

Toxicity Management

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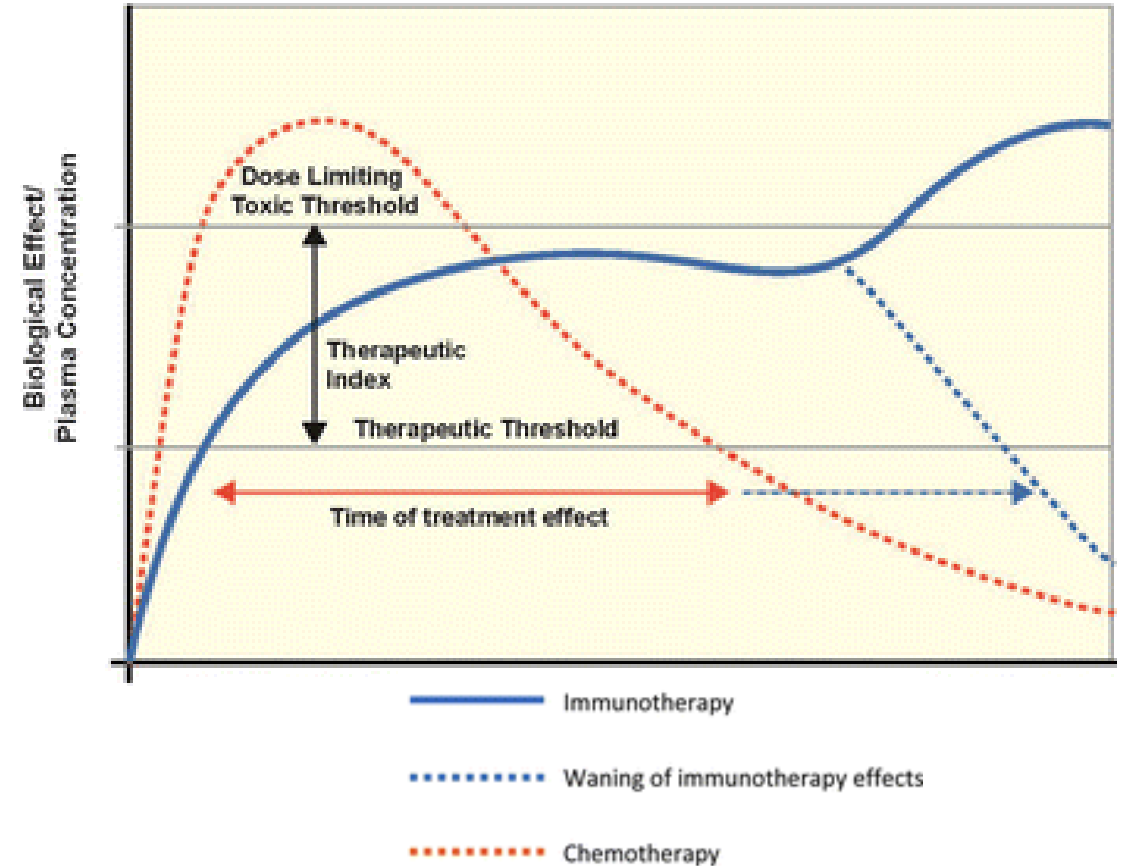
Lombardi Comprehensive Cancer Center

Disclosures

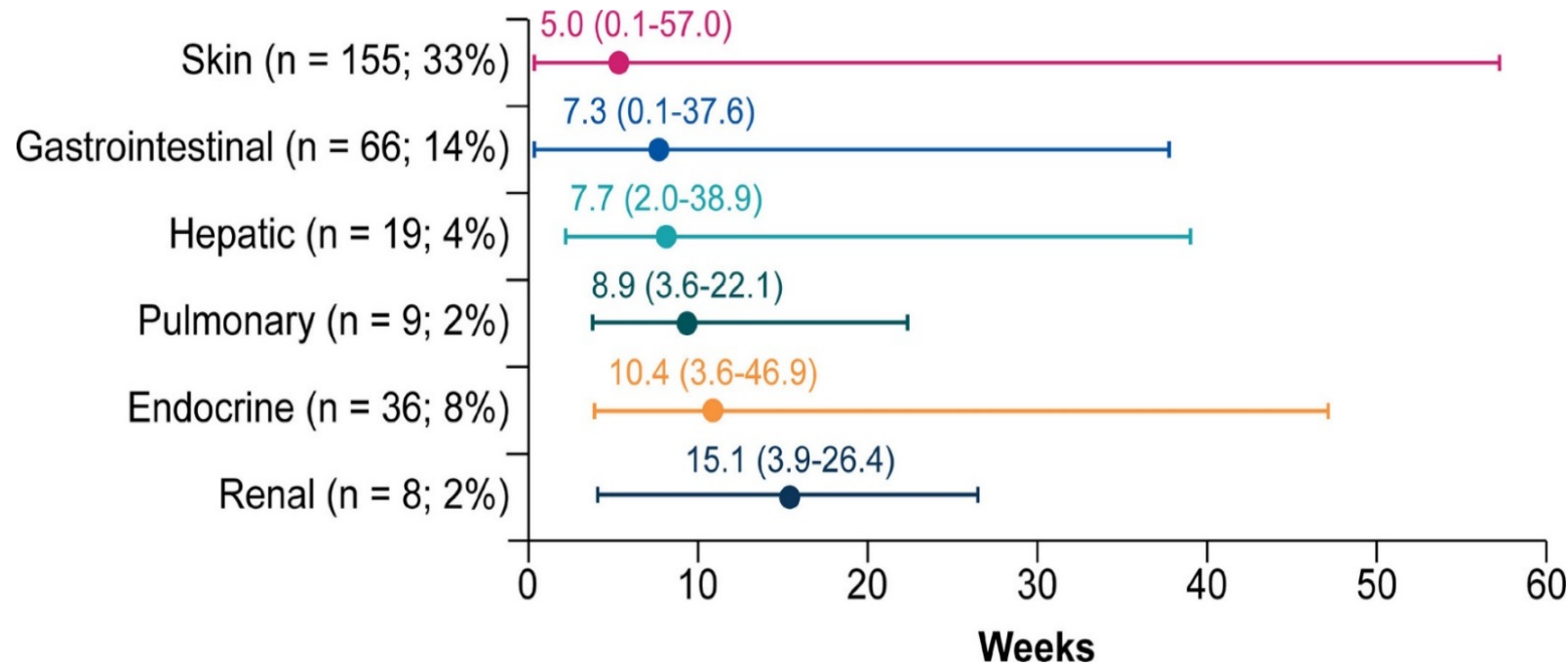
- Consulting Fees: Eisai, Exelixis
- I will be discussing non-FDA approved indications during my presentation.

Immune-related adverse events (irAEs)

- Immune checkpoint inhibitor (ICI) toxicities often have delayed onset and prolonged duration relative to chemotherapy toxicity
- Toxicities result from non-specific activation of the immune system and can mimic a number of other medical conditions



Onset of irAEs



- Can be days to months after therapy initiation
- May occur even after treatment is discontinued
- Important to identify patients who are currently **OR** previously on ICI treatment!

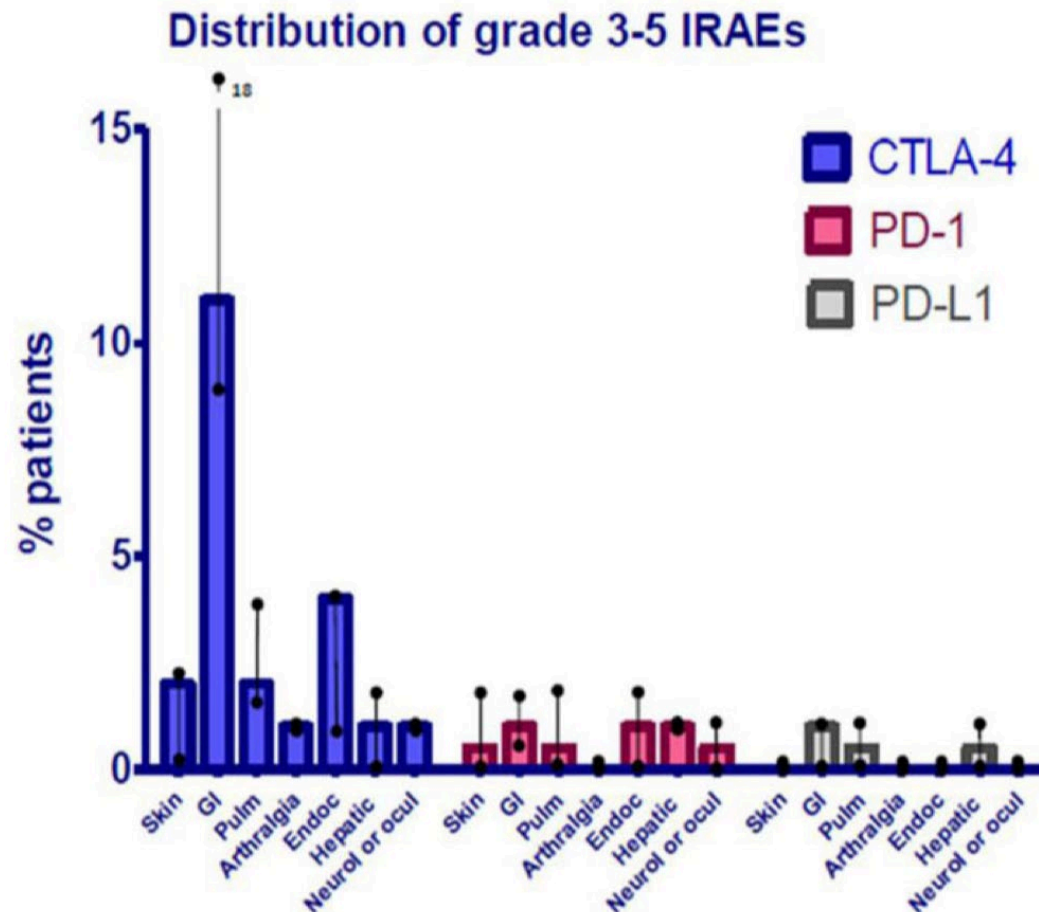
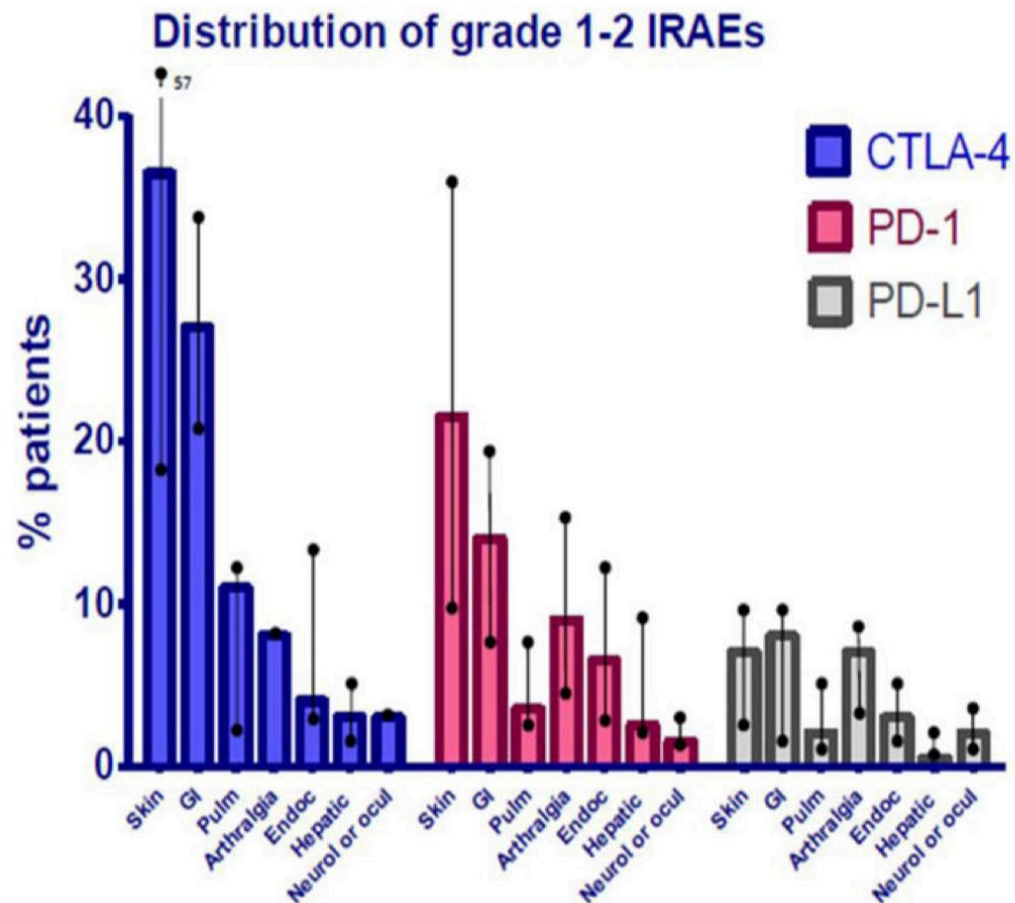
Incidence of irAEs

- Overall incidence of all-grade irAEs with single-agent ICI reported as 15-90% in studies
- Anti-CTLA-4 inhibitor (ipilimumab): dose-dependent toxicities
 - Any grade toxicity \leq 75% (**Grade 3+: \leq 43%**)
- PD-1/PD-L1 inhibitors: toxicities less dose-dependent
 - Any grade toxicity \leq 30% (**Grade 3+: \leq 20%**)
- Life-threatening irAEs are rare but treatment-related deaths reported in up to 2% of clinical trial patients

Incidence of specific irAEs by ICI

Drug	Dermatitis	Colitis	Hepatitis	Endocrinopathies	Pneumonitis
	All grades (grade 3-4)				
Ipilimumab	14.5 (12)	10 (7)	5 (2)	10 (3)	<1
Ipilimumab/Nivolumab	30 (3)	26 (16)	13 (6)	35 (4)	6 (2.2)
Nivolumab	28 (1.5)	2.9 (0.7)	1.8 (0.7)	12 (0)	3.1 (1.1)
Pembrolizumab	20 (0.5)	1.7 (1.1)	0.7 (0.4)	12.5 (0.3)	3.4 (1.3)
Atezolizumab	17 (0.8)	1 (<1)	1.3 (<1)	5.9 (<1)	2.6 (<1)
Avelumab	15 (0.4)	1.5 (0.4)	0.9 (0.7)	6.5 (0.3)	1.2 (0.5)
Durvalumab	11 (1)	1.3 (0.3)	1.1 (0.6)	16.2 (0.1)	2.3 (0.5)

Severity of irAEs by ICI



Common irAEs with ICI's

- 1 Dermatologic: maculopapular rash, dermatitis, pruritis
- 2 Gastrointestinal: diarrhea, colitis, hepatitis, gastritis
- 3 Rheumatologic: arthralgias, myositis, sicca symptoms
- 4 Pulmonary: pneumonitis, sarcoidosis
- 5 Endocrine: thyroid dysfunction, hypophysitis

Uncommon irAEs with ICI's

Cardiovascular:

Myocarditis, pericarditis,
arrhythmias

Renal:

Interstitial nephritis,
granulomatous nephritis

Endocrine:

Adrenal insufficiency,
pancreatitis, type 1
diabetes mellitus

Hematologic:

Hemolytic anemia, red
cell aplasia, neutropenia,
thrombocytopenia

Neurologic:

Myasthenia gravis,
Guillain-Barré syndrome,
peripheral neuropathies

Ophthalmologic:

Uveitis, episcleritis,
conjunctivitis

Pre-treatment screening

- Patient History
 - Autoimmune, infectious, endocrine, organ-specific diseases
 - Baseline bowel habits
- Dermatologic
 - Full skin and mucosal exam
- Pulmonary
 - Baseline O₂ saturation
- Cardiovascular
 - ECG
 - Troponin I or T
- Blood tests
 - CBC with diff
 - CMP
 - TSH and free T4
 - HbA1c
 - Total CK
 - Fasting lipid profile
 - Infectious disease screen:
 - Hepatitis serologies
 - CMV antibody
 - HIV antibody and antigen (p24)
 - TB testing (T-spot, quantiferon gold)

Additional screening for high-risk patients

- Endocrine tests
 - 8 am cortisol and ACTH
- Cardiac tests
 - Brain natriuretic peptide (BNP) or N-terminal pro B-type natriuretic peptide (NT pro-BNP)
- Pulmonary tests
 - PFTs
 - 6MWT

Approach to Treatment

- Treatment approach is guided by grading of specific toxicity
- Resources for grading:
 - SITC Toxicity Management Working Group
 - Common Terminology Criteria for Adverse Events
 - National Comprehensive Cancer Network
- 1st line for **MOST** irAE's is systemic high-dose corticosteroids
 - Endocrine toxicities managed with hormone replacement
 - Some grade 1-2 irAEs may respond to topical steroids (dermatologic, ophthalmologic)
- OTC drugs may not be appropriate for managing symptoms
 - i.e. loperamide for colitis may result in bowel perforation

General corticosteroid management

Grade of irAE	Corticosteroid Management	Additional Notes
1	Usually not indicated	Continue immunotherapy
2	<ul style="list-style-type: none"> Start prednisone 0.5-1 mg/kg/day (or equivalent dose of IV methylprednisolone) If no improvement in 2-3 days, increase dose to 2 mg/kg/day Once improved to ≤grade 1, 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

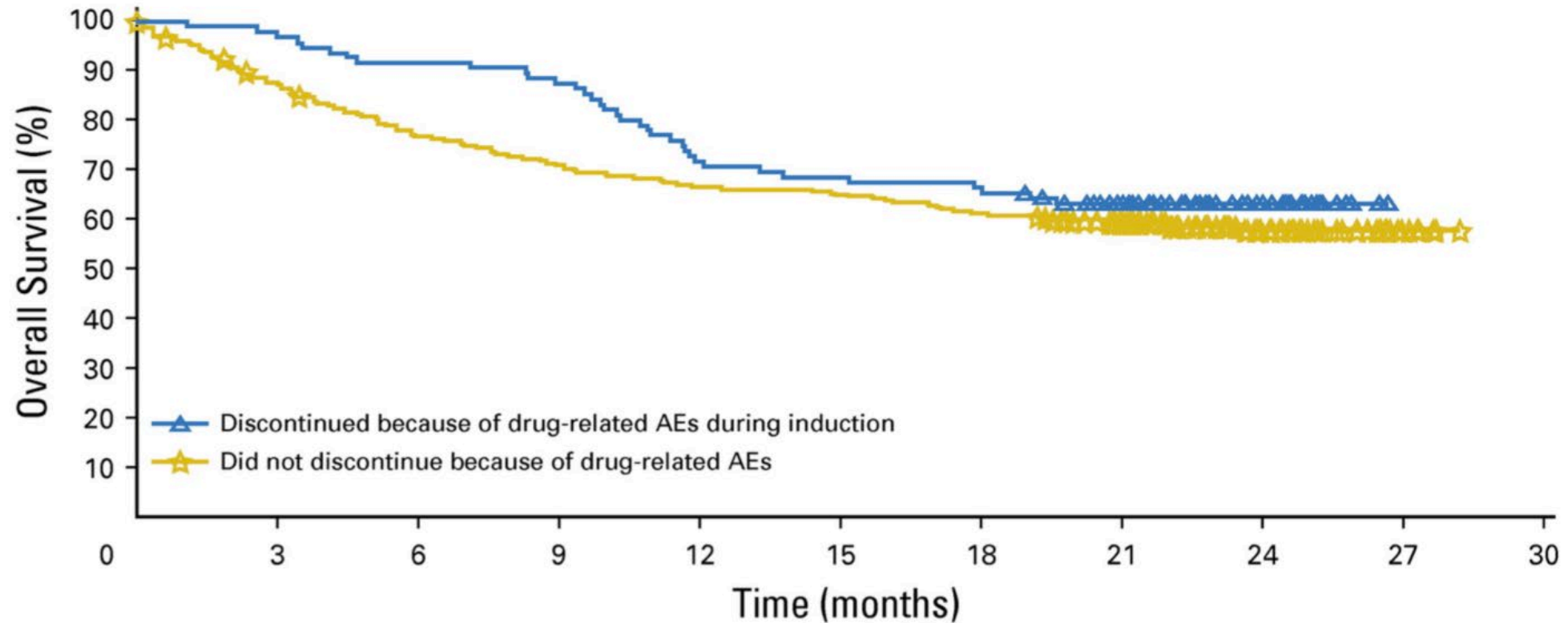
General corticosteroid management

Grade of irAE	Corticosteroid Management	Additional Notes
3	<ul style="list-style-type: none"> Start prednisone 1-2 mg/kg/day (or equivalent dose of IV methylprednisolone) If no improvement in 2–3 days, ADD additional immunosuppressant Once improved to \leq grade 1, start 4–6-week steroid taper 	<ul style="list-style-type: none"> Hold immunotherapy; if symptoms do not improve in 4–6 weeks, discontinue immunotherapy Start proton pump inhibitor for GI prophylaxis Add PJP prophylaxis if more than 3 weeks of immunosuppression expected (>30 mg prednisone or equivalent/day)
4		<ul style="list-style-type: none"> Discontinue immunotherapy Start proton pump inhibitor for GI prophylaxis Add PJP prophylaxis if more than 3 weeks of immunosuppression expected (>30 mg prednisone or equivalent/day)

Additional immunosuppressives

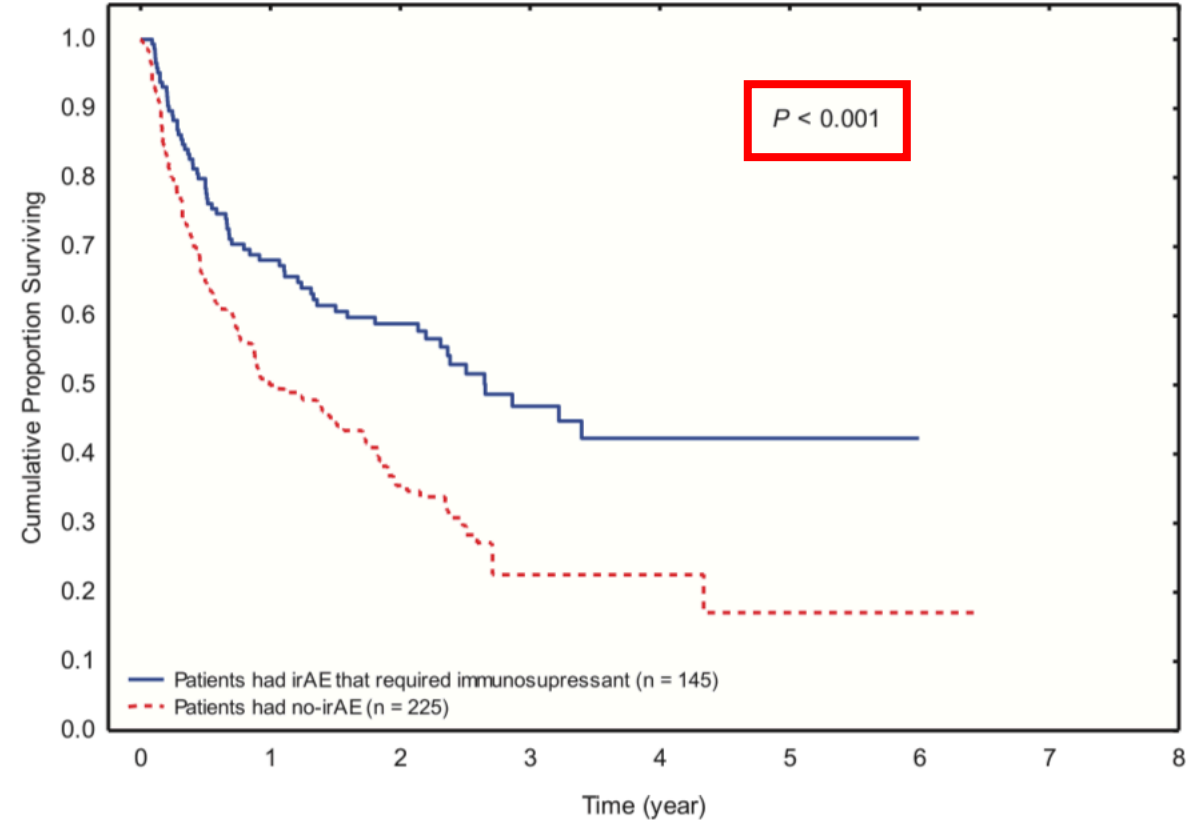
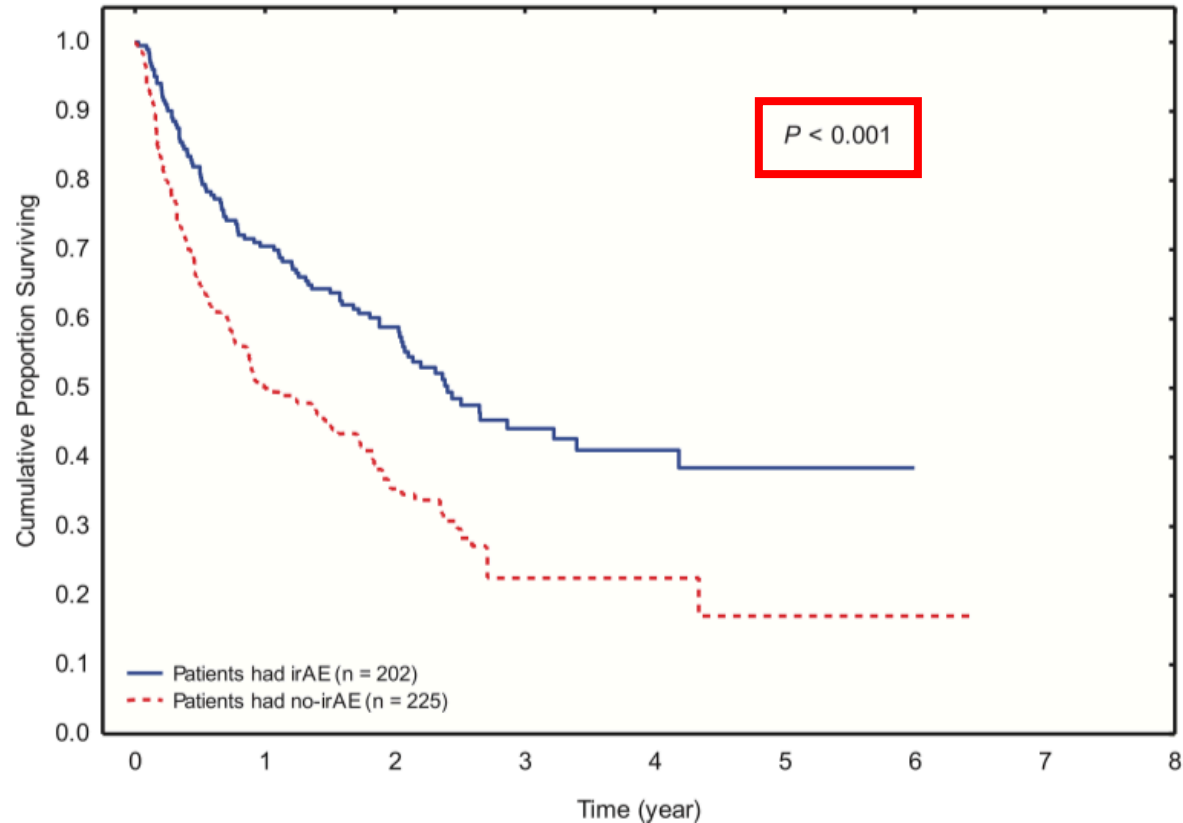
- **Infliximab: anti-TNF- α mAb**
 - Hepatotoxic so should NOT be used for immune-mediated hepatitis
 - Risk for hepatitis B and tuberculosis activation; obtain hepatitis serologies and TB testing prior to initiation
 - Dose: 5 mg/kg; 2nd dose may be administered after 2 weeks
- **Vedolizumab: $\alpha 4\beta 7$ integrin mAb**
 - **Selective GI immunosuppression** → inhibits migration of T cells across endothelium into inflamed GI tissues
 - Dose: 300 mg; repeat dose at 2 and 6 weeks
- **Others: mycophenolate, IVIG, tacrolimus**

Effect of irAEs on patient outcomes



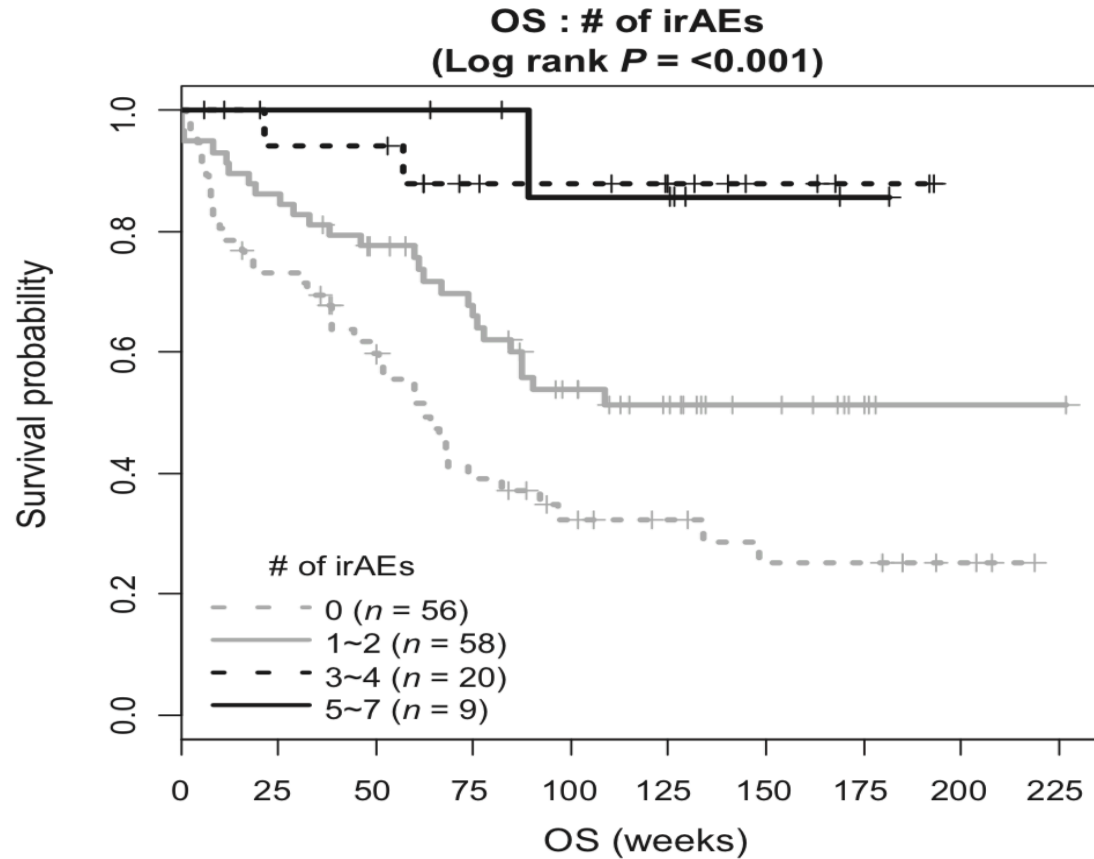
No significant difference in survival in melanoma patients who discontinued ipilimumab + nivolumab due to irAEs versus those who did not discontinue treatment

Autoimmunity as prognostic marker?

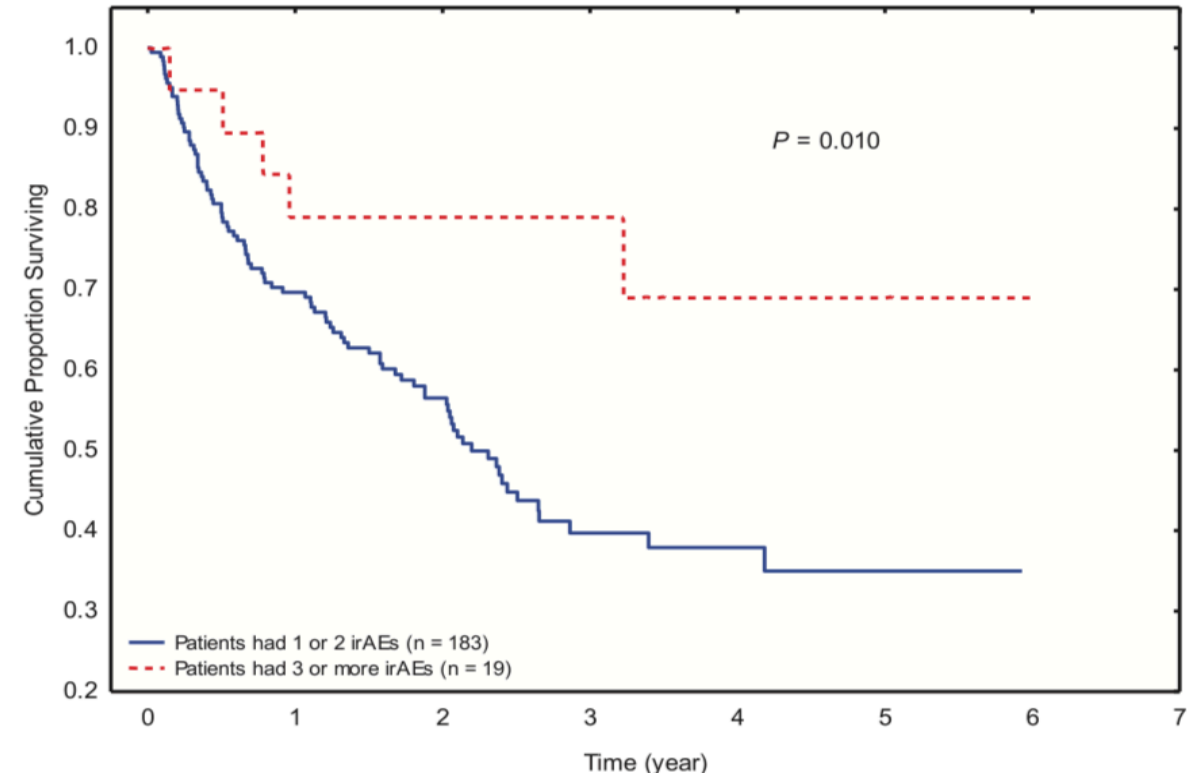


Based on **retrospective** data, patients who experience irAEs (regardless of needing treatment) may have better outcomes compared to patients who do not experience irAEs

Number of irAEs on patient outcomes

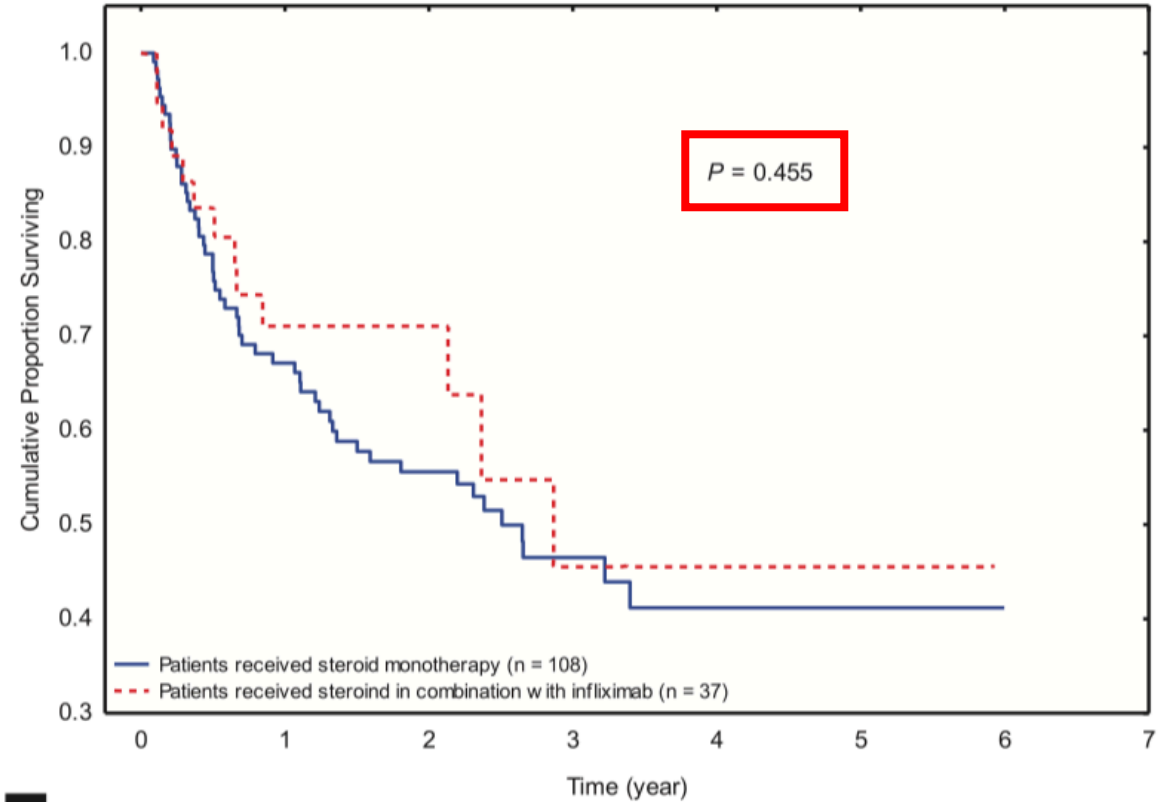
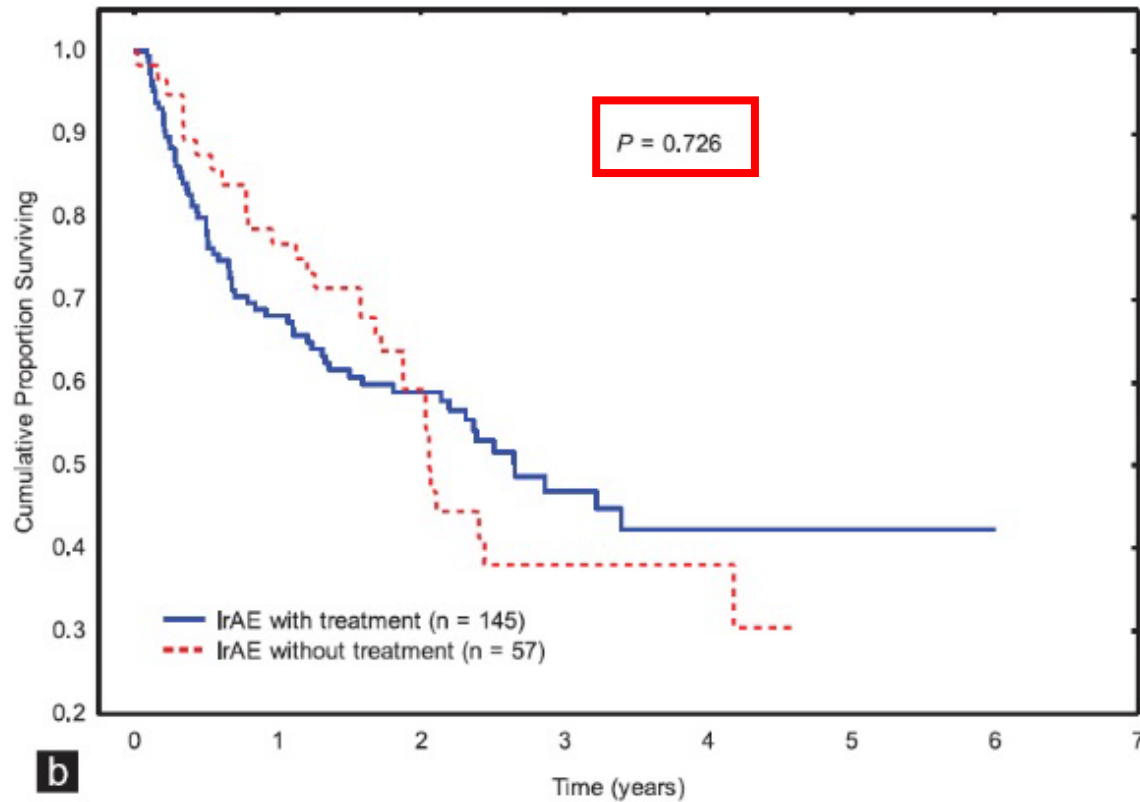


Nivolumab in metastatic melanoma: greater OS
 in patients with 3+ irAEs versus ≤ 1 irAE



Patients receiving ICI's for various malignancies:
 greater OS in those with 3+ irAEs versus ≤ 2 irAEs

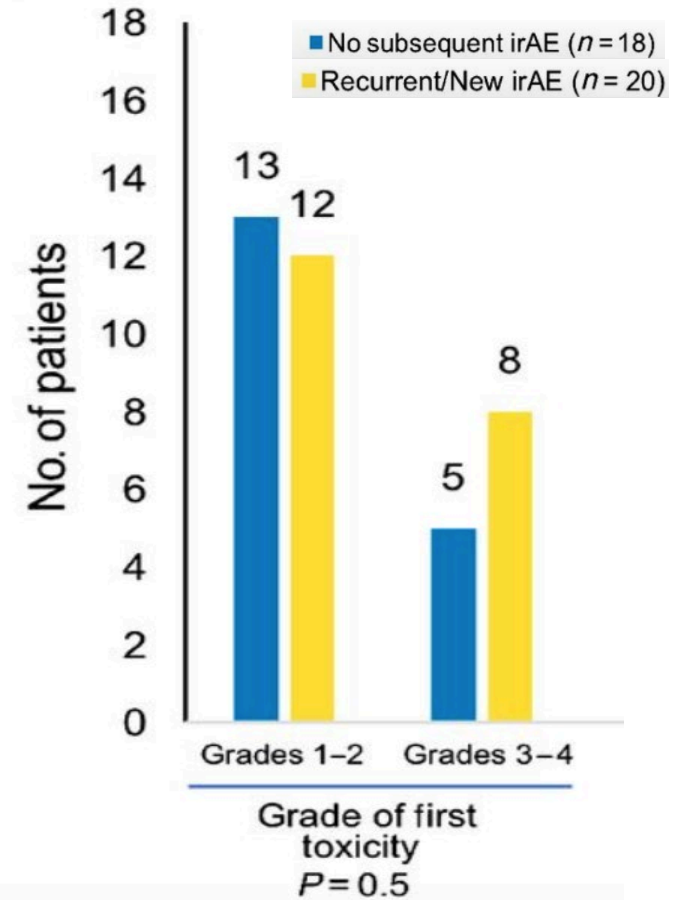
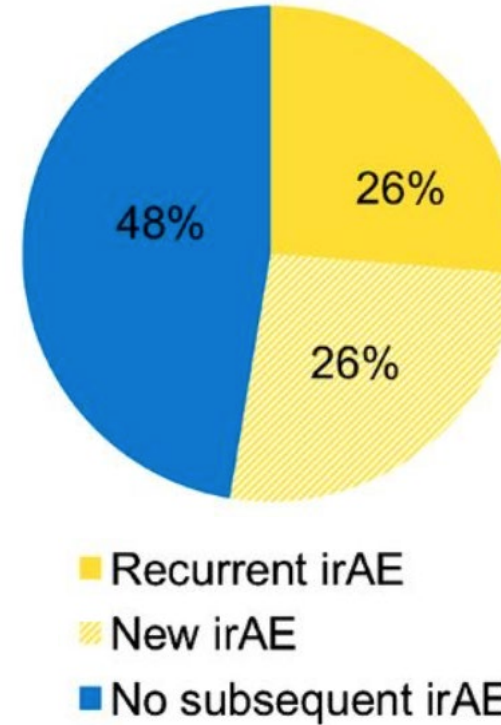
Impact of toxicity management on patient outcomes



While still under debate, the administration of immunosuppressive treatments NOR the type of immunosuppressant used for irAE management does not seem to impact cancer control

Rechallenging with ICI after irAEs

- Patients should not be rechallenged until irAE resolved to grade ≤ 1
- Re-challenge with anti-PD-1/L1 after anti-CTLA-4 \pm anti-PD-1 likely safe
- Caution in re-challenging with same ICI in patients who previously had grade 3-4 irAEs



Patients with autoimmune disorders

- Ipilimumab in melanoma patients
 - 29% experienced flare of pre-existing disorder; 29% experienced new irAEs
 - 56% experienced no flare OR additional irAEs
- PD-1 in melanoma patients
 - 38% experienced flare; 29% experienced new irAEs
 - Lower response rates in patients who remained on immunosuppressive treatment (15% vs 44%)
- Efficacy appears similar for patients with autoimmune disorders compared to those without

ICI use in SOT or SCT

- Patients who relapse after allogeneic SCT:
 - Ipilimumab: 32% response (10 mg/kg); 14% GVHD; 21% irAEs
 - Anti-PD-1: 77% response; 26% died due to new-onset GVHD
- Solid organ data is limited; most is in renal SOT patients
 - One retrospective study (n=39) reported graft loss in 81% and death in 46%
 - Also reported rapid time to rejection with median onset of 21 days
- PD-1 pathway appears to be more critical in allograft immune tolerance compared to CTLA-4 pathway

CAR T-cell related toxicities

More
Common

Cytokine release syndrome

Immune cell associated neurotoxicity syndrome (ICANS)

Less
Common

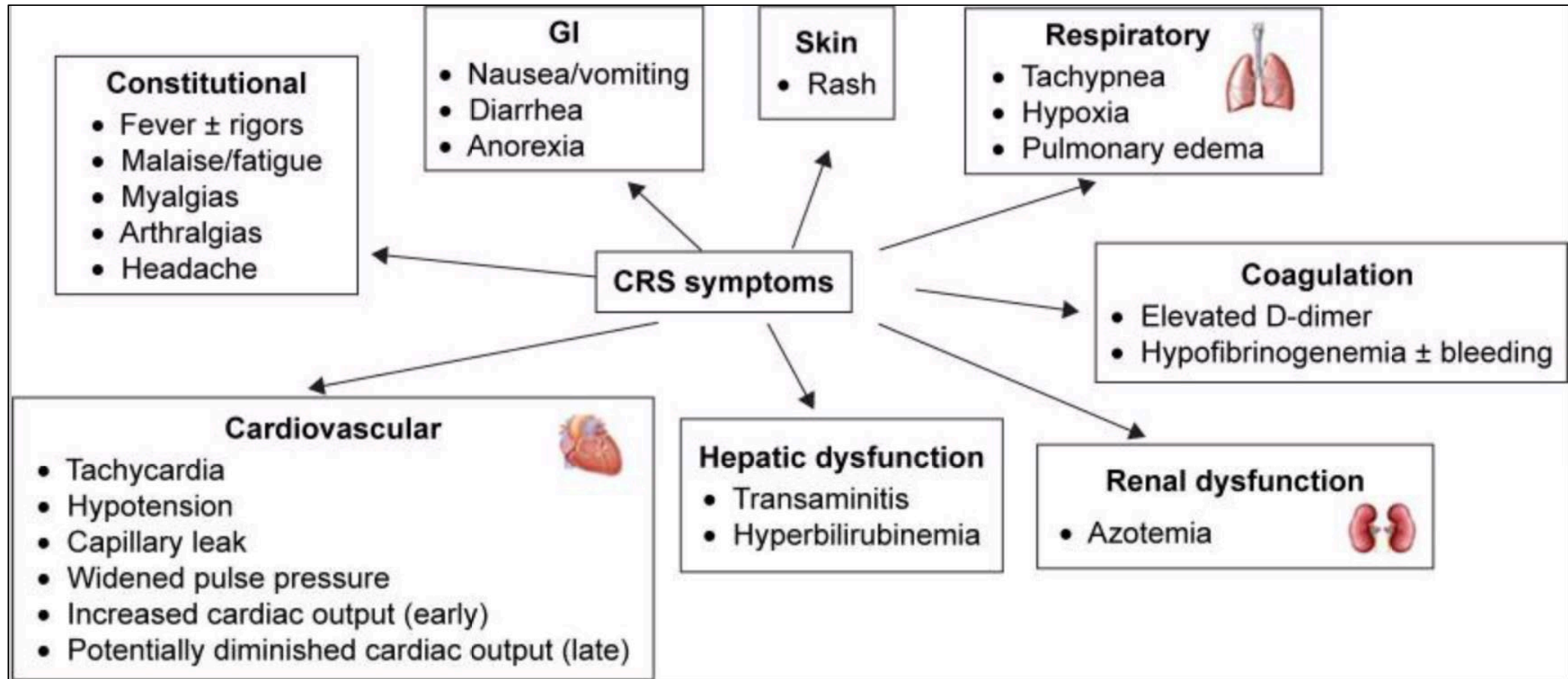
Hemophagocytic Lymphohistiocytosis/
Macrophage Activation Syndrome (HLH/MAS)

Anaphylaxis, B cell aplasia and
hypogammaglobulinemia

CRS and Neurotoxicity

- Should not be viewed as two unrelated adverse events
 - Overlapping toxicities from excessive immune activation
 - May occur together or exclusive of one another
 - However, they do have distinct timing and responses to treatment
- Risk factors for both include:
 - High disease burden
 - Higher infused CAR-T cell dose
 - High intensity lymphodepletion regimen
 - Pre-existing endothelial activation
 - Severe thrombocytopenia

Cytokine release syndrome



Cytokine release syndrome

- Occurs in ~70% of patients; severe = 12-47%
 - Median onset 2-3 days after infusion, typical duration 7-8 days
- Multiple grading systems exist (MSKCC, CarTox, ASTCT)
 - Hypotension and hypoxia are main drivers of CRS severity
- Tocilizumab approved for CRS treatment (blocks IL-6R)
 - Dose for patients >30 kg: 8 mg/kg (up to 800 mg/dose)
 - May be repeated every 8 hours up to 4 doses
- Consider adding dexamethasone 10 mg q6h for grade 3-4 CRS and/or refractory to tocilizumab

Neurotoxicity

- Also called CAR-T Related Encephalopathy Syndrome (CRES) or iIEC-associated neurologic syndrome (ICANS)
- Occurs in 20-64% of patients, \geq grade 3 in 11-42%
 - Onset 4-5 days after infusion, typical duration 5-12 days
- Common symptoms include encephalopathy, headache, delirium, anxiety, tremor, aphasia
 - Severe neurotoxicity: seizures, cerebral edema, hemi/paraparesis
- Diagnosis usually based on clinical symptoms
 - MRI/CT often negative although ~30% will have abnormal MRI (poorer outcome)
- Also has multiple grading systems which guide treatment
 - Usually includes early use of high-dose steroids (dexamethasone 10 mg IV q6h)

HLH/MAS

- Inflammatory syndrome caused by hyperactivation of macrophages and lymphocytes
- Rare; frequency reported to be as low as ~1%
- Should be managed with anti-IL-6 and corticosteroid therapy
- If no improvement after 48 hours, consider adding etoposide for additional immunosuppression
 - Dose: 75-100 mg/m²
 - May be repeated after 4-7 days

Box 5 | Diagnostic criteria for CAR-T-cell-related HLH/MAS

A patient might have HLH/MAS if he/she had a peak serum ferritin level of >10,000 ng/ml during the cytokine-release syndrome phase of CAR-T-cell therapy (typically the first 5 days after cell infusion) and subsequently developed any two of the following:

- Grade ≥3 increase in serum bilirubin, aspartate aminotransferase, or alanine aminotransferase levels*
- Grade ≥3 oliguria or increase in serum creatinine levels*
- Grade ≥3 pulmonary oedema*
- Presence of haemophagocytosis in bone marrow or organs based on histopathological assessment of cell morphology and/or CD68 immunohistochemistry

The importance of patient education

- Many immune-related adverse events can present in similar ways to other disease states, but the treatment of them is very different.
- Patients may not go back to their oncologist for treatment of irAEs and need to identify themselves as immunotherapy recipients
 - Emergency room & general practitioners need to understand the proper identification and management of irAEs
- Reassure patients that irAEs will likely resolve over time (except endocrinopathies)

Education along the healthcare continuum

- Patients may not go back to their original clinic for adverse event management
- Emergency departments and primary care physicians need to recognize and know how to manage irAEs
- For example, the most common irAE in emergency departments is diarrhea – recognize immune-related symptoms versus other causes

Case 1

- 50-year old male with metastatic melanoma and lung metastases
- PMH: Hypertension, well-controlled
- S/p 3 cycles of nivolumab + ipilimumab
- Presents to clinic reporting watery diarrhea over the past few days and abdominal discomfort
 - 8 episodes/24 hour period (grade 3)
 - No fevers

Case 1

Work up

- Differential diagnosis
 - Infectious colitis
 - Immunotherapy-induced colitis
- Work up
 - Stool sample: Stool culture, Stool WBC, C. Difficile toxin, Ova & parasites
 - CT scan A/P or colonoscopy
 - CT scan shows: Rectosigmoid colon is decompressed with new wall thickening, and mild adjacent pericolic fat stranding, consistent with distal colitis.

Case 1

Treatment

- Hospital Admission with IV hydration
- IV methylprednisone 2mg/kg day
 - Pt responds well to IV steroids with improvement in symptoms
 - Discharged after 2 days of IV steroids on oral prednisone steroid taper
- Permanently discontinue ipilimumab
- Can consider resuming anti PD-1 after resolution of colitis and steroid taper
- Early recognition/intervention is crucial to effective management of toxicities

Case 2

- 64-year old woman with Stage IIIB resected melanoma, now on adjuvant anti-PD1
- PMH: non-contributory
- s/p 2 cycles and presents to clinic for consideration of C3
- Notes mild fatigue since C1, now with mildly pruritic rash on forearms
- PE: Gr1 maculopapular rash on forearms, no blisters, afebrile, pt appears comfortable

How would you manage this patient?

Case 2

- 1) Hold treatment, bring patient back in one week to evaluate
- 2) Continue with C3 nivolumab, consider oral antihistamine and mild-moderate potency topical steroids for affected areas
- 3) Start oral steroids (0.5-1mg/kg) daily with steroid taper over 4 weeks

Case 2

- 1) Hold treatment, bring patient back in one week to evaluate
- 2) Continue with C3 nivolumab, consider oral antihistamine and mild-moderate potency topical steroids for affected areas prn**
- 3) Start oral steroids (0.5-1mg/kg) daily with steroid taper over 4 weeks

Case 2

- Rash was Gr1 by BSA and symptoms
 - Can proceed with immunotherapy with use of antihistamines and/or topical steroids prn
 - Close monitoring/reporting indicated due to risk of worsening symptoms
- Pt now presents prior to C4 complaining of worsening rash/pruritus
 - Rash affecting back/chest and forearms with moderate pruritus
 - Difficulty sleeping due to pruritus even with antihistamines/topical steroids
 - No blisters, mouth sores, fevers

How would you manage this patient?

Case 2

- 1) Continue moderate potency topical steroids and oral antihistamines, proceed with cycle 4
- 2) Hold immunotherapy, have patient return in one week for evaluation
- 3) Hold immunotherapy, start oral corticosteroids (0.5-1mg/kg daily), consider dermatology referral and hospitalization if symptoms fail to improve or worsen

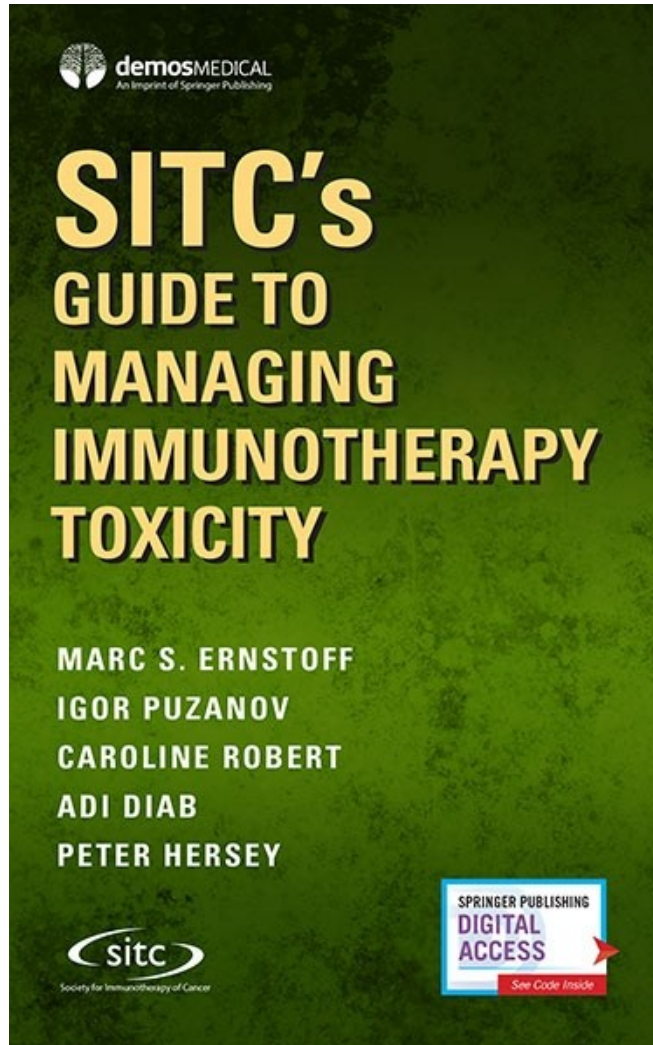
Case 2

- 1) Continue moderate potency and oral antihistamines, proceed with cycle 4
- 2) Hold immunotherapy, have patient return in one week for evaluation
- 3) Hold immunotherapy, start oral corticosteroids (0.5-1mg/kg daily), consider dermatology referral and hospitalizations if symptoms fail to improve or worsen**

Case 2

- Now Gr3 by BSA/symptoms
- Treatment should be held given worsening symptoms
- Start oral steroids at 0.5-1mg/kg daily, can increase to 2mg/daily if needed
- Dermatology referral
- Consider inpatient treatment if symptoms worsening or patient not responding to oral steroids/unable to tolerate oral medications


Additional Resources



Puzanov et al. *Journal for Immunotherapy of Cancer* (2017) 5:95
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
Journal for Immunotherapy of Cancer

POSITION ARTICLE AND GUIDELINES **Open Access**



Managing toxicities associated with immune checkpoint inhibitors: consensus recommendations from the Society for Immunotherapy of Cancer (SITC) Toxicity Management Working Group

I. Puzanov^{1†}, A. Diab^{2†}, K. Abdallah³, C. O. Bingham III⁴, C. Brogdon⁵, R. Dadu², L. Hamad¹, S. Kim², M. E. Lacouture⁶,
 N. R. LeBoeuf⁷, D. Lenihan⁸, C. Onofrei⁹, V. Shannon², R. Sharma¹, A. W. Silk¹², D. Skondra¹⁰, M. E. Suarez-Almazor²,
 Y. Wang², K. Wiley¹¹, H. L. Kaufman^{12†}, M. S. Ernstoff^{1††} and on behalf of the Society for Immunotherapy of Cancer
 Toxicity Management Working Group


 National Comprehensive Cancer Network®

NCCN Clinical Practice Guidelines in Oncology (NCCN Guidelines®)

Management of Immunotherapy-Related Toxicities