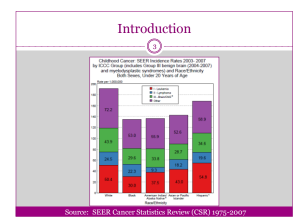
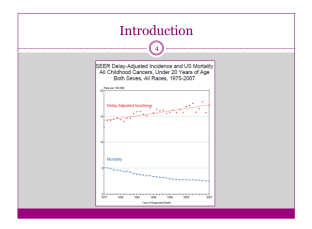


Program Outline

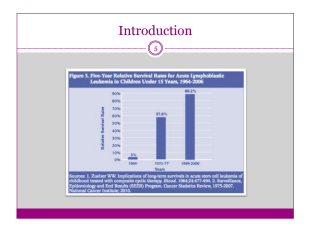
- Introduction
- Types of Pediatric Neoplasms
- Signs and Symptoms
- Causes and Risk Factors
- MPH Rules Solid Tumors
- MPH Rules Heme/Lymph Neoplasms
- Staging Pediatric Tumors
- Collaborative Stage Data Collection System
- Treatment Options
- Future Webcasts
- Q&A







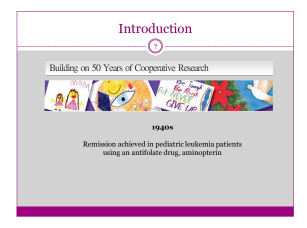


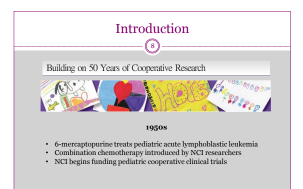


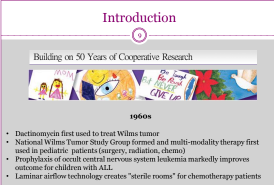


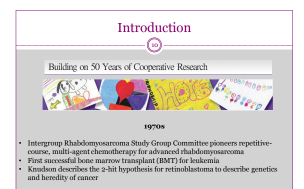






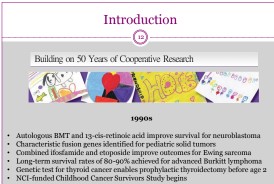




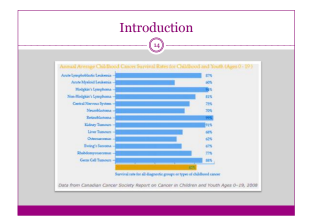




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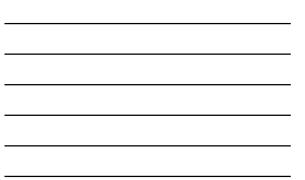


Pediatric Cancer Research Pediatric Cancer Registries



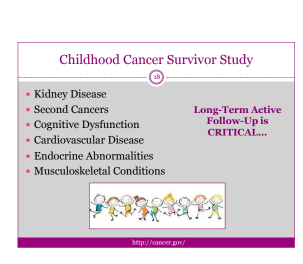






Childhood Cancer Survivor Study

- Diagnosis 1970-1986
- 20,000 person cohort
- Survival at least 5 years
- Chance for long-term effects increase over time
- > 70% at least 1 chronic illness related to treatment
- $\bullet>25\%$ have 3 or more chronic illnesses related to tx



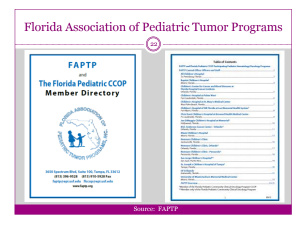
	((19))	
Health Effects	Predisposing Therapy	Clinical Manifestations
Oral/dental	Any chemotherapy in a patient who has not developed permanent dentition	Dental maldevelopment (tooth/root agenesis, microdontia, root thinning and shortening, enamel dysplasia)
	Radiation impacting oral cavity and salivary	Salivary gland dysfunction
	glands	Xerostomia
		Accelerated dental decay
		Periodontal disease
Thyroid	Radiation impacting thyroid gland	Hypothyroidism
		Hyperthyroidism
		Thyroid nodules
Cardiovascular	Radiation impacting cardiovascular structures	Subclinical left ventricular dysfunction
		Cardiomyopathy
		Pericarditis
		Heart valve dysfunction
		Conduction disorder
		Coronary, carolid, subclavian vascular diseas
		Myocardial infarction
		Stroke
	Anthracycline chemotherapy	Subclinical left ventricular dysfunction
		Cardiomyopathy
		Congestive heart failure

Pulmonary	Radiation impacting the lungs	Subclinical pulmonary dysfunction	
	Bleomycin	Pulmonary fibrosis	
Musculoskeletal	Radiation of musculoskeletal tissues in any patient who is not skeletally mature	Growth impairment	
	Glucocorticosteroids	Bone mineral density deficit	
		Osteonecrosis	
Reproductive	Alkylating agent chemotherapy	Hypogonadism	
	Gonadal irradiation	Infertility	
Immune	Splenectomy	Overwhelming post-splenectomy sepsis	
Subsequent neoplasm or disease	Alkylating agent chemotherapy	Myelodysplasia/acute myeloid leukemia	
	Epipodophyllotoxins	Myelodysplasia/acute myeloid leukemia	
	Radiation	Solid benign and malignant neoplasms	

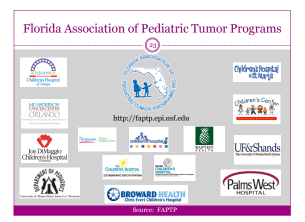


Pediatric Cancer Registries

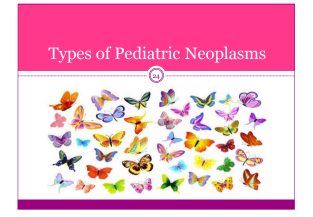
- COG Childhood Cancer Research Network
- CDC NPCR National Childhood Cancer Registry
- FAPTP Florida Consortia Pediatric Cancer Registry
- Cancer Site/Type Specific Registries
- Bone Marrow Donor Registries
- National Children's Study
- Other













	ediatric Neoplasm
	cidence Rates (SEER) by 004 – All Sex, All Race
Leukemia	44.2
Brain/CNS	27.4
Lymphoma	23.2
Soft Tissue	12.0
Germ Cell	11.8
Bone	8.9
Neuroblastoma	7.6
Renal	6.0
Retinoblastoma	3.0
Note: Rates are pe	r 1,000,000 population



o Acute Lymphocytic Leukemia

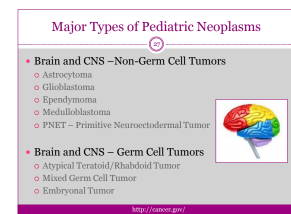


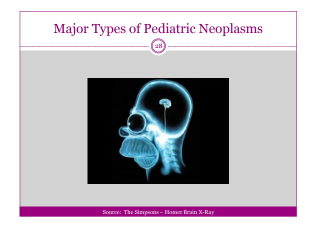
Myeloid Neoplasms

o Myeloid Leukemia Associated with Down Syndrome

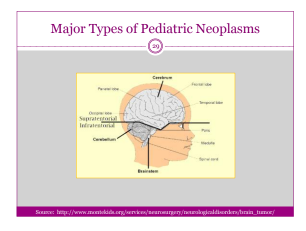
http://cancer.gov/

- o Chronic Myeloid Leukemia
- Acute Myeloid Leukemia

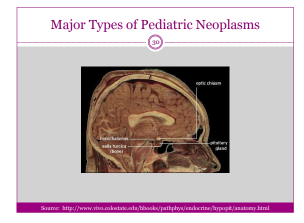










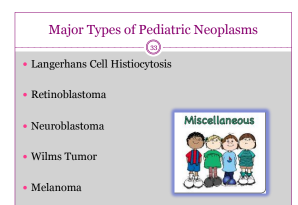


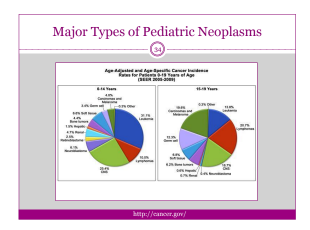


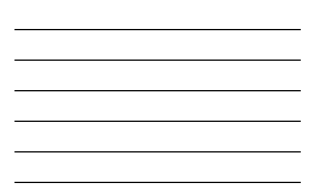
N	/lajor '		diatric I	Neoplasms	
	Sarcoma			a – Connective/Soft Tissu	ie
Ewing Sarcoma – Undifferentiated pPNET pPNET – PNET with Neural Differentiation		ion			
Osteosarco	ma		Rhabdomyosarcoma		
Odontogen	ic Sarcoma		Lymphangiosarcoma		
Chondrosa	rcoma		Fibrosarcoma		
		Ewing Fa	mily of Tumor	s	
	9473/3	PNET		Brain/CNS	
	9364/3	pPNET		Soft Tissue	
	9365/3	Askin Tumor		Soft Tissue	
	9260/3	Ewing Sarcoma		Bone	
	9260/3	Ewing Sarcoma - Ex	tra Osseous	Soft Tissue	
		http://c	ancer.gov/		

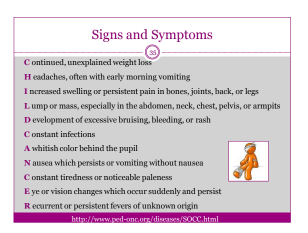
	-(32) .				
Childhood Soft T	issue S	arcoma	Treatm	ent (PDC	8)	
National Cancer		Age 5- 9 Y	Age 10- 14 y	Age 15- 19 y	% of	the Total Number of ST Cases <20 y
Other specified soft tissue sarcomas	198	220	512	856	38	
Ewing tumor and Askin tumor of soft tissue	22	::8	57	81		4
pPNET of soft tissue	21	19	29	42		2.4
Extrarenal rhabdoid tumar	37	3	8	3		1
Liposareomas	5	6	22	66		2
Fibrohistiocytic tumors *	53	69	171	293		12
Leionyosarcomas	13	19	22	57		2.4
Synovial sarcomas	12	39	133	204		8.3
Blood vessel tumors	15	7	11	33		1.4
Osseous and chondromatous neoplasms of soft tissue	1	5	9	36		0.6
Alveolar soft parts sarcoma	3	7	19	26		1
Miscellaneous soft tissue sarcomas	10	18	31	35		2
Unspecified soft tissue sarcomas	70	58	136	163	9	





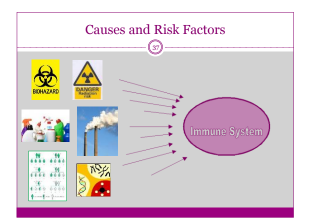


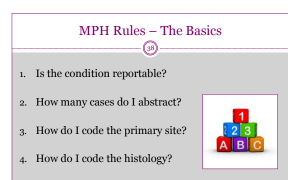




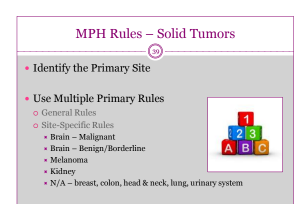


12





5. How do I code the grade?



MPH Rules – Solid Tumors

- Histology Coding Rules
- What Drives Treatment Decisions?
- Pediatric Pathology Characteristics and Terminology
- Tumor Characteristic Testing
- Tumor Marker Testing



• Profile

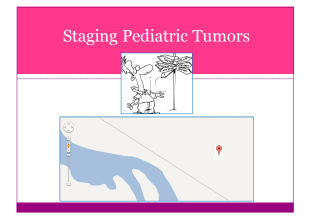


MPH Rules – Heme/Lymph Neoplasms

- Hodgkin Lymphoma
- Non-Hodgkin Lymphoma
- Acute Lymphoblastic Leukemia
- Acute Myeloid Leukemia
- Chronic Myeloid Leukemia
- Myeloid Leukemia Associated with Down Syndrome
- Langerhans Cell Histiocytosis solitary/multifocal

MPF	H Rules – Heme/Lymph Neopla	asms
For East View Faucultes Tauls Hey X Gangle	* • • • • • • • • • • • • • • • • • • •	Sipche Ma, e @i = Pape= Safety= Tools= @=
	National Cancer Institute U.L. National endoes of Health (News, cancer pre 2012 Hematlopolietic and Lymphoid Database Desire for service and access (New 20, 2012 Cancer of the Cancer of	į
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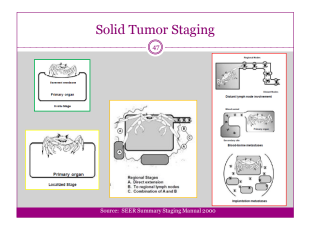
Staging Pediatric Tumors

- Wilms Tumor Study Group Staging
- International Neuroblastoma Staging System
- Children's Oncology Group Neuroblastoma Risk Grouping
- Intergroup Rhabdomyosarcoma Study Staging System
- Soft Tissue Sarcoma Tumor Pathological Grading System
- FNCLCC Grading System
- TNM Staging System

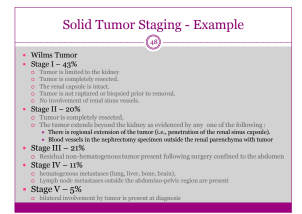


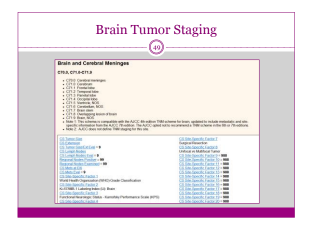




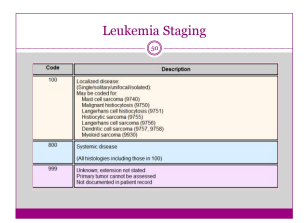


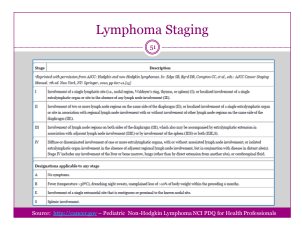


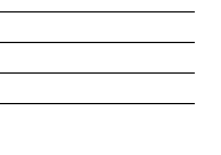












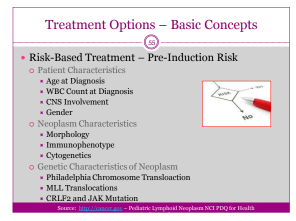


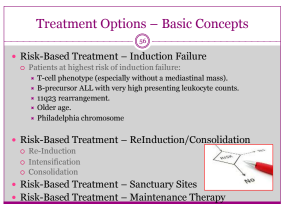




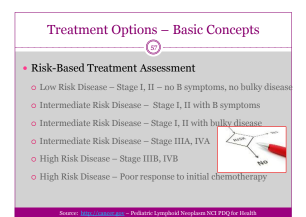
	reatment Options – Basic Concepts
	Pre-Induction Risk Assessment
	Induction Therapy
	Post-Induction Assessment
	Re-Induction Therapy
	Intensification/Consolidation Therapy
	Post-Consolidation Assessment
	BMT/Stem Cell Transplant
	Maintenance Therapy
Ĩ	Maintenance Assessment

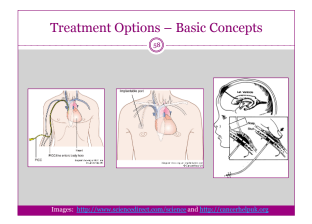














Treatment Options – Lymphoid Neoplasms

- Hodgkin Lymphoma
- Non-Hodgkin Lymphoma
- Chronic Lymphocytic Leukemia
- Acute Lymphocytic Leukemia
- Other Lymphoid Neoplasm



Table 5. Low-Risk Disease	(Stages I–IIA; N	(o Bulky Di	sease; No B Sy	mptoms)	Enla
Chemotherapy (No. of Cycles)*	Radiation (Gy)	Stage	No. of Patients	Event-Free Survival (No. of Years of Follow-up)	Survival (No. of Years o Follow-up)
VAMP (4) [38]	15-25-5, IFRT	CS I/IIb	110	89 (10)	96 (10)
VAMP (4) [44]	25-5, IFRT/None	CS I/II ^b	41/47	88/89 (5)	100/100 (5)
COPP/ABV (4) [14,17]	21, IFRT/None	CS IA/B, IIA:	94/113	100/89 (10) ⁱⁱ	97/96 (10) ¹
OEPA/OPPA (2) [18]	20-35, IFRT/None	I, IIA	281/113	94/97 (5)	N/A
ABVE (2-4) [47]	25.5, IFRT	IA, IIA, IIIAs	51	91(6)	98 (á)

Source: Mosaic Rainbow and Woodland Forest - http://www.et

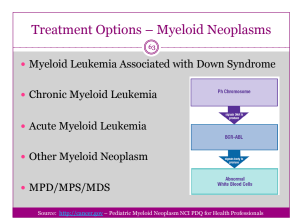


Table 6. Intermediate-Ris	ik Disease (All S	Stage I and Stage II	Patients Not	Classified as Early Stage; Stage III	A; Stage IVA) Enlarge
Chemotherapy (No. of Cycles)*	Radiation (Gy)	Stage	No. of Patients	Event-Free Survival (No. of Years of Follow-up)	Survival (No. of Years of Follow-up)
COPP/ABV (6) [17]	21, IFRT/None	CS I/II ^b , CS IIB, CS III	103/122	84/78 (10)	100 (3)
OEPA/OPPA (2) + COPP (2) [18]	20-35, IFRT	$\mathrm{II}_{1}\mathrm{A},\mathrm{IIB},\mathrm{IIIA}$	212	92 (5)	N/A
OEPA/OPPA (2) + COPDAC (2) [37]	20–35, IFRT	II ₂ B, III ₈ A/B, IIIB, IVA/B	139	88.3 (5)	98.5 (5)
ABVE-PC (3-5) [32]	21, IFRT	IB, IIA, IIIA	53	84 (5)	95 (5)



Table 7. High-Risk Disease (St	ages IIIB, IVB)			Enlare
Chemotherapy (No. of Cycles)*	Radiation (Gy)	Stage	No. of Patients	Event-Free Survival (No. of Years of Follow-up)	Survival (No. of Years of Follow-up)
OEPA/OPPA (2) + COPP (4) [18]	20-35, IFRT	II18B, III8A/B, IIIB, IVA/B	265	91 (5)	N/A
DEPA/OPPA (2) + COPDAC (4) [37]	20-35, IFRT	II18, III8A/B, IIIB, IVA/B	239	Bá.9 (5)	94-9 (5)
ABVE-PC (3-5) [32]	21, IFRT	IB, IIA, IIIA	163	85 (5)	95 (5)
BEACOPP (4); COPP/ABV (4) (RER; girls) [40]	None	IIB, IIIB, IV	38	94 (5)	97 (5)
BEACOPP (4); ABVD (2) (RER; boys) [40]	21, IFRT	IIB, IIIB, IV	34		
BEACOPP (8) (SER) [40]	21, IFRT	IIB, IIIB, IV	25		



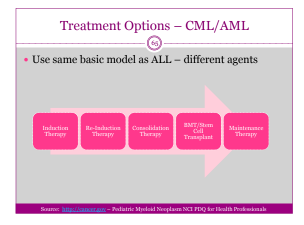




Treatment Options – CML/AML

- About 85% of patients are diagnosed in the chronic phase
- The accelerated phase of CML is characterized by 10% -19% blasts in the WBC of peripheral blood (WHO)
- Note: There are alternative ways to define the accelerated phase proposed by Sokal et al., the International Bone Marrow Transplant Registry, and MD Anderson.
- The **blast phase**, also referred to as blast crisis, is most often defined as >20% blasts WBC of peripheral blood

Source: http://cancer.gov - Pediatric Myeloid Neoplasm NCI PDQ for Health Professionals



Treatment Options – CML/AML

- Ph⁺ chronic phase CML is typically treated with a tyrosine kinase inhibitor (TKI).
- TKIs include imatinib, nilotinib or dasatinib.
- All TKIs are given orally so there will be no "administration" documentation rather the patient will be given prescriptions

Source: http://cancer.gov – Pediatric Myeloid Neoplasm NCI PDQ for Health Professionals

• Other treatment options include clinical trial or Hematopoietic Stem Cell Transplant [HSCT].

Treatment Options – CML/AML

- Patients are evaluated for a cytogenetic response
- Treatment is based on response
- o Complete Response no Ph⁺ positive metaphases
- o Partial Response 1 %-35% $\rm Ph^+$ positive metaphases
- o Major Response o %-35% Ph⁺ positive metaphases

Source: http://cancer.gov - Pediatric Myeloid Neoplasm NCI PDQ for Health Professionals

- Complete + Partial
- o Minor > 35% Ph⁺ positive metaphases

Treatment Options – CML/AML

- Patients achieving a complete hematologic response are continued on their current medication at the same dose.
- Patients who fail to achieve a complete hematologic response are evaluated for compliance, drug-drug interaction and possibly mutational status but generally are switched to an alternate TKI as second line treatment.
- Other treatment options include evaluation and discussion of HSCT and clinical trial.

Treatment Options – CML/AML

ncer.gov – Pediatric Myeloid Neoplasm NCI PDQ for Health Professionals

- Growth Factors
 - o filgrastim (Neupogen®)
 - o pegfilgrastim (Neulasta ®)
- Diuretics (aldactone, hydrochlorothiazide [HCTZ]), steroids (prednisone 20mg/day x 3 for effusions)
- Topical steroids (hydrocortisone cream)
- Antidiarrheal agents (loperamide [Imodium®])

Source: http://cancer.gov - Pediatric Myeloid Neoplasm NCI PDQ for Health Professionals

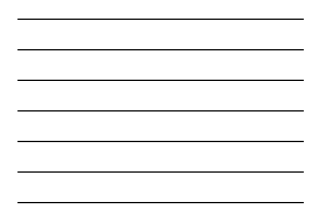
· Analgesics (acetaminophen, ibuprofen)

Treated Bas	ed on Histolog	y and Location	
Tumor Type	Pathologic Subtype	Staging and Treatment of Newly Diagnose and Recurrent Disease	
CNS = central nervous system.			
Astrocytomas and Other Tumors of Glial Origin			
	Diffuse fibrillary astrocytoma	Childhood Astrocytomaa Treatment	
	Gemistocytic astrocytoma		
	Oligoastrocytoma		
	Oligodendroglioma		
- Low-Grade Astrocytomas	Pilocytic astrocytoma		
	Pilomyzoid astrocytoma		
	Pleomorphic xanthoastrocytoma		
	Protoplasmic astrocytoma		
	Subependymal giant cell astrocytoma		
- High-Grade Astrocutomas	Anaplastic astrocytoma	Childhood Astrocytomas Treatment	

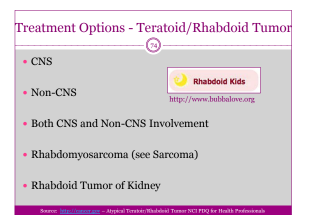
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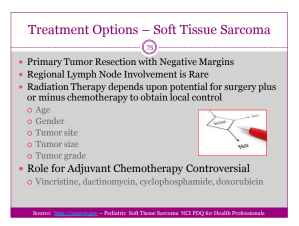
Treated B	ased on Histolo	gy and Location
Tumor Type	Pathologic Subtype	Staging and Treatment of Newly Diagnos and Recurrent Disease
	Anaplastic oligoastrocytoma	
	Anaplastic oligodendroglicena	
	Giant cell glioblastoma	
	Glioblastoma	
	Gliomatosis eerebri	
	Gliosarcoma	
Brain Stem Glioma		
	Diffuse intrinsic pontine gliomas	Childhood Brain Stem Glioma Treatment
	Focal or low-grade brain stem gliomas	Chaidhood Brain Stem Ghoma Treatment
CNS Embryonal Tumors		
	Anaplastic	
	Classie	
- Medulloblastoma	Desmoplastic/nodular	Childhood CNS Embryonal Tumors Treatment
	Large cell	

Treated Ba	sed on Histolo	gy and Location
Tumor Type	Pathologic Subtype	Staging and Treatment of Newly Diagnoses and Recurrent Disease
	Medulloblastoma with extensive nodularity	
	CNS ganglioneuroblastoma	
- CNS Primitive Neuroectodermal	CNS neuroblastoma	
Tumors (PNETs)	Ependymoblastoma	
	Medulloepithelioma	
	Pineal parenchymal tumor of intermediate differentiation	
- Tumors of the Pineal Region	Pineoblastoma	
, ,	Pineocytoma	
	Papillary tumor of the pineal region	
– CNS Atypical Teratoid/Rhabdoid Tumor		Childhood CNS Atypical Teratoid/Rhabdoid Tumo Treatment
CNS Germ Cell Tumors		
- Germinamas		Childhood CNS Germ Cell Tumors Treatment



T	(73)	
Tumor Type	Pathologic Subtype	Staging and Treatment of Newly Diagnosed and Recurrent Disease
	Immature teratomas	
- Teratomas	Mature teratomas	
	Teratomas with malignant transformation	
	Chorioearcinoma	
– Non-Germinomatous Germ Cell	Embryonal careinoma	
Tumors	Mixed germ cell tumors	
	Yolk sac tumor	
Craniopharyngioma		Childhood Craniopharyngioma Treatment
Ependymoma		Childhood Ependymoma Treatment





Treatment Options – Ewing Sarcoma

• Pre-Treatment Factors

- o Site of Tumor
- o Tumor Size or Volume
- o Age of Patient
- Gender (favorable girls)
- o Metastases
- Standard Cytogenetics
- o Detectable Fusion Transcripts
- Mon Ho

- Surgery
- Chemo vincristine, doxorubicin, cyclophosphamide
- Radiation Therapy

Treatment Options – Wilms Tumor

- Preoperative chemotherapy prior to nephrectomy is indicated in the following situations:[10,17,20-23]
 - o Metachronous bilateral Wilms tumor.
 - Wilms tumor in a solitary kidney.
 - ${\rm \circ}\,$ Extension of tumor thrombus above the level of the hepatic veins.
 - Tumor involves contiguous structures whereby the only means of removing the kidney tumor requires removal of the other structures (e.g., spleen, pancreas, colon but excluding the adrenal gland).
- Pulmonary compromise due to extensive pulmonary metastases.
- Patients with massive, nonresectable unilateral tumors, bilateral tumors, or venacaval tumor thrombus are candidates for preoperative chemotherapy

Treatment Options – Wilms Tumor

- Pre-Surgical Chemotherapy for High Risk Group
- Nephrectomy
- Chemotherapy
- Radiation Therapy

Table 2. Standard Chemotherapy Regimens for Wilms Tumor		
Regimen Name	Regimen Description	
Regimen EE-4A [1]	Vincristine, dactinomycin x 18 weeks postnephrectomy	
Regimen DD-4A [1]	Vineristine, dactinomycin, doxorubicin x 24 weeks postnephrectomy	
Regimen I [2]	Vincristine, doxorubicin, cyclophosphamide, etoposide x 24 weeks	

Treatment Options – Neuroblastoma

- Low-Risk Neuroblastoma
 - o Surgery
 - Chemo carboplatin, cyclophoasphamide, doxorubicin, etoposide
- Intermediate-Risk Neuroblastoma
 - o Surgery
 - Chemo as above x 2 cycles
 - o Dose Intensive Multi-Agent Chemo
- High-Risk Neuroblastoma
- o Dose Intensive Multi-Agent Chemo as above plus ifosfamide, cisplatin o Surgery
- Response Assessment then next steps

Treatment Options - Retinoblastoma

- Goals of Treatment
- ${\rm \circ}~{\rm Eradicate}$ the disease to save the patient's life.
- Preserve as much vision as possible.
- Decrease risk of late sequelae from treatment, particularly subsequent neoplasms.
- Enucleation
- Radiation Therapy (beam or brachytherapy)
- Local Treatments (Cryotherapy/Laser Therapy)
- Chemo carboplatin, etoposide, vincristine
- Subteton (subconjunctival) Chemo carboplatin
- Opthalmic Artery Infusion Chemo topotecan, carboplatin

Late Effects of Treatment

Childhood Cancer Survivors Study

- Chance for long-term effects increase over time
- o > 70% at least 1 chronic illness related to treatment
- 0 > 25% have 3 or more chronic illnesses related to tx
- Kidney Disease
- Second Cancers
- Cognitive Dysfunction
- Cardiovascular Disease
- Endocrine Abnormalities
- Musculoskeletal Conditions

References and Resources



- o Childhood Cancers Fact Sheet
- o Childhood Acute Lymphoblastic Leukemia
- o Childhood Acute Myeloid Leukemia
- o Childhood Brain and Spinal Cord Tumors Overview
- o Childhood Astrocytoma
- Childhood CNS Embryonal Tumors
- Childhood CNS Atypical Teratoid/Rhabdoid Tumors
- Childhood Hodgkin Lymphoma
- o Childhood Non-Hodgkin Lymphoma
- o Ewing Sarcoma
- o Childhood Soft Tissue Sarcoma
- o Childhood Rhabdomyosarcoma
- NeuroblastomaWilms Tumor



- Florida Association of Pediatric Tumor Programs (FAPTP)
- Children's Oncology Group (COG)







