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CAR = Chimeric Antigen Receptor CRS = cytokine release syndrome ICANS = immune effector cell-associated neurotoxicity syndrome IEC = immune effector cells

GVHD = graft versus host disease

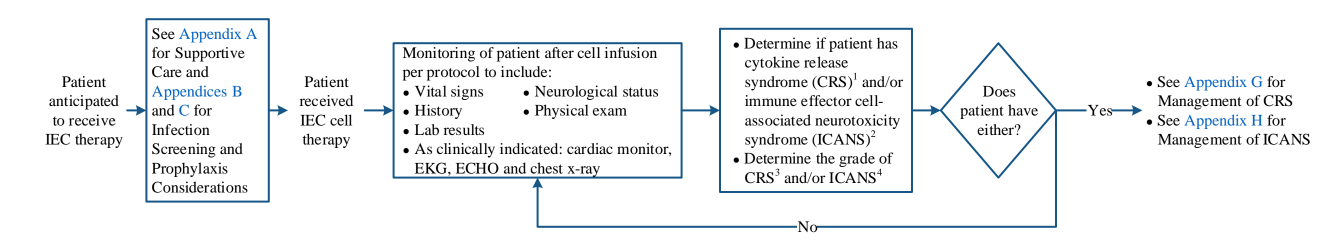
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INITIAL EVALUATION

MANAGEMENT



¹ If the subject has fever with or without hypotension or hypoxia within the first 4 weeks of engineered immune effector cell (IEC) therapy, the subject <u>may have</u> CRS if the symptoms or signs are not attributable to any other cause

- Fever should be present at onset of CRS (temperature $\geq 38^{\circ}$ C)
- Hypotension (requiring IV fluids or vasopressors to maintain normal blood pressure)
- Hypoxia (requiring supplemental oxygen to correct a deficit in oxygenation)

²If the subject has any of the following within the first 8 weeks of engineered IEC- therapy, the subject <u>may have</u> ICANS if the symptoms or signs are not attributable to any other cause

- IEC-Associated Encephalopathy (ICE) Score of less than 10 (Appendix F)
- Depressed level of consciousness
- Convulsive or non-convulsive seizures (can be focal or generalized)
- Motor weakness (can be focal motor weakness, hemiparesis, paraparesis)
- Focal / diffuse cerebral edema on imaging or signs of raised intracranial pressure including decerebrate or decorticate posturing, cranial nerve VI palsy, papilledema, or Cushing's triad

³See Appendix D for Grading of CRS ⁴See Appendix E for Grading of ICANS

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APPENDIX A: Checklist / Supportive Care Considerations for Managing Patients Receiving IEC Therapy

For Inpatients or Outpatients:

Before and During IEC Infusion

- Imaging of the brain prior to IEC infusion (preferably MRI with and without contrast but CT without contrast is acceptable if MRI cannot be performed) to rule out any central nervous system disease and also to serve as a baseline for comparison in case the patient develops ICANS
- For patients with known history of seizures, migraines and/or other CNS disorders including malignant disease, consider Neurology consult prior to IEC infusion
- Central venous access with port-a-cath or double/triple lumen catheter is recommended for IEC infusion as well as for IV fluids and other infusions in case of toxicities
- IEC infusion may be administered either in the ambulatory unit or in the inpatient unit
- If the median time to onset of CRS is expected to be < 48 hours, hospitalization should be considered for IEC infusion
- When hospitalized, admission to an IEC-designated unit with capability for cardiac monitoring by telemetry is recommended
- Tumor lysis precautions for patients with high tumor burden, as per standard guidelines
- Seizure prophylaxis with levetiracetam 500-750 mg PO every 12 hours for 30 days, starting on the day of infusion for IEC therapies associated with a high incidence of ICANS, in patients with history of seizures or brain metastases
- Consider filgrastim products if patient is neutropenic and concern for infection (if not already receiving)
- Ensure appropriate documentation in EHR regarding IEC therapy and "conditional" corticosteroid contraindication

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APPENDIX A: Checklist / Supportive Care Considerations for Managing Patients Receiving IEC Therapy- continued

For Outpatients:

Patient Monitoring After IEC infusion (for at least 14 days post-IEC infusion)

- Assess and record vital signs at least once daily in clinic
- Daily weights
- Daily review of patient history and physical examination
- Daily complete blood count with differential and complete metabolic profile
- Coagulation profile at least twice weekly
- Consider monitoring C-reactive protein (CRP) and ferritin levels daily during the phase when CRS is likely to occur and then as needed thereafter
- Consider cytokine panel if clinically indicated
- Assessment and grading of CRS (document in CARTOX flowsheet) at least daily and if a change in patient status while in clinic
- Assessment and grading for ICANS (document in CARTOX flowsheet) at least daily including 10-point ICE score assessment

Supportive Care

- Encourage oral fluid intake to ensure adequate hydration
- IV fluids as needed

Patient Home Monitoring (provide patient with a log to document and bring daily to clinic visits and dictate the findings from home log in each clinic note)

- Provide patient with self-care instructions and team contact information
- Provide patient with guidance for when to report to the emergency center
- Oral temperature every evening
- ICE-score with sentence writing every evening

Considerations for Admission

- Temperature $\geq 38^{\circ}C$
- SBP < 90 mmHg
- New arrhythmia
- Upward trend in liver function tests and/or creatinine

- Oxygen saturation < 92% on room air
- Tremors or jerky movements in extremities
- Grade 1 CRS or greater
- Grade 1 ICANS or greater

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APPENDIX A: Checklist / Supportive Care Considerations for Managing Patients Receiving IEC Therapy - continued

For Inpatients:

Patient Monitoring After IEC infusion

- Assess vital signs every 4 hours (inpatient encounter)
- Strict monitoring of oral and IV fluid input and output (including urine and stool)
- Daily measurement of body weight
- Daily review of patient history and physical examination
- Daily complete blood count with differential and complete metabolic profile
- Coagulation profile at least twice weekly or more frequently if clinically indicated
- Consider monitoring C-reactive protein (CRP) and ferritin levels daily during the phase when CRS is likely to occur and continue to monitor until CRS and/or ICANS resolves (if present). Monitor as needed thereafter.
- Consider cytokine panel if clinically indicated
- Assessment and grading of CRS (document in CARTOX flowsheet) should be completed at least every 12 hours and whenever there is a change in patient's status
- Assessment and grading for ICANS (document in CARTOX flowsheet) should be completed at least every 12 hours including the 10-point ICE score assessment
- Maintenance IV fluids with normal saline to ensure adequate hydration
- Cardiac monitoring by telemetry is recommended for \geq Grade 1 CRS and continued until CRS resolves
- For post-IEC infusion headache that is unresponsive to analgesics, consider brain imaging and lumbar puncture
- Neurology consult recommended for patients who develop Grade 1 or higher ICANS
- Critical Care and/or MERIT team will follow patients on an as-needed basis
- Infectious Diseases team will follow patients on an as-needed basis
- Consult should be performed early for patients with positive infectious disease screening or for persistent fevers

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APPENDIX A: Checklist / Supportive Care Considerations for Managing Patients Receiving IEC Therapy - continued

For Inpatients:

- Notifications and contingency orders
- Notify primary physician on detection of any of the following:
- \circ SBP > 140 or < 90 mmHg
- \circ Heart rate > 120 or < 60 beats per minute or arrhythmia
- \circ Respiratory rate > 25 or < 12 breaths per minute
- \circ Oxygen saturation < 92% on room air
- \circ Urine output < 1,500 mL/24 hours or 60 mL/hour
- \circ Upward trends in creatinine or liver function tests
- \circ Tremors or jerky movements in extremities
- Change in mental status (alertness, orientation, speech, ability to write a sentence, or ICE score of < 10)
- For temperature \geq 38°C, send blood cultures (central and peripheral) and urine for urinalysis and culture, obtain portable chest x-ray, and notify physician
- For patients with neutropenia and fever, start empiric broad-spectrum antibiotics
- Do not administer corticosteroids unless approved by physician
- If patient develops ICANS, withhold oral intake of food, fluids, and medicines, and notify physician
- PRN medications
- \circ Acetaminophen (1st choice) or ibuprofen (2nd choice, if not contraindicated) for fever \geq 38.3°C
- \circ Cooling blanket for fever \geq 38.3°C
- Normal saline 500 1000 mL bolus prn for hypotension; may repeat once if patient remains hypotensive after 1st bolus
- \circ Transfuse packed red blood cells (PRBC) to maintain hemoglobin > 8 gm/dL
- Transfuse platelets to maintain > 10 K/microliter; for patients with abnormal brain imaging, see recommendations as in Grades 3 and 4 ICANS
- PRN tocilizumab to be activated only on physician order ("ok to give tocilizumab" order should be placed if dose approved by physician)

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APPENDIX B: Infectious Disease Screening (within 30 days prior to apheresis is recommended)

Required Infectious Disease Screening ¹	Optional Infectious Disease Screening (as clinically indicated) ²
• Hepatitis B surface antigen (HBsAg)	Anti-human T-cell lymphotrophic virus (HTLV) antibody (HTLV I/II Ab)
• Anti-hepatitis B core antibody (HBcAb)	• Rapid Plasma Reagin (RPR) – syphilis
• Anti-hepatitis C virus antibody (HCVAb)	West Nile Virus nucleic acid test
• Anti-human immunodeficiency virus (HIV) antibody (HIV type 1 / 2 type O Ab)	• T Cruzi antibody
• HIV-1 / HCV / HBV Nucleic Acid Test	• Strongyloides antibody to assess for previous infection or exposure
• HHV-6 IgG (Herpesvirus 6 Ab panel)	• T-spot to assess for exposure or history of tuberculosis
• Cytomegalovirus (CMV) IgG and IgM	

¹Primary team should follow up on all testing and order follow up testing and consults as indicated prior to proceeding to IEC therapy ²Patients with recent travel out of the country should be considered for some/all of these additional tests

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APPENDIX C: Infection Prophylaxis Considerations for IEC Therapy

Most patients who receive immune effector cells receive a fludarabine-based chemotherapy regimen prior to IEC infusion. Fludarabine causes immunosuppression and can increase the risk of opportunistic infections. Additionally, patients who receive IEC therapies targeting B-cell are at increased risk of infection due to B-cell aplasia.

Prophylaxis	Preferred Medication	Alternative Medication(s)	Start	Stop	Comment
Viral • Herpes simplex • Varicella zoster	Valacyclovir ¹ 500 - 1,000 mg PO daily	Acyclovir ¹ 400 - 800 mg PO twice daily	IEC infusion day	At least 1 year post IEC infusion; may stop after 1 year if CD4 count > 200 cells/microliter	-
Hepatitis B (only for patients who are positive for HBsAg or HBcAb)	Entecavir ¹ 0.5 mg PO daily	Tenofovir alafenamide 25 mg PO daily or Tenofovir disoproxil fumarate ¹ 300 mg PO daily	2 weeks before IEC	12-24 months post IEC	Consider Infectious Disease and/or Hepatology consult if not already following. Monitor HBV DNA PCR once a month while on prophylaxis and for a year after stopping. Consult Infectious Diseases if entecavir cannot be used or if DNA PCR detectable.
Bacterial (if neutropenia with ANC < 1 K/microliter is expected to last \geq 7 days)	Levofloxacin ¹ 500 mg PO or IV daily	Cefpodoxime ^{1,2} 200 mg PO twice daily or Ciprofloxacin ¹ 500 mg PO twice daily	IEC infusion day or when ANC ≤ 0.5 K/microliter	Continue until ANC > 0.5 K/microliter for 3 consecutive days without growth factor support	Consult Infectious Diseases if patient is allergic to quinolones and cephalosporins

ANC = absolute neutrophil count

¹Adjust for renal function

² Cefpodoxime does not cover pseudomonas

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APPENDIX C: Infection Prophylaxis Considerations for IEC Therapy - continued

Most patients who receive immune effector cells receive a fludarabine-based chemotherapy regimen prior to IEC infusion. Fludarabine causes immunosuppression and can increase the risk of opportunistic infections. Additionally, patients who receive IEC therapies targeting B-cell are at increased risk of infection due to B-cell aplasia.

Prophylaxis	Preferred Medication(s)	Alternative Medication(s)	Start	Stop	Comment
Pneumocystis jiroveci	jiroveci infusion and then transition to sulfamethoxazole/ trimethoprim (SMZ/TMP) (preferred post-IEC infusion) by 3-4 weeks if counts have recovered: • 1 double strength tablet PO every M, W, F or • 1 single strength tablet PO daily or • 1 double strength tablet PO twice daily for two consecutive days/week	-	Within 1 week prior to IEC infusion	At least 1 year post IEC infusion; may stop after 1 year if CD4 count > 200 cells/microliter	SMZ/TMP also has activity against toxoplasma and nocardia
		Pentamidine inhaled 300 mg flat dose every 28 days	Within 1 week prior to IEC infusion	At least 1 year post IEC infusion; may stop after 1 year if CD4 count > 200 cells/microliter	Albuterol nebulizer premedication encouraged
		Pentamidine ¹ IV 4 mg/kg (max 300 mg) every 21 days	Within 1 week prior to IEC infusion	At least 1 year post IEC infusion; may stop after 1 year if CD4 count > 200 cells/microliter	Can cause pancreatitis
		Dapsone 100 mg PO daily or 50 mg PO every 12 hours	3-4 weeks post IEC infusion	At least 1 year post IEC infusion; may stop after 1 year if CD4 count > 200 cells/microliter	Check G6PD level Use caution if patient has sulfa allergy Can cause hemolytic anemia
		Atovaquone 1,500 mg PO daily	Within 1 week of IEC infusion	At least 1 year post IEC infusion; may stop after 1 year if CD4 count > 200 cells/microliter	Must take with a fatty meal. Also has activity against toxoplasma, but inferior to SMZ/TMP.

¹Adjust for renal function

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APPENDIX C: Infection Prophylaxis Considerations for IEC Therapy - *continued*

Prophylaxis	Preferred Medication	Alternative Medication	Start	Stop	Comment	
Fungal (low risk)	Fluconazole ¹ 200 - 400 mg PO or IV daily	Caspofungin 50 mg IV daily ²	IEC infusion day	Continue until ANC > 0.5 K/microliter for 3 consecutive days without growth factor support	-	
Fungal (high risk) ³	Posaconazole 300 mg PO (as tablets) or $IV \text{ daily}^2$	Caspofungin 50 mg IV daily ²	IEC infusion day or when high-risk criteria are met	Continue as clinically indicated ³	-	
HIV	Antiretroviral Therapy (ART) and monitoring	ng per ID recommendations. Obtain	an ID consult on any patient	with HIV.		
HHV-6	Monitor HHV-6 by quantitative PCR from I if patient receives \geq 3 days of corticosteroid			-		
CMV	Routine CMV prophylaxis is not required b CRS/ICANS, if patient receives \geq 3 days of corticosteroids.	Routine CMV prophylaxis is not required but CMV monitoring by PCR is recommended 1-2 times a week if neutropenia lasts \geq 14 days, if patient experiences Grade 3 or 4 CRS/ICANS, if patient receives \geq 3 days of corticosteroids, or if patient develops HLH ⁴ . CMV monitoring is recommended for at least 30 days after completion of				
Immunoglobulin replacement therapy	Hypogammaglobulinemia may be observed after IEC therapies that target B-cells and IgG levels should be checked in such patients when they develop respiratory infections. Immunoglobulin replacement therapy and/or prophylaxis is only indicated for patients who develop hypogammaglobulinemia and recurrent infections.					
Prolonged cytopenias	Grade 3 or 4 cytopenias lasting beyond day 30 have been reported in approximately 30% of patients after IEC therapies. Cytopenias may be managed with filgrastim products; monitor blood counts at least weekly. Continue appropriate prophylactic antimicrobials as described above. Diagnostic bone marrow may be performed to rule out other causes such as myelodysplasia, malignancy, HLH ⁴ , or infection.					
HHV-6 = Herpesvirus 6	HIV = Human Immunodeficiency Virus HLH =	hemophagocytic lymphohistic vtosis				

HHV-6 = Herpesvirus 6 HIV = Human Immunodeficiency Virus HLH = hemophagocytic lymphohistiocytosis

¹Adjust for renal function

²Loading dose of antifungals is not needed if it is being used for prophylaxis

³Posaconazole prophylaxis is recommended for HIGH RISK patients with leukemia, recent allogenic stem cell transplant, prior history of mold infection, neutropenia lasting ≥ 14 days, Grade 3 or 4 CRS/ICANS, those who receive ≥ 3 days of corticosteroids, or those who develop hemophagocytic lymphohistiocytosis (HLH) (see Appendix M). If corticosteroids are given, continue posaconazole for at least 1 month AFTER COMPLETION of corticosteroids. Do not stop posaconazole prophylaxis if ANC < 1 K/microliter. Voriconazole or isavuconazole may be used if the patient had previously been taking them or if posaconazole is not covered by insurance. In the event posaconazole, voriconazole, isavuconazole, or an echinocandin are contraindicated or pose affordability/access issues, then use fluconazole for prophylaxis and consider aspergillus antigen testing at least once a week DURING corticosteroids and for at least a month AFTER completion of corticosteroids. Patients not meeting high risk definitions will be considered to be at LOW RISK for fungal infections and receive prophylaxis as detailed above.

⁴ See Appendix M for Diagnostic Criteria for IEC-associated Fulminant Hemophagocytic Lymphohistiocytosis (HLH) or Macrophage Activation Syndrome (MAS)

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APPENDIX D: ASTCT Grading for CRS¹ (Note: CRS grade should be determined at least twice daily and any time there is a change in patient's status)

CRS Parameter	CRS Grade 1	CRS Grade 2	CRS Grade 3	CRS Grade 4
Fever ²	Yes	Yes	Yes	Yes
		With		
Hypotension ³	No	Requiring IV fluids but not requiring vasopressors	Requiring one vasopressor with or without vasopressin	Requiring multiple vasopressors (excluding vasopressin)
		And/Or		
Hypoxia ³	No	Requiring low-flow O ₂ via nasal cannula ⁴ or blow-by	Requiring O ₂ via high-flow nasal cannula ⁴ , facemask, non-rebreather mask, or Venturi mask	Requiring O ₂ via positive pressure (<i>e.g.</i> , CPAP, BiPAP, and mechanical ventilation)

CPAP = continuous positive airway pressure

BiPAP = bilevel positive airway pressure

¹Organ toxicities associated with CRS may be graded according to CTCAE v5.0 but they do not influence CRS grading

² Fever is defined as temperature \geq 38°C not attributable to any other cause. In patients who have CRS then receive antipyretics or anti-cytokine therapy such as tocilizumab or corticosteroids, fever is no longer required to grade subsequent CRS severity. In this case, CRS grading is driven by hypotension and/or hypoxia.

³ CRS grade is determined by the more severe event: hypotension or hypoxia not attributable to any other cause. For example, a patient with temperature of 39.5°C, hypotension requiring one vasopressor and hypoxia requiring low-flow nasal cannula is classified as having Grade 3 CRS.

⁴Low-flow nasal cannula is defined as oxygen (O₂) delivered at less than or equal to 6 liters/minute. Low flow also includes blow-by oxygen delivery, sometimes used in pediatrics.

High-flow nasal cannula is defined as oxygen delivered at greater than 6 liters/minute.

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APPENDIX E: ASTCT Grading of ICANS¹

Symptom/Sign	Grade 1	Grade 2	Grade 3	Grade 4
ICE Score ²	7-9	3-6	0^{3} -2	0^3 (patient is unarousable and unable to perform ICE)
Depressed level of consciousness ⁴	Awakens spontaneously	Awakens to voice	Awakens only to tactile stimulus	Patient is unarousable or requires vigorous or repetitive tactile stimuli to arouse; stupor or coma
Seizure	-	-	Any clinical seizure (focal or generalized) that resolves rapidly (< 5 minutes) or non-convulsive seizures on EEG that resolve with intervention	Life-threatening prolonged seizure (≥ 5 minutes) or repetitive clinical or electrical seizures without return to baseline in between
Motor findings ⁵	-	-	-	Deep focal motor weakness such as hemiparesis or paraparesis
Raised intracranial pressure ⁶ / cerebral edema	-	-	Focal/local edema on neuroimaging ⁷	Diffuse cerebral edema on neuroimaging, decerebrate or decorticate posturing, cranial nerve VI palsy, papilledema, or Cushing's triad

EEG = electroencephalogram

¹ICANS grade is determined by the most severe event (ICE score, level of consciousness, seizure, motor findings, raised intracranial pressure/cerebral edema) not attributable to any other cause). For example, a patient with an ICE score of 3 who has a generalized seizure is classified as having Grade 3 ICANS.

² See Appendix F for Immune Effector Cell-associated Encephalopathy (ICE) Score

³ A patient with an ICE score of 0 may be classified as having Grade 3 ICANS if the patient is awake with global aphasia or Grade 4 ICANS if the patient is unarousable

⁴Depressed level of consciousness should not be attributable to any other cause (*e.g.*, sedating medication)

⁵Tremors and myoclonus associated with immune effector cell therapies may be graded according to CTCAE v 5.0 but they do not influence ICANS grading

⁶Ophthalmology may be consulted to assess for papilledema if concern for elevated ICP, but otherwise not needed for all patients

⁷ Intracranial hemorrhage with or without associated edema is not considered a neurotoxicity feature and is excluded from ICANS grading. It may be graded according to CTCAE v5.0.

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APPENDIX F: Immune Effector Cell-associated Encephalopathy (ICE) Score

- Orientation: Orientation to year, month, city, hospital: 4 points (1 point each)
- Naming: Name 3 objects (e.g., clock, pen, button): 3 points (1 point each)
- Following commands: (e.g., Show me 2 fingers or close your eyes and stick out your tongue): 1 point
- Writing: Ability to write a standard sentence (*e.g.*, Our national bird is the bald eagle): 1 point
- Attention: Count backwards from 100 by 10: 1 point

Score 10: No impairment

Score 7-9: Grade 1 ICANS

Score 3-6: Grade 2 ICANS

Score 0-2: Grade 3¹ ICANS

Score 0 due to patient unarousable and unable to perform ICE assessment: Grade 4 ICANS

¹ A patient with an ICE score of 0 may be classified as having Grade 3 ICANS if the patient is awake with global aphasia or may be classified as having Grade 4 ICANS if the patient is unarousable

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APPENDIX G: Management of CRS

CRS Grade	CRS Parameter	Management				
CKS Graue	CKS rarameter	Diagnostic Work-Up	Supportive Care	Anti-IEC Therapies		
Grade 1	Fever	 Assess for infection with blood and urine cultures, and chest radiography Cardiac telemetry and pulse oximetry 	 Acetaminophen and hypothermia blanket as needed for the treatment of fever Ibuprofen if fever is not controlled with above; use with caution or avoid with thrombocytopenia or renal dysfunction Empiric broad-spectrum antibiotics and consider filgrastim products if neutropenic Maintenance IV fluids for hydration Symptomatic management of constitutional symptoms and organ toxicities as per standard guidelines If not on seizure prophylaxis, initiate levetiracetam 500 mg PO twice daily 	• Consider tocilizumab ¹ for 1 dose for persistent fever lasting greater than 3 days		

¹See Appendix I for Dosing of IL-6 Antagonists and Alternative Agents

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APPENDIX G: Management of CRS - continued

CRS Grade	CRS Parameter		Management	
CKS Graue	CNS rarailleter	Diagnostic Work-up	Supportive Care	Anti-IEC Therapies
 Fever work-up if not previously performed Assess for infection with blood and urine cultures, and chest radiography Fever work-up if not previously performed If hypotension persists after I dexamethasone, start vasopre transfer patient to ICU, obtain management as in Grade 3 or Symptomatic management of Symptomatic management of Symptomatic management of the symptomatic m		 IV fluid bolus of 500 – 1,000 mL normal saline; repeat once as needed to maintain normal BP If hypotension persists after IV fluids, tocilizumab, and dexamethasone, start vasopressors, transfer patient to ICU, obtain ECHO, and refer to further management as in Grade 3 or 4 CRS Symptomatic management of fever as in Grade 1 CRS Symptomatic management of constitutional symptoms and organ toxicities as per standard guidelines 	 Administer tocilizumab¹ for 1 dose <u>and</u> consider dexamethasone 4 - 10 mg IV for 1 dose (or methylprednisolone equivalent) and reassess in 6 hours or earlier if clinically indicated Tocilizumab may be repeated every 8 hours for up to 3 doses in a 24-hour period 	
Grade 2	Hypoxia	 Pulse oximetry Fever work-up if not previously performed Assess for infection with blood and urine cultures, and chest radiography 	 Use supplemental oxygen as needed If hypoxia persists after above interventions, but oxygen requirement is stable with low-flow nasal cannula, continue close monitoring. If oxygen requirement increases to high-flow nasal cannula, face mask, or positive pressure ventilation, refer to further management as in Grade 3 or 4 CRS Symptomatic management of fever as in Grade 1 CRS Symptomatic management of constitutional symptoms and organ toxicities as per standard guidelines 	 Administer tocilizumab¹ for 1 dose and consider dexamethasone 4 - 10 mg IV for 1 dose (or methylprednisolone equivalent) and reassess in 6 hours or earlier if clinically indicated Tocilizumab may be repeated every 8 hours for up to 3 doses in a 24-hour period

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APPENDIX G: Management of CRS - continued

CRS Grade	CRS Parameter		Ma	nagement
CR5 Graue	CKS Faranieter	Diagnostic Work-up	Supportive Care	Anti-IEC Therapies
	Hypotension	 Obtain ECHO if not performed already Cardiac telemetry Fever work-up if not previously performed Assess for infection with blood and urine cultures, and chest radiography 	 Transfer patient to ICU IV fluid boluses as needed as in Grade 2 CRS Use vasopressors as needed Symptomatic management of fever as in Grade 1 CRS Symptomatic management of constitutional symptoms and organ toxicities as per standard guidelines 	 Tocilizumab¹ as in Grade 2 CRS if not administered previously; tocilizumab may be repeated every 8 hours for up to 3 doses in a 24-hour period If on one vasopressor: tocilizumab as in Grade 2 CRS and dexamethasone 10 mg IV every 6 hours (or methylprednisolone equivalent) If on two vasopressors: tocilizumab as in Grade 2 CRS and dexamethasone 20 mg IV every 6 hours (or methylprednisolone equivalent) If vasopressin and norepinephrine equivalent² is ≥ 15 mcg/minute, follow as in Grade 4 CRS Once CRS improves to Grade 1 or less, taper and/or stop corticosteroids depending on clinical situation
Grade 3	Hypoxia	 Pulse oximetry Fever work-up if not previously performed Assess for infection with blood and urine cultures, and chest radiography 	 Supplemental oxygen including high- flow nasal cannula, face mask, non- rebreather mask, or Venturi mask as needed Symptomatic management of fever as in Grade 1 CRS Symptomatic management of constitutional symptoms and organ toxicities as per standard guidelines 	 Tocilizumab¹ and dexamethasone 10 mg IV every 6 hours (or methylprednisolone equivalent) if not administered previously; tocilizumab may be repeated every 8 hours for up to 3 doses in a 24-hour period If there is no improvement in hypoxia within 24 hours or there is rapid progression of pulmonary infiltrates or sharp increase in FiO₂ requirements, increase dexamethasone to 20 mg IV every 6 hours (or methylprednisolone equivalent) Once CRS improves to Grade 1 or less, taper and/or stop corticosteroids depending on clinical situation

¹See Appendix I for Dosing of IL-6 Antagonists and Alternative Agents

²VASST Trial vasopressor equivalent equation: norepinephrine equivalent dose = [norepinephrine (mcg/minute)] + [dopamine (mcg/kg/minute) / 2] + [epinephrine (mcg/minute)] + [phenylephrine (mcg/minute) / 10]

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APPENDIX G: Management of CRS - *continued*

CRS Grade	CRS Parameter		Ma	nagement		
CKS Graue	CKS rarameter	Diagnostic Work-up	Supportive Care	Anti-IEC Therapies		
Grade 4	Hypotension	 Obtain ECHO if not performed already Cardiac telemetry Fever work-up if not previously performed Assess for infection with blood and urine cultures, and chest radiography 	 Transfer patient to ICU IV fluid boluses as needed as in Grade 2 CRS Vasopressors as in Grade 3 CRS Use vasopressors as needed Symptomatic management of fever as in Grade 1 CRS Symptomatic management of constitutional symptoms and organ toxicities as per standard guidelines 	 Tocilizumab¹ as in Grade 2 CRS if not administered previously; tocilizumab may be repeated every 8 hours for up to 3 doses in a 24-hour period Methylprednisolone 1,000 mg/day in divided doses IV for 3 days followed by rapid taper as per clinical situation If hypotension is refractory for > 24 hours or if patient is deteriorating rapidly, consider additional therapies (see Appendix I) including activation of safety switches if applicable 		
	Hypoxia	 Monitor oxygen saturation while on mechanical ventilation Fever work-up if not previously performed Assess for infection with blood and urine cultures, and chest radiography 	 Positive pressure ventilation including CPAP, BiPAP, mechanical ventilation Symptomatic management of fever as in Grade 1 CRS 	 Tocilizumab¹ as in Grade 2 CRS if not administered previously; tocilizumab may be repeated every 8 hours for up to 3 doses in a 24-hour period Methylprednisolone 1,000 mg/day in divided doses IV for 3 days followed by rapid taper as per clinical situation If hypoxia is refractory for > 24 hours or if patient is deteriorating rapidly, consider additional therapies (see Appendix I) including activation of safety switches if applicable 		

¹See Appendix I for Dosing of IL-6 Antagonists and Alternative Agents

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APPENDIX H: Management of ICANS

ICANS	Sign or		Management				
Grade	symptom	Diagnostic Work-up	Supportive Care	Anti-IEC Therapies			
Grade 1	Encephalopathy and/or depressed le vel of consciousness	 MRI imaging of the brain with and without contrast; CT of brain without contrast may be performed if MRI is not feasible; MRI spine if focal deficits are noted Neurology consultation ICE Score assessment every 6 hours or more frequently if clinically indicated EEG Consider diagnostic lumbar puncture if other causes of encephalopathy are suspected (<i>e.g.,</i> infections, autoimmune, leptomeningeal disease) Add a meningitis-encephalitis panel from CSF in patients with neurologic symptoms that persist or worsen after ICANS therapy and/or if symptoms start after corticosteroids 	 Vigilant supportive care; aspiration precautions; IV hydration Withhold oral intake of food/medications/fluids and assess swallowing; convert all oral medications and/or nutrition to IV if swallowing is impaired Avoid medications that cause central nervous system depression Low doses of lorazepam after EEG is performed (0.25-0.5 mg IV every 8 hours) or haloperidol (0.5 mg IV every 6 hours) may be used with careful monitoring for agitated patients If no seizures on EEG, continue prophylactic levetiracetam If EEG shows focal or generalized convulsive or non- convulsive seizure or convulsive status epilepticus, refer to further management as in Grade 3 or 4 ICANS 	 Dexamethasone 10 mg IV for 1 dose (or methylprednisolone equivalent) and reassess in 6 hours or earlier if clinically indicated If associated with concurrent CRS, add tocilizumab¹ 			
Grade 2	Encephalopathy and/or depressed level of consciousness	• Neurological work-up as in Grade 1 ICANS	• Supportive care as in Grade 1 ICANS	 Dexamethasone 10 mg IV every 12 hours (or methylprednisolone equivalent) If associated with concurrent CRS, add tocilizumab¹ Once ICANS improves to Grade 1 or less, taper and/or stop corticosteroids depending on clinical situation 			

¹See Appendix I for Dosing of IL-6 Antagonists and Alternative Agents

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APPENDIX H: Management of ICANS - continued

ICANS	Sign or	Management		
Grade	symptom	Diagnostic Work-up	Supportive Care	Anti-IEC Therapies
	Encephalopathy and/or depressed level of consciousness	 Neurological work-up as in Grade 1 ICANS Consider repeat neuro-imaging (CT or MRI) every 2-3 days for persistent ≥ Grade 3 encephalopathy Consider diagnostic lumbar puncture if Grade 3 encephalopathy persists ≥ 2 days or earlier if other causes are suspected (<i>e.g.</i>, infections, autoimmune, leptomeningeal disease) Add a meningitis-encephalitis panel from CSF in patients with neurologic symptoms that persist or worsen after ICANS therapy and/or if symptoms start after corticosteroids 	(transfuse to keep platelets > 20-50 K/microliter, fibrinogen > 200 mg/dL and INR < 1.5)	 Dexamethasone 10 mg IV every 6 hours (or methylprednisolone equivalent) If associated with concurrent CRS, add tocilizumab² If Grade 3 encephalopathy is persistent for > 24 hours, increase dexamethasone to 20 mg IV every 6 hours (or methylprednisolone equivalent) Once ICANS improves to Grade 1 or less, taper and/or stop corticosteroids depending on clinical situation
Grade 3	Seizure	 Neurological work-up as in Grade 1 ICANS EEG if clinically indicated (<i>e.g.</i>, ongoing seizures, depressed level of consciousness) Rule out other potential causes of seizure (i.e., beta-lactams, etc.) 	 Transfer to ICU Supportive care as in Grade 1 ICANS For focal or generalized convulsive seizures, or non- convulsive seizures, treat as per Appendix J 	 Dexamethasone 20 mg IV every 6 hours (or methylprednisolone equivalent) If associated with concurrent CRS, add tocilizumab² Once ICANS improves to Grade 1 or less, taper and/or stop corticosteroids depending on clinical situation
	Focal cerebral edema	 Neurological work-up as in Grade 1ICANS Consider repeat neuro-imaging (CT or MRI) every 24 hours until edema resolves or more frequently if clinically indicated 	 Transfer to ICU Supportive care as in Grade 1 ICANS 	 If focal edema is in brain stem or thalamus, methylprednisolone 1,000 mg/day in divided doses IV for 3 days followed by taper depending on clinical situation o If associated with concurrent CRS, add tocilizumab² If focal edema is in other areas of brain, methylprednisolone 1,000 mg/day in divided doses IV for 1 day; assess daily and continue or taper depending on clinical situation o If associated with concurrent CRS, add tocilizumab²

¹Abnormal findings on imaging where correction of hypertension, uremia, and/or coagulopathy should be performed include changes suggestive of typical or atypical posterior reversible encephalopathy syndrome (PRES), temporal lobe and limbic system encephalitis (autoimmune or infection), acute disseminated encephalomyelitis, emboli, vasculitis, strokes, and/or seizure-related changes

²See Appendix I for Dosing of IL-6 Antagonists and Alternative Agents Copyright 2020 The University of Texas MD Anderson Cancer Center Department of Clinical Effectiveness V4

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APPENDIX H: Management of ICANS - *continued*

ICANS	Sign or	Management		
Grade	symptom	Diagnostic Work-Up	Supportive Care	Anti-IEC Therapies
	Encephalopathy and/or depressed level of consciousness	 Neurological work-up as in Grade 1 ICANS Repeat neuro-imaging and lumbar puncture as in Grade 3 ICANS 	 Transfer to ICU Supportive care as in Grade 1 ICANS Consider mechanical ventilation for airway protection If there are new abnormal findings on brain imaging¹ not related to primary malignancy, control hypertension with the goal of maintaining MAP within 20-25 mmHg of baseline MAP; correct any uremia (dialysis if needed) and/or coagulopathy (transfuse to keep platelets > 20 - 50 K/microliter, fibrinogen > 200 mg/dL and INR < 1.5) 	 Methylprednisolone 1,000 mg/day in divided doses IV for 3 days followed by taper as clinically indicated; if associated with concurrent CRS, add tocilizumab² Continue corticosteroids until improvement to less than or equal to Grade 1 ICANS and then taper and stop corticosteroids depending on clinical situation If Grade 4 ICANS is refractory for > 24 hours or if patient is deteriorating rapidly, consider additional therapies (see Appendix I) including activation of safety switches if applicable
Grade 4	Seizure	 Neurological work-up as in Grade 1 ICANS Rule out other potential causes of seizure (i.e., beta-lactams, etc.) 	 Transfer to ICU Supportive care as in Grade 1 ICANS For focal or generalized convulsive or non-convulsive seizure or convulsive status epilepticus, treat as in Appendix J For convulsive status epilepticus, treat as in Appendix K 	 Methylprednisolone 1,000 mg/day in divided doses IV for 3 days followed by taper as clinically indicated; if associated with concurrent CRS, add tocilizumab² If Grade 4 ICANS is refractory for > 24 hours or if patient is deteriorating rapidly, consider additional therapies (see Appendix I) including activation of safety switches if applicable
	Motor Weakness	 Neurological work-up as in Grade 1 ICANS MRI with and without contrast of the spine 	 Transfer to ICU Supportive care as in Grade 1 ICANS 	 Methylprednisolone 1,000 mg/day in divided doses IV for 3 days followed by taper as clinically indicated; if associated with concurrent CRS, add tocilizumab² If Grade 4 ICANS is refractory for > 24 hours or if patient is deteriorating rapidly, consider additional therapies (see Appendix I) including activation of safety switches if applicable
	Diffuse cerebral edema or raised intracranial pressure	 Neurological work-up as in Grade 1 ICANS Consider repeat neuro- imaging as in focal cerebral edema from Grade 3 ICANS 	 Transfer to ICU Supportive care as in Grade 1 ICANS For diffuse cerebral edema or signs of raised intracranial pressure, treat as in Appendix L 	 Methylprednisolone 1,000 mg/day in divided doses IV for 3 days followed by taper as clinically indicated; if associated with concurrent CRS, add tocilizumab² If Grade 4 ICANS is refractory for > 24 hours or if patient is deteriorating rapidly, consider additional therapies (see Appendix I) including activation of safety switches if applicable

¹Abnormal findings on imaging where correction of hypertension, uremia, and/or coagulopathy should be performed include changes suggestive of typical or atypical posterior reversible encephalopathy syndrome (PRES), temporal lobe and limbic system encephalitis (autoimmune or infection), acute disseminated encephalomyelitis, emboli, vasculitis, strokes, and/or seizure-related changes

² See Appendix I for Dosing of IL-6 Antagonists and Alternative Agents

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APPENDIX I: Recommendations for Use of IL-6 Antagonists and Alternative Agents for Management of CRS and ICANS

Drug	Recommended Dose for CRS and/or ICANS	Maximum Dose	Mechanism of Action	Comments
Tocilizumab ¹	8 mg/kg IV	Maximum 800 mg per dose	IL-6 receptor antagonist	 Maximum of 4 doses total over the entire course of CRS and ICANS Dose may be repeated every 8 hours for up to three doses in a 24-hour period
Siltuximab ²	11 mg/kg IV once	-	IL-6 antibody	 Recommended primarily for patients who are intolerant to tocilizumab No more than 1 dose in a 3 week period
Anakinra ³	100 mg subcutaneously daily for 7 days	-	IL-1 receptor antagonist	• Renal dose adjustment may be needed for creatinine clearance < 30 mL/minute
Cyclophosphamide	1,500 mg/m ² IV for one dose	-	Alkylating agent	• Give with mesna 1500 mg/m ² IV over 24 hours for one dose
Anti-thymocyte globulin (rabbit)	1-2 mg/kg IV daily for 3 days	-	Immunosuppressant	 Hypersensitivity reactions can occur; premedicate with diphenhydramine and scheduled dose of corticosteroid Infuse over a minimum of 6 hours
Safety switches	-	_	-	• If the IEC product contains a safety switch (<i>e.g.</i> , iCapsase-9 or EGFRt-positive), the corresponding drug to eliminate those cells can be considered in doses according to manufacturer Examples include rimiducid to eliminate iCaspase-9 or cetuximab to eliminate EGFRt-positive cells

¹ MD Anderson formulary restricted for use in CRS/ICANS and for use in hemophagocytic lymphohistiocytosis (HLH), see Appendix M

² MD Anderson formulary restricted for use in CRS/ICANS

³ Not on MD Anderson formulary; use at MD Anderson is based on internal data in patients with tocilizumab and/or siltuximab failure

MDAnderson IEC Therapy Toxicity Assessment and Management Page 22 of 28 **Cancer** Center (also known as CARTOX) – Adult

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APPENDIX J: Management of Focal or Generalized Convulsive or Non-Convulsive Seizures

- Assess CAB / consider airway protection / check blood glucose
- Consult Neurology
- For focal and generalized convulsive seizures, lorazepam 1-2 mg IV and repeat as needed (to a maximum cumulative dose of 4 mg)
- For electrographical seizures, including non-convulsive status epilepticus, lorazepam 0.5 mg IV and repeat every 5 minutes as needed (to a maximum cumulative dose of 2 mg)
- Levetiracetam 500-1,500 mg IV bolus (in addition to maintenance dose)
- Replete with magnesium as needed to maintain magnesium level > 2 mg/dL
- Thiamine 100 mg IV every 8 hours for 5 days
- If non-convulsive seizures persist, transfer to ICU and add phenobarbital loading dose of 60 mg IV (monitor for respiratory depression, bradycardia and hypotension)
- Maintenance doses after resolution of non-convulsive status epilepticus
- Lorazepam 0.5 mg IV every 8 hours for 3 doses
- Levetiracetam 1,000-1,500 mg IV every 12 hours
- Phenobarbital 30 mg IV every 12 hours (~0.5 mg/kg every 12 hours)
 - Monitor for respiratory depression, bradycardia and hypotension
 - Assess for drug-drug interactions (*i.e.*, may induce metabolism of azole antifungals or other CYP3A4 substrates) and consider alternative therapy if drug interactions are significant
 - Target serum trough levels 15-40 mcg/mL

APPENDIX K: Management of Convulsive Status Epilepticus

- Assess CAB / consider airway protection / check blood glucose
- Transfer to ICU
- Consult Neurology
- Lorazepam 0.1 mg/kg (maximum 4 mg/dose) given at a maximum rate of 2 mg/minute; may repeat in 5 to 10 minutes
- Levetiracetam 500-1,500 mg IV bolus (in addition to maintenance dose)
- Replete with magnesium as needed to maintain magnesium > 2 mg/dL
- Thiamine 100 mg IV every 8 hours for 5 days
- If seizures persist, add phenobarbital loading dose of 15 mg/kg IV (monitor for respiratory depression, bradycardia and hypotension)
- If refractory, consider additional therapies (see Appendix I) including activation of safety switches if applicable
- Maintenance doses after resolution of convulsive status epilepticus
- Levetiracetam 1,000-1,500 mg IV every 12 hours
- Phenobarbital 0.5 mg/kg IV every 12 hours
 - Monitor for respiratory depression, bradycardia and hypotension
 - Assess for drug-drug interactions (*i.e.*, may induce metabolism of azole antifungals or other CYP3A4 substrates) and consider alternative therapy if drug interactions are significant
 - Target serum trough levels 15-40 mcg/mL
- Continuous EEG monitoring if seizures are refractory to treatment

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APPENDIX L: Management of Diffuse Cerebral Edema and/or Raised Intracranial Pressure

For papilledema without diffuse cerebral edema or other signs of raised intracranial pressure	 Acetazolamide 1,000 mg IV followed by 250-1,000 mg IV every 12 hours (monitor renal function and acid/base balance once or twice daily and adjust dose accordingly) Dexamethasone 20 mg IV every 6 hours (or methylprednisolone equivalent) and start taper after resolution of papilledema
For diffuse cerebral edema on neuroimaging or signs of raised intracranial pressure such as decerebrate or decorticate posturing, cranial nerve VI palsy, or Cushing's triad	 Methylprednisolone 1,000 mg/day in divided doses IV for 3 days followed by taper as clinically indicated Elevate head end of patient's bed to an angle of 30 degrees Hyperventilation to achieve target PaCO₂ of 28-30 mmHg, but maintained for no longer than 24 hours Hyperosmolar therapy with either mannitol (20 g/dL solution) or hypertonic saline (3% or 23.4% as detailed below) Mannitol: initial dose 0.5-1 g/kg IV; maintenance dose 0.25-1 g/kg IV every 6 hours while monitoring metabolic profile and serum osmolality every 6 hours; and withhold mannitol if serum osmolality is ≥ 320 mOsm/kg or osmolality gap is ≥ 400 Hypertonic 3% saline: initial dose 250 mL IV over 15 minutes, maintenance dose of 50-75 mL/hour IV while monitoring electrolytes every 4 hours; withhold infusion if serum sodium levels reach ≥ 155 mEq/L) Hypertonic 23.4% saline (for patients with imminent herniation): dose to be administered by physician; initial dose of 30 mL IV; repeat after 15 minutes, if needed If patient has ommaya reservoir, drain CSF to target OP < 20 mmHg Control hypertension with the goal of maintaining mean arterial pressure (MAP) within 20-25 mmHg of baseline MAP; correct any uremia (dialysis if needed) and/or coagulopathy (transfuse to keep platelets > 20-50 K/microliter, fibrinogen > 200 mg/dL and INR < 1.5) Consider neurosurgery consultation and IV anesthetics for burst-suppression pattern on EEG; transfuse to keep platelets ≥ 100 K/microliter if possible and correct coagulopathy in case of surgical intervention Consider additional therapies (see Appendix I) including activation of safety switches if applicable Metabolic profile every 6 hours and daily CT scans of head without contrast, with adjustments in usage of aforementioned medications to prevent rebound cerebral edema, renal failure, electrolyte abnormalities, hypovolemia and hypotension

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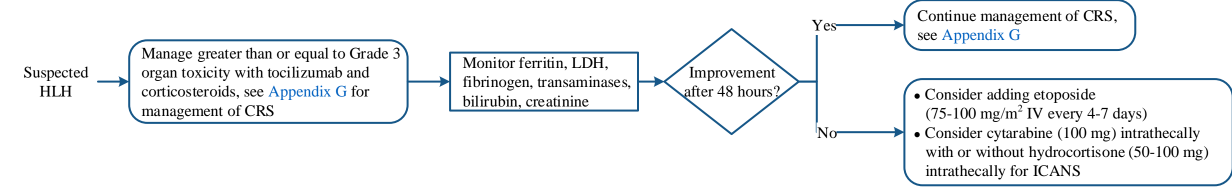
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APPENDIX M: Diagnostic Criteria for IEC-associated Fulminant Hemophagocytic Lymphohistiocytosis (HLH) or Macrophage Activation Syndrome (MAS)

- Consider HLH/MAS if a patient has a peak ferritin > 10,000 ng/mL during the CRS phase and develops any two of the following organ toxicities after IEC therapy
- $\circ \geq$ Grade 3 increase in bilirubin, aspartate transaminase, or alanine transaminase¹
- $\circ \ge$ Grade 3 oliguria or increase in creatinine¹
- $\circ \ge$ Grade 3 pulmonary edema¹
- Presence of hemophagocytosis by morphology and/or CD68 immunohistochemistry in bone marrow or organs
- If HLH/MAS is suspected, obtain baseline fasting triglyceride level and serum soluble IL-2 receptor

¹Grading as per Common Terminology Criteria for Adverse Events, version 5.0

Management of IEC-associated Fulminant HLH / MAS



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APPENDIX N: Determine if the Subject Has Allogeneic IEC-associated Acute Graft-Versus-Host Disease (GVHD)

If a subject has any of the following symptoms or signs within the first 3 months after allogeneic IEC therapy, the subject <u>may have</u> acute GVHD if the symptoms or signs are not attributable to any other cause.

1. Skin rash

2. Diarrhea (may also be associated with nausea, vomiting, and/or anorexia due to upper GI GVHD)¹

3. Total bilirubin $\geq 2 \text{ mg/dL}$

¹ Isolated upper GI GVHD with nausea, vomiting, and/or anorexia is uncommon following allogeneic lymphocyte infusion

APPENDIX O: Determine the Grade of IEC-associated Acute GVHD GVHD Target Organ Staging

Stage	Skin	Liver (bilirubin)	Lower GI (stool output/day)
0	No active (erythymatous) GVHD rash	< 2 mg/dL	< 500 mL/day or < 3 episodes/day
1	Maculopapular rash < 25% BSA	2 - 3 mg/dL	500-999 mL/day or 3-4 episodes/day
2	Maculopapular rash 25 – 50% BSA	3.1 - 6 mg/dL	1000 -1500 mL/day or 5-7 episodes/day
3	Maculopapular rash > 50% BSA	6.1 - 15 mg/dL	> 1500 mL/day or > 7 episodes/day
4	Generalized erythroderma (> 50% BSA) plus bullous formation and desquamation > 5% BSA	> 15 mg/dL	Severe abdominal pain with or without ileus or grossly bloody stool (regardless of stool volume)
Overall Clinical Grade (based on most severe organ involvement) BSA = body surface area			

Grade	Comment	
0	No stage 1-4 of any organ	
Ι	Stage 1-2 skin without liver, upper GI ² , or lower GI involvement	
II	Stage 3 rash and/or stage 1 liver and/or stage 1 upper GI ² and/or stage 1 lower GI	
III Stage 2-3 liver and/or stage 2-3 lower GI, with stage 0-3 skin and/or stage 0-1 upper C		
IV	Stage 4 skin, liver, or lower GI involvement, with stage 0-1 upper GI ²	

² Isolated upper GI GVHD with nausea, vomiting, and/or anorexia is uncommon following allogeneic lymphocyte infusion

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APPENDIX P: Manage IEC-associated Acute GVHD

Grade	Sign or Symptom	Management
Grade I	Skin rash	 Skin biopsy, preferably non-sun exposed site Hydrocortisone cream 1% twice daily to face Triamcinolone cream 0.1% three times daily to affected body area If patient fails triamcinolone, may consider clobetasol cream 0.05% twice daily to body; limit use to no longer than 1-2 weeks All corticosteroid creams should be followed by an emollient such as CeraVe, Aquaphor or Eucerin (creams not lotions) 20-40 minutes after application of corticosteroid
Grade II-IV	 Skin rash > 50% BSA and/or Total bilirubin > 2 mg/dL and/or Diarrhea > 500 mL/day 	 At onset of symptoms that are grade II or higher, consult Stem Cell Transplant team for GVHD workup and management Skin biopsy as above for rash Gastrointestinal consult for flexible sigmoidoscopy with or without upper GI endoscopy with duodenal biopsy¹ DO NOT give GI prep (GoLytely, etc.) unless full colonoscopy ordered Stool culture for <i>C. difficile</i> and GI multiplex panel DO NOT wait for completion of these procedures to start systemic therapy Start prednisone 2 mg/kg/day orally or methylprednisolone equivalent in divided doses

BSA = body surface area

¹Upper GI endoscopy may be considered if patient has nausea, vomiting, and/or anorexia and upper GI GVHD is suspected

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Disclaimer: This algorithm has been developed for MD Anderson using a multidisciplinary approach considering circumstances particular to MD Anderson's specific patient population, services and structure, and clinical information. This is not intended to replace the independent medical or professional judgment of physicians or other health care providers in the context of individual clinical circumstances to determine a patient's care. This algorithm should not be used to treat pregnant women.

DEVELOPMENT CREDITS

This practice consensus algorithm is based on majority expert opinion of the CAR Cell Therapy Toxicity Assessment and Management work group at the University of Texas MD Anderson Cancer Center. It was developed using a multidisciplinary approach that included input from the following providers:

Sherry Adkins, MSN, ANP-C (Lymphoma/Myeloma) Ella Ariza Heredia, MD (Infectious Diseases) Anne Rain Brown, PharmD (Pharmacy Clinical Programs) Aman Buzdar, MD (Clinical Research Administration) Wendy Covert, PharmD (Pharmacy Clinical Programs) Naval Daver, MD (Leukemia) John de Groot, MD (Neurology Oncology) Alison Gulbis, PharmD (Pharmacy Clinical Programs) Cristina Gutierrez, MD (Critical Care & Respiratory Care) Sandra Horowitz, PharmD (Pharmacy Clinical Programs) Patrick Hwu, MD (Cancer Medical Administration) Partow Kebriaei, MD (Stem Cell Transplantation) Monica Elena Loghin, MD (Neurology Oncology) Victor Mulanovich, MD (Infectious Diseases) Sattva Neelapu, MD (Lymphoma/Myeloma)^T Christina Perez[•] Corey Russell, MS, RN (Clinical Nursing) Elizabeth Shpall, MD (Stem Cell Transplantation)^T Sudhakar Tummala, MD (Neurology Oncology) William Wierda, MD, PhD (Leukemia)^T Milena Zhang, PharmD[•]

^TCore Development Team
Clinical Effectiveness Development Team