# Supportive Care

Keith Eaton, MD, PhD September 2020

# What is supportive care?

- Encompasses significant amount of what an oncologist does widely applicable
- Not specific to any oncologic disease
- Aimed at improving symptoms and tolerance of therapy
- Multiple topics
- Guidelines by NCCN, MASCC, ASCO, and others
- Supportive Care, Survivorship, and Communication = 10% boards

# Topics – covered today

**Antiemesis** 

Anemia

Myeloid growth factors

Skeletal

**Fatigue** 

Brief reviews – neuropathy, cachexia

**Not covered:** pain, mucositis, GI, distress, palliative care, infections, survivorship, chemotherapy dosing, IV access, immunotherapy toxicity management, communication ...

# Antiemesis

## **CINV** Introduction

- N/V are the most common and feared symptoms of cancer chemotherapy
- Management of these symptoms is the most important determinant of the patient experience
- Innovation in this area has undoubtedly improved QOL and likely survival though improved adherence

## Potential problems due to N/V:

- Metabolic disturbances
- Dehydration
- Anorexia
- Decline in PS
- Wound complications, esophageal tears
- Withdrawal from treatment

## **Definitions**

 <u>Acute onset N/V</u> usually occurs within minutes to hours after chemotherapy administration and , it peaks after ~ 6 hours and commonly resolves within 24 hours

# **Delayed CINV**

- Delayed = (>24hrs)
- Common with platins, cyclophosphamide, doxorubicin
- Cisplatin peaks at 48-72 hours, can last up to a week
- The risk of N/V extends to at least 4 days after drug is given for agents of moderate to high emetogenic potential and patients should be protected through this period

# **Anticipatory CINV**

- N/V before next chemotherapy
- a conditioned response
- estimates range from 20-60%
- main indication for benzodiazepines (lorazepam) in CINV

# Refractory/Breakthrough CINV

 Breakthrough emesis occurs despite prophylactic treatment and/or requires "rescue" antiemetics

 <u>Refractory</u> emesis may occur during subsequent cycles following ineffective treatment in earlier cycles

## **CINV Risk Factors**

#### **Acute**

Patient-related factors

- Age
- Gender
- Alcohol use
- previous CINV
- History of anxiety
- Prone to motion sickness
- Morning sickness during pregnancy

Chemotherapy-related factors

- Emetogenicity
- Combination regimens, dose
- Number of cycles
- Unfractionated regimens
- Infusion time

#### <u>Delayed</u>

Any predictive factor for acute CINV

Poor control of acute CINV

Concomitant drugs after chemotherapy

(i.e. opioids, antibiotics)

Low QOL score

## **Emetogenicity of Chemotherapy**

- No universal classification system, NCCN guidelines
- High (> 90%) of patients experience emesis
- Moderate (30-90%)
- Low (10-30%)
- Minimal (< 10%)</li>



#### EMETOGENIC POTENTIAL OF PARENTERAL ANTICANCER AGENTS<sup>a</sup>

LEVEL	AGENT		
High emetic risk (>90% frequency of emesis) <sup>b,c,d</sup>	<ul> <li>AC combination defined as any chemotherapy regimen that contains an anthracycline and cyclophosphamide</li> <li>Carboplatin AUC ≥4</li> </ul>	<ul> <li>Carmustine &gt;250 mg/m²</li> <li>Cisplatin</li> <li>Cyclophosphamide &gt;1,500 mg/m²</li> <li>Dacarbazine</li> <li>Doxorubicin ≥60 mg/m²</li> </ul>	<ul> <li>Epirubicin &gt;90 mg/m²</li> <li>Ifosfamide ≥2 g/m² per dose</li> <li>Mechlorethamine</li> <li>Streptozocin</li> </ul>
Moderate emetic risk (>30%–90% frequency of emesis) b,c,d	<ul> <li>Aldesleukin &gt;12–15 million IU/m²</li> <li>Amifostine &gt;300 mg/m²</li> <li>Azacitidine</li> <li>Bendamustine</li> <li>Busulfan</li> <li>Carboplatin AUCe &lt;4</li> <li>Carmustinee ≤250 mg/m²</li> <li>Clofarabine</li> <li>Cyclophosphamidee ≤1500 mg/m²</li> <li>Cytarabine &gt;200 mg/m²</li> <li>Dactinomycine</li> </ul>	<ul> <li>Daunorubicin<sup>e</sup></li> <li>Dual-drug liposomal encapsulation of cytarabine and daunorubicin</li> <li>Dinutuximab</li> <li>Doxorubicin<sup>e</sup> &lt;60 mg/m²</li> <li>Enfortumab vedotin-ejfv</li> <li>Epirubicin<sup>e</sup> ≤90 mg/m²</li> <li>Fam-trastuzumab deruxtecan</li> <li>Idarubicin<sup>e</sup></li> <li>Ifosfamide<sup>e</sup> &lt;2 g/m² per dose</li> <li>Interferon alfa ≥10 million IU/m²</li> </ul>	<ul> <li>Irinotecan<sup>e</sup></li> <li>Irinotecan (liposomal)</li> <li>Melphalan</li> <li>Methotrexate<sup>e</sup> ≥250 mg/m²</li> <li>Oxaliplatin<sup>e</sup></li> <li>Temozolomide</li> <li>Trabectedin<sup>e</sup></li> </ul>

NCCN Guidelines Index
Table of Contents
Discussion

#### **EMETOGENIC POTENTIAL OF PARENTERAL ANTICANCER AGENTS**<sup>a</sup>

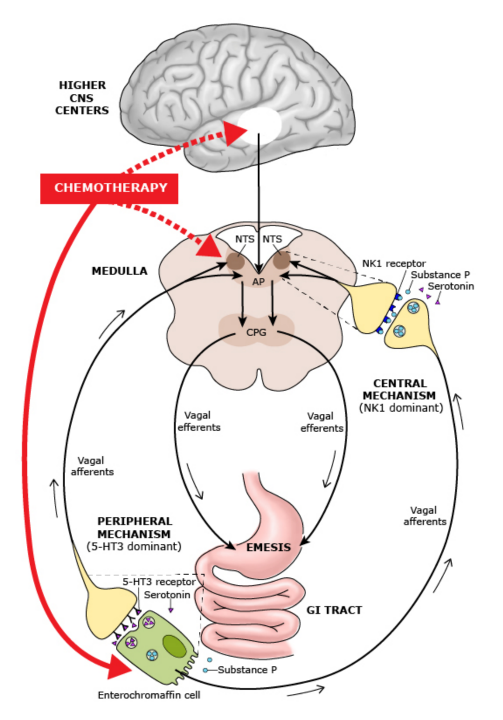
LEVEL	AGENT			
Low emetic risk (10%–30% frequency of emesis) <sup>b,d,f</sup>	<ul> <li>Ado-trastuzumab emtansine</li> <li>Aldesleukin ≤12 million IU/m²</li> <li>Amifostine ≤300 mg/m²</li> <li>Arsenic trioxide</li> <li>Axicabtagene ciloleucel<sup>g</sup></li> <li>Belinostat</li> <li>Brentuximab vedotin</li> <li>Cabazitaxel</li> <li>Carfilzomib</li> <li>Copanlisib</li> </ul>	Cytarabine (low dose) 100–200 mg/m² Docetaxel Doxorubicin (liposomal) Eribulin Etoposide 5-Fluorouracil (5-FU) Floxuridine Gemcitabine Gemtuzumab ozogamicin Inotuzumab ozogamicin	<ul> <li>Methotrexate &gt;50 mg/m² - &lt;250 mg/m²</li> <li>Mitomycin</li> <li>Mitoxantrone</li> <li>Mogamulizumab</li> <li>Moxetumomab</li> <li>Necitumumab</li> <li>Olaratumab</li> <li>Omacetaxine</li> <li>Paclitaxel</li> <li>Paclitaxel-albumin</li> </ul>	<ul> <li>Pemetrexed</li> <li>Pentostatin</li> <li>Polatuzumab vedotin</li> <li>Pralatrexate</li> <li>Romidepsin</li> <li>Tagraxofusp</li> <li>Talimogene laherparepvec</li> <li>Thiotepa</li> <li>Tisagenlecleucel<sup>g</sup></li> <li>Topotecan</li> <li>Ziv-aflibercept</li> </ul>
Minimal emetic risk (<10% frequency of emesis) b,d,f	<ul> <li>Alemtuzumab</li> <li>Atezolizumab</li> <li>Avelumab</li> <li>Asparaginase</li> <li>Bevacizumab</li> <li>Bleomycin</li> <li>Blinatumomab</li> <li>Bortezomib</li> <li>Cetuximab</li> <li>Cemiplimab</li> <li>Cladribine</li> <li>Cytarabine &lt;100 mg/m²</li> <li>Daratumumab</li> </ul>	<ul> <li>Decitabine</li> <li>Denileukin diftitox</li> <li>Dexrazoxane</li> <li>Durvalumab</li> <li>Elotuzumab</li> <li>Fludarabine</li> <li>Ipilimumab</li> <li>Methotrexate ≤50 mg/m²</li> <li>Nelarabine</li> <li>Nivolumab</li> </ul>	<ul> <li>Obinutuzumab</li> <li>Ofatumumab</li> <li>Panitumumab</li> <li>Pegaspargase</li> <li>Peginterferon</li> <li>Pembrolizumab</li> <li>Pertuzumab</li> <li>Ramucirumab</li> <li>Rituximab</li> <li>Rituximab and hyaluronidase human injection, for subcutaneous use</li> </ul>	<ul> <li>Siltuximab</li> <li>Temsirolimus</li> <li>Trastuzumab</li> <li>Trastuzumab and hyaluronidase injection, for subcutaneous use</li> <li>Valrubicin</li> <li>Vinblastine</li> <li>Vincristine</li> <li>Vincristine (liposomal)</li> <li>Vinorelbine</li> </ul>

Adapted with permission from: Hesketh PJ, Kris MG, Grunberg SM, et al. Proposal for classifying the acute emetogenicity of cancer chemotherapy. J Clin Oncol 1997;15:103-109. Grunberg SM, Warr D, Gralla RJ, et al. Evaluation of new antiemetic agents and definition of antineoplastic agent emetogenicity--state of the art. Support Care Cancer 2011;19:S43-S47.

NCCN Guidelines Index
Table of Contents
Discussion

#### EMETOGENIC POTENTIAL OF ORAL ANTICANCER AGENTS<sup>a</sup>

LEVEL	AGENT			
Moderate to high emetic risk <sup>b,z</sup> (≥30% frequency of emesis)	<ul> <li>Altretamine</li> <li>Avapritinib</li> <li>Binimetinib</li> <li>Busulfan (≥4 mg/d)</li> <li>Ceritinib</li> <li>Crizotinib</li> </ul>	<ul> <li>Cyclophosphamide (≥100 mg/m²/day)</li> <li>Dabrafenib</li> <li>Enasidenib</li> <li>Encorafenib</li> <li>Estramustine</li> </ul>	<ul><li>Etoposide</li><li>Lenvatinib</li><li>Lomustine (single day)</li><li>Midostaurin</li><li>Mitotane</li></ul>	<ul> <li>Niraparib</li> <li>Olaparib</li> <li>Procarbazine</li> <li>Rucaparib</li> <li>Selinexor</li> <li>Temozolomide (&gt;75 mg/m²/day)</li> </ul>
Minimal to low emetic risk <sup>b</sup> (<30% frequency of emesis)	Abemaciclib     Acalabrutinib     Afatinib     Alectinib     Alpelisib     Axitinib     Bexarotene     Brigatinib     Bosutinib     Busulfan (<4 mg/day)     Cabozantinib     Capecitabine     Chlorambucil     Cobimetinib     Cyclophosphamide     (<100 mg/m²/day)     Dacomitinib     Dasatinib	<ul> <li>Duvelisib</li> <li>Entrectinib</li> <li>Erdafitinib</li> <li>Erlotinib</li> <li>Everolimus</li> <li>Fludarabine</li> <li>Gefitinib</li> <li>Gilteritinib</li> <li>Glasdegib</li> <li>Hydroxyurea</li> <li>Ibrutinib</li> <li>Idelalisib</li> <li>Imatinib</li> <li>Ixazomib</li> <li>Ivosidenib</li> <li>Lapatinib</li> <li>Larotrectinib</li> <li>Lenalidomide</li> </ul>	<ul> <li>Lorlatinib</li> <li>Melphalan</li> <li>Mercaptopurine</li> <li>Methotrexate</li> <li>Nilotinib</li> <li>Neratinib</li> <li>Osimertinib</li> <li>Palbociclib</li> <li>Panobinostat</li> <li>Pazopanib</li> <li>Pomalidomide</li> <li>Ponatinib</li> <li>Regorafenib</li> <li>Ribociclib</li> <li>Ruxolitinib</li> <li>Sonidegib</li> <li>Sorafenib</li> </ul>	<ul> <li>Sunitinib</li> <li>Talazoparib tosylate</li> <li>Temozolomide (≤75 mg/m²/day)<sup>aa</sup></li> <li>Thalidomide</li> <li>Thioguanine</li> <li>Topotecan</li> <li>Trametinib</li> <li>Tretinoin</li> <li>Trifluridine/tipiracil</li> <li>Vandetanib</li> <li>Vemurafenib</li> <li>Venetoclax</li> <li>Vismodegib</li> <li>Vorinostat</li> <li>Zanubrutinib</li> </ul>



# **Biology of CINV**

NTS: nucleus tractus solitarius

AP: area postrema

CPG: central pattern generator

From: UpToDate

# Pharmacologic options for CINV

- 5HT<sub>3</sub> antagonists (ondansetron, dolasetron, granisetron, palonosetron)
- Corticosteroid (dexamethasone)
- Benzodiazepines (lorazepam)
- Phenothiazines\*\*
   (prochlorperazine, promethazine)
- Butyrophenones\*\* (droperidol, haloperidol)
- Olanzapine

- Cannabinoids (dronabinol)\*\*
- Substituted benzamides (metoclopramide)\*\*
- Antihistamine/Anticholinerg ics (diphenhydramine, scopolamine)\*\*
- Substance P/NK<sub>1</sub> receptor antagonist (aprepitant, netupitant)

\*\* low therapeutic index agents not discussed in this lecture

- No final common pathway has been discovered
- Current agents act on different receptor families (M1, D2, H1, 5-HT3, NK1)
- No single agent expected to provide complete protection

# Serotonin (5HT<sub>3</sub>) in CINV

- Closely associated with acute phase CINV
- Chemotherapy administration causes release of serotonin from the GI tract, thereby stimulating emesis via vagus and greater splanchnic nerves, as well as the area postrema of the brain
- In early trials, 5HT<sub>3</sub> release was not found in delayed phase of CINV
- Palonosetron has efficacy for prevention of delayed emesis, but role of other 5HT<sub>3</sub> is debated

# 5-HT3 receptor antagonists

- ondansetron (1991), granisetron, dolasetron, palonsetron (2003)
- Numerous studies have demonstrated the 5-HT3 agents have same
   SE profile and efficacy\*
- SE are mild HA, constipation counsel patients
- Steroids improve efficacy
- QTc prolongation (except palonosetron and ER formulations)
- Limited role in treatment of delayed phase N/V

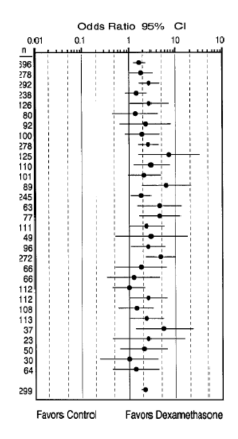
## Palonsetron

- pharmacologically distinct
- 100-fold higher binding affinity for 5-HT3R
- T ½ ~ 40 hours
- As effective as traditional 5-HT3 agents for acute CINV (single dose)
- Superior in preventing delayed emesis (single dose)

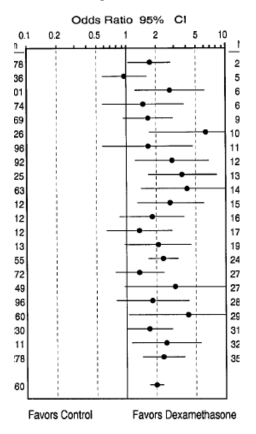
### Dexamethasone addition to 5HT3

Meta-analysis of 32 studies showing OR of 2 vs 5HT3 monotherapy for acute and delayed phase

#### **Acute Phase**



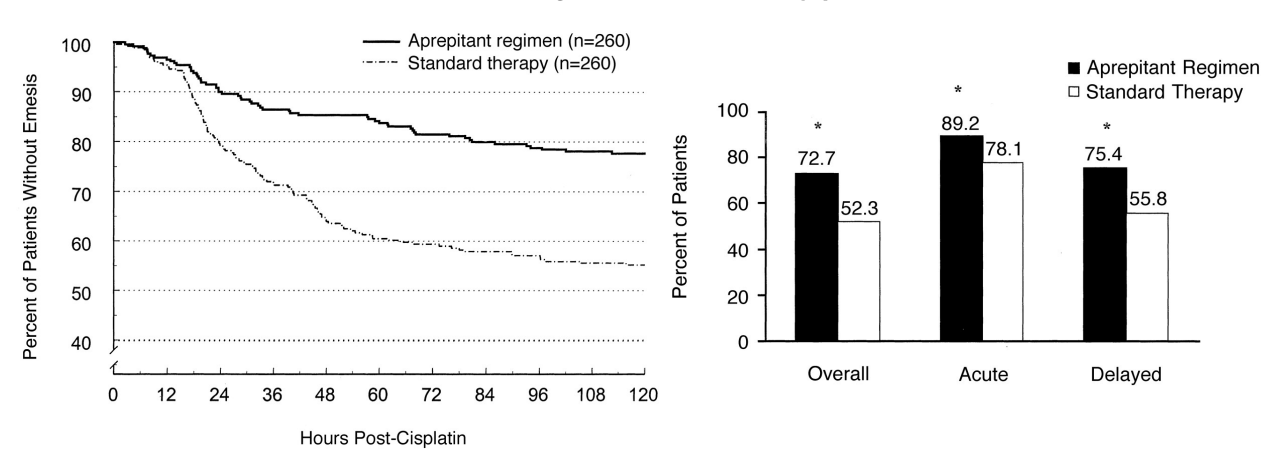
#### **Delayed Phase**



# Substance P / Neurokinin Receptors Aprepitant/fosaprepitant

- <u>Substance P</u>: a member of the tachykinin family of neuropeptides
- Biological activity mediated by neurokinin (NK-1) receptor
- Substance P and NK-1 receptors located in brain stem dorsal vagal complex – nucleus tractus solitarus (NTS) and area postrema
- Also located in the GI tract
- Beneficial in delayed > acute CINV, but use is in prevention
- New non-polysorbate-80 IV formulation for aprepitant

# Kaplan-Meier curves demonstrating percentages of patients without emesis during the 120-hour study period.



Standard therapy – ondansetron d1, dexamethasone d1-4

Hesketh et al. JCO 2003;21:4112-4119

# RCT: (<u>olanzapine</u>10mg vs placebo) + fosaprepitant, 5HT3, Dex

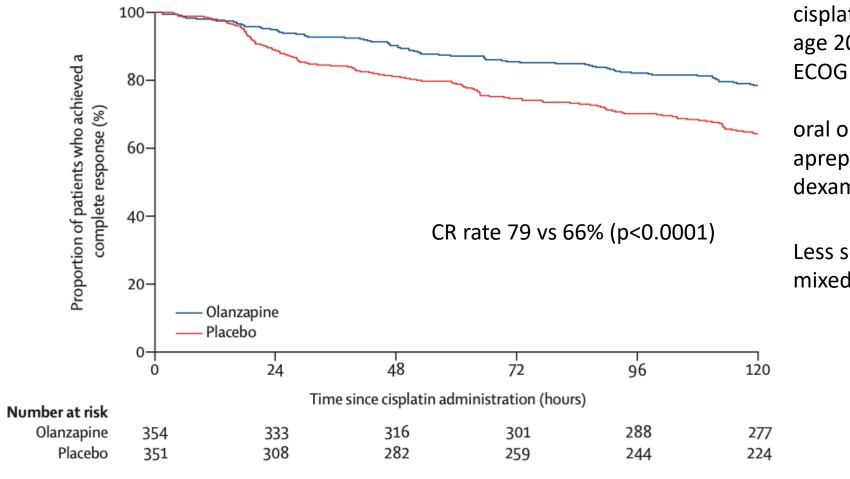
	CR Rate=no emesis or rescue(%)		No nausea (%) = primary endpt	
	Olanz	PCO	Olanz	PCO
0-24 hr	86	65	74	55
0-120 hr	64	40	37	22

All P < 0.01, N = 380

Side Effects: mild increase in **sedation** at day 2 (2/10 vs. 1/10) and increased appetite

Navari RM et al. NEJM 2016: 375: 134-142

# Olanzapine 5 mg plus standard antiemetic therapy for the prevention of chemotherapy-induced nausea and vomiting (J-FORCE):



cisplatin (≥50 mg/m2) age 20 - 75 years, ECOG 0–2.

oral olanzapine 5 mg or placebo d1–4 aprepitant, palonosetron, and dexamethasone

Less sedation than 10mg mixed effect on sleep

# Principles

- Prophylactic therapy should be given before chemotherapy to prevent adverse outcomes
- Routes of administration: PO, PR, IV, IM
- PO route is preferred as it is most convenient /cost effective
- Often IV is needed due to inability to take PO
- Lowest maximally effective dose should be used
- Once daily dosing
- Delayed N/V therapy incorporated proactively
- Avoid using concomitant drugs in same class



 In contrast to other guidelines that are often based on expert opinion there is a significant amount of clinical trials data supporting the recommendations

• USE THE GUIDELINES



Dexamethasone 12 mg PO/IV once<sup>u,v</sup>

#### NCCN Guidelines Version 2.2020 Antiemesis

NCCN Guidelines Index
Table of Contents
Discussion

#### HIGH EMETIC RISK PARENTERAL ANTICANCER AGENTS — ACUTE AND DELAYED EMESIS PREVENTION<sup>h,i,j,k,l</sup> **DAYS 2, 3, 4**: **DAY 1:** Select treatment option A, B, or C All treatment options are category 1 and should be started before chemotherapy Treatment option A (preferred), use the following combination: Treatment option A: Olanzapine 5–10 mg PO once<sup>m,n</sup> Olanzapine 5–10 mg PO daily on days 2, 3, 4<sup>m</sup> • NK1 RA (choose one): Aprepitant 80 mg PO daily on days 2, 3 ▶ Aprepitant 125 mg PO once (if aprepitant PO used on day 1) ▶ Aprepitant injectable emulsion 130 mg IV once<sup>o</sup> • Dexamethasone 8 mg<sup>u,v</sup> PO/IV daily on days 2, 3, 4 Fosaprepitant 150 mg IV once Netupitant 300 mg / palonosetron 0.5 mg (available as fixed combination product only) PO once Fosnetupitant 235 mg / palonosetron 0.25 mg (available as fixed combination product only) IV oncep ▶ Rolapitant 180 mg PO once<sup>q</sup> 5-HT3 RA (choose one):<sup>r,s</sup> Dolasetron 100 mg PO once ▶ Granisetron 10 mg SQ once, t or 2 mg PO once, or 0.01 mg/kg (max 1 mg) IV once, or 3.1 mg/24-h transdermal patch applied 24-48 h prior to first dose of chemotherapy. → Ondansetron 16–24 mg PO once, or 8–16 mg IV once ▶ Palonosetron 0.25 mg IV once Dexamethasone 12 mg PO/IV once<sup>u,v</sup> Treatment option B, use the following combination: Treatment option B: Olanzapine 5–10 mg PO daily on days 2, 3, 4<sup>m</sup> Olanzapine 5–10 mg PO once<sup>m</sup> Palonosetron 0.25 mg IV once Dexamethasone 12 mg PO/IV once<sup>u,v</sup> Treatment option C, use the following combination: Treatment option C: Aprepitant 80 mg PO daily on days 2, 3 NK1 RA (choose one): → Aprepitant 125 mg PO once (if aprepitant PO used on day 1) ▶ Aprepitant injectable emulsion 130 mg IV once<sup>o</sup> Dexamethasone 8 mg<sup>u,v</sup> PO/IV daily on days 2, 3, 4 ▶ Fosaprepitant 150 mg IV once Netupitant 300 mg / palonosetron 0.5 mg (available as fixed combination product only) PO once Fosnetupitant 235 mg / palonosetron 0.25 mg (available as fixed combination product only) IV oncep ▶ Rolapitant 180 mg PO once<sup>q</sup> 5-HT3 RA (choose one):<sup>r,s</sup> ▶ Dolasetron 100 mg PO once → Granisetron 10 mg SQ once, to 2 mg PO once, or 0.01 mg/kg (max 1 mg) IV once, or 3.1 mg/24-h transdermal patch applied 24-48 h prior to first dose of chemotherapy. ▶ Ondansetron 16-24 mg PO once, or 8-16 mg IV once ▶ Palonosetron 0.25 mg IV once

#### Comprehensive NCCN Guidelines Version 2.2020 **Antiemesis**

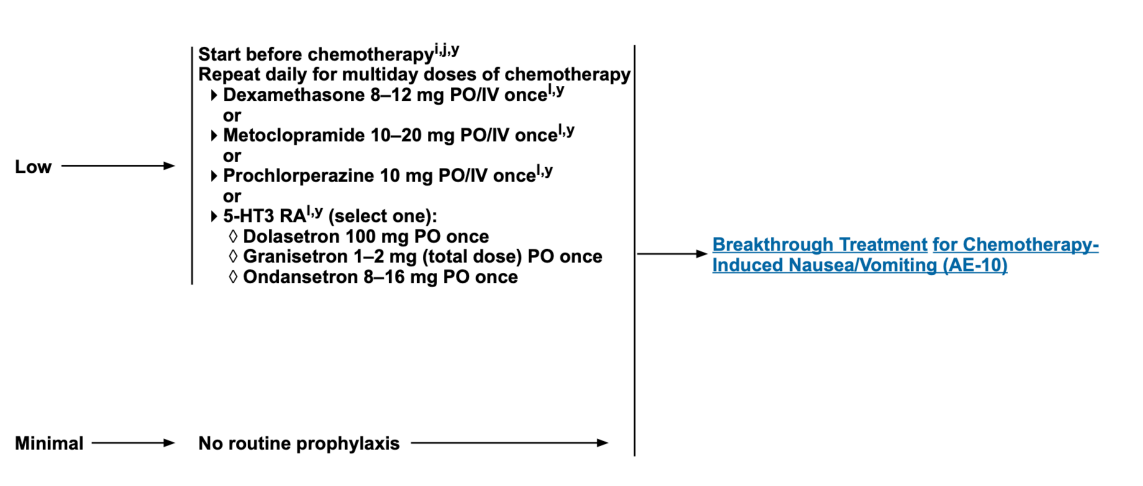
**NCCN Guidelines Index Table of Contents Discussion** 

#### ${\tt MODERATE\ EMETIC\ RISK\ PARENTERAL\ ANTICANCER\ AGENTS\ --\ ACUTE\ AND\ DELAYED\ EMESIS\ PREVENTION^{h,i,j,k,l}}$

<u>DAY 1</u> : Select treatment option D, E, or F. All treatment options are category 1 and should be started before chemotherapy:	<b>DAYS 2, 3</b> :
Treatment option D, use the following combination:  • 5-HT3 RA (choose one):  • Dolasetron 100 mg PO once  • Granisetron 10 mg SQ once <sup>t</sup> (preferred), or 2 mg PO once, or 0.01 mg/kg (max 1 mg) IV once, or 3.1 mg/24-h transdermal patch applied 24–48 h prior to first dose of chemotherapy.  • Ondansetron 16–24 mg PO once, or 8–16 mg IV once  • Palonosetron 0.25 mg IV once (preferred)  • Dexamethasone 12 mg PO/IV once <sup>u,v</sup>	Treatment option D:  • Dexamethasone 8 mg <sup>u,v</sup> PO/IV daily on days 2, 3  OR  • 5-HT3 RA monotherapy <sup>w</sup> :  • Granisetron 1–2 mg (total dose) PO daily or 0.01 mg/kg (max 1 mg) IV daily on days 2 and 3  • Ondansetron 8 mg PO twice daily or 16 mg PO daily or 8–16 mg IV daily on days 2, 3  • Dolasetron 100 mg PO daily on days 2, 3
Treatment option E, use the following combination: <sup>x</sup> • Olanzapine 5–10 mg PO once <sup>m</sup> • Palonosetron 0.25 mg IV once • Dexamethasone 12 mg PO/IV once <sup>u,v</sup>	Treatment option E: • Olanzapine 5–10 mg PO daily on days 2, 3 <sup>m</sup>
Treatment option F, use the following combination: <sup>x</sup> NK1 RA (choose one): Aprepitant 125 mg PO once Aprepitant injectable emulsion 130 mg IV once <sup>o</sup> Fosaprepitant 150 mg IV once <sup>p</sup> Netupitant 300 mg / palonosetron 0.5 mg (available as fixed combination product only) PO once <sup>p</sup> Fosnetupitant 235 mg / palonosetron 0.25 mg (available as fixed combination product only) IV once <sup>p</sup> Rolapitant 180 mg PO once <sup>q</sup> 5-HT3 RA (choose one): <sup>r,s</sup> Dolasetron 100 mg PO once Granisetron 10 mg SQ once, <sup>t</sup> or 2 mg PO once, or 0.01 mg/kg (max 1 mg) IV once, or 3.1 mg/24-h transdermal patch applied 24–48 h prior to first dose of chemotherapy. Ondansetron 16–24 mg PO once, or 8–16 mg IV once Palonosetron 0.25 mg IV once Dexamethasone 12 mg PO/IV once <sup>u,v</sup>	Treatment option F: • Aprepitant 80 mg PO daily on days 2, 3 (if aprepitant PO used on day 1) • ± Dexamethasone 8 mg <sup>u,v</sup> PO/IV daily on days 2, 3

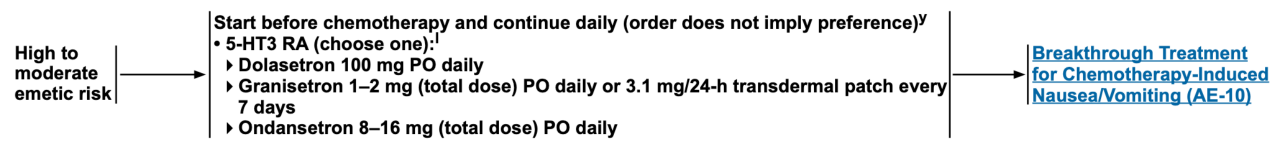
NCCN Guidelines Index
Table of Contents
Discussion

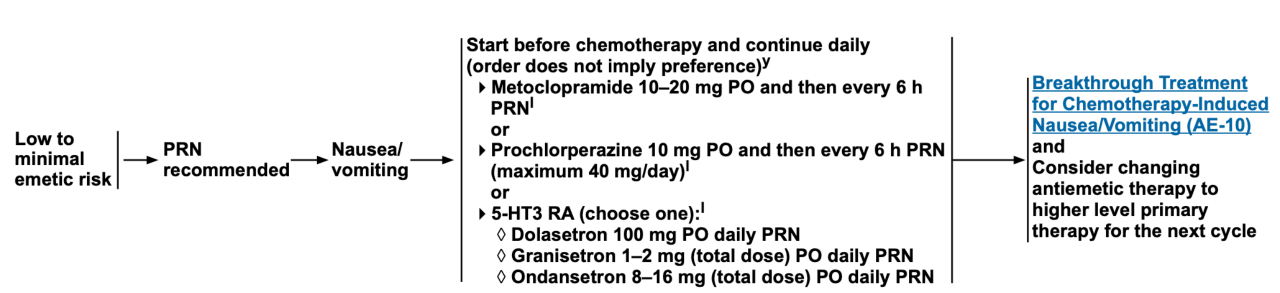
#### LOW AND MINIMAL EMETIC RISK PARENTERAL ANTICANCER AGENTS - EMESIS PREVENTION<sup>h,i,j,l</sup>





#### ORAL CHEMOTHERAPY - EMESIS PREVENTION<sup>i,j,bb,cc</sup>





# Breakthrough Treatment

- assess what was actually taken (medication reconciliation)
- add agents from a different drug class
  - Additional steroid for prolonged nausea in delayed phase
  - (don't use additional 5HT3 for 3 days post-palonosetron)
  - (5HT3 likely minimally effective in delayed phase)
- use multiple concurrent agents
- IV therapy often needed (drugs, IVF)
- round-the-clock administration
- remember this for the next cycle, assess for other causes

### Consider non-CINV causes

- bowel obstruction
- constipation
- vestibular dysfunction
- brain metastases
- electrolytes, dehydration
- uremia
- other drugs (opiates)
- gastro paresis (tumor or vincristine)
- anxiety, anticipatory N/V

## Take Home Points

- 5-HT3 agents are the mainstay for the prevention of acute CINV in moderate to highly emetogenic regimens
- The benefit of the 5-HT3 agents (except palonset.) in delayed CINV is debated
- Steroids significantly augment 5-HT3s and should almost always be used
- NCCN recommends avoiding steroids in immunotherapy
- Aprepitant and/or olanzapine (~5mg) are indicated for highly emetogenic chemotherapy
- High therapeutic index agents: 5HT3, NK1, olanzapine
- CW: Don't give patients starting on a chemotherapy regimen that has a low or moderate risk of causing nausea and vomiting antiemetic drugs intended for use with a regimen that has a high risk of causing nausea and vomiting.

# Erythropoiesis-Stimulating Agents (ESA)

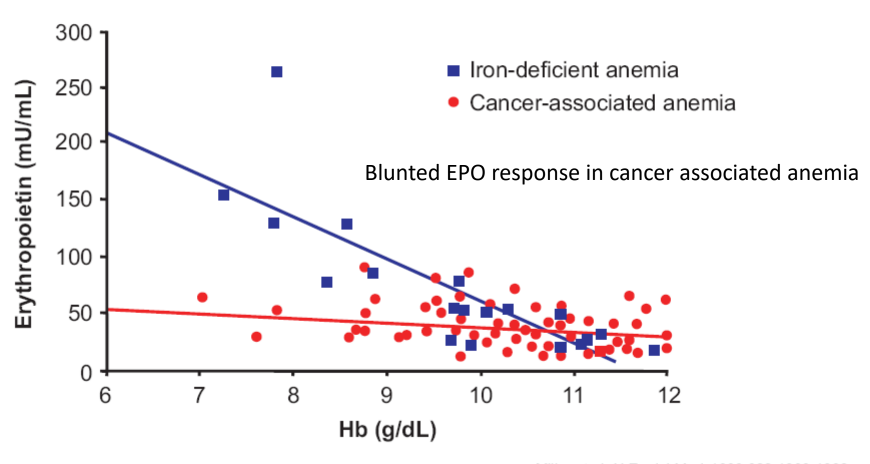
#### Cancer Related Anemia

- High prevalence among cancer patients
- Multifactorial
  - Inflammatory state related to cancer
  - Treatment related myelosuppression
  - BM infiltration
  - Paraneoplastic
  - Other (bleeding, nutritional, hemolysis, congenital,...)

### Workup of Anemia in Ca Patients

- Screen for anemia in cancer patients
- Complete workup not always needed
- Consider: smear, BM, B12, folate, guaiac, Creat, retics
- EPO levels not recommended as they are not predictive of response
- Screening iron studies: ferritin, Fe, TIBC, TSAT

#### **Erythropoietin Response to Anemia**



# ESAs in solid tumor oncology

- Anemia is very common in cancer
- Linked to worse prognosis
- Worse outcomes with radiation
  - hypoxia leads to radio-resistance
- ESAs initially used in CRF, use extended to oncology
  - Reduction of transfusions, HR = 0.64 in chemo patients
  - Difference between placebo was ~1 unit, NNT = 6
  - Marginal effects on QOL and fatigue
- Utilization was quite high, but has decreased due to safety concerns

This is a controversial subject, with a vast literature

#### ESAs: Risks/Benefits National NCCN Guidelines Version 2.2019 Management of Cancer- and Chemotherapy-Induced Anemia

NCCN Guidelines Index Table of Contents Discussion

#### COMPARISON OF RISKS AND GOALS OF ESA USE VERSUS RBC TRANSFUSION<sup>h</sup>

Discuss the following risks and goals with patients when considering anemia treatment options:

	ESA in the Cancer Setting	RBC Transfusion
Risks	Increased thrombotic events     Possible decreased survival     Time to tumor progression shortened	Transfusion reactions (eg, hemolytic, febrile, non-hemolytic, lung injury) Transfusion-associated circulatory overload (TACO) Virus transmission (eg, hepatitis, HIV) Bacterial contamination Iron overload Increased thrombotic events Possible decreased survival Alloimmunization Increased risk of poor response to future platelet transfusions due to HLA immunization
Goals	Transfusion avoidance     Gradual improvement in anemia- related symptoms	Rapid increase of Hb and hematocrit levels     Rapid improvement in anemia-related symptoms

See Erythropoietic Therapy - Dosing, Titration, and Adverse Effects (ANEM-A)

#### When considering ESAs:

- Discuss the risks of ESAs with patients including the potential for tumor growth, death, blood clots, and serious heart problems.
- Refer patients to the following medication guides for more information on the benefits and risk of ESAs: Epoetin Alfa Medication Guide, Epoetin Alfa-epbx Medication Guide and Darbepoetin Alfa Medication Guide

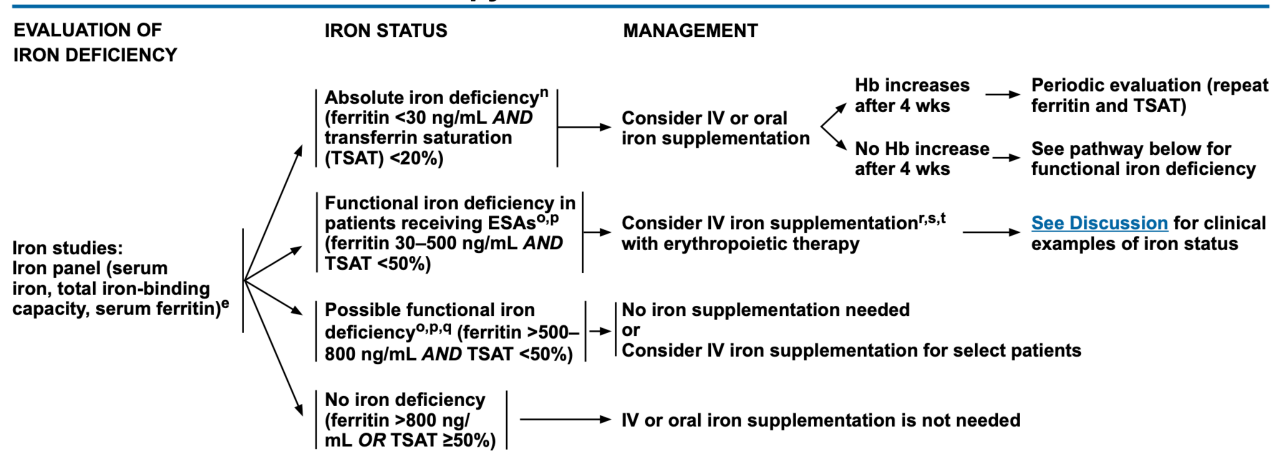
When considering RBC transfusion, see AABB Clinical Practice Guidelines: Tobian AA, Heddle NM, Wiegmann TL, Carson JL. Red blood cell transfusion: 2016 clinical practice guidelines from AABB. Transfusion 2016;56:2627-2630.

### Iron deficiency



#### NCCN Guidelines Version 2.2020 Management of Cancer- and Chemotherapy-Induced Anemia

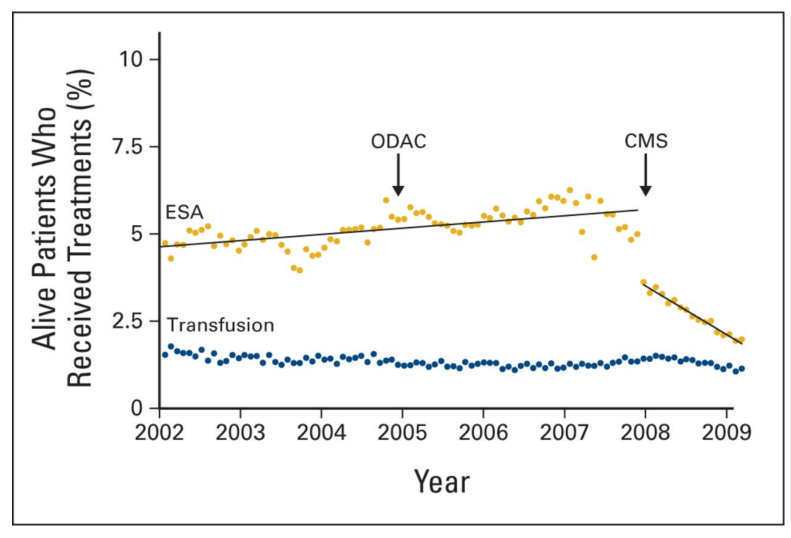
NCCN Guidelines Index
Table of Contents
Discussion



#### Recommendations

- No use of ESA for anemia not associated with chemotherapy
- For chemotherapy related anemia, weigh risks/benefits
- Start Hgb < 10, goal = avoid transfusion, increase < 1gm/2w</li>
- CMS start Hgb < 10, DC for >10
- FDA indications and dosing should be used, NCCN lists alternative regimens

Percentage of patients with cancer who received erythropoiesis-stimulating agents (ESAs) per month in relation to regulatory changes.



Hershman D L et al. JOP 2014;10:264-269

# **Myeloid Growth Factors**

### Myeloid Growth Factors

- Neutropenia is a common DLT of chemotherapy
- Febrile neutropenia (FN) results in hospitalization, IV antibiotic use, decreased QOL, and morbidity
- FN risk is highest with first two cycles of a regimen
- Neutropenia may result in reductions in dose-density and intensity which can compromise outcomes
- This all can be reduced with use of myeloid CSFs

# Myeloid CSFs

- Reduce risk (by ~50% for FN), severity and duration of neutropenia
- Cost-benefit threshold is now at 20% risk of FN, previously was at 40%
- Many common regimens have 25-40% FN risk in treatment naïve patients

# Risk of FN – chemotherapy

- Risk is hard to define precisely
- Published trials are informative
- Guidelines (NCCN) have been published which estimate risk for regimens

### Patient risk factors for neutropenia

#### Treatment-related

- Previous history of severe neutropenia with similar chemotherapy
- Type of Chemotherapy (anthracyclines)
- Planned relative dose intensity > 80%
- Preexisting neutropenia (< 1000) or lymphocytopenia
- Extensive prior chemotherapy
- Concurrent or prior radiation therapy to marrow containing bone

#### Patient-related

- Age (> 65 y)
- Female gender
- Poor performance status (ECOG ≥ 2)
- Poor nutritional status (eg, low albumin)
- Decreased immune function

#### Cancer-related

- Bone marrow involvement with tumor
- Advanced or uncontrolled cancer
- Elevated Lactate Dehydrogenase (Lymphoma)
- Leukemia
- Lymphoma
- Lung cancer

#### Conditions associated with risk of serious infection

- Open wounds
- Active tissue infection

#### Comorbidities

- COPD
- Cardiovascular disease
- Liver disease (elevated bilirubin, alkaline phosphatase)
- Diabetes mellitus
- Low baseline hemoglobin

From NCCN guidelines

# Use of myeloid CSFs

- Risk of FN
  - >20% recommended
  - 10-20% consider
  - <10% generally not recommended</p>
  - CW: Don't use white cell stimulating factors for primary prevention of febrile neutropenia for patients with less than 20 percent risk for this complication.
- Also consider intent of treatment: curative, adjuvant, palliative
- Prior FN is an indication for CSFs
- Prior FN w/CSF-> dose reduction or change regimen
- Do not use with chemoradiation
- (antibiotics not recommended)

# Myeloid CSF regimens

- Filgrastim
  - 5mcg/kg/d rounded to 300 or 480mcg
  - Start 1-3 days after chemo
  - Treat through post-nadir recovery
- Tbo-filgrastim, filgrastim-sndz, other biosimilars
- Pegfilgrastim, other biosimilars, OnPro®
  - 6mg/cycle
  - Start 1-3 days after chemo
  - Data for q3wk regimens, phase II data for q2wk
  - Dosing on day 1 safe, but less efficacious\*
  - NCCN recommends administration on day 2

#### **Adverse Effects**

- Bone pain (common)
- Allergic reactions
- ARDS
- Splenic rupture (transplant setting)
- Precipitate sickle cell crisis
- MDS/AML\* (increased AR 0.4%, RR 1.9)
- Cutaneous vasculitis (Sweet's syndrome)

# **Bone Supportive Care**

# Skeletal Morbidity

- Cancer treatment induced bone loss
  - Androgen deprivation
  - Estrogen deprivation
  - Corticosteroids, TSH suppression
  - These will not be discussed further
- Bone metastases
  - Common in many cancer
  - Lung, breast, and prostate are most common

# Measuring Skeletal Morbidity

- "Skeletal related event" SRE
  - Fracture, spinal cord compression
  - Need for surgery or radiation
  - (some definitions) hypercalcemia
- QOL and pain are other outcomes of interest
- SREs are quite common, estimates are > 50% of metastatic breast cancer patients will have a SRE

# Bisphosphonates

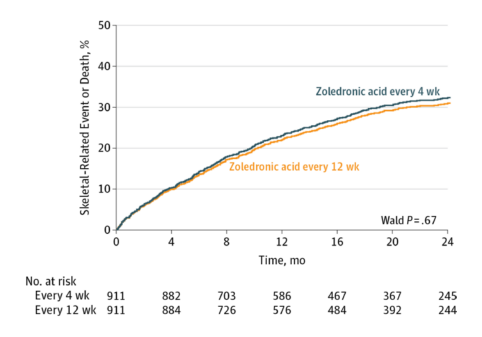
- Analogs of pyrophosphate a major constituent of bone
- Decrease bone resorption and increase mineralization by inhibiting osteoclast activity
- Induce apoptosis in osteoclasts
- Zoledronic acid (ZA) and pamidronate are potent bisphosphonates

# Bisphosphonates in solid tumors with established bone metastases

- Positive data is primarily for zoledronic acid (ZA)
- ZA vs. placebo in AR-prostate cancer
  - Incidence of SRE 38% vs. 49% median FU 2yrs
  - TTE was 488 vs. 321 days, benefit in pain control
- ZA vs. placebo in solid tumor
  - (no breast/prostate, mostly NSCLC)
  - Incidence of SRE 38% vs. 47%
  - TTE was 230 vs. 163 days

From: Effect of Longer-Interval vs Standard Dosing of Zoledronic Acid on Skeletal Events in Patients With Bone Metastases A Randomized Clinical Trial

JAMA. 2017;317(1):48-58. doi:10.1001/jama.2016.19425



Cause-Specific Cumulative Incidence of Skeletal-Related EventsThere were 256 patients with skeletal-related events in the zoledronic acid every 4-week dose group and 246 patients in the every 12-week dose group (hazard ratio, 0.96 [95% CI, 0.81-1.15]).

#### Denosumab

- Monoclonal antibody targeting the RANKL which is involved in osteoclast formation and activation
- Has indications for osteoporosis and prevention of SREs in solid tumors
- Denosumab does not have renal toxicity
- Given as 120mg SQ injection q 4 weeks
- Goodrx: \$2400 vs \$33 for ZA

# Denosumab efficacy

- All have ZA as comparator arm
- Three positive trials: breast, AR-prostate, "other"
- Denosumab vs. ZA
- Other (N=1176) MM and solid tumors (not breast or prostate), 40% were NSCLC)
  - TTE 20.6m vs. 16.3 mo.
  - P=0.03, but 0.06 after correction for multiple comparisons

# ONJ - osteonecrosis of the jaw

- Presents as infection with exposed necrotic maxillary or mandibular bone
- Risks: poor dental hygiene, dental extractions/implants, chemotherapy?, anti-angiogenics?
- Incidence is ~2% for both ZA and denosumab
- Most patients who get ONJ have a risk factor (~80%)
- "Dental" exam prior to initiation
- Avoid invasive dental procedures

### Comparison

#### Denosumab

- Expensive
- Monthly
- Ok in renal dysfunction
- Mildly improved SRE
- Rebound vertebral fractures after DC
- Hypersensitivity, neutralizing Abs
- Mild increase in infections (skin, UTI)

#### ZA

- Cheap
- Q 3 month
- Avoid if CrCl < 30, dose adjust; potential for renal injury
- Acute phase reaction flu like ~50%
- conjunctivitis, uveitis, scleritis, and orbital inflammation
- Afib/flutter , stroke RR~1.3 in SEER
- MSK pain

Common to both: hypocalcemia, ONJ, atypical fractures

#### Conclusions

- Use agents in patients with established bone metastases
- Aggregate data favors denosumab over ZA, but cost is high
- Among bisphosphonates ZA is the preferred agent
- Screen for ONJ risk factors prior to use
- Adverse events are similar between agents
- Supplement Ca, D, replete if deficient prior to therapy

### Fatigue

- High symptom burden among cancer patients
- Some nihilism regarding treatment
- I will focus on NCCN guidelines and trials data
- "Cancer-related fatigue is a distressing, persistent, subjective sense of physical, emotional, and/or cognitive tiredness or exhaustion related to cancer or cancer treatment that is not proportional to recent activity and interferes with usual functioning."

Sources – NCCN Guidelines and Cancer-related fatigue; UpToDate

# **Fatigue Evaluation**

- Medications
- Pain
- Emotional distress -depression
- Anemia
- Sleep disturbance
- Comorbidities endocrine disorders, organ dysfunction
- Assessment on 0-10 scale

### Interventions: non – pharmacologic

- Different interventions for different milestones in cancer treatment
- Management strategies (delegation, prioritize)
- physical activity (cat 1)
- massage therapy, CBT, educational therapies (cat 1)
- Sleep hygiene structure, naps

# Pharmacologic Interventions

- Stimulants methylphenidate
- Modafinil
- Corticosteroids
- Ginseng

 Overall evidence is weak or mixed for these interventions, but downside is relatively low

# Methylphenidate

- Of 8 RCTs, only 2 have demonstrated benefit
- Most rigorous studies were negative
- Trials were small and populations heterogeneous
- Suggestion of greater benefit with
  - Higher levels of fatigue
  - More advanced disease
  - Opioid related fatigue
  - Higher dose

#### Modafinil

- "Wake-promoting" agent for narcolepsy
- Initial pilot studies were encouraging
- Subsequent studies did not show overall benefit
  - N=631 evaluable, any level of fatigue, only patients with score ≥ 7 showed benefit
  - N=160 in ITT, NSCLC no benefit over placebo

#### Corticosteroids

- Studied in terminal stage of cancer
- Long-term side effects limit utility in patients with longer life expectancy
- N=84 RCT of advanced cancer patients with fatigue(≥4) and high symptom burden, dexamethasone 4mg bid vs. placebo
- Improved QOL and fatigue scores
- J Clin Oncol. 2013 Sep 1;31(25):3076-82.

# Ginseng

- N=364 cancer patients with curative intent therapy and fatigue(≥4), RCT of ginseng 2000mg vs. PCO
- Improved fatigue at 8 week (but not 4 week)
- No discernable toxicities
- Potential for drug interactions, inhibitor of CYP3A4
- J Natl Cancer Inst. 2013 Aug 21;105(16):1230-8.

# Chemotherapy Induced Peripheral Neuropathy (CIPN)

- Common side effect of many agents
  - Most common in breast and colon cancer
  - Platins, taxanes, vincas, bortezomib
- Can be dose-limiting
- Potential for significant impact on QOL

#### **CIPN**

- Prevention despite some reports demonstrating benefit, <u>NO agent has</u> been useful for prevention of CIPN
- Preliminary data suggests possible beneficial effect of limb cooling
- Prevention strategies are dose reduction, dose delays, and treatment breaks
- Bortezomib: Weekly vs. 2x/week and SQ vs IV is preferred
- Treatment the <u>only agent that has demonstrated efficacy is duloxetine</u>
- 59% vs 38% (PCO) reported pain decrease
- Difference in decrease of pain was modest: 0.7 on a 1-10 scale
- RCT: Smith EM. JAMA. 2013 Apr 3;309(13):1359-67. PMID: 23549581

#### Cancer Cachexia

- Pharmacologic interventions:
- Only corticosteroids and progesterone analogs have demonstrated benefit
- Increased appetite, modest weight gain
- No effect on survival or overall QOL

#### Treatment of Cancer Cachexia

For patients with short life expectancy (~weeks) dexamethasone (4mg daily)

Side effects: myopathy, Cushingoid, PUD

Megestrol 400-800mg daily for longer term

- Side effects: edema, VTE, increased mortality with doses >800mg/d
- Effect is weak, 16% of patients with >15# gain

No benefit of dronabinol in RCTs

# Sources for further study

- ASCO Guidelines: Supportive Care and Treatment Related Issues; Patient and Survivor Care
- NCCN Guidelines for Supportive Care
- ESMO Clinical Practice Guidelines: Supportive and Palliative Care
- MASCC, Multinational Association for Supportive Care in Cancer
- UpToDate multiple topics covered

thank you!