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## Pediatric Hematology Oncology and Vascular Anomalies/Malformations Consult and referral guidelines

*Pediatric Hematology Oncology Clinic referring provider line: 616.267.1908 (business hours)*

*Helen DeVos Children's Hospital  
100 Michigan Street NE*

*Traverse City  
217 South Madison Street*

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### About Pediatric Hematology Oncology

We care for children and teens from birth to age 21.

#### Most common referrals

- Abnormal CBC results such as neutropenia, thrombocytopenia or anemia
- Abnormal WBC differential
- Refractory iron deficiency
- Hemolytic anemia, such as congenital spherocytosis
- Hemoglobinopathies including sickle cell disease and thalassemia
- Bone marrow failure conditions
- Cancers of all types
- Conditions predisposing to cancer including Fanconi anemia, von Hippel Lindau, hemihypertrophy, Li-Fraumeni, neurofibromatosis and others
- Lymphadenopathy
- Abnormal coagulation tests
- Bleeding disorders including hemophilia, other factor deficiencies, von Willebrand disease
- Hereditary thrombophilia including factor V Leiden mutation
- Hemangiomas: infantile and congenital
- Vascular anomalies and malformations
- Lymphedema

### Pediatric Hematology Oncology Appointment Priority Guide

<b>Immediate</b>	During business hours, call referring provider line at 616.267.1908. After hours and on weekends, contact HDVCH Direct at 616.391.2345 and ask to speak to the on-call pediatric hematology oncologist. We will help decide if your patient should go to the emergency department or be seen immediately in our clinic.
<b>Urgent</b>	Likely to receive an appointment within 2 days. During business hours, call referring provider line at 616.267.1908. After hours and on weekends, call HDVCH Direct (616.391.2345) and ask to speak to the on-call pediatric hematology oncologist regarding an urgent referral.
<b>Routine</b>	Likely to receive an appointment within 10 days to 6 weeks. Send referral via Epic Care Link, fax completed referral form to 616.267.1005, or send referral through Great Lakes Health Connect.

Diagnosis/Symptom	Suggested Workup/Initial Management	When to Refer	Information Needed
<b>Enlarged Lymph Node</b>	<p><b><i>DO NOT, for any reason, give systemic steroids.</i></b></p> <p>Detailed history paying attention to constitutional symptoms, weight loss/failure to thrive, musculoskeletal pain and exposure to cats</p> <p>Physical exam paying attention to weight/growth curve, all lymph node regions of neck, axilla and inguinal areas, abdominal exam for hepatosplenomegaly and bruising on skin exam</p> <p>If suspicion for malignancy is high, send to a hospital laboratory (not satellite lab): CBC with manual differential, CMP, LDH, phosphorus, uric acid. Obtain chest X-ray</p>	<ul style="list-style-type: none"> <li>• <i>Urgent referral:</i> Patient with large lymph nodes (&gt;2 cm) without known infectious etiology, or firm, non-tender or matted lymph nodes. <i>Those with abnormal labs may need to be seen immediately.</i></li> </ul>	<ul style="list-style-type: none"> <li>• Office notes</li> <li>• Any prior work-up including all laboratory studies and chest X-ray (if performed)</li> </ul>
<b>Concern for Acute Leukemia</b>	<p><b><i>DO NOT, for any reason, give systemic steroids.</i></b></p> <p>Detailed history paying attention to constitutional symptoms, weight loss/failure to thrive, musculoskeletal pain and complaints of enlarged lymph nodes.</p> <p>Physical exam paying attention to weight/growth curve, all lymph node regions of neck, axilla and inguinal areas, abdominal exam for hepatosplenomegaly, and bruising or pallor on skin exam.</p> <p>If suspicion is high, send to a hospital laboratory (not satellite lab): CBC with manual differential, CMP, LDH, phosphorus, uric acid. Consider chest X-ray</p>	<ul style="list-style-type: none"> <li>• <i>Immediate referral:</i> If there is concern for acute leukemia based on physical exam or laboratory findings. We will be glad to provide consultation and interpretation of tests and management guidance.</li> </ul>	<ul style="list-style-type: none"> <li>• Office notes</li> <li>• Any prior work-up, including all laboratory studies and chest X-ray (if performed)</li> </ul>

Diagnosis/Symptom	Suggested Workup/Initial Management	When to Refer	Information Needed
<b>Abdominal Mass</b>	<p>Detailed history paying attention to constitutional symptoms, weight loss/failure to thrive, abdominal pain and chronic constipation</p> <p>Physical exam paying attention to blood pressure, weight/growth curve, all lymph node regions of neck, axilla and inguinal areas, abdominal exam for hepatosplenomegaly and abdominal mass</p> <p>Abdominal ultrasound, CBC and CMP are good screening tests</p>	<ul style="list-style-type: none"> <li>• We will be glad to provide guidance about the most efficient and safest way to work up your patient (prior to referral)</li> <li>• If your index of suspicion is high for an abdominal mass, and your patient is being worked up as an outpatient, notify us prior to scheduling diagnostic imaging or triaging to the ER so we are prepared to act upon the results or ER notification. <i>These patients are seen on an immediate or urgent referral basis.</i></li> </ul>	<ul style="list-style-type: none"> <li>• Office notes</li> <li>• Any prior work-up, including all laboratory studies</li> </ul>
<b>Extremity Mass</b>	<p>Detailed history paying attention to constitutional symptoms, weight loss/failure to thrive, pain and inability to bear weight</p> <p>Physical exam paying attention to weight/growth curve, all lymph node regions and extremity exam</p> <p>Plain X-ray films of limbs above and below the area of pain are a good initial step. This should be followed up with MRI of the extremity with and without contrast when index of suspicion is high. If sedation is required, consider chest X-ray PA/lateral to evaluate airway and look for lung disease prior to MRI.</p>	<ul style="list-style-type: none"> <li>• We will be glad to provide guidance about the most efficient and safest way to work up your patient (prior to referral)</li> <li>• Pediatric patients with an extremity mass are often referred to Spectrum Health Orthopedic Oncology at Lemmen-Holton Cancer Pavilion. We can help facilitate that referral. <i>These patients are seen on an urgent referral basis.</i></li> </ul>	<ul style="list-style-type: none"> <li>• Office notes</li> <li>• Any prior work-up including all laboratory studies</li> </ul>

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<b>New Cranial Nerve Palsy, Onset of Weakness, Lethargy</b> (concern for hydrocephalus)	<p>Detailed history paying attention to constitutional symptoms, weight loss/failure to thrive, headache, seizures, decline in school performance and history of NF-1.</p> <p>Physical exam paying attention to weight/growth curve, complete neurologic exam and vision</p>	<ul style="list-style-type: none"> <li>We will be glad to provide guidance about the most efficient and safest way to work up your patient (prior to referral). In many cases, referral to the emergency department is most appropriate.</li> <li>If your index of suspicion is high for an intracranial mass, and your patient is being worked up as an outpatient, we would be grateful for advance notification prior to scheduled radiology imaging or triage to the ER. This helps us to make quick schedule changes in those challenging situations where we are asked to meet a patient in the ER or radiology waiting room. <i>These patients are seen on an immediate or urgent referral basis.</i></li> </ul>	<ul style="list-style-type: none"> <li>Office notes</li> <li>Any prior work-up including all laboratory studies</li> </ul>
<b>Overgrowth Syndromes</b> including hemihypertrophy (hemihyperplasia), Beckwith Weidemann, Sotos syndrome, Megalencephaly Capillary Malformation syndrome, others	<p>These children have a small increased risk for embryonal tumors of childhood such as Wilms tumor, neuroblastoma, hepatoblastoma and adrenal corticocarcinoma</p> <p>Consider ordering abdominal ultrasound prior to consultation visit</p>	<ul style="list-style-type: none"> <li><i>Routine referral:</i> We will generally follow these children until 8 years of age</li> </ul>	<ul style="list-style-type: none"> <li>Office notes</li> <li>Any prior work-up</li> </ul>
<b>Familial Cancer Syndromes</b> including Li-Fraumeni, von Hippel Lindau, Lynch syndrome, Familial Adenomatous Polyposis syndrome	<p>Referral to Medical Genetics for appropriate counseling and screening test</p>	<ul style="list-style-type: none"> <li><i>Routine referral:</i> After confirmed diagnosis of familial cancer syndrome (even if the child has no personal history of cancer) we will discuss and provide a cancer screening regimen specific to the familial cancer syndrome.</li> </ul>	<ul style="list-style-type: none"> <li>Office notes</li> <li>Any prior work-up</li> </ul>

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<b>Conditions Associated with Bone Marrow Failure</b>	<p>Detailed history</p> <p>Physical exam paying particular attention to microcephaly, features of VACTERL association, thumb anomalies and short stature</p> <p>Obtain CBC with manual differential, reticulocyte count</p>	<ul style="list-style-type: none"> <li>• <i>Immediate or urgent referral:</i> Those with significant pancytopenia as leukemia is also in the differential diagnosis.</li> </ul>	<ul style="list-style-type: none"> <li>• Office notes including growth charts</li> <li>• Any prior work-up</li> </ul>
<b>Normocytic Anemia</b> (low hemoglobin with normal MCV)	<p>Detailed history</p> <p>Peripheral smear (manual differential acceptable too), reticulocyte count, direct Coombs, CMP <i>Note: retic count, and direct Coombs can be added to specimen in lab when CBC results show normocytic anemia.</i></p>	<ul style="list-style-type: none"> <li>• We will be glad to provide consultation and interpretation of tests and management guidance</li> <li>• <i>Immediate referral:</i> Patients with positive Coombs test</li> </ul>	<ul style="list-style-type: none"> <li>• Office notes</li> <li>• Any prior work-up including all laboratory studies</li> </ul>
<b>Macrocytic Anemia</b> (low hemoglobin with high MCV)	<p>Detailed history including diet</p> <p>Peripheral smear (manual differential acceptable too), reticulocyte count, TSH with reflexive T4, RBC folate, B12 level, CMP</p>	<ul style="list-style-type: none"> <li>• We will be glad to provide consultation and interpretation of tests and management guidance</li> <li>• All patients should be referred for routine consultation. Some patients may require bone marrow aspiration and biopsy to evaluate for marrow failure or myelodysplastic syndromes.</li> </ul>	<ul style="list-style-type: none"> <li>• Office notes including growth charts</li> <li>• Any prior work-up including all laboratory studies</li> </ul>

Diagnosis/Symptom	Suggested Workup/Initial Management	When to Refer	Information Needed
<b>Microcytic Anemia</b> (low hemoglobin with low MCV)	<p>Detailed history, especially diet history (quantitation of cow's milk), menstrual history and any GI symptoms</p> <p>Labs: Ferritin, TIBC and serum iron. Consider stool for hemocult if appropriate.</p> <p>Trial of oral iron replacement 3 mg/kg of elemental iron given once daily; discontinue cow's milk</p>	<ul style="list-style-type: none"> <li>We will be glad to provide consultation and interpretation of tests and management guidance</li> <li>Immediate or urgent referral: Patients with hemoglobin less than 7g/dL, depending on patient factors</li> <li><i>Routine referral:</i> Patients with lack of response to oral iron supplementation after 2 weeks, ensuring medication adherence</li> <li>Males and non-menstruating females above 2 years with iron deficiency anemia without excessive milk intake but with normal iron absorption will likely need referral to Pediatric Gastroenterology for possible inflammatory bowel disease</li> </ul>	<ul style="list-style-type: none"> <li>Office notes</li> <li>Any prior work-up including all laboratory studies</li> </ul>
<b>Sickle Cell Disease</b> (Hemoglobin SS, hemoglobin SC or sickle beta thalassemia)	<p>Send confirmatory hemoglobin fractionation (not electrophoresis)</p> <p>Immediately start penicillin VK:</p> <ul style="list-style-type: none"> <li>125 mg twice daily for child &lt;3 years</li> <li>250 mg twice daily for child &gt;3 years</li> </ul>	<ul style="list-style-type: none"> <li>All patients should be referred and will be seen for routine consultation</li> </ul>	<ul style="list-style-type: none"> <li>Office notes</li> <li>Newborn screening results</li> </ul>
<b>Hemoglobinopathy Trait including Sickle Cell Trait; Isolated Hemoglobin C Trait or hemoglobin E Trait</b> (without concomitant thalassemia or sickle cell)	<p>Patients with hemoglobinopathy trait do not need ongoing care from a hematologist</p>	<ul style="list-style-type: none"> <li>One-time routine consultation to discuss inheritance, etc., is offered</li> </ul>	<ul style="list-style-type: none"> <li>Office notes</li> <li>Newborn screening results and/or hemoglobin fractionation</li> </ul>
<b>Alpha Thalassemia Trait or Beta Thalassemia Trait</b>	<p>Patients with thalassemia trait do not need ongoing care from a hematologist</p>	<ul style="list-style-type: none"> <li>One-time routine consultation to discuss laboratory findings, inheritance and potential confusion with iron deficiency anemia is offered</li> </ul>	<ul style="list-style-type: none"> <li>Office notes</li> <li>Newborn screening results and/or hemoglobin fractionation</li> </ul>

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<b>Newborn with Rh Sensitization</b>	No additional work up needed beyond usual CBC/retic and bilirubin	<ul style="list-style-type: none"> <li>We like to see these patients prior to hospital discharge; <i>inpatient consult should be requested.</i></li> <li><i>Urgent referral:</i> If born outside of HDVCH, refer so that monitoring plan can be put in place</li> </ul>	<ul style="list-style-type: none"> <li>Office notes</li> <li>Birth records, including laboratory studies</li> </ul>
<b>Newborn with ABO Incompatibility</b>	No additional work up needed beyond usual CBC/retic and bilirubin	<ul style="list-style-type: none"> <li><i>At birth:</i> Hemoglobin less than 12 g/dL</li> <li><i>After discharge:</i> Hemoglobin less than 10 g/dL</li> </ul>	<ul style="list-style-type: none"> <li>Office notes</li> <li>Birth records, including laboratory studies</li> </ul>
<b>Newborn with Family History of Hereditary Spherocytosis (HS)</b>	<p>CBC with manual differential, reticulocyte count. Consider bilirubin if jaundiced</p> <p>Osmotic fragility should not be sent in newborn period.</p>	<ul style="list-style-type: none"> <li>We will be glad to provide consultation and interpretation of tests.</li> <li>Newborn with anemia and hyperbilirubinemia and family history of HS should be referred within 2 weeks of hospital discharge. <i>Patient will be seen for urgent or routine consultation depending on patient factors.</i></li> </ul>	
<b>Isolated Thrombocytopenia</b>	<p>Detailed history including maternal/gestational history if patient is a newborn, recent medication changes or immunizations.</p> <p>Physical exam paying attention to weight/growth curve, oral exam for petechiae, all lymph node regions of neck, axilla and inguinal areas, abdominal exam for hepatosplenomegaly and bruising/petechiae on skin exam.</p> <p>If no bruising and incidentally found, consider repeating CBC in case platelets were clumped.</p> <p>If patient has any associated symptoms, obtain CBC with manual differential, CMP, uric acid, LDH, Coombs test, blood type</p>	<ul style="list-style-type: none"> <li>Platelets &lt;20,000 <i>will require immediate referral/probable hospitalization</i></li> <li>Platelets <math>\geq</math>20,000 but &lt;50,000 without other cytopenias <i>will be seen on an urgent or routine basis depending on patient factors</i></li> <li>Platelets <math>\geq</math>50,000 <i>will be seen on a routine basis</i></li> </ul>	<ul style="list-style-type: none"> <li>Office notes</li> <li>Any prior work-up</li> </ul>



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<b>Isolated Neutropenia</b>	<p>Detailed history including recent illness, history of infections/recurrent fevers, mouth sores, diarrhea, autoimmune disease, race/ethnicity</p> <p>Physical exam paying attention to weight/growth curve, any signs of infection, all lymph node regions of neck, axilla and inguinal areas, abdominal exam for hepatosplenomegaly</p> <p>CBC with manual differential. Consider repeating 1-2 weeks later to see if low absolute neutrophil count persists.</p>	<ul style="list-style-type: none"> <li>• Patients with neutrophils &lt;500/uL (severe neutropenia) <i>with fever <b>should be sent immediately to the ER, and will be hospitalized.</b></i> Contact us to assist with fast-tracking these patients in the ER.</li> <li>• Patients with neutrophils <math>\geq</math>500/uL but &lt; 1000/uL <i>with fever <b>should be sent immediately to the ER for treatment of fever and neutropenia, but will not necessarily be hospitalized.</b></i> Contact us to assist with fast-tracking these patients in the ER.</li> <li>• <i>Routine referral:</i> Patients with persistent neutropenia with ANC &lt;1000/uL over at least 3-6 weeks may need referral. <i>Phone consultation should be first step.</i></li> <li>• Patients with persistent neutropenia with ANC <math>\geq</math>1000/uL but &lt;1500/uL may not necessarily require referral. <i>Phone consultation should be first step.</i></li> </ul>	<ul style="list-style-type: none"> <li>• Office notes including growth charts</li> <li>• Any prior work-up</li> </ul>
<b>Excessive Bruising or Bleeding</b>	<p>Detailed bleeding history (bleeding with surgery including circumcision, epistaxis, bleeding gums, prolonged bleeding with tooth loss, excessive bruising, heavy menses); family history of bleeding</p> <p>Must assess for non-accidental trauma/need for CPS referral as appropriate</p> <p>Obtain PT/INR, PTT, fibrinogen, CBC with manual differential, von Willebrand antigen, von Willebrand ristocetin cofactor activity, factor 8 activity level</p>	<ul style="list-style-type: none"> <li>• <i>Routine referral:</i> child with bleeding history and prolonged PT, PTT; low fibrinogen, von Willebrand antigen, or von Willebrand ristocetin cofactor activity</li> </ul>	<ul style="list-style-type: none"> <li>• Office notes</li> <li>• Any prior work up</li> </ul>

Diagnosis/Symptom	Suggested Workup/Initial Management	When to Refer	Information Needed
<b>Isolated Prolonged PTT</b> in non-hospitalized patient	<p>Detailed bleeding history (bleeding with surgery including circumcision, epistaxis, bleeding gums, prolonged bleeding with tooth loss, excessive bruising, heavy menses); family history of bleeding</p> <p>Obtain repeat PTT as result could be spurious. If abnormal, and no bleeding history, consider repeating in 3-4 weeks as transient anti-phospholipid antibodies are common in children.</p>	<ul style="list-style-type: none"> <li>Note that normal ranges differ for newborns and infants. We will be glad to provide consultation and interpretation of tests and management guidance.</li> <li><i>Routine referral:</i> Child with bleeding history and prolonged PTT confirmed on repeat measurement.</li> </ul>	<ul style="list-style-type: none"> <li>Office notes</li> <li>Any prior work-up</li> </ul>
<b>Isolated Prolonged PT or Combined Prolonged PT &amp; PTT</b> in non-hospitalized patient	<p>Detailed bleeding history (bleeding with surgery including circumcision, epistaxis, bleeding gums, prolonged bleeding with tooth loss, excessive bruising, heavy menses); family history of bleeding</p> <p>Obtain repeat PT and aPTT as result could be spurious.</p>	<ul style="list-style-type: none"> <li>Note that normal ranges differ for newborns and infants. We will be glad to provide consultation and interpretation of tests and management guidance</li> <li><i>Routine referral:</i> Child with bleeding history and prolonged PT or PT and aPTT confirmed on repeat measurement</li> </ul>	<ul style="list-style-type: none"> <li>Office notes</li> <li>Any prior work-up</li> </ul>
<b>Acute Thrombosis</b>	<p>Detailed family history of thrombosis, risk factors for thrombosis (modifiable and unmodifiable).</p> <p>If you are considering initiating anticoagulation prior to referral, please obtain the following labs: PT, aPTT, protein C activity, protein S activity and an antithrombin III activity level</p>	<ul style="list-style-type: none"> <li><i>Immediate referral:</i> Child with acute thrombosis should be directed to emergency department for further management. We will either consult (on a critically ill child) or admit the patient to our service</li> </ul>	<ul style="list-style-type: none"> <li>Office notes</li> <li>Any prior work-up</li> </ul>

Diagnosis/Symptom	Suggested Workup/Initial Management	When to Refer	Information Needed
<p><b>History of Familial Thrombophilia WITHOUT Active Thrombosis</b> (factor V Leiden mutation, prothrombin G20210A mutation, antithrombin III deficiency, protein S deficiency, protein C deficiency)</p>	<p>Detailed family history of thrombosis, risk factors for thrombosis (modifiable and unmodifiable).</p> <p>Testing for thrombophilia is controversial but could be considered in high-risk patients (obesity, tobacco use, immobilization due to surgery) and those in whom oral contraceptives are being considered</p> <p>Avoidance of oral contraceptive pills is strongly recommended in patients with family history of thrombosis. Non-estrogen alternative should be considered.</p>	<ul style="list-style-type: none"> <li>• <i>Routine referral:</i> Child with familial risk factor for thrombosis can be seen for one time consultation to discuss laboratory findings, inheritance and risk reduction</li> </ul>	<ul style="list-style-type: none"> <li>• Office notes</li> <li>• Any prior work-up</li> </ul>
<p><b>Hemangioma</b></p>	<p>Birth history, time course in terms of initial appearance and growth pattern</p> <p>Physical exam paying particular attention to size (documenting dimensions), location, potential for organ compromise, ulceration and presence of petechiae or bruising</p> <p>If concern exists for bruises or petechiae, or non-traumatic bleeding from the vascular tumor, immediate evaluation for Kasabach-Merritt Syndrome should occur (CBC with manual differential, fibrinogen, PT, PTT) as KMS can be life threatening</p> <p>If electing to perform ultrasound of area prior to visit, please order ultrasound with doppler to assess blood flow</p> <p>For small, flat (&lt;1 cm), non-ulcerated, superficial hemangiomas that are not near mucus membranes, topical Timolol may be an option. Please call to discuss.</p>	<ul style="list-style-type: none"> <li>• <i>Urgent referral:</i> child with vascular tumor in organ-threatening location (e.g., on face near eye, nose, mouth) or with significant risk or presence of ulceration. <i>Note: infants with "port-wine stain" of face may have other associated anomalies that will require additional work-up.</i></li> <li>• <i>Routine referral:</i> child with vascular tumor in non-threatening location and without presence of ulceration. <i>Note: infants with segmental hemangioma involving lower body may have other associated anomalies that will require additional work-up.</i></li> <li>• Not all children with infantile hemangiomas need to be referred. We will be glad to provide phone guidance to determine if consultation is warranted.</li> </ul>	<ul style="list-style-type: none"> <li>• Office notes</li> <li>• Any prior work-up</li> </ul>

Diagnosis/Symptom	Suggested Workup/Initial Management	When to Refer	Information Needed
<b>Capillary Malformation on Face in V1, V2 Distribution, High Risk for Sturge-Webber Syndrome</b>	MRI brain should be ordered	<ul style="list-style-type: none"> <li>• We do not generally care for these patients unless they have a capillary malformation in the context of an overgrowth syndrome. In the case of an overgrowth syndrome, routine referral should be made.</li> <li>• Otherwise, these patients should be referred to Pediatric Dermatology, Pediatric Ophthalmology and potentially Pediatric Neurology. If there is uncertainty in terms of diagnosis, we are happy to see the patient and make recommendations for referral.</li> </ul>	
<b>Vascular Anomaly, Vascular Malformation, Lymphedema</b>	<p>Birth history, time course in terms of initial appearance and growth pattern</p> <p>Physical exam paying particular attention to size (documenting dimensions), location, potential for organ or airway compromise, pain or acute swelling.</p> <p>Presence of petechiae, bruising, bleeding.</p> <p>Presence of limb length or girth discrepancy.</p> <p>Ultrasound with doppler of lesion prior to referral is helpful.</p>	<ul style="list-style-type: none"> <li>• <i>Urgent referral:</i> Infant or child with vascular malformation or lymphedema causing pain, with acute swelling or with signs/symptoms of cellulitis.</li> <li>• <i>Routine referral:</i> Infant or child with vascular malformation of limb without pain or acute growth.</li> </ul>	<ul style="list-style-type: none"> <li>• Office notes</li> <li>• Any prior work-up</li> </ul>