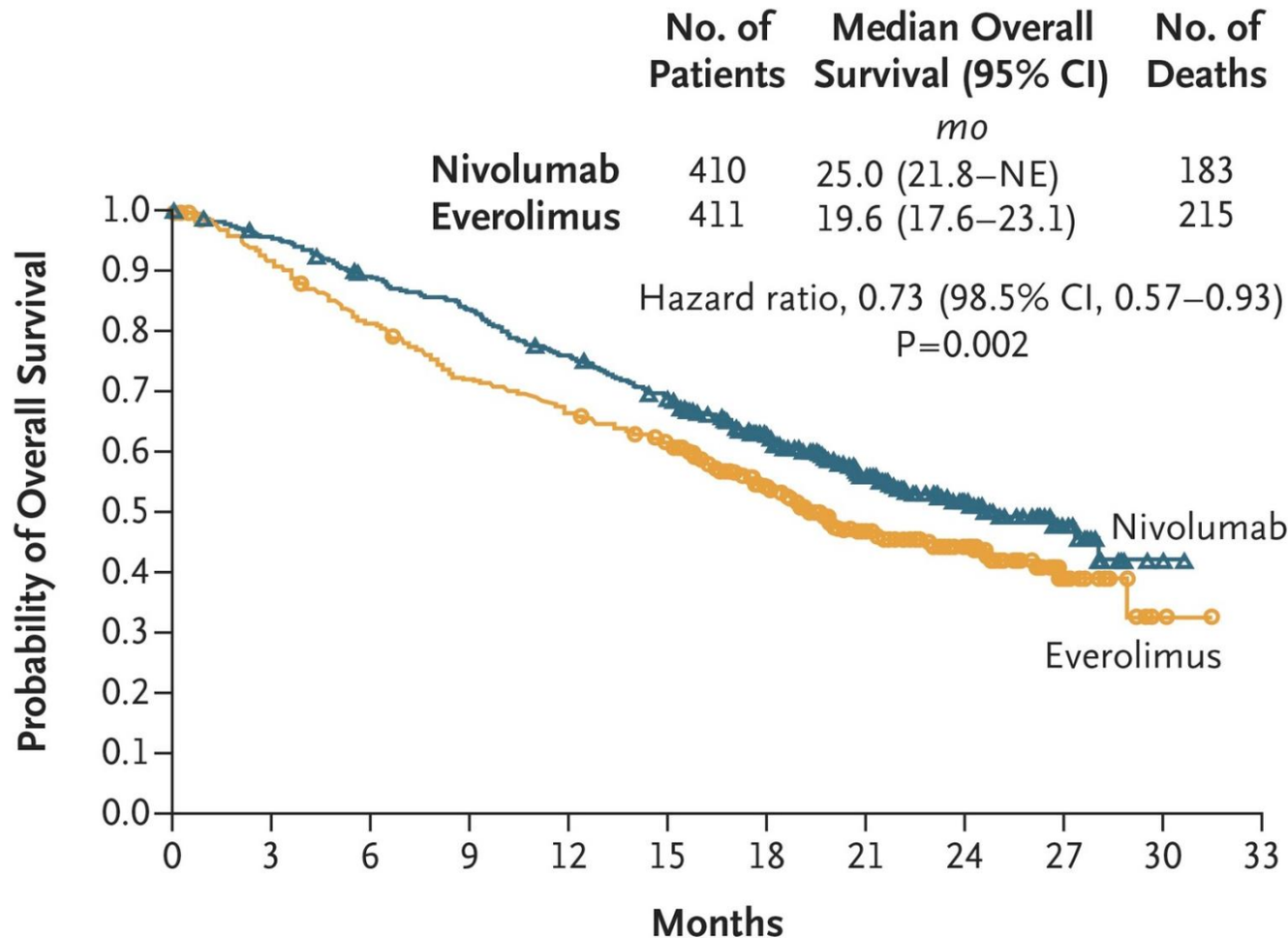
- 20 year analysis of 259 patients
- ORR = 20%
 - 9% CR (n = 23)
 - 12% PR (n = 30)
- Median duration of response = 15.5 months
- Median OS = 19 months



Klapper et al. Cancer 2008

Second-Line Nivolumab in mRCC

- CheckMate 025 Phase III trial
- Nivolumab = anti-PD-1 antibody
- Metastatic, clear-cell disease
- One or two previous antiangiogenic treatments
- Nivolumab (3 mg/kg IV Q2W) vs everolimus (10 mg daily)

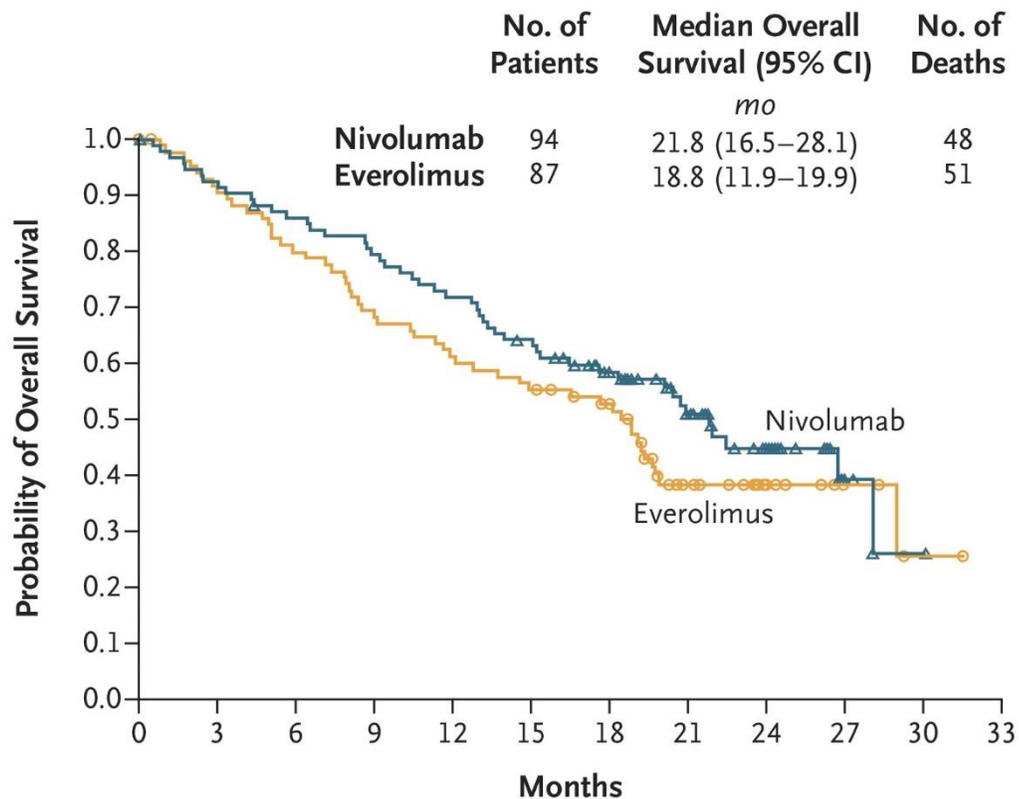


Motzer et al. NEJM 2015

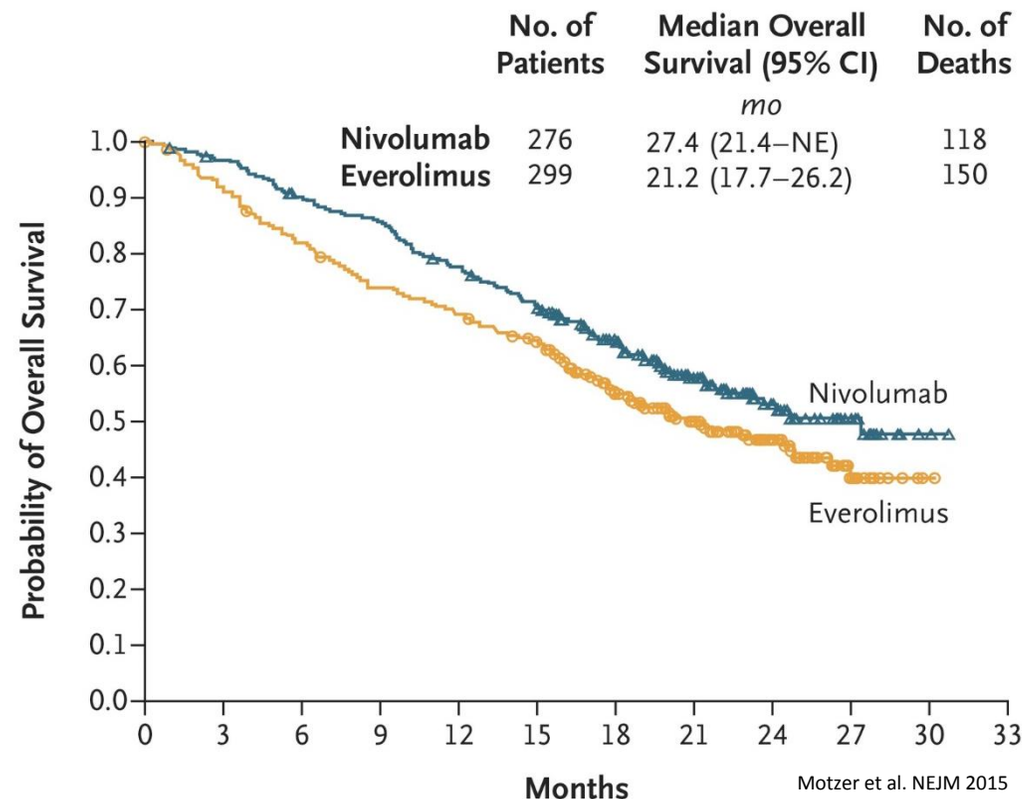
Second-Line Nivolumab in mRCC

PD-L1 subgroups

PD-L1 ≥ 1%

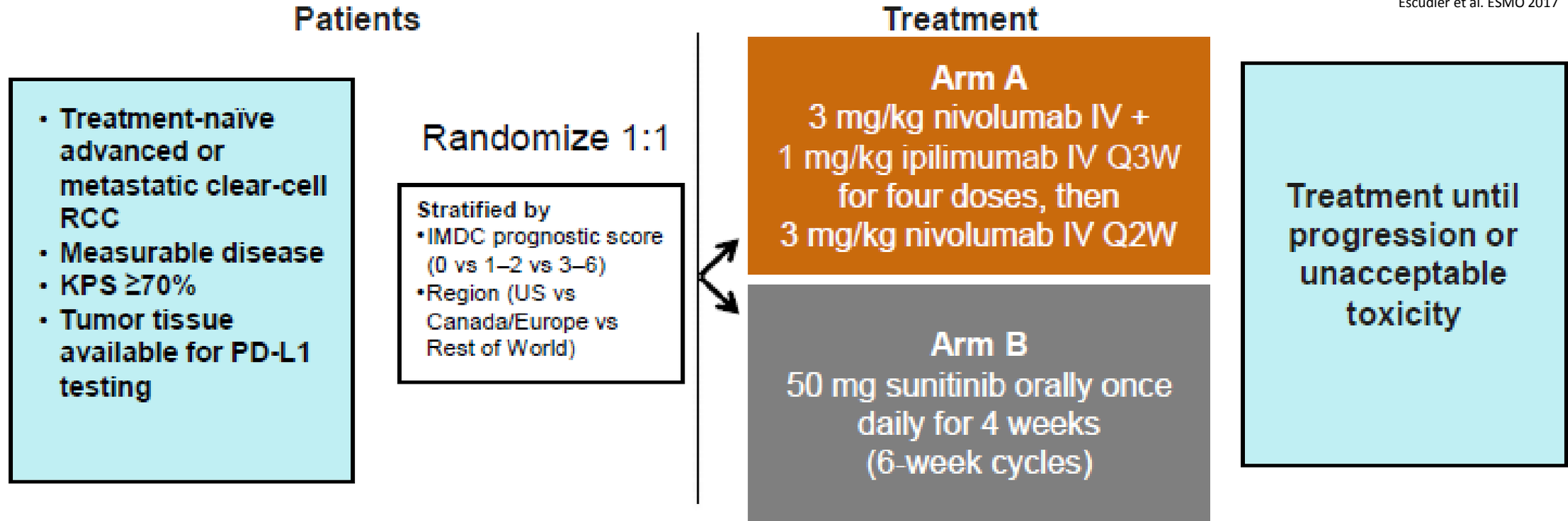


PD-L1 < 1%



First-line Nivolumab + Ipilimumab in mRCC

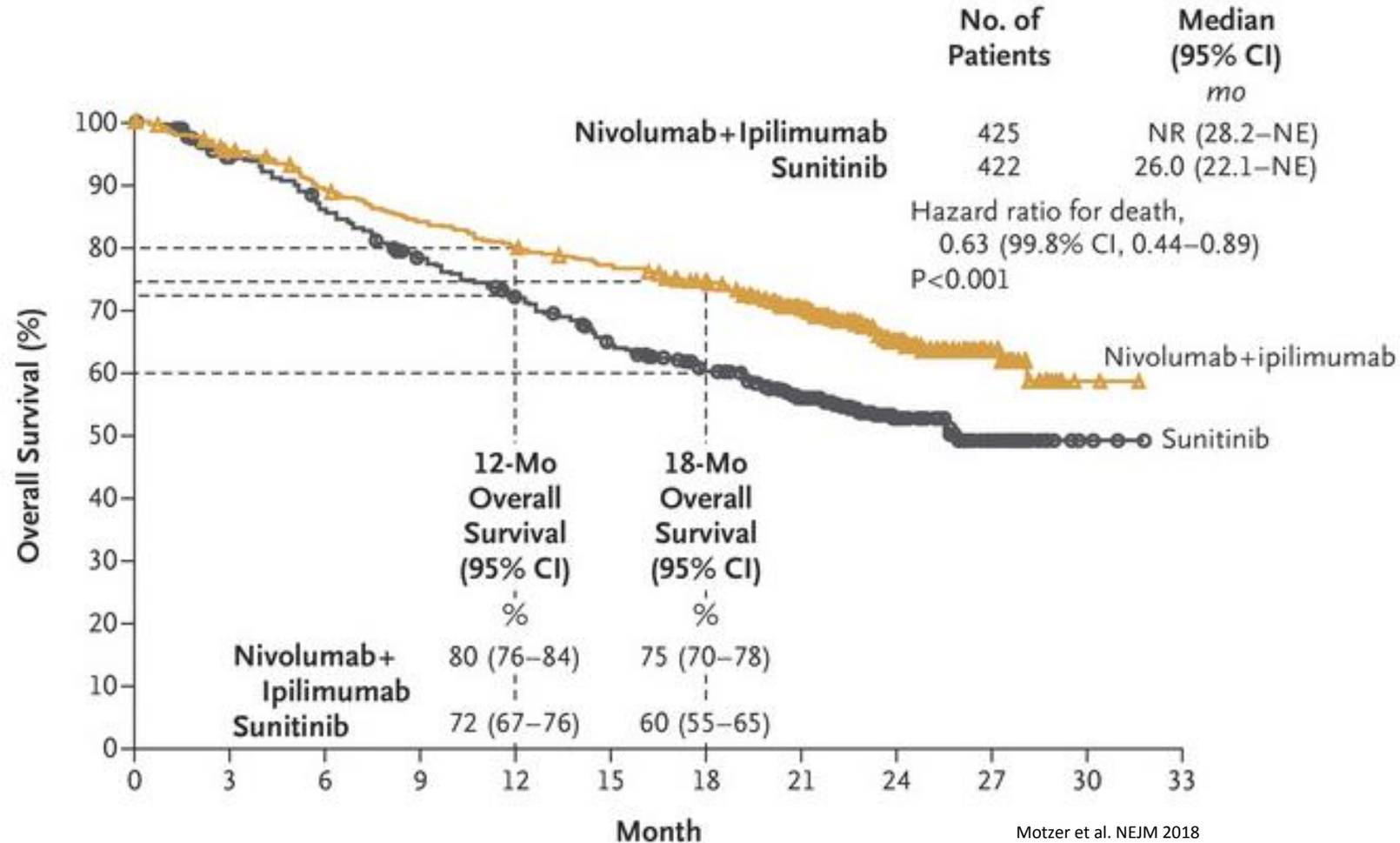
Escudier et al. ESMO 2017



Nivolumab = anti-PD-1 antibody

Ipilimumab = anti-CTLA-4 antibody

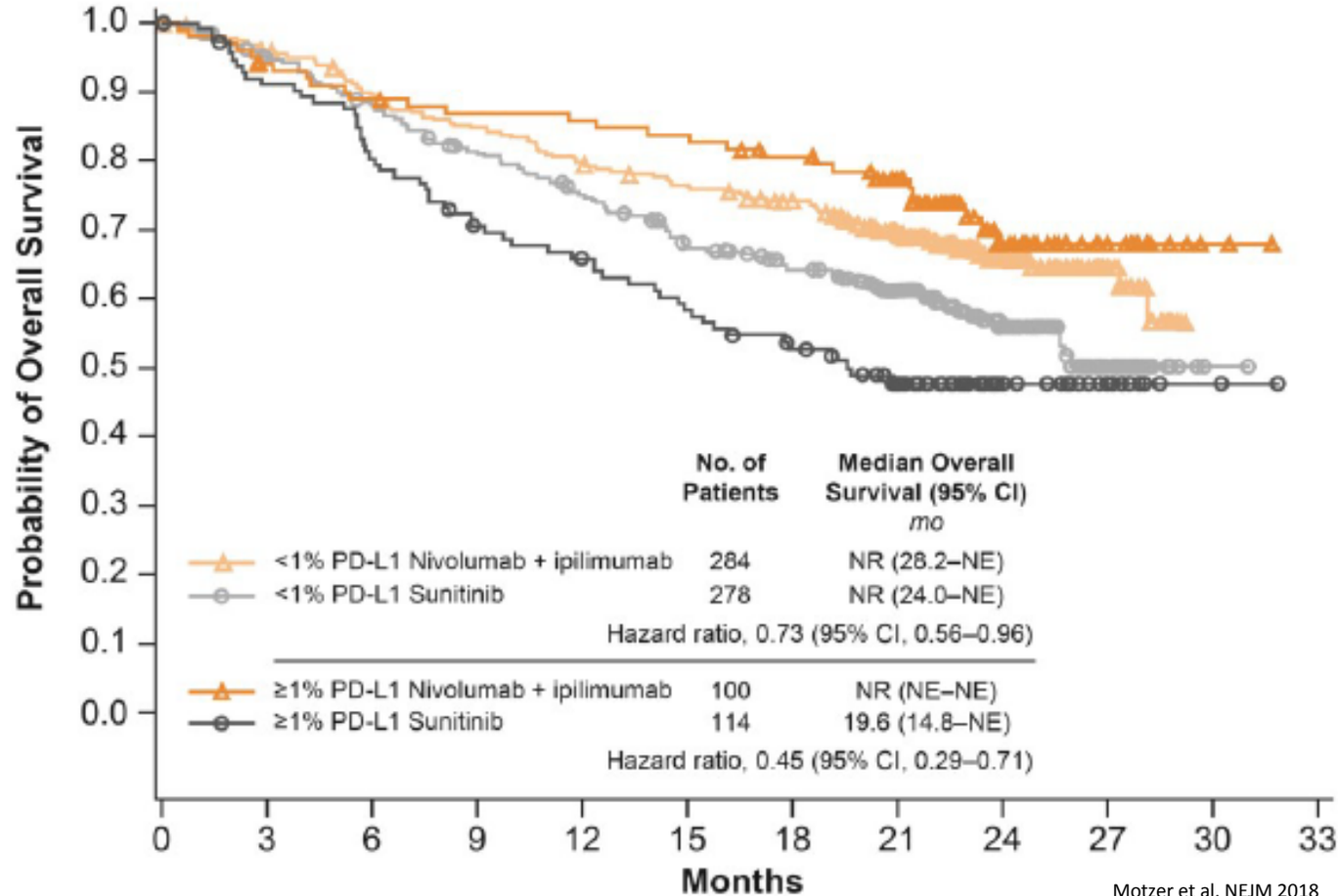
First-line Nivolumab + Ipilimumab in mRCC



Motzer et al. NEJM 2018

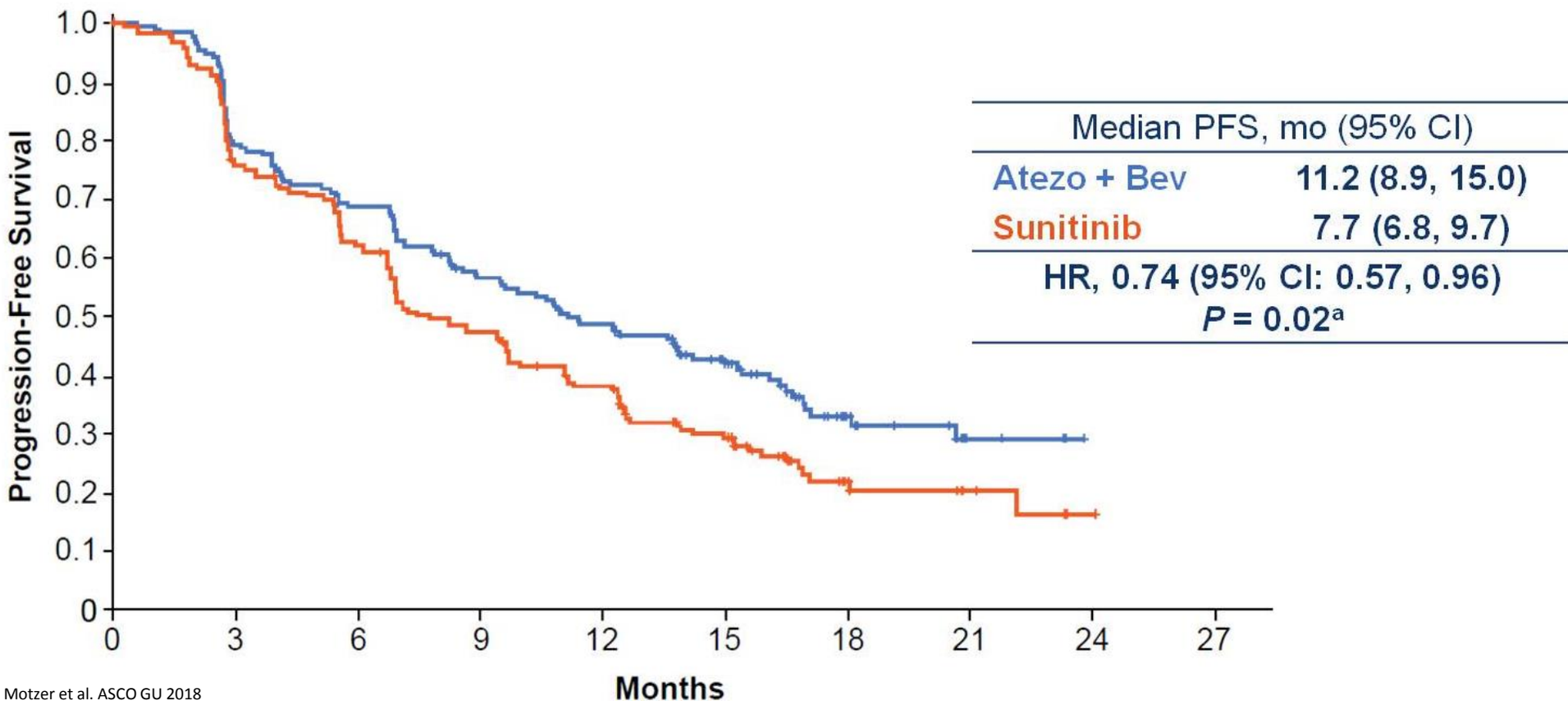
First-line Nivolumab + Ipilimumab in mRCC

PD-L1 Subgroups



Motzer et al. NEJM 2018

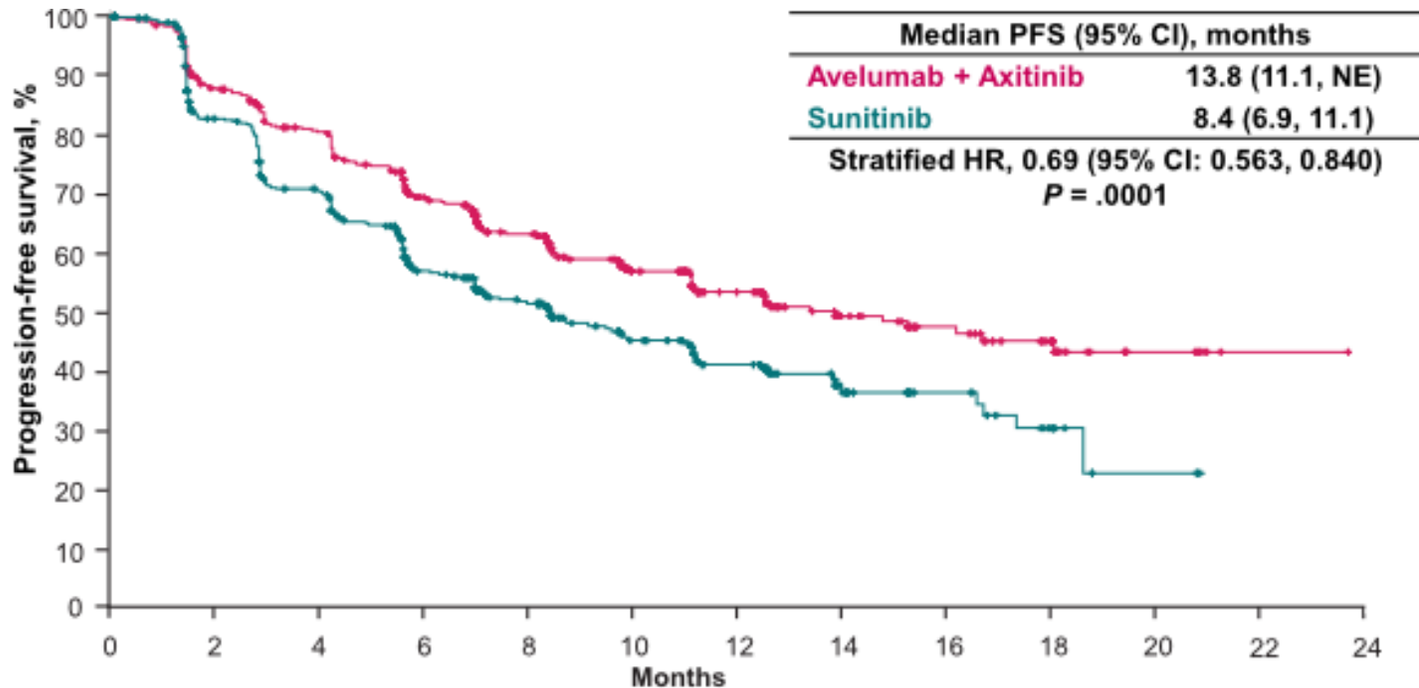
In Development: First-line Atezolizumab + Bevacizumab in PD-L1+ mRCC



Motzer et al. ASCO GU 2018
 Escudier et al. ASCO 2018

In Development: First-line Checkpoint Inhibitors + Axitinib in mRCC

JAVELIN Renal 101

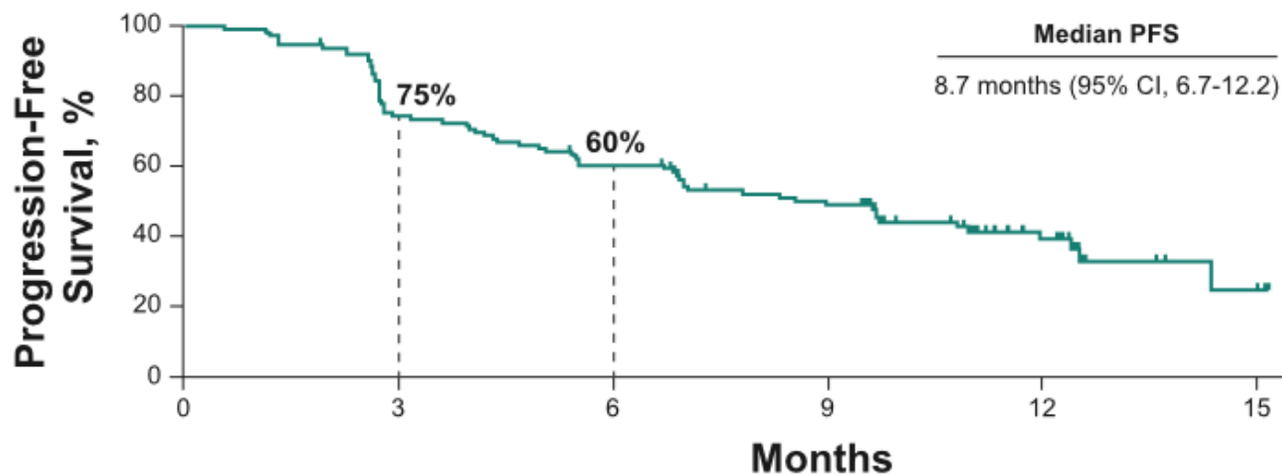


- KEYNOTE-426
 - Pembrolizumab + axitinib in mRCC
 - Positive for OS and PFS (10/18/2018)

Motzer et al. ESMO 2018

In Development: First-line Pembrolizumab in mRCC

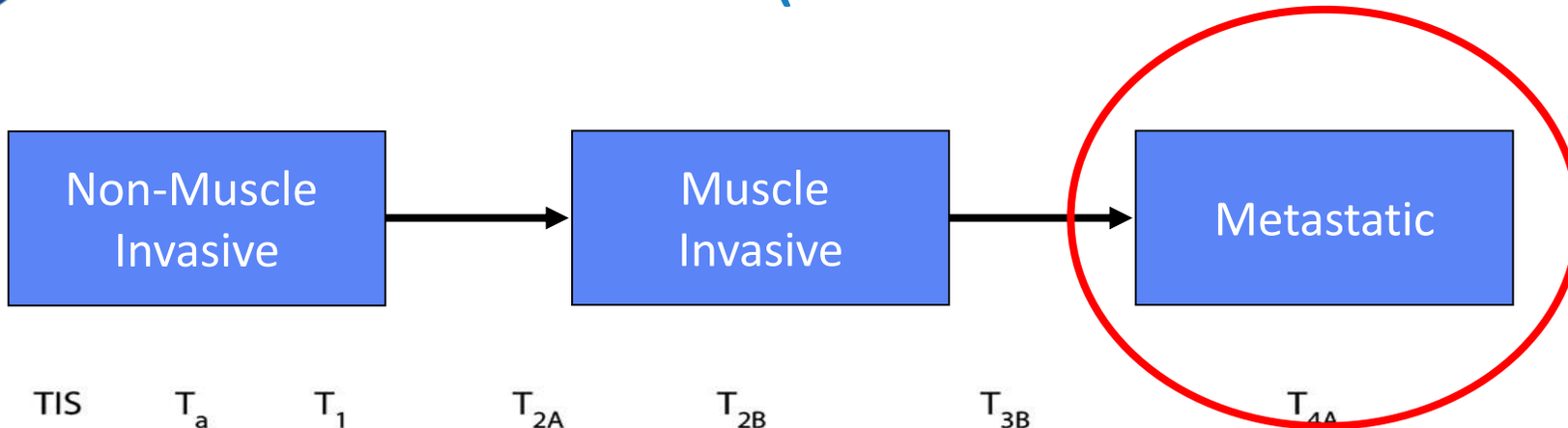
KEYNOTE - 427



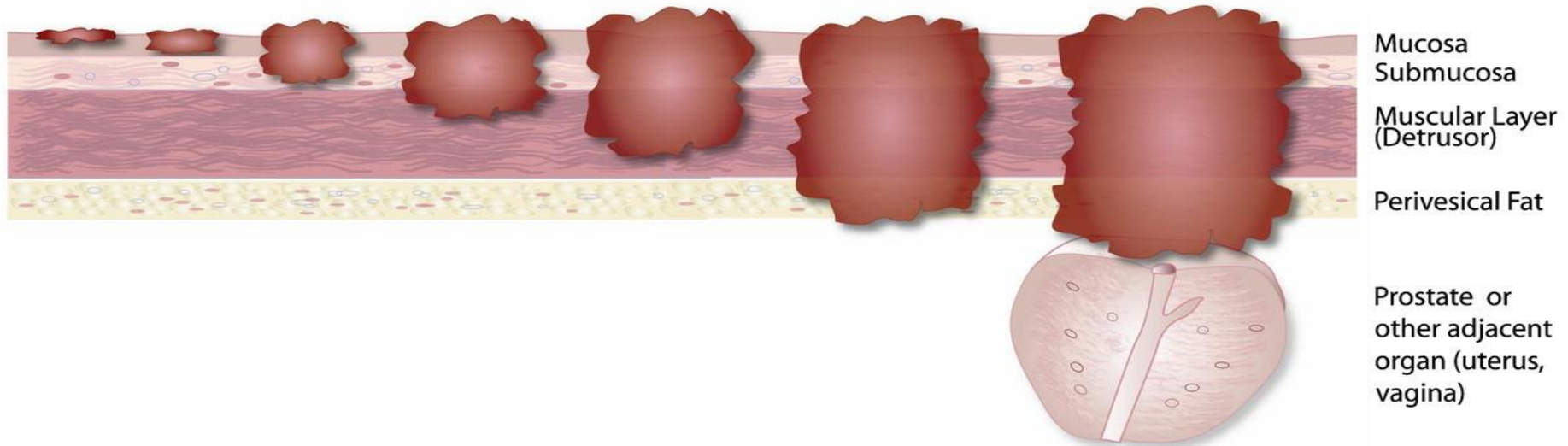
	N = 110
Confirmed ORR, % (95% CI)	38 (29 – 48)
Confirmed BOR, n (%)	
CR	3 (3)
PR	39 (35)
SD	35 (32)
PD	31 (28)
No assessment	2 (2)

Donskov et al. ESMO 2018

Immunotherapy for Metastatic Bladder Cancer (Urothelial Carcinoma; UC)



TNM	TIS	T _a	T ₁	T _{2A}	T _{2B}	T _{3B}	T _{4A}
JSM	0	0	A	B ₁	B ₂	C	D ₁



Approved Checkpoint Inhibitors for mUC

Platinum Refractory

Drug/Trial name	Phase	No. of patients	ORR	PFS	OS	Duration of response	Grade 3/4 AE (treatment related deaths)	Maximal duration of treatment
CISPLATIN REFRACTORY								
Atezolizumab IMvigor210 cohort 2	II	310	16% (6% CR)	2.1 mo	7.9 mo (1yr 29%)	22.1 mo	18% (0 deaths)	NR
Atezolizumab IMvigor211	III	931	13%	NR	8.6 mo	21.7 mo	20%	NR
Pembrolizumab KEYNOTE-045	III	542	21%	2.1 mo	10.3 mo	NR	14% (4 deaths)	2 years
Nivolumab CheckMate275	II	265	19.6% (2% CR)	2 mo	8.7 mo	NR	18% (3 deaths)	NR
Avelumab JAVELIN	Ib	242*	17% (6% CR)	6.6 weeks	6.5 mo	NR	10% (1 death)	NR
Durvalumab	I/II	191	17.8% (4% CR)	1.5 mo	18.2 mo	NR	7% (2 deaths)	1 year

Anti-PD-L1 Antibodies

- 1) Atezolizumab
- 2) Avelumab
- 3) Durvalumab

Anti-PD-1 Antibodies

- 1) Nivolumab
- 2) Pembrolizumab

In development: Combinations

- 1) IO + IO
- 2) IO + Chemotherapy

Approved Checkpoint Inhibitors for mUC

Cisplatin Ineligible

CISPLATIN INELIGIBLE								
Atezolizumab IMvigor210 cohort 1	II	119	23% (9% CR)	2.7 mo	15.9 mo, 1yr 57%	NR	16% (1 death)	NR
Pembrolizumab KEYNOTE-052	II	370	29% (7% CR)	6mo 30%	6 mo 67%	NR	19% (1 death)	2 years

Anti-PD-L1 Antibodies

- 1) Atezolizumab
 - PD-L1 stained tumor-infiltrating immune cells [IC] covering $\geq 5\%$ of the tumor area

Anti-PD-1 Antibodies

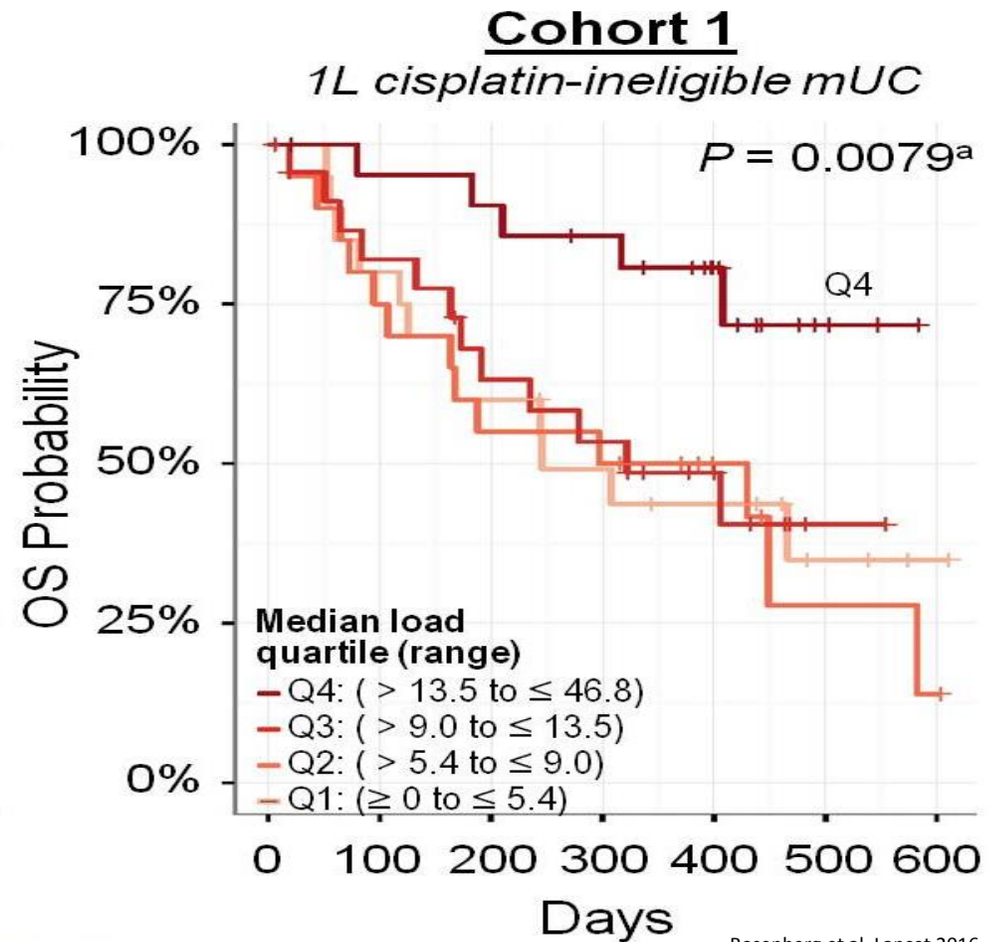
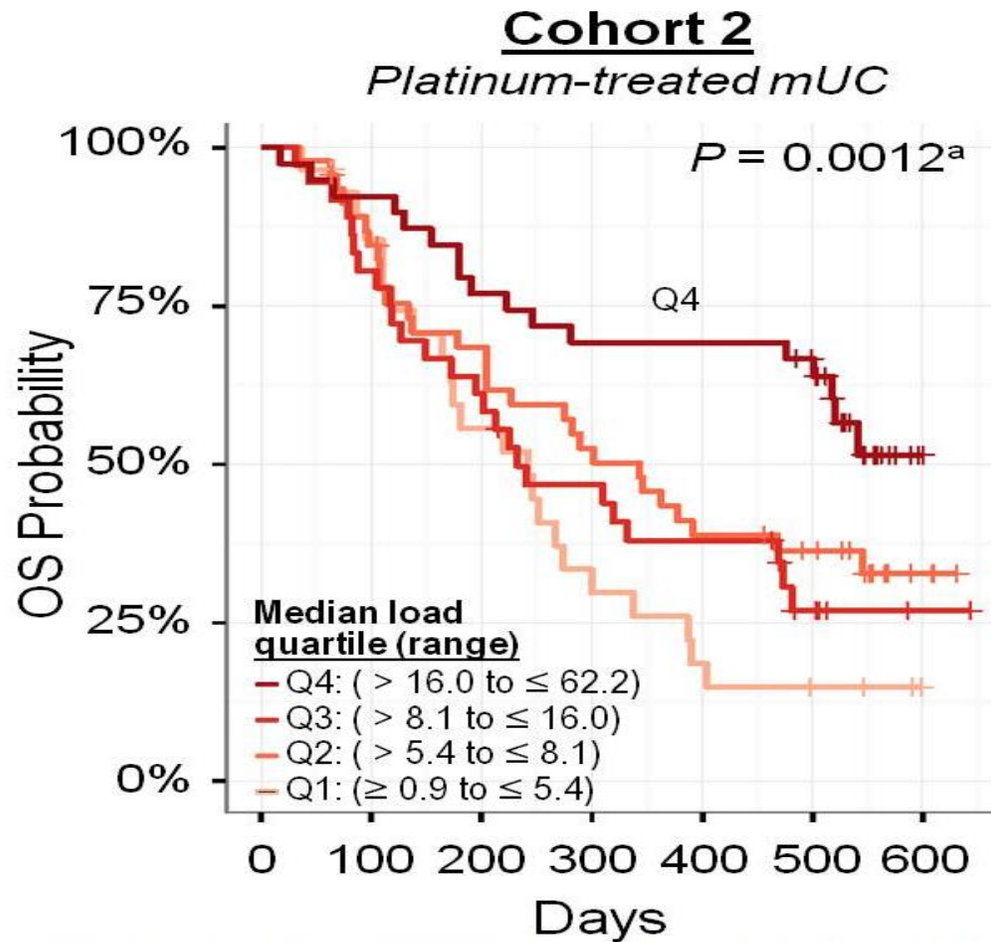
- 1) Pembrolizumab
 - PD-L1 CPS ≥ 10

In development: Combinations

- 1) IO + IO
- 2) IO + Chemotherapy

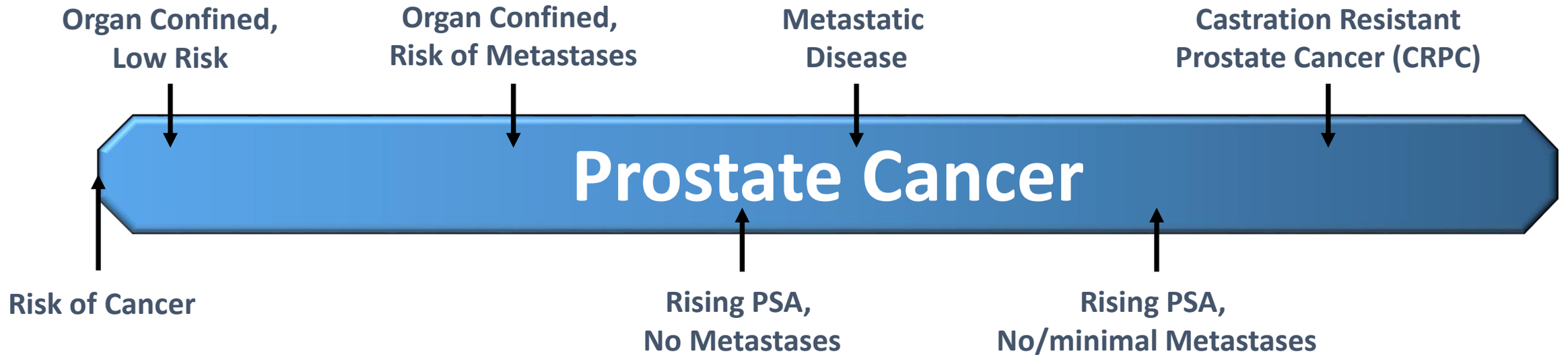
Tumor Mutational Burden (TMB) May Signal Responses with PD-1 Blockade

Atezolizumab in mUC



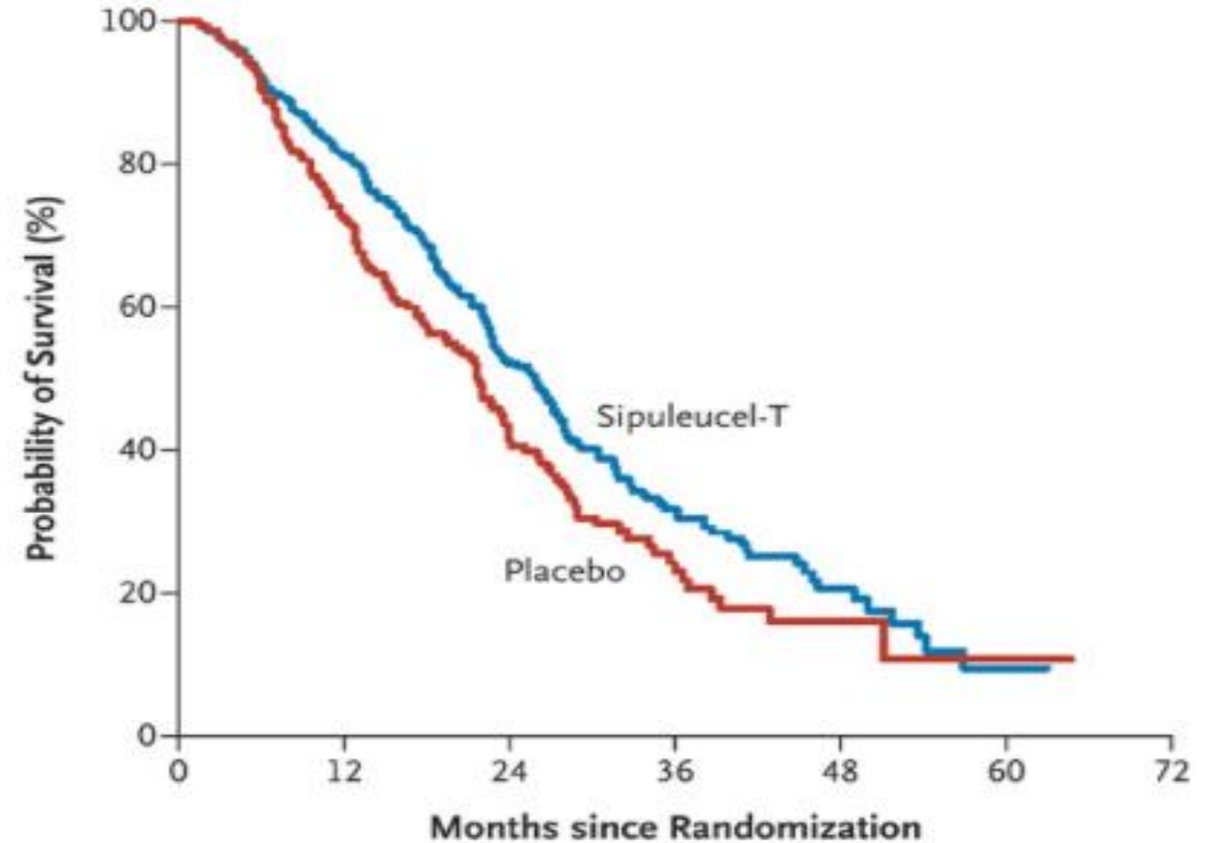
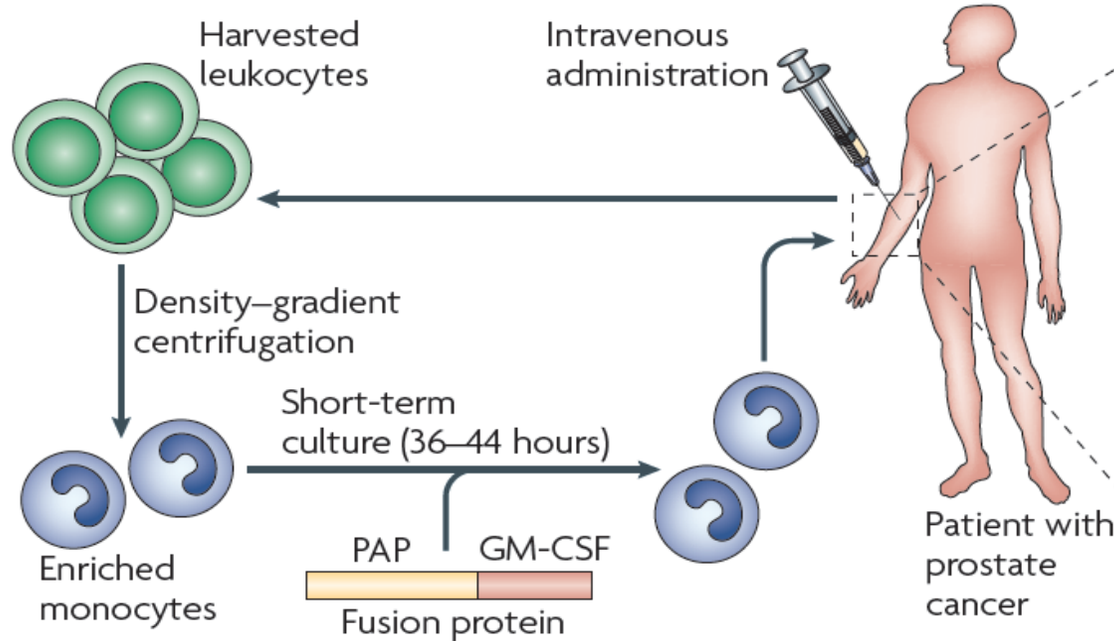
Rosenberg et al. Lancet 2016

The Spectrum of Prostate Cancer



Sipuleucel-T in mCRPC

- First anticancer therapeutic vaccine



Kantoff et al. NEJM 2010

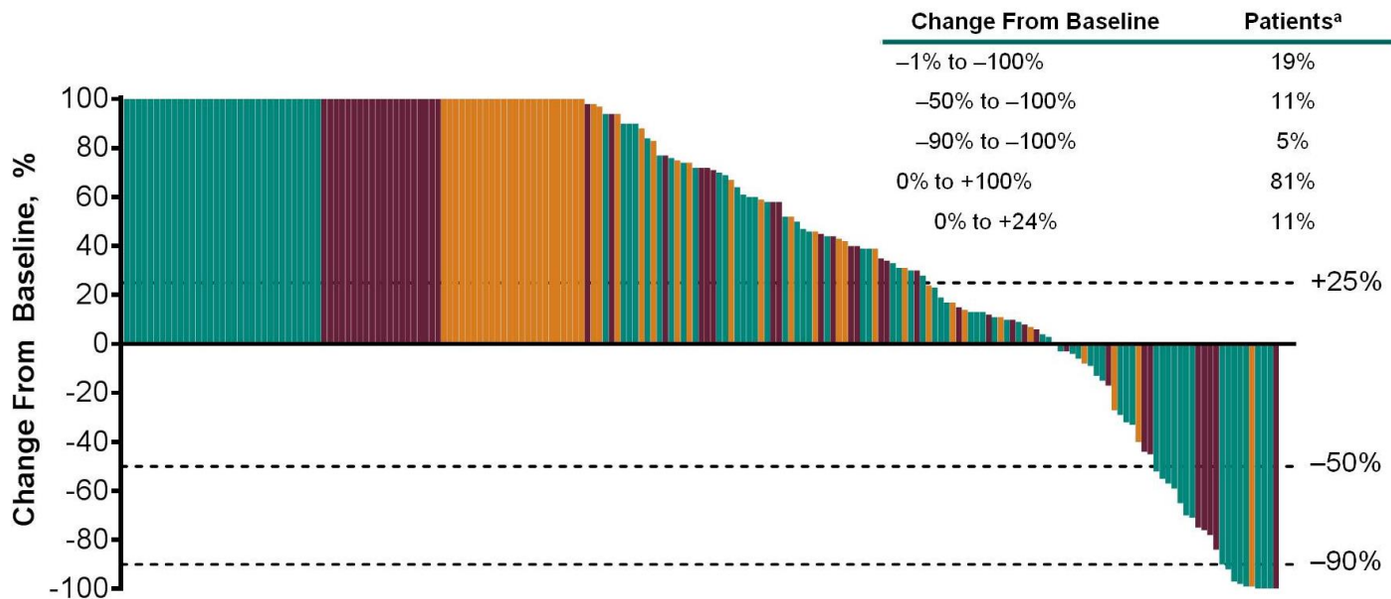
Drake et al. Curr Opin Urol 2010

Limited efficacy of Checkpoint Inhibitors in mCRPC

No FDA-approved CIs for mCRPC

Ex. – KEYNOTE-199 (Pembrolizumab)

■ Cohort 1 (PD-L1+)
 ■ Cohort 2 (PD-L1-)
 ■ Cohort 3 (Any PD-L1; Bone)

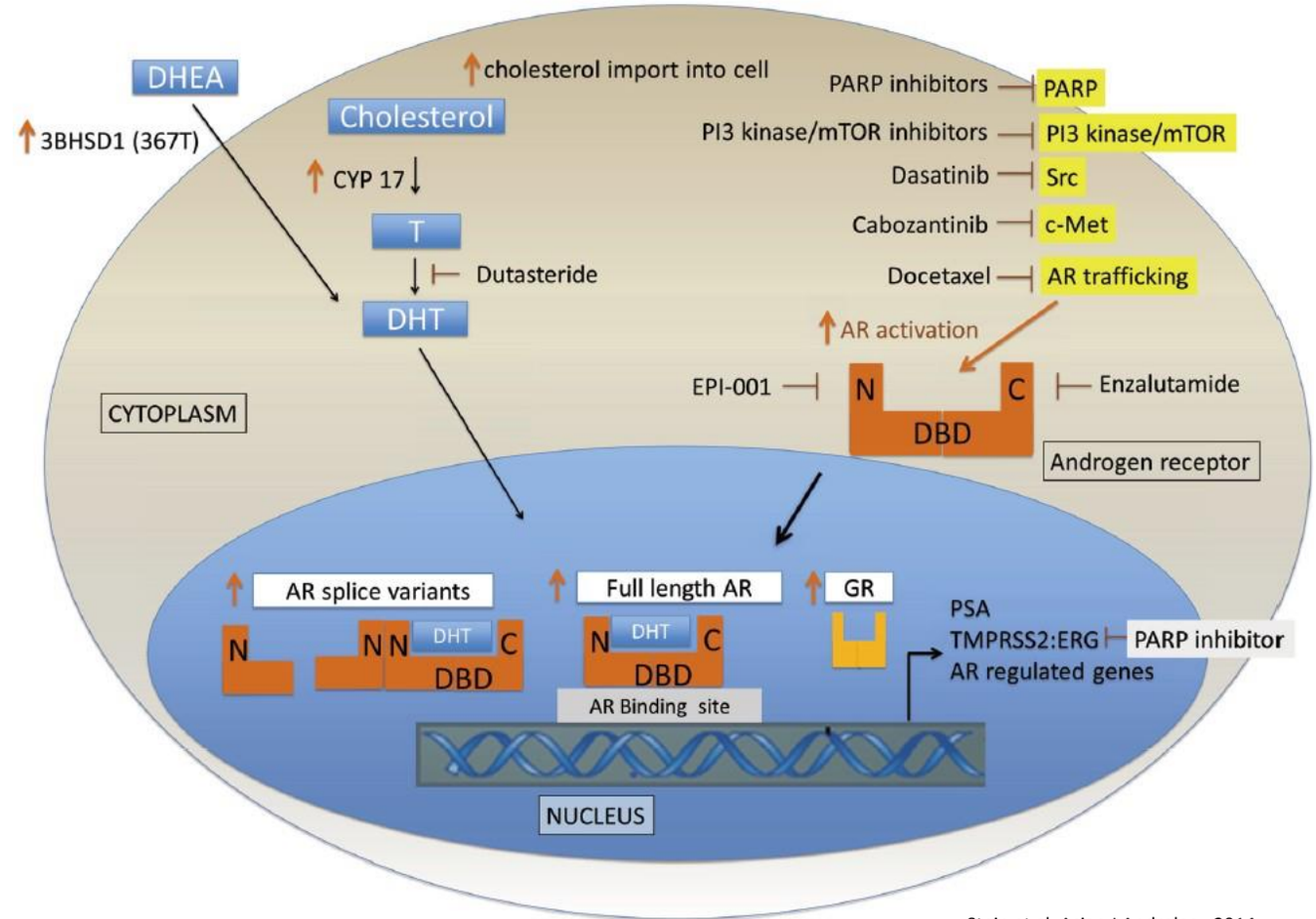


DeBono et al. ASCO 2018

- Pembrolizumab is approved for all Microsatellite Instability-High (MSI-H) solid tumors
- MSI-H incidence is low in PC
 - Localized PC ~2%
 - Autopsy series of mCRPC ~12%
- MSI testing may offer pembrolizumab as an option

Future Combinations in mCRPC to Engage Immune System

- Hormonal therapy
- Radiation
- Radium-223
- PARP inhibitors
- Chemotherapy
- New targets



Stein et al. Asian J Andrology 2014

irAEs with Immune Checkpoint Inhibitors in GU Cancers

Meta-analysis of 8 studies

- Similar incidence overall

Adverse event	Incidence, any grade (GU only trials) (%)	Incidence, grades 3–5 (GU only trials) (%)	Incidence any grade (non-GU clinical trials) (%)	Incidence, grades 3–5 (non-GU clinical trials) (%)
Hypothyroid/thyroiditis	0.8–9	0–0.6	3.9–12	0–0.1
Diabetes/DKA	0–1.5	0–0.7	0.8–0.8	0.4–0.7
LFT changes/hepatitis	1.5–5.4	1–3.8	0.3–3.4	0.3–2.7
Pneumonitis	2–4.4	0–2	1.8–3.5	0.25–1.9
Encephalitis	NR	NR	0.2–0.8	0.0–0.2
Colitis/diarrhea	1–10	1–10	2.4–4.1	1.0–2.5
Hypophysitis	0–0.5	0–0.2	0.2–0.9	0.2–0.4
Renal Dysfunction/nephritis	0.3–1.6	0–1.6	0.3–4.9	0.0–0.5
Myositis	0.8–5	0–0.8	NR	NR

Maughan et al. Front Oncol 2017

Immune-related Adverse Events

Table 2 General guidance for corticosteroid management of immune-related adverse events

Grade of immune-related AE (CTCAE/equivalent)	Corticosteroid management	Additional notes
1	<ul style="list-style-type: none"> Corticosteroids not usually indicated 	<ul style="list-style-type: none"> Continue immunotherapy
2	<ul style="list-style-type: none"> If indicated, start oral prednisone 0.5-1 mg/kg/day if patient can take oral medication. If IV required, start methylprednisolone 0.5-1 mg/kg/day IV If no improvement in 2–3 days, increase corticosteroid dose to 2 mg/kg/day Once improved to ≤grade 1 AE, start 4–6 week steroid taper 	<ul style="list-style-type: none"> Hold immunotherapy during corticosteroid use Continue immunotherapy once resolved to ≤grade 1 and off corticosteroids Start proton pump inhibitor for GI prophylaxis
3	<ul style="list-style-type: none"> Start prednisone 1-2 mg/kg/day (or equivalent dose of methylprednisolone) If no improvement in 2–3 days, add additional/alternative immune suppressant Once improved to ≤ grade 1, start 4–6-week steroid taper Provide supportive treatment as needed 	<ul style="list-style-type: none"> Hold immunotherapy; if symptoms do not improve in 4–6 weeks, discontinue immunotherapy Consider intravenous corticosteroids Start proton pump inhibitor for GI prophylaxis Add PCP prophylaxis if more than 3 weeks of immunosuppression expected (>30 mg prednisone or equivalent/day)
4	<ul style="list-style-type: none"> Start prednisone 1-2 mg/kg/day (or equivalent dose of methylprednisolone) If no improvement in 2–3 days, add additional/alternative immune suppressant, e.g., infliximab Provide supportive care as needed 	<ul style="list-style-type: none"> Discontinue immunotherapy Continue intravenous corticosteroids Start proton pump inhibitor for GI prophylaxis Add PCP prophylaxis if more than 3 weeks of immunosuppression expected (>30 mg prednisone or equivalent/day)

Puzanov Journal for ImmunoTherapy of Cancer 2017

Additional Resources

Rini et al. *Journal for Immunotherapy of Cancer* (2016) 4:81
DOI 10.1186/s40425-016-0180-7

Journal for Immunotherapy
of Cancer

POSITION ARTICLE AND GUIDELINES **Open Access**

 CrossMark

Society for Immunotherapy of Cancer consensus statement on immunotherapy for the treatment of renal cell carcinoma

Brian I. Rini¹, David F. McDermott², Hans Hammers³, William Bro⁴, Ronald M. Bukowski⁵, Bernard Faba⁶, Jo Faba⁶, Robert A. Figlin⁷, Thomas Hutson⁸, Eric Jonasch⁹, Richard W. Joseph¹⁰, Bradley C. Leibovich¹¹, Thomas Olencki¹², Allan J. Pantuck¹³, David I. Quinn¹⁴, Virginia Seery², Martin H. Voss¹⁵, Christopher G. Wood⁹, Laura S. Wood¹ and Michael B. Atkins^{16*}

Kamat et al. *Journal for Immunotherapy of Cancer* (2017) 5:68
DOI 10.1186/s40425-017-0271-0

Journal for Immunotherapy
of Cancer

POSITION ARTICLE AND GUIDELINES **Open Access**

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Society for Immunotherapy of Cancer consensus statement on immunotherapy for the treatment of bladder carcinoma

Ashish M. Kamat^{1*}, Joaquim Bellmunt², Matthew D. Galsky³, Badrinath R. Konety⁴, Donald L. Lamm⁵, David Langham⁶, Cheryl T. Lee⁷, Matthew I. Milowsky⁸, Michael A. O'Donnell⁹, Peter H. O'Donnell¹⁰, Daniel P. Petrylak¹¹, Padmanee Sharma¹², Ella C. Skinner¹³, Guru Sonpavde¹⁴, John A. Taylor III¹⁵, Prasanth Abraham¹⁶ and Jonathan E. Rosenberg¹⁷

McNeel et al. *Journal for Immunotherapy of Cancer* (2016) 4:92
DOI 10.1186/s40425-016-0198-x

Journal for Immunotherapy
of Cancer

POSITION ARTICLE AND GUIDELINES **Open Access**

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The Society for Immunotherapy of Cancer consensus statement on immunotherapy for the treatment of prostate carcinoma

Douglas G. McNeel¹, Neil H. Bander², Tomasz M. Beer³, Charles G. Drake⁴, Lawrence Fong⁵, Stacey Harrelson⁶, Philip W. Kantoff⁷, Ravi A. Madan⁸, William K. Oh⁹, David J. Peace¹⁰, Daniel P. Petrylak¹¹, Hank Porterfield¹², Oliver Sartor¹³, Neal D. Shore⁶, Susan F. Slovin⁷, Mark N. Stein¹⁴, Johannes Vieweg¹⁵ and James L. Gulley^{16*}

Case Study 1

- 60 yo F cyclist who noted increasing fatigue after a hysterectomy. She was quite anemic pre and post-operatively and was started on oral iron yet continued to feel weak and to lose weight. An abdominal ultrasound was ordered to evaluate her weight loss and a 10 cm renal mass was noted. She underwent a left radical nephrectomy revealing a T3aNxMx Clear Cell RCC, Fuhrman grade $\frac{3}{4}$.
- Imaging at the time revealed a 1.8 cm lower lobe pulmonary nodule.
- Follow up imaging 2 months post operatively revealed multiple new pulmonary lesions, bilaterally which were biopsy proven RCC.
- She began sunitinib with improvement in fatigue, anemia, and appetite. Her restaging imaging at 3 months revealed new liver lesions and enlarging pulmonary lesions.

Case Study 1

- Second line nivolumab was chosen
- Tolerated well with some mild GI symptoms.
- Pulmonary nodules resolved
- Liver lesions appeared less active
- Remained on therapy for 15 months and then developed a new small liver lesion in the setting of otherwise stable PR

Case Study 1

- Underwent Radiofrequency ablation to the solitary new liver lesion with good results.
- Follow up imaging revealed a good response with scarring at the RFA site and virtual resolution of her additional sites of disease
- Treatment holiday after 2 years of nivolumab and stable after 13 months of follow up

Case Study 1

Discussion topics

- focal therapy of oligometastatic disease or focal progression
- treatment holidays after prolonged response to checkpoint inhibition

Case study 2

73 yo F PMHx of COPD, tobacco use, uncontrolled diabetes who lives with family members in Colorado and Texas presented with intermittent hematuria underwent cystoscopy and TURBT revealing a large friable bladder tumor. Pathology was T2 high grade urothelial carcinoma

Imaging at the time revealed a mass with perivesicular stranding, non avid mildly enlarged lymph nodes.

ECOG is 1; no neuropathy, GFR is

She underwent 4 cycles of neoadjuvant cisplatin, gemcitabine and 4 weeks post-chemotherapy had a cystectomy, lymph node dissection and ileal conduit creation.

ypT0N0 (0/15 nodes)

Case study 2

She was on surveillance and approximately 9 months post-operatively her images revealed a 3 cm soft tissue nodule along the right pelvic sidewall which was PET avid, as well as intensely avid, non-enlarged right common iliac node in the mid-upper pelvis

- Initiated therapy with atezolizumab given every 21 days
- Cycle 6 she was clinically stable. Scans with mixed response of mildly enlarging pelvic sidewall node and a new PET avid node in the pre-aortic region of 1.5 mm in short axis. Therapy continued

Case study 2

By cycle 9 she had increasing difficulties ambulating without a prop and general arthralgia/myalgia. Complaining of hip and shoulder girdle stiffness, and weight loss

ESR 69 CRP 87; CPK <10; LFTs normal; xray with some mild healed erosions in the carpometacarpal articulations

Received 1mg/kg steroids with rapid improvement in mobility and markers. Thought to be PMR.

ESR 10 CRP 16 on follow up

Atezolizumab discontinued

Follow up imaging 3 months later with resolution of pelvic mass and RP lymphadenopathy. No new sites of disease

Case study 2

Discussion points

- sequencing of immunotherapy
- pseudoprogression
- inflammatory toxicities
- continued response