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What's Next for Cancer Immunotherapy?

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Society for Immunotherapy of Cancer



Disclosures

- Nektar Therapeutics Advisory Board; Honorarium
- By virtue of discussing the future, I will be discussing non-FDA approved indications during my presentation.







Improvements in Staging and Immunotherapy Biomarkers

• Immunoscore

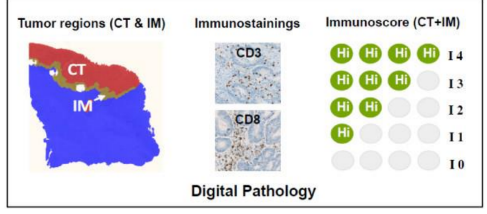
- CD3, CD8, memory markers
- PD-L1, PD-L2 and other checkpoint ligands
- Next generation sequencing
 - MSI-high, MMR defects, etc.
- Gut microbiome
- Tumor microenvironment/metabolomics







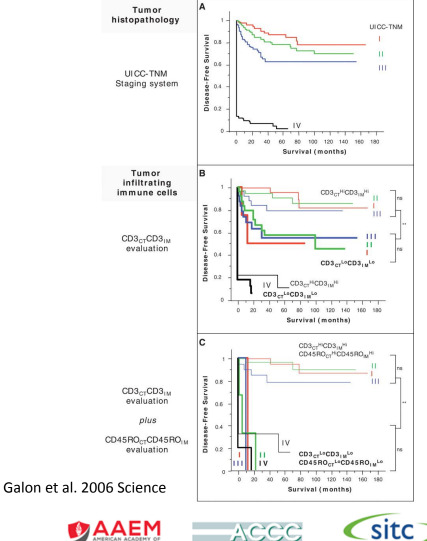
Immunoscore will become part of standard pathologic reports for all tumors, used as a biomarkers for responses and correlate with survival



CT = center of tumor; IM = invasive margin; Galon et al. 2012 J Transl Med

Cancers where immunoscore correlates with outcome

Adult tumors	Hepatocellular carcinoma
Colorectal cancer	Breast cancer
Melanoma	Ovarian cancer
Renal cell carcinoma	Spinal chordoma
Non-small cell lung cancer	Pediatric tumors
Head and neck cancer	Neuroblastoma
Gastric cancer	Osteosarcoma



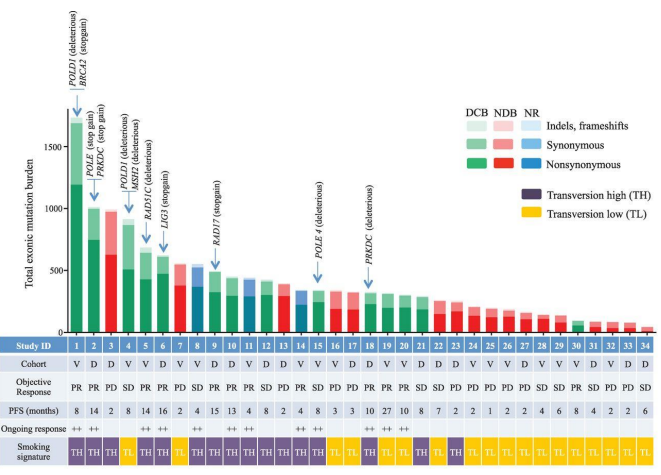
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Better intersection of next generation sequencing with predicting immunotherapy responses

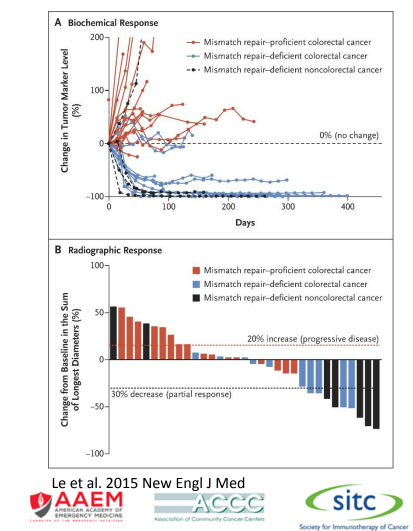
Tumor mutational burden

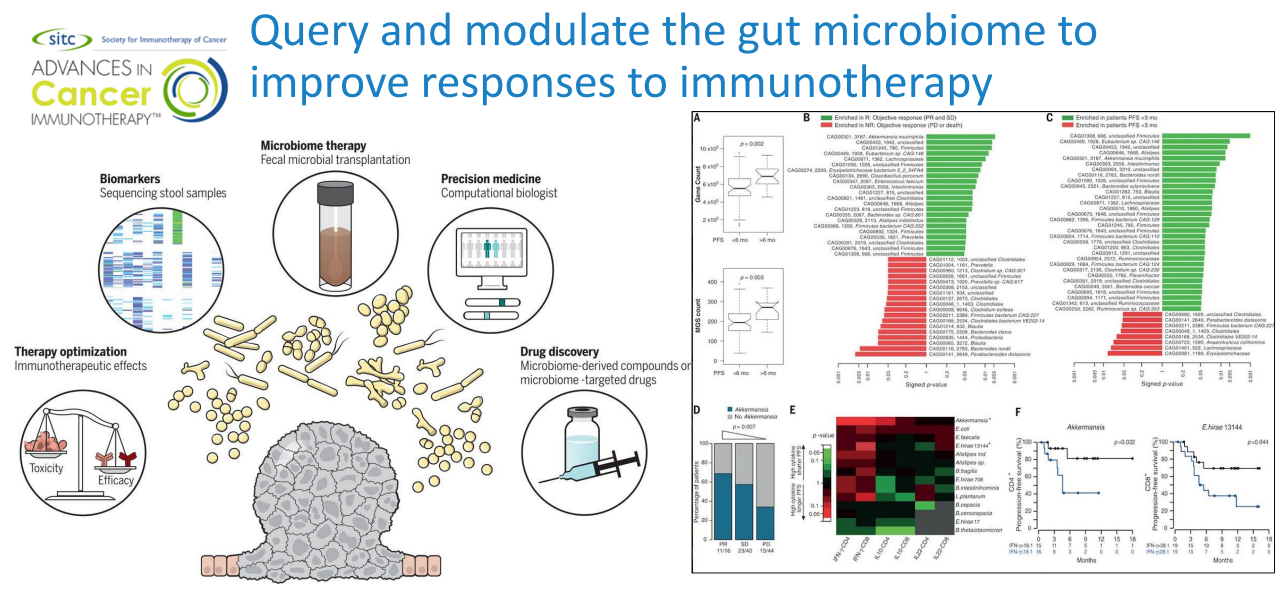


Rizvi et al. 2016 Science

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Mismatch repair defects





Zitvogel et al. 2018 Science

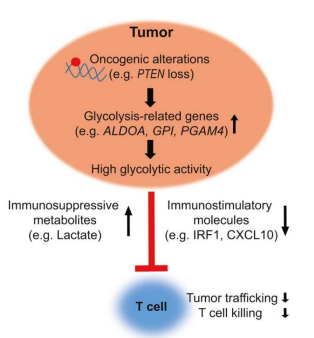
Routy et al. 2018 Science







Target the tumor metabolic environment to enhance immunotherapy responses



GENE	FC (N/R)	P value	В			С			
ALDOA	1.697746	0.136385					1		
ALDOC	1.758665	0.186307	⁵⁰]	P=0.011		_ 800	Aerobic	T	Energetic
ENO2	1.257908	0.685237	ъ ⁴⁰ -			ation			
ENO3	2.365525	0.205193			<u> </u>	spire		YUN I	
GAPDH	1.733471	0.143722	c ger c ger			ol/n			
GPI	1.700951	0.015443	erall expression glycolytic genes	r st n	••	drial Respira (pmol/min)			
LDHA	1.429014	0.302088	Overall expression glycolytic genes 0 0 00			Mitrochondrial Respiration OCR (pmol/min)			
LDHB	1.511258	0.152462	Ó		'	C			
PFKM	1.152264	0.57808	0 +	CR/PR	SD/PD	200			
PFKP	1.232823	0.463708				0-	Quiescen	3-	Glycolytic
PGAM1	1.421356	0.108866				C	5	10 1	5 20
PGAM4	1.5305	0.033143						R (mpH/min) Iycolysis	
PGK1	1.636341	0.087465						, ,	

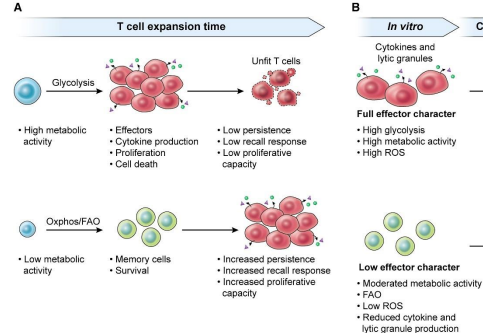
Cascone et al. 2018 Cell Metab



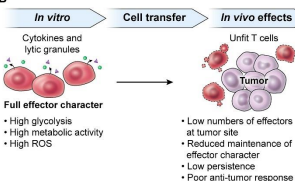


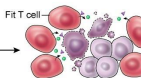


Target T cell metabolism to enhance immunotherapy responses



Kishton et al. 2017 Cell Metab





Increased numbers of effectors at tumor site
Effector activity against tumor
Increased persistence
High anti-tumor activity

Molecule	Target	Metabolic outcome	Clinical (C), pre-clinical (P)
2-DG	Hexokinase	↓Glycolysis	Ρ
Mdivi-1	Drp-1	\downarrow Mitochondrial fission	Ρ
JQ1	c-Myc	↓Glycolysis	Р
STF-31	GLUT1	↓Glycolysis	Р
WZB117	GLUT1	↓Glycolysis	Р
Rapamycin	mTOR	↓Glutamine metabolism	С
Metformin	АМРК, ЕТС	个FAO, others	С
Fenofibrate	ΡΡΑRα	个Fatty acid catabolism	Ρ
ugnani et al 2017 (ancer l ett		

Dugnani et al. 2017 Cancer Lett







Expansion of immunotherapy therapeutics

- Antibody therapy
 - Checkpoint agonists/inhibitors
 - Antibody-drug conjugates
 - Bispecifics
- Oncolytic viral therapy
- Radiotherapy/Immunotherapy

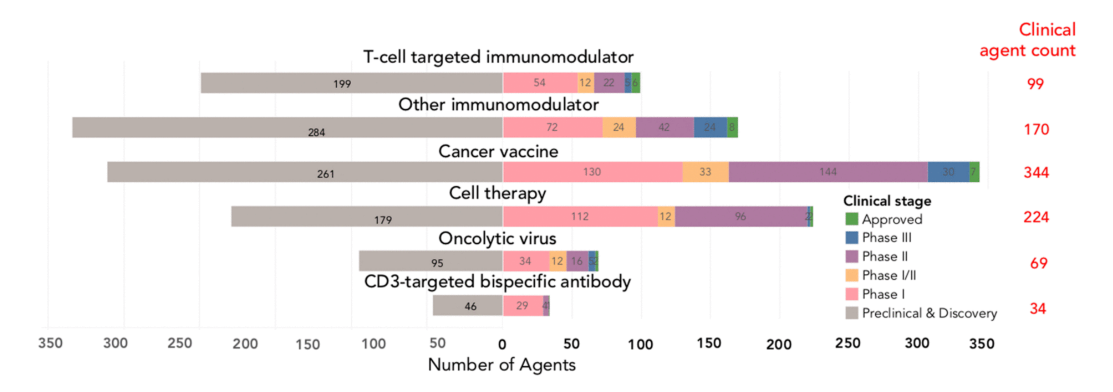
- Cellular therapy
 - CAR T cells and CAR NK cells
 - Bispecific
 - Bicistronic
 - Armored
 - TCR transduced T cells





2,004 IO AGENTS IN DEVELOPMENT

940 AGENTS ARE IN CLINICAL STAGES, AND 1,064 IN PRECLINICAL





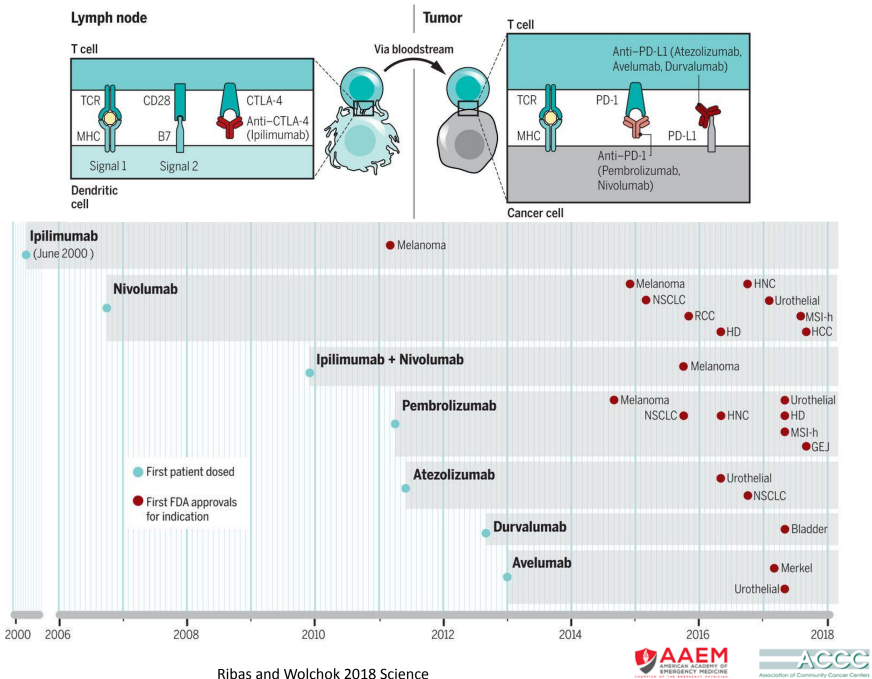
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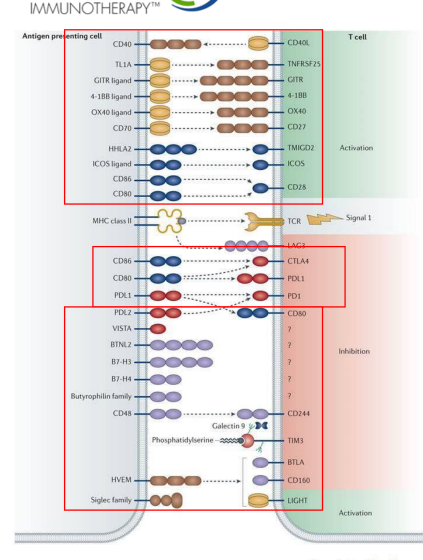




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Checkpoint agonists and antagonists will expand Society for Immunotherapy of Cancer and be used in combination



Appendix: Immune checkpoint modulators in combination clinical trials (August 2017



Checkpoint modulator name	Target	Checkpoint modulator name	Target	Checkpoint modulator name	Target
PF-05082566 / Utomilumab	4-1BB	GWN323	GITR	MOXR0916	OX-40
Urelumab	4-1BB	INCAGN01876	GITR	PF-04518600	OX-40
AZD4635	ADORA2A	MK-1248	GITR	AMP-224	PD-1
CPI-444	ADORA2A	MK-4166	GITR	BGB-A317	PD-1
NIR178	ADORA2A	GSK3359609	ICOS	IBI308	PD-1
PBF-509	ADORA2A	JTX-2011	ICOS	JS001	PD-1
Preladenant / MK-3814 /	4000434	Epacadostat	IDO	MED10680	PD-1
SCH420814	ADORA2A	Indoximod	IDO	Nivolumab	PD-1
Enoblituzumab	B7-H3	KHK2455	IDO	PDR001	PD-1
Varlilumab	CD27	NLG919 / GDC-0919	IDO	Pembrolizumab	PD-1
APX005M	CD40	BMS-986205	IDO	PF-06801591	PD-1
CP-870,893 / RO7009789	CD40	Lirilumab	KIR	REGN2810	PD-1
Dacetuzumab	CD40	BMS-986016	LAG-3	SHR-1210	PD-1
Lucatumumab	CD40	LAG525	LAG-3	Atezolizumab	PD-L1
SEA-CD40	CD40	MK-4280	LAG-3	Avelumab	PD-L1
ISF35 / rAd-CD40L	CD40	REGN3767	LAG-3	Durvalumab	PD-L1
MEDI5083	CD-40L	IMP321	MHC II	FAZ053	PD-L1
ARGX-110	CD70	Monalizumab	NKG2A	LY3300054	PD-L1
Galiximab	CD80	ABBV-368	OX-40	CX-072	PD-L1
BMS-986218	CTLA-4	BMS-986178	OX-40	BMS-986207	TIGIT
Ipilimumab	CTLA-4	GSK3174998	OX-40	MTIG7192A	TIGIT
MK-1308	CTLA-4	MEDI0562	OX-40	LY3321367	TIM-3
Tremelimumab	CTLA-4	MEDI6383	OX-40	MBG453	TIM-3
BMS-986156	GITR	MEDI6469	OX-40	TSR-022	TIM-3

Copyright: Hanson Wade, August 2017

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Nature Reviews | Drug Discovery

Mahoney KM et al. 2015 Nat Rev Drug Discovery

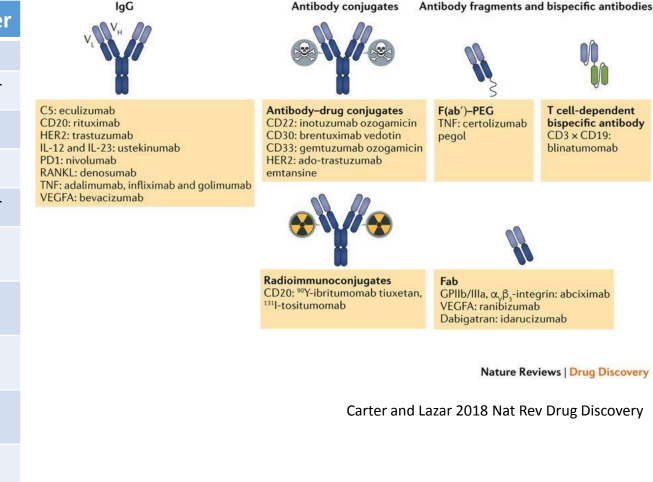
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More development and potential approvals of antibody conjugates vs. cancer

Emerging antibody-drug conjugates	Target cancer
Sacituzumab govitecan	Breast cancer
Mirvetuximab canavanine	Ovarian cancer
Rovalpituzumab tesirine	Lung cancer
Depatuxizumab mafodotin	Glioblastoma
Oportuzumab monatox	Bladder cancer
Denintuzumab mafodotin	B cell malignancies
Indatuximab ravtansine	Multiple myeloma
Lorvotuzumab mertansine	Small cell lung cancer
Moxetumomab pasudotox	B cell malignancies
Pinatuzumab vedotin	B cell malignancies
Polatuzumab vedotin	B cell malignancies



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Oncolytic viral therapy will continue to expand viral agents, biologics, and target tumors

	Adenovirus ^a	Herpes simplex virus ^b	Pox virus	Coxsackie virus	Maraba virus	Poliovirus	Measles virus	Newcastle Disease virus
Genome	dsDNA	dsDNA	dsDNA	ssRNA	ss (–) RNA	ss (+) RNA	ss (–) RNA	ss (–) RNA
Genome size	Moderate (32 kb)	Large (152 kb)	Large (130– 375 kb)	Small (~8 kb)	Small (11– 15 kb)	Small (7.5 kb)	Small (~16 kb)	Small (~15 Kb)
Cell entry mechanism	Endocytosis	Endocytosis; penetration	Membrane penetration and fusion	Micropinocytos is via epithelial tight junctions	Endocytosis; pH-dependent fusion activation	Receptor- mediated endocytosis	Membrane fusion	Endocytosis; pH- independent direct fusion
Cell entry receptors	hCAR; VCAM1; CD46	HVEM; nectin 1; nectin 2	GAGs; EFC	CAR; DAF	Unknown	CD155	CD46; SLAM	Neuraminidase receptor; sialoglyco- conjugates

^aE1B-55 K/E3B-deleted adenovirus in combination with chemotherapy was approved for the treatment of late-stage refractory nasopharyngeal cancer by the Chinese State Food and Drug Administration in 2005. ^bHerpes simplex virus type 1 (HSV-1) with ICP34.5 deletion and encoding granulocyte–macrophage colony-stimulating factor (GM-CSF) was approved for stage III–IV melanoma treatment by the US Food and Drug Administration in 2015 and by Australia and the European Medicines Agency (EMA) in 2016.

Adapted from Bommareddy et al. 2018 Nat Rev Immunol

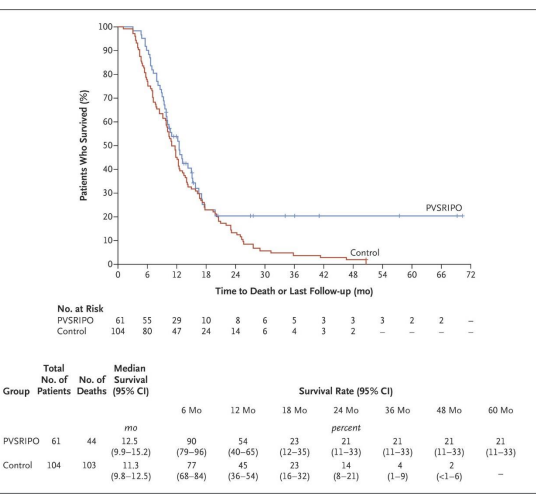






Oncolytic viral therapy will continue to expand viral agents, biologics, and target tumors

PVSRIPO for Glioblastoma multiforme



Desjardins et al. 2018 New Eng J Med

A Study of Intratumoral CAVATAK[™] in Patients With Stage IIIc and Stage IV Malignant Melanoma (VLA-007 CALM) NCT01227551

Outcome	Result
Percentage of Participants With Immune-related Progression- Free Survival (irPFS) at 6 Months	38.6 (26.0 to 52.4)
Percentage of Participants With Durable Response Rate of 6 months or more	21.1

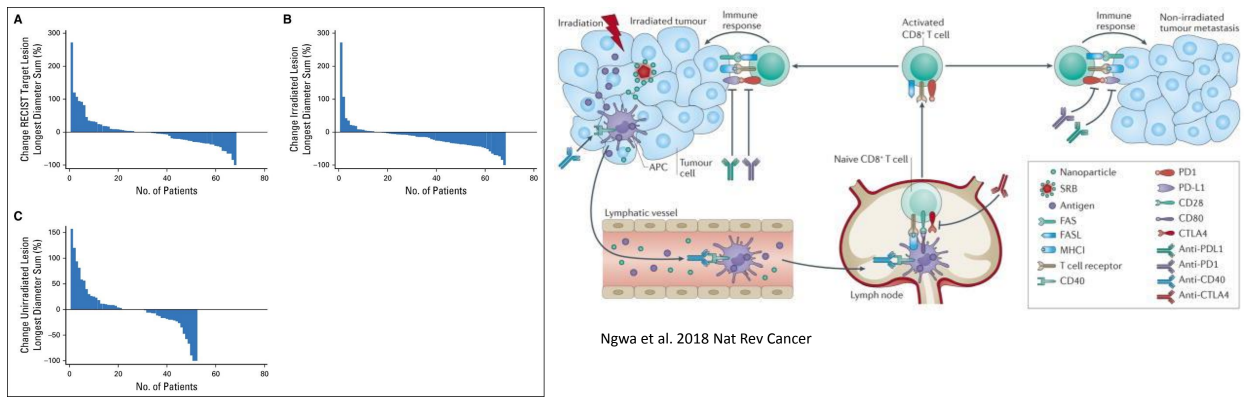




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Radiation therapy will be increasingly used as a means of enhancing immunotherapy



Luke et al. 2018 J Clin Oncol







Radiation therapy can be safely combined with immunotherapy

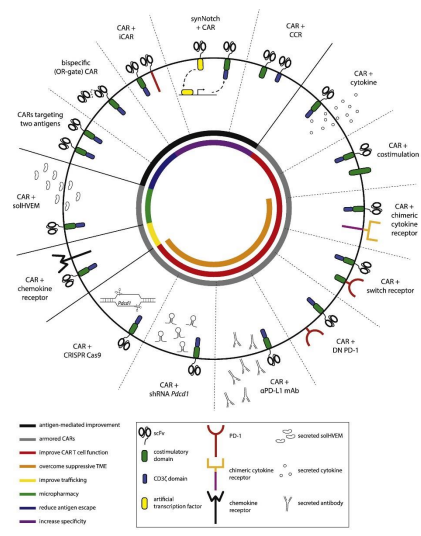
Primary site	n	Radiotherapy	Immunotherapy	Schedule	Nonirradia CR	ted lesions PR	SD	Grade 3+ toxicities
Melanoma	22	 6 Gy × 2–3 or 8 Gy × 2–3 1 site 	Ipilimumab 3 mg/kg every 3 weeks × 4	First ipilimumab 3–5 days after RT	0/22 (0%)	4/22 (18%)	4/22 (18%)	 Number of patients with any grade 3 toxicity not reported Grade 3 anemia (4/22; 18%) most common No grade 4–5 No DLT
Melanoma	22	 Multiple dose-fx regimens (BED10 range 28.0–112.5 Gy) 1–2 sites 	Ipilimumab 3 mg/kg every 3 weeks × 4	RT within 5 days of first ipilimumab	3/22 (14%)	3/22 (14%)	5/22 (23%)	 2/22 (9%) grade 3 1/22 (5%) grade 4 No grade 5
NSCLC, CRC, sarcoma, RCC, and others	35	 50 Gy/4 fx or 60 Gy/10 fx 1 site 	Ipilimumab 3 mg/kg every 3 weeks × 4	RT 1 day after first ipilimumab or 1 week after second ipilimumab	0/31 (0%)	3/31 (10%)	4/31 (13%)	 12/35 (34%) grade 3 No grade 4–5 2/35 (6%) with DLT
	Melanoma Melanoma NSCLC, CRC, sarcoma, RCC, and others	Melanoma 22 Melanoma 22 NSCLC, CRC, sarcoma, RCC, and 35	Melanoma226 Gy × 2–3 or 8 Gy × 2–3 • 1 siteMelanoma22• Multiple dose-fx regimens (BED10 range 28.0–112.5 Gy) • 1–2 sitesNSCLC, CRC, sarcoma, RCC, and 35 others• 50 Gy/4 fx or 60 Gy/10 fx • 1 site	Melanoma22•6 Gy × 2–3 or 8 Gy × 2–3 •Ipilimumab 3 mg/kg every 3 weeks × 4Melanoma22•Multiple dose-fx regimens (BED10 range 28.0–112.5 Gy) •Ipilimumab 3 mg/kg every 3 weeks × 4NSCLC, CRC, sarcoma, RCC, and 35 others•50 Gy/4 fx or 60 Gy/10 fx •Ipilimumab 3 mg/kg every 3 weeks × 4	Melanoma226 Gy × 2–3 or 8 Gy × 2–3 1 siteIpilimumab 3 mg/kg every 3 weeks × 4First ipilimumab 3–5 days after RTMelanoma22• Multiple dose-fx regimens (BED10 range 28.0–112.5 Gy) • 1–2 sitesIpilimumab 3 mg/kg every 3 weeks × 4RT within 5 days of first ipilimumabNSCLC, CRC, sarcoma, RCC, and 35 others• 50 Gy/4 fx or 60 Gy/10 fx • 1 siteIpilimumab 3 mg/kg every 3 weeks × 4RT 1 day after first ipilimumab or 1 week after second ipilimumab	Primary sitenRadiotherapyImmunotherapyScheduleCRMelanoma22 $\begin{array}{c} 6 \ Gy \times 2 - 3 \ or \ 8 \ Gy \times 2 - 3 \ 1 \ siteIpilimumab 3 mg/kgevery 3 weeks \times 4First ipilimumab3 - 5 days afterRT0/22 (0%)Melanoma22\begin{array}{c} Multiple \ dose \ fx \ regimens \ (BED10 \ range 28.0 - 112.5 \ Gy) \ 1 - 2 \ sitesIpilimumab 3 mg/kg \ of \ first \ ipilimumabof \ first \ ipilimumab3/22 (14%)NSCLC, CRC, sarcoma, RCC, and others35\begin{array}{c} 50 \ Gy/4 \ fx \ or \ 60 \ Gy/10 \ fx \ 1 \ siteIpilimumab 3 mg/kg \ every 3 weeks \times 4RT 1 \ day after first \ ipilimumab or 1 \ week \ after \ second \ ipilimumab0/31 (0%)$	Primary sitenRadiotherapyImmunotherapyScheduleCRPRMelanoma226 Gy × 2-3 or 8 Gy × 2-3 1 siteIpilimumab 3 mg/kg every 3 weeks × 4First ipilimumab 3-5 days after RT0/22 (0%)4/22 (18%)Melanoma22CMultiple dose-fx regimens (BED10 range 28.0-112.5 Gy) 1 -2 sitesIpilimumab 3 mg/kg every 3 weeks × 4RT within 5 days of first ipilimumab3/22 (14%)3/22 (14%)NSCLC, CRC, sarcoma, RCC, and others3550 Gy/4 fx or 60 Gy/10 fx 1 siteIpilimumab 3 mg/kg every 3 weeks × 4RT 1 day after first ipilimumab or 1 week after or 1 week aft	Melanoma22• 6 Gy × 2-3 or 8 Gy × 2-3 • 1 siteIpilimumab 3 mg/kg every 3 weeks × 4First ipilimumab 3-5 days after RT0/22 (0%)4/22 (18%)4/22 (18%)Melanoma22• Multiple dose-fx regimens (BED10 range 28.0-112.5 Gy) • 1-2 sitesIpilimumab 3 mg/kg every 3 weeks × 4RT within 5 days of first ipilimumab3/22 (14%)3/22 (14%)5/22 (23%)NSCLC, CRC, sarcoma, RCC, and 35 others• 50 Gy/4 fx or 60 Gy/10 fx • 1 siteIpilimumab 3 mg/kg every 3 weeks × 4RT 1 day after first ipilimumab or 1 week after second ipilimumab0/31 (0%)3/31 (10%)4/31 (13%)

Ko and Formenti 2018 Ther Adv Med Oncl

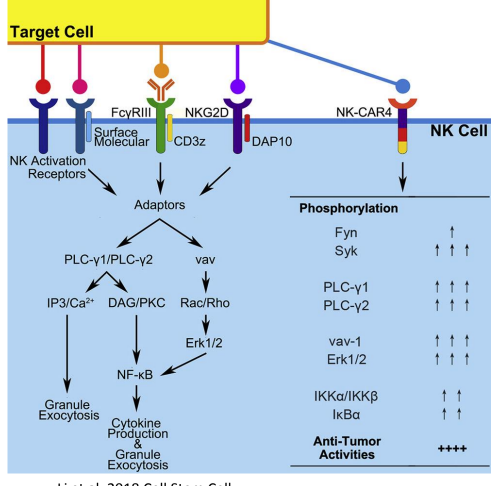




Enhanced engineering of CAR T and CAR NK cells may Society for Immunotherapy of Cancer help reduce side effects while improving efficacy



Jaspers and Brentjens 2017 Pharmacol Ther



Li et al. 2018 Cell Stem Cell







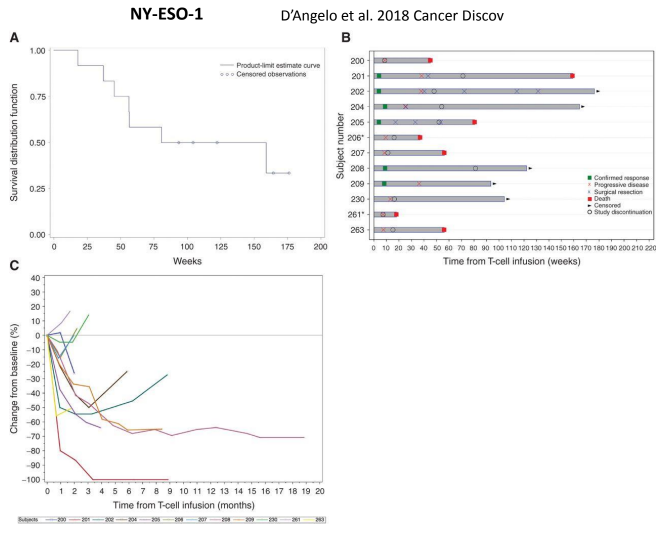
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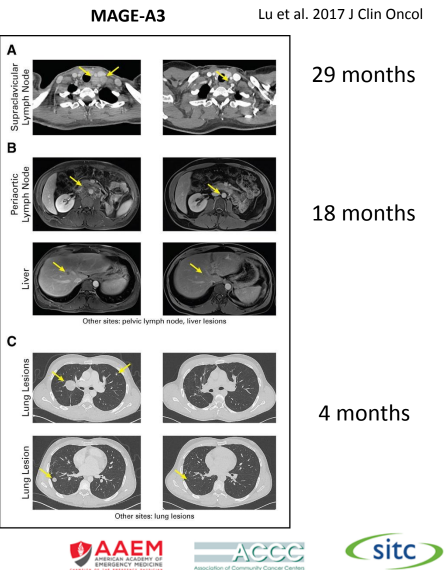
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TCR transduced T cells will provide durable responses in solid tumors





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Conclusions

- The future of cancer immunotherapy is bright, with advances occurring in both diagnostics and therapeutics at a rapid pace
- We will see improvements in our ability to distinguish immunologically "hot" vs. "cold" tumors, and potentially be able to convert "cold" into "hot" tumors
- Advances in genetic engineering and biomanufacturing will permit development of "next generation" antibodies, viruses, targeted radiotherapies and cellular therapies for cancer.

