

# NCG GUIDELINES 2019



PAEDIATRIC HEMATOLYMPHOID AND SOLID TUMOURS  
- 2019

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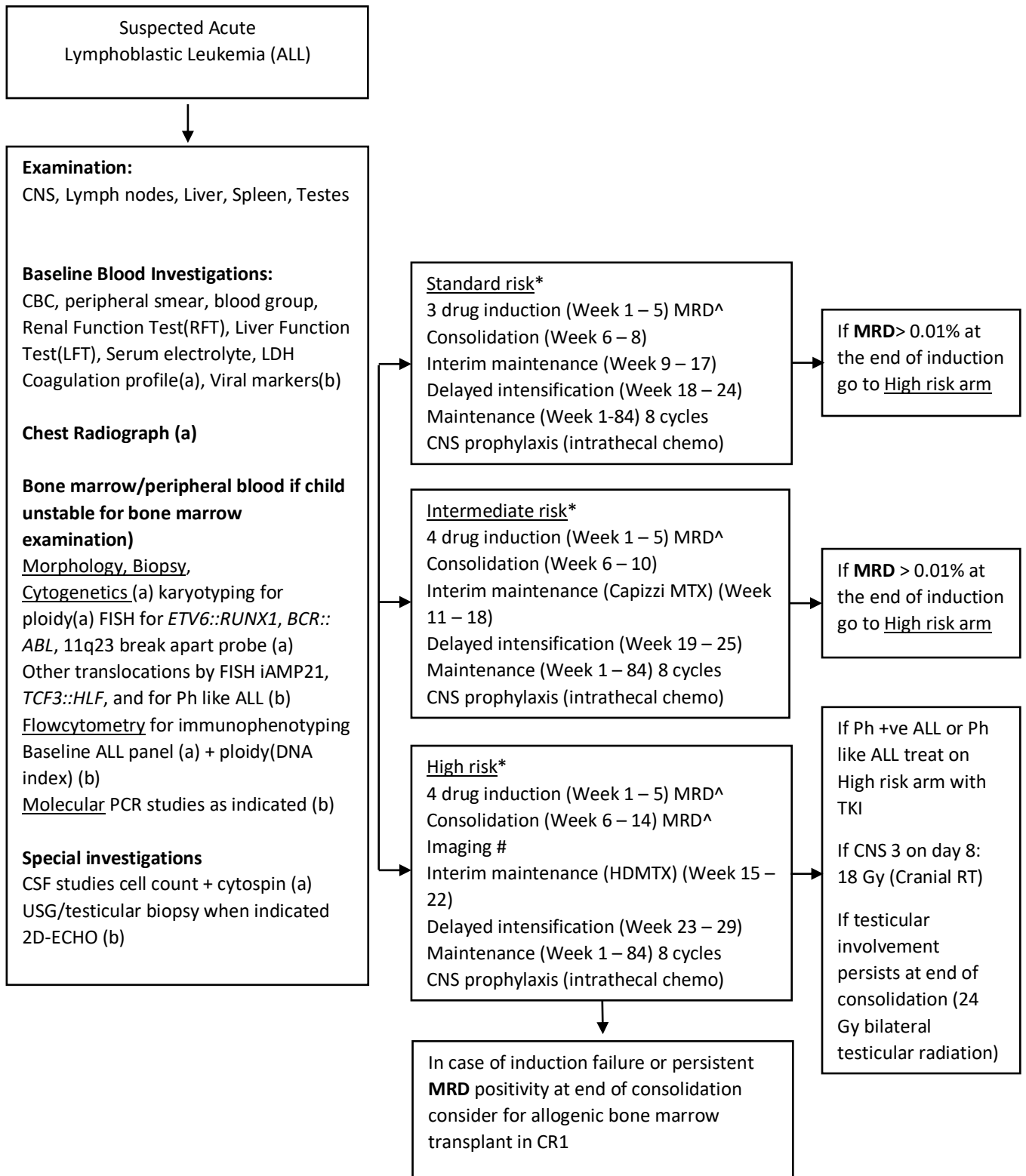
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## PEDIATRIC HEMATOLYMPHOID TUMORS

## PEDIATRIC ACUTE LYMPHOBLASTIC LEUKEMIA

### Treatment Algorithm: Newly diagnosed ALL



<sup>^</sup> MRD = Minimal residual disease by flow cytometry or PCR to assess response to treatment. Risk stratification depends upon MRD assessment. To be done at centers with expertise in flowcytometry and/PCR MRD detection. # Imaging like

CXR for patients with T-ALL or LBL with mediastinal mass at the end of consolidation at the end of consolidation to assess response to treatment.



## Risk Stratification

### Standard Risk

B lineage ALL **and**  
Age > 1 and <10 years **and**  
WBC <50,000/mm<sup>3</sup> **and**  
Prednisolone good responder **and**  
No testicular or bulky disease **and**  
No high-risk cytogenetics **and**  
MRD <10<sup>-4</sup> at week 5

### Intermediate risk

B lineage ALL **and**  
Age ≥ 10 years  
**or** Presenting WBC ≥  
50,000/mm<sup>3</sup>  
**or** testicular/bulky disease **and**  
Prednisolone good response **and**  
No high-risk cytogenetics **and**  
MRD <10<sup>-4</sup> at week 5

### High risk

T lineage ALL **or**  
B lineage ALL **and**  
High risk cytogenetics  
BCR-ABL, iAMP21, MLL rearranged,  
t(17;19), Hypodiploidy (< 45  
chromosomes or DNA index <0.81)  
**or** Prednisolone poor response  
**or** MRD positive (>10<sup>-4</sup>) at week 5  
**or** No CR at the end of induction  
**or** CNS disease

### Protocol options:

- ICICLE-2014
- BFM 90
- BFM 95 (ALL)
- UK ALL protocols
- COG protocols
- MCP 841
- Any others with similar backbone

### Standard risk B ALL (ICICLE-2014/BFM protocols/UK ALL protocols/MCP 841)

#### 3 drug Induction

Vincristine 1.5mg/m<sup>2</sup> IV bolus day 8,15,22,29

L Asparaginase 10,000units/m<sup>2</sup> IM Day 18, 21, 24, 27 or PEG Asparaginase 1,000units/m<sup>2</sup> x 1 dose IM Day 16

Prednisolone 60mg/m<sup>2</sup> PO x 3 to 5 weeks

Intrathecal Methotrexate on Day 8, 15, 35

#### Consolidation

6MP 60mg/m<sup>2</sup> PO Day 1 - 21

Intrathecal Methotrexate Day 8, 15

Some protocols also will give

Cyclophosphamide 1000mg/m<sup>2</sup> IV infusion with or without MESNA Days 1, 15

Cytarabine 75mg/m<sup>2</sup> IV bolus or Subcutaneously on Days 2-5, 8-11, 16-19, 23-26

6 MP 60mg/m<sup>2</sup>/day PO Day 1 - 28

#### Interim maintenance

Dexamethasone 6mg/m<sup>2</sup>/day PO Day 1-5 and Day 29-33

Vincristine 1.5mg/m<sup>2</sup> IV bolus Day 1, 29

6 MP 60mg/m<sup>2</sup>/day PO Day 1 - 49

Methotrexate 20mg/m<sup>2</sup> PO once a week x 7 doses

Intrathecal Methotrexate Day 1, 31

#### Delayed intensification

Dexamethasone 10mg/m<sup>2</sup> PO Day 1-5 and Day 15-19

Vincristine 1.5mg/m<sup>2</sup> IV bolus Day 1,8,15

Doxorubicin 25mg/m<sup>2</sup> IV bolus or short infusion day 1, 8, 15 or Mitoxantrone 10mg/m<sup>2</sup> IV bolus Day 1

L asparaginase 10,000units/m<sup>2</sup> IM on Day 4, 7, 10, 13 or PEG asparaginase 1000units/m<sup>2</sup> IM on Day 4

Intrathecal Methotrexate Day 1, 15

Cyclophosphamide 1000mg/m<sup>2</sup> IV over 30 min with or without MESNA on day 29

Cytarabine 75mg/m<sup>2</sup> IV bolus or subcutaneous Days 30-33 and day 37 - 40

6 Mercaptopurine 60mg/m<sup>2</sup> PO Day 29 - 42

#### Maintenance phase

Every 3 months x 8 such cycles

Intrathecal Methotrexate on day 1

6MP 60mg/m<sup>2</sup> PO Days 1 – 90

Methotrexate 20mg/m<sup>2</sup> PO every week x 12 weeks

#### **Intermediate risk B ALL (ICICLE-2014/BFM protocols/UK ALL protocols/MCP 841)**

##### Intermediate risk 4 drug induction

Vincristine 1.5mg/m<sup>2</sup> IV bolus Day 8,15,22,29

Daunorubicin 25-30mg/m<sup>2</sup> IV Bolus or infusion Day 8, 15

L Asparaginase 10,000units/m<sup>2</sup> IM Day 9, 12, 15, 18, 21, 24, 27, 30 or PEG Asparaginase 1,000units/m<sup>2</sup> IM Day 9, 23

Prednisolone 60mg/m<sup>2</sup> PO x 3 to 5 weeks

Intrathecal Methotrexate on Day 8, 15, 35

##### Consolidation

Cyclophosphamide 1000mg/m<sup>2</sup> IV infusion with or without MESNA Days 1, 15

Cytarabine 75mg/m<sup>2</sup> IV bolus or subcutaneously on Days 2-5, 8-11, 16-19, 23-26

6MP 60mg/m<sup>2</sup> PO daily Day 1 - 28

Intrathecal Methotrexate Day 8, 15

##### Consolidation with protocol I2A (MCP 841 only)

Cytarabine 2000mg/m<sup>2</sup> infusion 12 hrly on day 1,2 and 15,16

##### Interim maintenance

Vincristine 1.5mg/m<sup>2</sup> IV push Day 2,12,22,32,42

Methotrexate 100mg/m<sup>2</sup> IV bolus with +50mg/m<sup>2</sup> dose escalation every 10 days Day 2, 12, 22, 32, 42

Intrathecal Methotrexate Day 1, 31

Some protocols will administer L Asparaginase or PEG Asparaginase

##### Delayed intensification

Dexamethasone 10mg/m<sup>2</sup> PO Day 1-5 and Day 15-19

Vincristine 1.5mg/m<sup>2</sup> IV bolus Day 1, 8, 15

Doxorubicin 25 – 30 mg/m<sup>2</sup> IV bolus or infusion day 1, 8, 15 or Mitoxantrone 10mg/m<sup>2</sup> IV bolus Day 1

L Asparaginase 10,000units/m<sup>2</sup> IM on day 4, 7, 10, 13 or PEG Asparaginase 100units/m<sup>2</sup> IM on day 4

Intrathecal Methotrexate Day 1,15

Cyclophosphamide 1000mg/m<sup>2</sup> IV over 30 min with or without MESNA on day 29  
Cytarabine 75mg/m<sup>2</sup> IV bolus or subcutaneous Day 30-33 and Day 37-40  
6 Mercaptopurine 60mg/m<sup>2</sup> PO Day 29 - 42

Maintenance phase  
Every 3 months x 8 such cycles  
Intrathecal Methotrexate on day 1 or Day 15  
6MP 60mg/m<sup>2</sup> PO Days 1 – 90  
Methotrexate 20mg/m<sup>2</sup> PO every week x 12 weeks

**High risk B ALL or T ALL or B LBL or T LBL (ICICLE-2014/BFM protocols/UK ALL protocols/MCP 841)**

High risk 4 drug induction  
Vincristine 1.5mg/m<sup>2</sup> IV day 8,15,22,29  
Daunorubicin 25-30mg/m<sup>2</sup> IV slow push Day 8, 15, 22, 29  
L Asparaginase 10,000units/m<sup>2</sup> IM every 3 days x 8 doses or PEG Asparaginase 1,000units/m<sup>2</sup> IM Day 9, 23  
Prednisolone 60mg/m<sup>2</sup> PO x 3 to 5 weeks  
Intrathecal Methotrexate x 3 doses on Day 8,15,35

Consolidation  
Cyclophosphamide 1000mg/m<sup>2</sup> IV over 30 min x 2 doses with or without MESNA Days 1, 29  
Cytarabine 75mg/m<sup>2</sup> IV push or Subcutaneously on Days 2-5, 9-12, 30-33, 37-40  
6MP 60mg/m<sup>2</sup> PO daily Day 1-14 and Day 29-42  
Intrathecal Methotrexate x 2 doses  
Vincristine 1.5mg/m<sup>2</sup> IV Day 16, 23, 44, 51  
L asparaginase 10,000units/m<sup>2</sup> IM Day 16 and Day 44

Interim maintenance  
IV Methotrexate 2 to 5 gram/m<sup>2</sup> Day 1,15,30,45  
Folinic acid rescue  
Intrathecal Methotrexate Day 1,15,30,45  
6MP 25mg/m<sup>2</sup> PO Day 1 to 49  
Some protocols might use Capizzi MTX with Asparaginase

Delayed intensification  
Dexamethasone 10mg/m<sup>2</sup> Day 1-5 and Day 15-19  
Vincristine 1.5mg/m<sup>2</sup> IV Day 1,8,15  
Doxorubicin 25mg/m<sup>2</sup> IV bolus or infusion Day 1,8,15 or Mitoxantrone 10mg/m<sup>2</sup> IV Day 1  
L Asparaginase 10,000units/m<sup>2</sup> IM on day 4,7,10,13 or PEG Asparaginase 100units/m<sup>2</sup> IM on day 4  
Intrathecal Methotrexate x 2 doses  
Cyclophosphamide 1000mg/m<sup>2</sup> IV infusion over 30 min with or without MESNA on day 29  
Cytarabine 75mg/m<sup>2</sup> IV bolus or subcutaneous Day 30-33 and day 37-40  
6 Mercaptopurine 60mg/m<sup>2</sup> PO Day 29 - 42

Maintenance phase  
Every 3 months x 8 such cycles

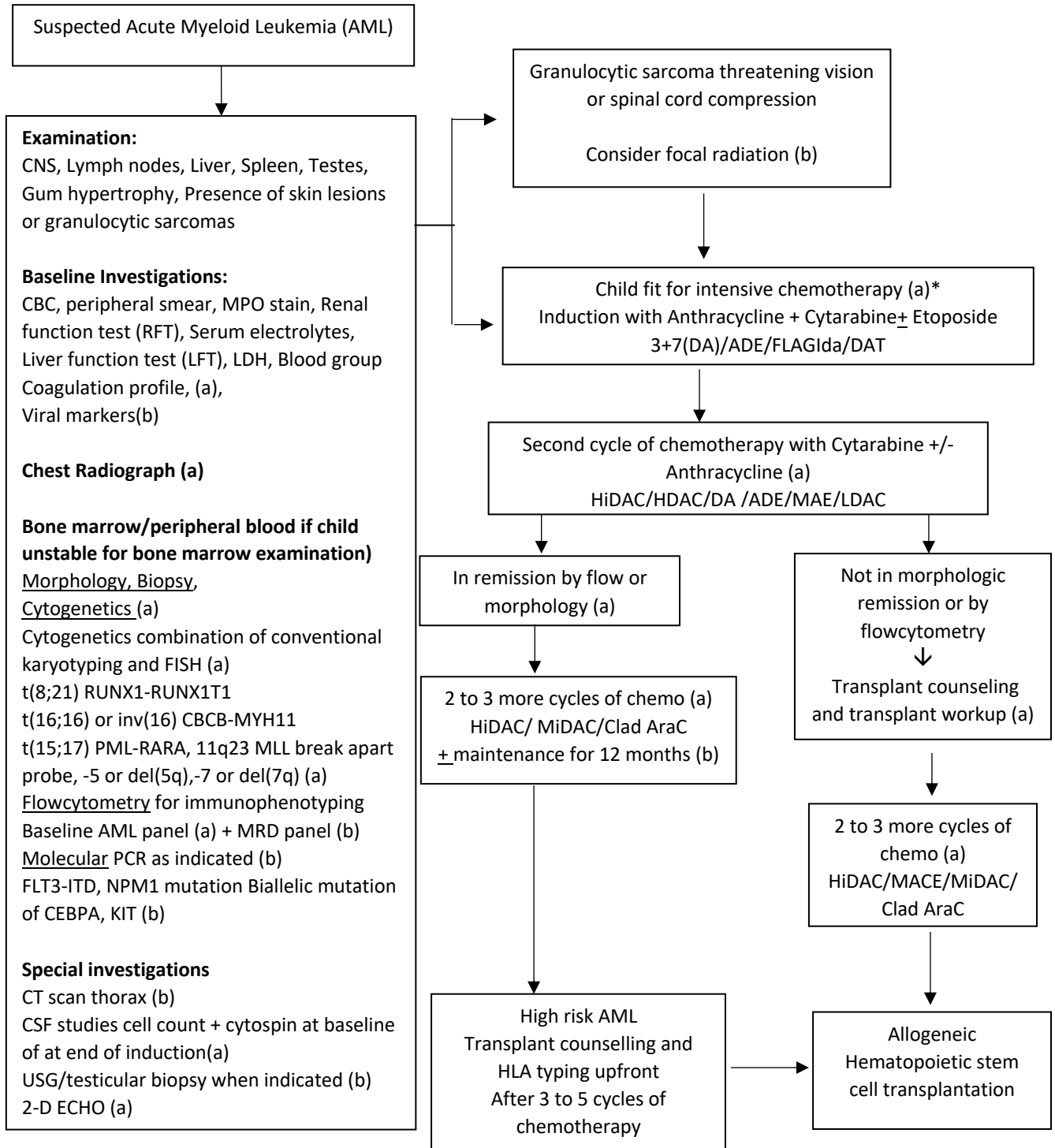
Intrathecal Methotrexate on day 1  
6MP 60mg/m<sup>2</sup> PO Days 1 – 90  
Methotrexate 20mg/m<sup>2</sup> PO every week x 12 weeks

**Radiation**

Therapeutic cranial irradiation 18 Gy for CNS 3 disease  
Testicular radiation 24 Gy if indicated

## PEDIATRIC ACUTE MYELOID LEUKEMIA

### Newly diagnosed AML



\* In case patient has active infection or malnourished and child is not fit for intensive chemotherapy, may consider low dose chemotherapy for 1 to 3 months

3+7(DA) Daunorubicin + Cytarabine/AraC  
DAE Daunorubicin AraC Etoposide  
FLAG Ida Fludarabine AraC GCSF Idarubicin

HiDAC High dose AraC  
LDAC Low dose AraC  
MAE Mitoxantrone AraC Etoposide  
Clad AraC Cladarabine AraC

1	Standard risk	Based on cytogenetics and M1 marrow or MRD (if available) negative at end of induction II	t(8;21) RUNX1-RUNX1T1 t(16;16) or inv(16) CBCB-MYH11
2	Intermediate risk		If not satisfying criteria for high risk or standard risk
3	High risk	Based on cytogenetics and positive MRD or >5% blasts on morphology in the bone marrow at end of Chemotherapy cycle II	Complex karyotype -5 or del(5q) -7 FLT3 ITD with allelic ratio > 0.4

### Treatment Algorithm: Newly diagnosed AML

#### Newly diagnosed AML (AML-1)

#### Protocol options

UK MRC series of protocols including Myechild-1 protocol  
St Jude AML 08  
BFM AML protocol  
Any other AML protocol with a combination of Anthracycline and Cytarabine

#### 3 + 7 induction

Daunorubicin 60mg/m<sup>2</sup> x 3 doses IV over 1 to 6 hrs Day 1-3  
Cytarabine 100mg – 200mg/m<sup>2</sup> IV over 24 hr continuous infusion or 100mg/m<sup>2</sup> bolus 12 hrly Day 1 - 7  
Some protocols add Etoposide 100mg/m<sup>2</sup> IV infusion Day 1 - 5

#### HiDAC

Cytarabine 2000 to 3000mg/m<sup>2</sup> IV infusion 12 hrly on day 1, 3, and 5

#### Clad AraC

Cladarabine 9mg/m<sup>2</sup> IV infusion Day 2 - 6  
Cytarabine 500mg/m<sup>2</sup> IV infusion Day 1 - 5

#### Triple Intrathecal chemotherapy x 4 to 5 doses once a month

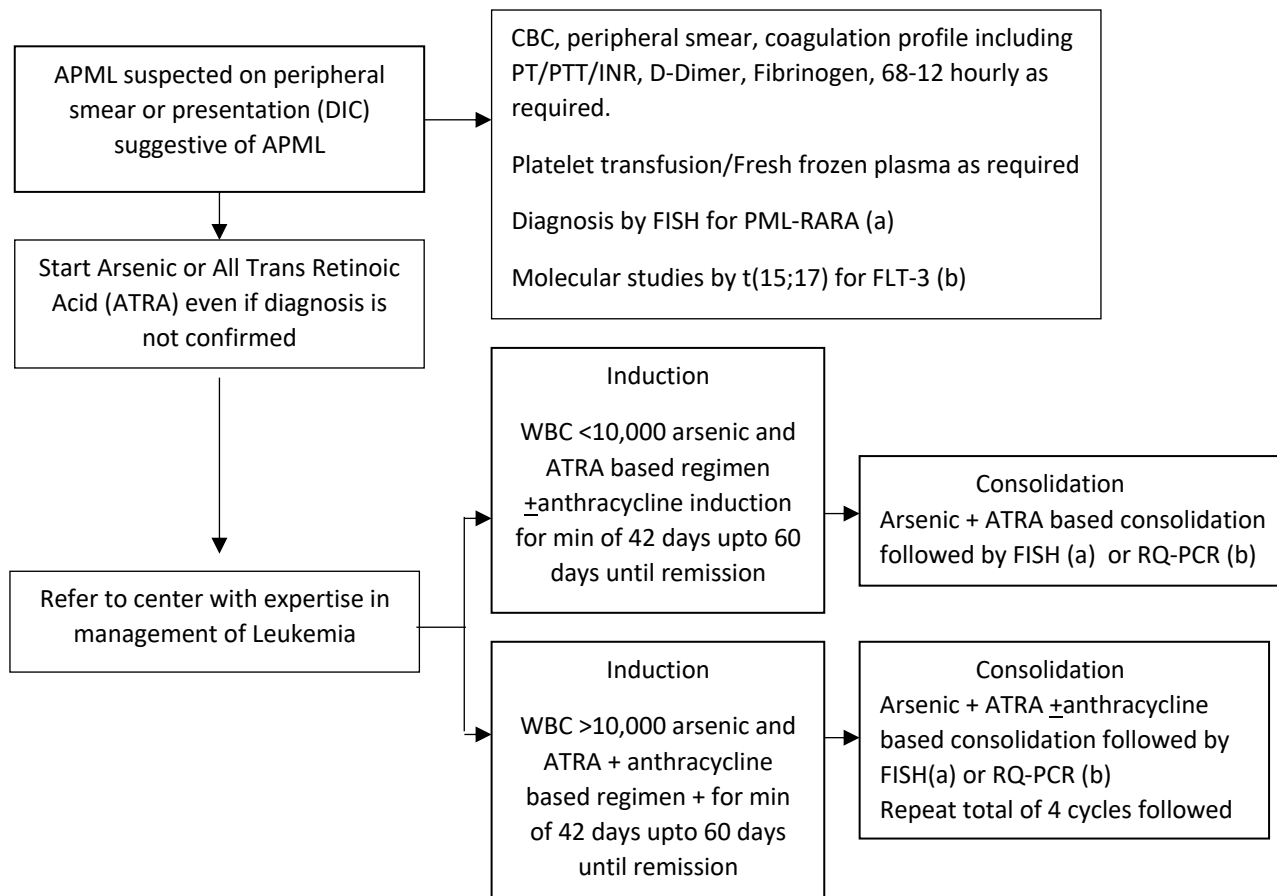
Methotrexate + Cytarabine + Hydrocortisone

#### Oral Maintenance chemotherapy 12 months in select protocols

6MP/6TG 50mg/40mg/m<sup>2</sup> PO Day 1 -21  
Etoposide 50mg/m<sup>2</sup> PO daily Day 1 - 21

## PEDIATRIC ACUTE PROMYELOCYTIC LEUKEMIA (APML)

### Treatment Algorithm: Newly diagnosed APML (start from AML pathway)



#### Supportive care

1. Platelet transfusion to keep platelets >30,000/cumm until coagulation parameters are stable
2. FFP/Cryoprecipitate for Hyperfibrinolysis to keep Fibrinogen >150 mg/dL
3. Hydroxyurea/Anthracyclines for WBC >10,000 or rising counts
4. Steroids Dexamethasone for Differentiation syndrome (DS) 10mg/m<sup>2</sup> in 2 divided doses
5. Hold Arsenic and ATRA for severe DS only

#### Induction

Arsenic 0.15mg/kg IV infusion day 1 - 60

ATRA 25 to 45mg/m<sup>2</sup> PO daily Days 1 – 60

Idarubicin/Daunorubicin/Mitoxantrone up to 4 doses for hyperleukocytosis

#### Other protocol options as below

Arsenic 0.15mg/kg IV infusion day 1 - 60

6MP 50mg/m<sup>2</sup> PO on Day 1 – 21 and Day 29 – 50

Etoposide 50mg/m<sup>2</sup> PO Day 1 – 21 and Day 29 -50

#### Consolidation x 4 cycles

Arsenic 0.15mg/kg IV daily day 1 -15 day 29 – 43  
ATRA 25 to 45mg/m<sup>2</sup> daily x day 1 to 60

**Other protocol options as below with 1 consolidation and 4 maintenance cycles**

ATRA 25 to 45mg/m<sup>2</sup> daily x day 1 to 60 (one cycle only)

Daunorubicin 45mg/m<sup>2</sup> IV bolus day 1-3, day 22-24, day 42-44 (one cycle only)

Followed by

**Maintenance 4 cycles (1-month cycle)**

ATRA Inj. ATO 0.15mg/kg (d1-10)

Tab. 6-TG 40mg/m<sup>2</sup> (d1-21)

Cap. Etoposide 50mg/m<sup>2</sup>(d1-21)

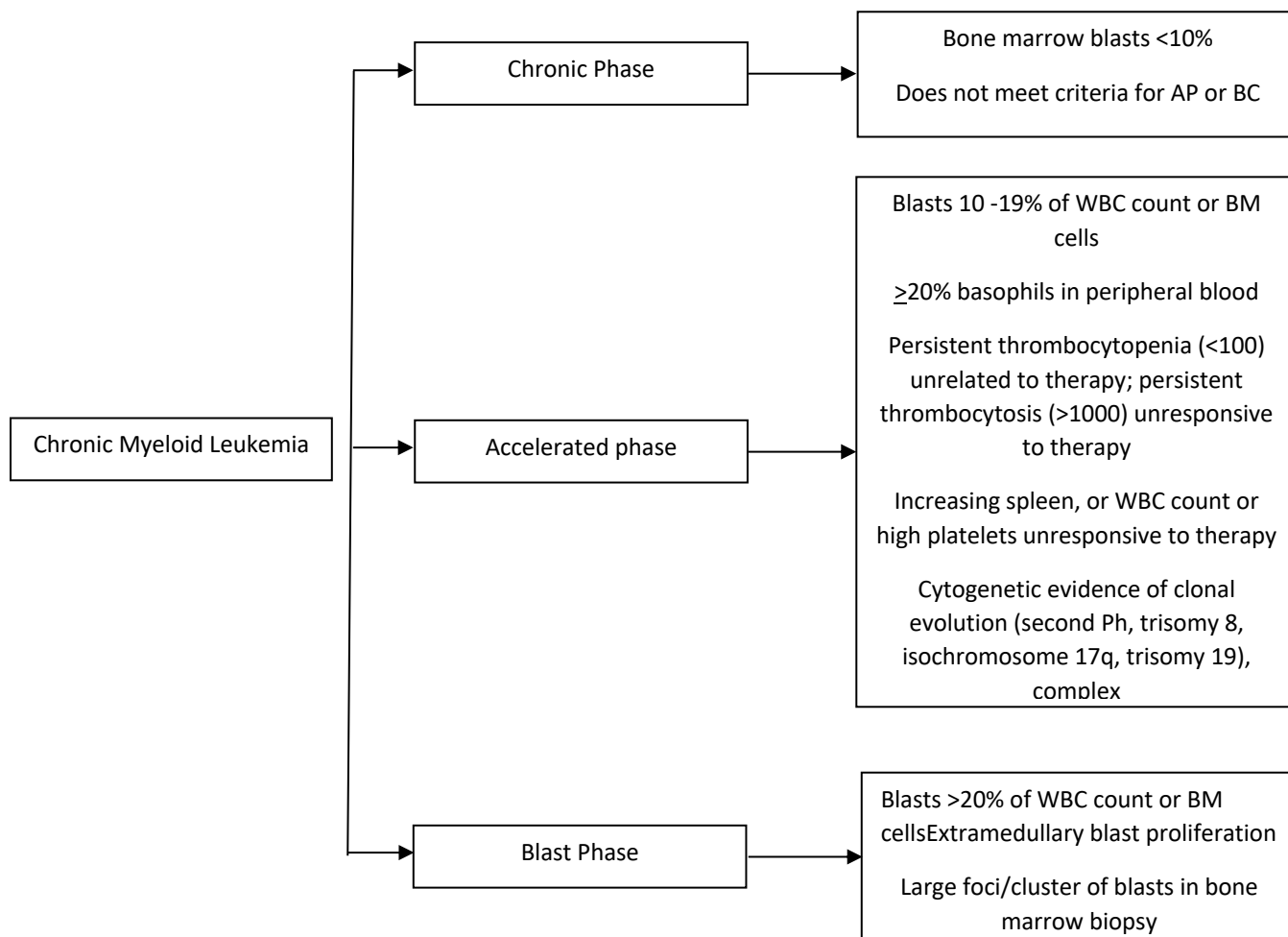


## PEDIATRIC CHRONIC MYELOID LEUKEMIA

### Treatment Algorithm: Pediatric CML



### Risk Stratification



Monitoring and response guidelines in CML CP			
Time (months) {Investigation}	Failure	Warning	Optimal Response
0 {FISH+ RTPCR} (a)	NA	High risk; Additional cytogenetic abnormalities	NA
3 {FISH} (a)	No HR; stable disease or disease progression	NA	CHR PCyR; Ph+ <35% BCR-ABL < 10 % IS
6 {FISH} (a)	Less than CHR; no CyR: Ph+ >95%	NA	CCyR; Ph+ 0% BCR-ABL < 1 % IS
12 {RQPCR} (a)	Less than PCyR: Ph+ >35%	Less than MMR	MMR BCR-ABL <0.1% IS
18 {RQPCR} (a)	Less than CCyR	NA	MMR or better
After 18 months {RQPCR} (a)	Loss of CHR; loss of CCyR; TKI resistant mutation	Loss of MMR; TKI resistant mutation	

**FISH**      Fluorescent in-situ hybridization  
**RQ-PCR**   Real-time Quantitative PCR  
**CHR**      Complete Hematological response

**PCyR** Partial Cytogenetic response  
**CCyR** Complete cytogenetic response  
**MMR** Major molecular response

**CML CP**

Imatinib 340mg/m<sup>2</sup> daily lifelong rounded up or down to nearest tablet strength  
 Or  
 Dasatinib 70mg/m<sup>2</sup> daily lifelong rounded up or down to nearest tablet strength

**CML AP**

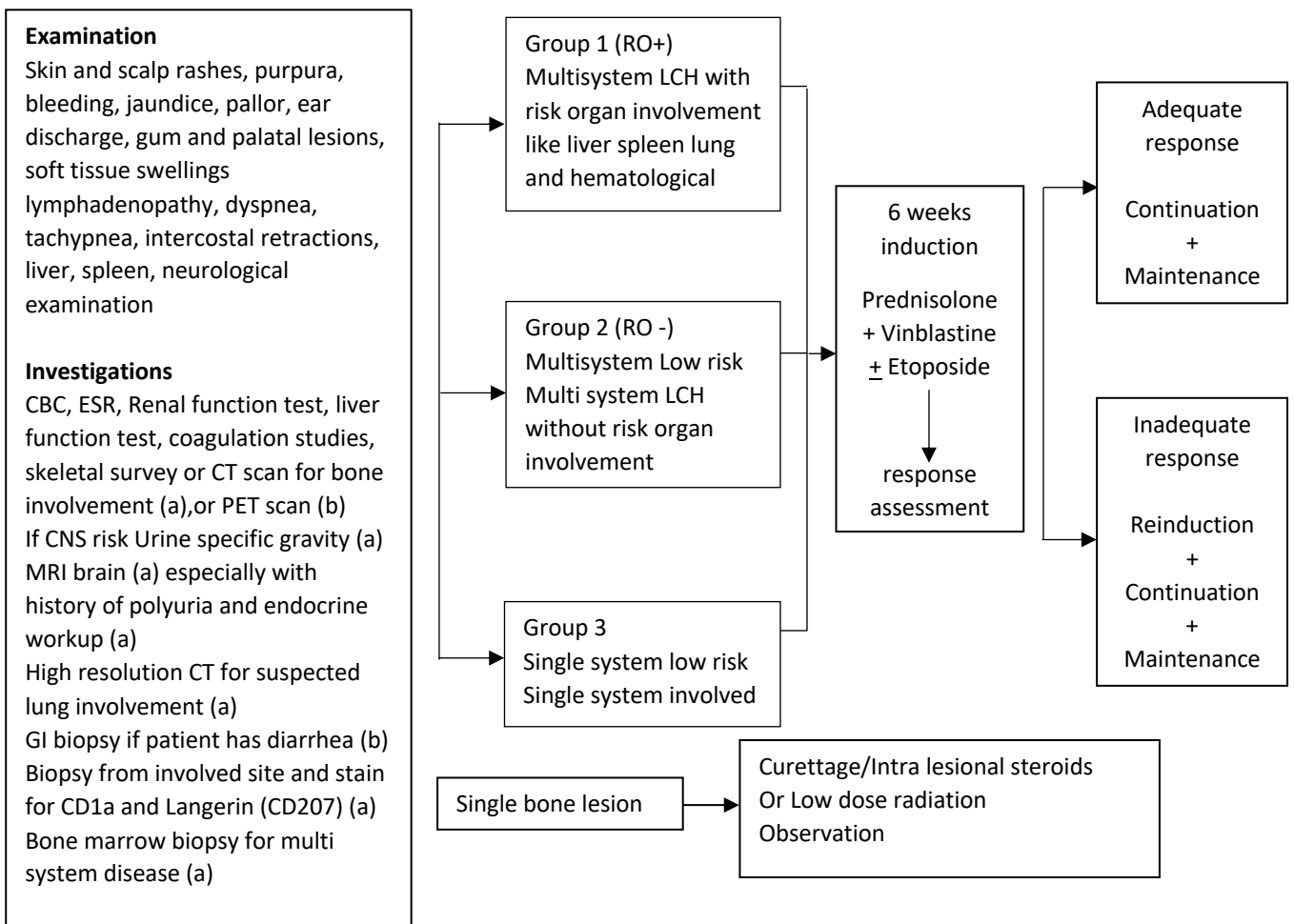
Imatinib 340mg/m<sup>2</sup> daily or Dasatinib 70mg/m<sup>2</sup> daily until CML CP followed by bone marrow transplant

**CML BC**

Imatinib 340mg/m<sup>2</sup> daily or Dasatinib 70mg/m<sup>2</sup> daily  
 ALL or AML like treatment  
 Bone marrow transplant

**LANGERHANS CELL HISTIOCYTOSIS**

**Treatment Algorithm: Newly diagnosed case of Langerhans cell Histiocytosis**



**AD better = Active disease better with reduction in size of lesion or SUV uptake on PET/Based on CT/Xray**

**AD worse = Active disease worse**

**RO+ Risk organ involved**

**RO- No involvement of Risk Organs**

### Chemotherapy

#### Induction

Vinblastine 6mg/m<sup>2</sup> IV bolus weekly x Week 1 - 6

Prednisolone 40mg/m<sup>2</sup> Day 1 - 42 with gradual tapering of dose

Some protocols add Etoposide 50mg/m<sup>2</sup> daily

#### Re-Induction as needed

Vinblastine 6mg/m<sup>2</sup> IV bolus weekly x Week 7 - 12

Prednisolone 40mg/m<sup>2</sup> Day 1 - 42 with gradual tapering of dose

Some protocols add Etoposide 50mg/m<sup>2</sup> daily

#### Continuation phase

Vinblastine 6mg/m<sup>2</sup> IV bolus every 3 weeks until end of 6 months

Prednisolone 40mg/m<sup>2</sup> Day 1 – 5 every 3 weeks until end of 6 months for low risk and 12 months for high risk disease

#### Maintenance 90 days 3 cycles for Group 2 and 6 cycles for Group 1

Vinblastine 6mg/m<sup>2</sup> every 3 weeks

Prednisolone 40mg/m<sup>2</sup> x 5 days every 3 weeks

6MP 50mg/m<sup>2</sup> daily x 3 months

Methotrexate 20mg/m<sup>2</sup>/week (added in some protocols)

Etoposide 50mg/m<sup>2</sup>/day for 21 days in a 4 week cycle (added in some protocols)

#### Chemotherapy options for salvage (recurrent or refractory LCH)

- Cladribine/Cytarabine
- Lenalidomide and Dexamethasone
- Vincristine/Cytarabine
- BRAF inhibitors (for BRAF V600E positive refractory or recurrent LCH)

	Induction (Vinblastine+ Prednisolone)	Reassessment imaging Or clinical assessment or compare imaging done at baseline	Post induction Treatment	Maintenance (3month/cycle)	Total duration of treatment
<b>Group 1</b>	High Risk (+Etoposide)	NAD/NED #	Continuation (+Etoposide)	6 cycles of maintenance	24 months  (in good responders)
		AD Better/Intermediate	Reinduction> Continuation	(Etoposide in first 2 cycles)	

		AD Worse	Salvage		
<b>Group 2</b>	Low Risk (-Etoposide)	NAD/NED	Continuation	3 cycles of maintenance (Etoposide)	15 months  (in good responders)
		AD Better/Intermediate	Reinduction> Continuation		
		AD Worse	Salvage		
<b>Group 3</b>	Low Risk (-Etoposide)	NAD/NED	Continuation	None  Stop treatment after Week 25 (End of continuation)	6 months  (in good responders)
		AD Better/Intermediate	Reinduction> Continuation		
		AD Worse	Salvage		

# AD Active disease  
 NAD No active disease  
 NED No evidence of disease

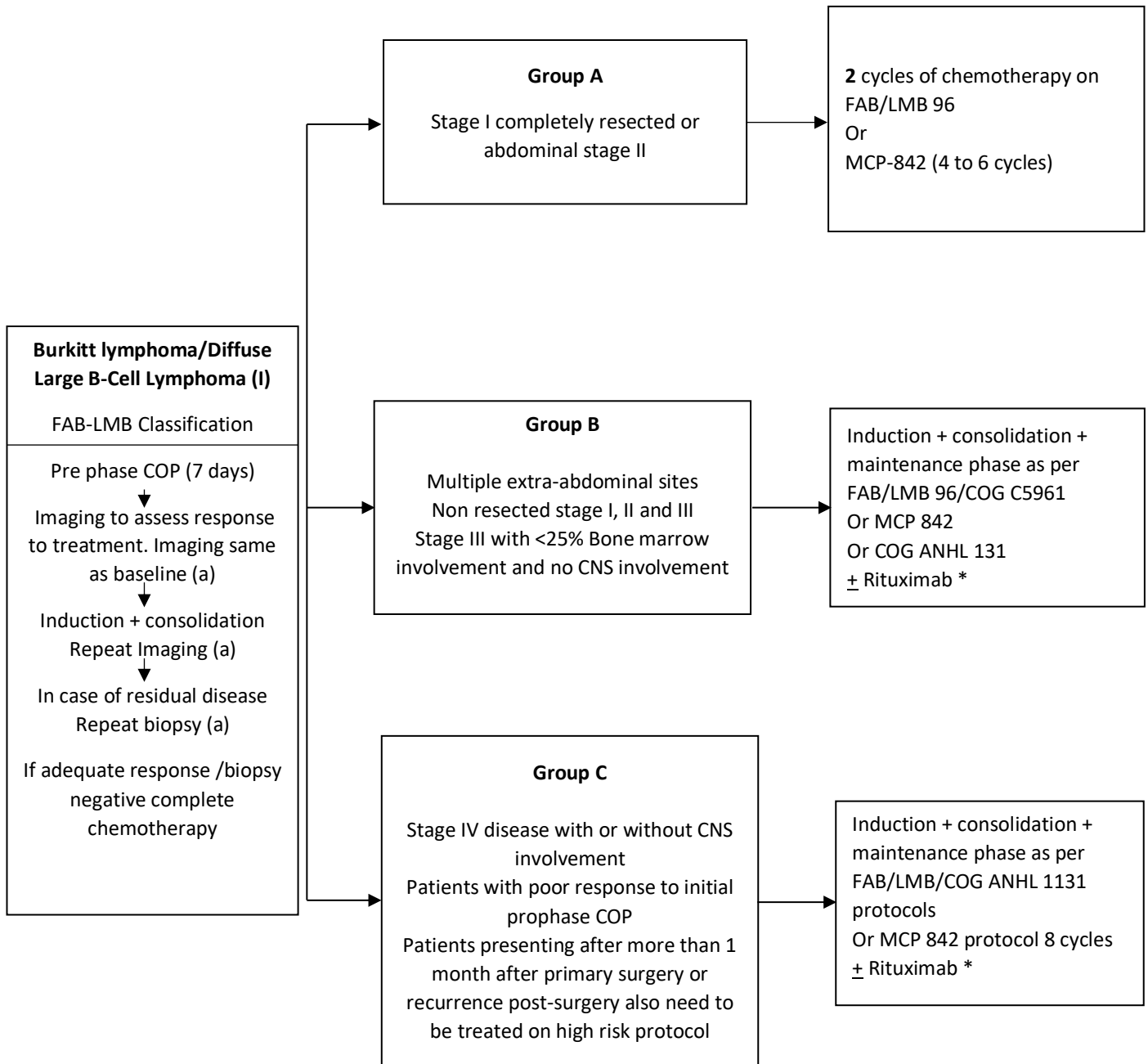
## NON-HODGKIN LYMPHOMA (NHL)

### Diagnostic workup for Suspected NHL

1	History and physical exam	Symptomatic large mediastinal masses, epidural or paraspinal tumors, should be recognized on examination and dealt with as an emergency. Steroids can be initiated before diagnostic confirmation but biopsy should not be delayed.	
2	Laboratory	Baseline labs CBC, Renal function test, Liver function test, serum electrolytes LDH Coagulation profile, (a) Viral Markers(b) Bone marrow aspirate and biopsy (a) CSF cytospin and morphology (a)	
3	Imaging	CXR (a) CT scan neck chest abdomen and pelvis (a) OR PET scan whole body (b) Ultrasound for certain intra-abdominal masses (b) 2 D ECHO (a)	For initial staging of tumor and response assessment (Treatment modality is based on site of lesion and best suited modality at that particular center on a case by case basis)
4	Histology	Biopsy (a) Excision biopsy or core biopsy Histopathology(a) Immunohistochemistry (a) as in appendix  Flowcytometry on pleural effusion or ascitic fluid where possible (b)	NHL is a heterogenous group of disorders. Common Pediatric NHLs include <ul style="list-style-type: none"> <li>• Aggressive mature B cell NHLs (Burkitt lymphoma and Diffuse large B cell lymphoma)</li> <li>• Lymphoblastic lymphoma T/B cell type</li> <li>• Primary mediastinal large B cell lymphoma (PMBCL)</li> <li>• Anaplastic large cell lymphoma (ALCL)</li> </ul> Miscellaneous NHL
5	Molecular studies (b)	Burkitt lymphoma t(8;14)(q24;q32) t(2;8) or t(8;22)(b) Anaplastic large cell lymphoma t(2;5)(p23;q35)(b)	
Staging (Due to high incidence of extra-nodal disease Murphy's staging is used for NHL) Burkitt's and DLBCL are stratified as mentioned subsequently.			
Stage I	Involvement of a single tumor or nodal area excluding the abdomen and mediastinum		
Stage II	Disease extent is limited to a single tumor with regional node involvement, two or more tumors or nodal areas involved on one side of the diaphragm, or a primary gastrointestinal tract tumor (completely resected) with or without regional node involvement.		
Stage III	Tumors or involved lymph node areas occur on both sides of the diaphragm. Stage III NHL also includes any primary intrathoracic (mediastinal, pleural, or thymic) disease, extensive primary intra-abdominal disease, or any paraspinal or epidural tumors.		
Stage IV	In stage IV childhood NHL, tumors involve the bone marrow and/or CNS, regardless of other sites of involvement.		
<ul style="list-style-type: none"> <li>• Bone marrow involvement is defined as 5% or more malignant cells the bone marrow, with normal peripheral blood counts and smears.</li> <li>• Patients with lymphoblastic lymphoma with more than 25% malignant cells in the bone marrow are considered to have Acute Lymphoblastic Leukaemia and to be treated on ALL protocols.</li> <li>• Patients with Burkitt's lymphoma with &gt;25% blasts are considered as Burkitt's leukaemia but treated on Burkitt's lymphoma protocol as stage IV disease with higher intensity of chemotherapy</li> </ul>			

- CNS disease is any malignant cell present in the CSF regardless of cell count.

**Treatment Algorithm: Newly diagnosed Non-Hodgkin lymphoma**



\* ± Rituximab can be given in following cases (b)

- Group B with Bone marrow involvement / Group C / inadequate initial response
- R2 disease Stage II stage IV as in the BFM classification below

## **FAB LMB regimen for Burkitt lymphoma and DLBCL**

### **Group A**

#### **COPAD x 2**

Vincristine 2mg/m<sup>2</sup> IV bolus on Day 1  
Prednisolone 60mg/m<sup>2</sup> PO day 1 – 5  
Cyclophosphamide 250mg/m<sup>2</sup> IV infusion every 12 hrs x 6 doses Day 2 – 4  
Doxorubicin 60mg/m<sup>2</sup> IV infusion Day 2  
Intrathecal Methotrexate + Hydrocortisone Day 2 and Day 6

#### **Pre phase COP Day 1 for Group B and Group C**

Vincristine 1mg/m<sup>2</sup> IV bolus Day 1  
Cyclophosphamide 300mg/m<sup>2</sup> IV infusion Day 1  
Prednisolone 60mg/m<sup>2</sup> PO Day 1 – 7  
Intrathecal Methotrexate + Hydrocortisone on Day 1

**\*Rituximab 375mg/m<sup>2</sup> day 0 before starting each cycle as per institutional guidelines/if clinically indicated  
(Total 4 to 6 doses throughout the treatment protocol)**

### **Group B**

#### **COPADM x 2**

Vincristine 2mg/m<sup>2</sup> IV bolus on Day 1  
Prednisolone 60mg/m<sup>2</sup> PO day 1 – 5  
Methotrexate 3g/m<sup>2</sup> IV infusion over 3 hrs on Day 1  
Folinic acid 15mg/m<sup>2</sup> IV/PO every 6 x 4 doses on Day 2 or until MTX level < 0.15Um/L  
Cyclophosphamide 250mg/m<sup>2</sup> IV infusion every 12 hrs x 6 doses Day 2 – 4  
Doxorubicin 60mg/m<sup>2</sup> IV infusion Day 2  
Intrathecal Methotrexate + Hydrocortisone Day 2 and Day 6

#### **CYM x 2**

Methotrexate 3g/m<sup>2</sup> IV infusion over 3 hrs on Day 1  
Folinic acid 15mg/m<sup>2</sup> IV/PO every 6 hrs x 4 doses on Day 2 or until MTX level < 0.15Um/L  
Cytarabine 100mg/m<sup>2</sup> over 24 hr continuous infusion from Day 2 - 6  
Intrathecal Methotrexate + Hydrocortisone Day 2  
Intrathecal Cytarabine + Hydrocortisone Day 7

### **Group C**

#### **COPADM1**

Vincristine 2mg/m<sup>2</sup> IV bolus on Day 1  
Prednisolone 60mg/m<sup>2</sup> PO day 1 – 5  
Methotrexate 8g/m<sup>2</sup> IV infusion over 4 hrs Day 1  
Folinic acid 15mg/m<sup>2</sup> IV/PO every 6 x 4 doses Day 2 or until MTX level < 0.15Um/L



Cyclophosphamide 250mg/m<sup>2</sup> infusion every 12 hrs x 6 doses Day 2 – 4  
Doxorubicin 60mg/m<sup>2</sup> IV infusion Day 2  
Intrathecal Methotrexate + Hydrocortisone + Cytarabine Day 2, 4 and 6

### **COPADM2**

Vincristine 2mg/m<sup>2</sup> IV bolus on Day 1  
Prednisolone 60mg/m<sup>2</sup> PO day 1 – 5  
Methotrexate 8g/m<sup>2</sup> IV infusion over 4 hrs on Day 1 (Infusion over 24hrs in case of CSF positive)  
Folinic acid 15mg/m<sup>2</sup> IV/PO every 6 x 4 doses on Day 2 or until MTX level < 0.15Um/L (12 doses for 24hr infusion of Methotrexate)  
Cyclophosphamide 500mg/m<sup>2</sup> IV infusion every 12 hrs x 6 doses Day 2 – 4  
Doxorubicin 60mg/m<sup>2</sup> IV infusion Day 2  
Intrathecal Methotrexate + Hydrocortisone + Cytarabine Day 2, 4 and 6

### **CYVE x 2**

Cytarabine 50mg/m<sup>2</sup> IV bolus over 12 hrs Day 1 - 5  
Cytarabine 3g/m<sup>2</sup> IV infusion over 3 hrs D2 to D5  
Etoposide 200mg/m<sup>2</sup> IV infusion Day 2 - 4  
For CSF pos  
Intrathecal Methotrexate + Hydrocortisone + Cytarabine Day 2, 4 and 6  
Methotrexate 8g/m<sup>2</sup> IV infusion over 4 hrs on Day 18  
Folinic acid 15mg/m<sup>2</sup> IV/PO every 6 x 4 doses on Day 2 or until MTX level < 0.15Um/L

### **Maintenance 1**

Vincristine 2mg/m<sup>2</sup> IV bolus on Day 1  
Prednisolone 60mg/m<sup>2</sup> PO day 1 – 5  
Methotrexate 8g/m<sup>2</sup> IV infusion over 4 hrs on Day 1 (For CSF pos 24 hr infusion)  
Folinic acid 15mg/m<sup>2</sup> IV/PO every 6 x 4 doses on Day 2 or until MTX level < 0.15Um/L ( 12 doses for 24 hr infusion of Methotrexate)  
Cyclophosphamide 500mg/m<sup>2</sup> IV infusion Day 2, 3  
Doxorubicin 60mg/m<sup>2</sup> IV infusion Day 2  
Intrathecal Methotrexate + Hydrocortisone + Cytarabine Day 2

### **Maintenance 2**

Cytarabine 50mg/m<sup>2</sup> IV bolus ever 12 hrs Day 1 – 5  
Etoposide 150mg/m<sup>2</sup> IV infusion Day 1 – 3

### **MCP 842 for all Non Hodgkin lymphoma (Vincristine replaced by Vinblastine for ALCL)**

#### **Total 6 to 8 cycles alternating A and B cycles**

#### **Regimen A**

Cytarabine 500mg/m<sup>2</sup> IV infusion 12hrly x 2 doses Day 1  
Cyclophosphamide 800mg/m<sup>2</sup> IV inifusion Day 1  
Cyclophosphamide 200mg/m<sup>2</sup> IV bolus Day 2 – 4  
Adriamycin 20mg/m<sup>2</sup> IV bolus Day 1, 2  
Vincristine 1.5mg/m<sup>2</sup> IV bolus Day 1, 8, 15

Intrathecal Cytarabine Day1, Day 4  
 Intrathecal Methotrexate Day 15, 18

**Regimen B**

Ifosfamide 1200mg IV infusion Day 1 – 5  
 Mesna 400mg/m<sup>2</sup> 0, 4, 8 hrs Day 1 – 5  
 Etoposide 60mg/m<sup>2</sup> IV infusion Day 1 – 3  
 Methotrexate 15mg/m<sup>2</sup> IV bolus Day 1 – 3  
 Vincristine 1.5mg/m<sup>2</sup> IV bolus Day 1, 8, 15  
 Intrathecal Cytarabine Day 1, Day 4  
 Intrathecal Methotrexate Day 15, 18

**DA-R EPOCH for Primary Mediastinal B cell lymphoma**

**Rituximab 375mg/m<sup>2</sup> can be administered before each cycle for Primary Mediastinal B cell lymphoma**

Etoposide 50mg/m <sup>2</sup> /day IV infusion over 24 hr Day 1 – 4	]	Dose escalation every cycle as tolerated based on count recovery
Doxorubicin 10mg/m <sup>2</sup> /day IV infusion over 24hr Day 1 – 4	]	
Vincristine 0.4mg/m <sup>2</sup> /day IV infusion over 24hr Day 1 – 4	]	
Cyclophosphamide 750mg/m <sup>2</sup> IV infusion Day 5	]	
Prednisolone 60mg/m <sup>2</sup> PO BD Day 1 – 5	]	

Intrathecal Methotrexate Day 1 and 5 Cycles 3 - 6

**ALCL 99 protocol for ALCL**

Pre-phase all patients  
 Dexamethasone 5mg/m<sup>2</sup> IV bolus or PO Day 1, 2  
 Dexamethasone 10mg/m<sup>2</sup> IV bolus or PO Day 3 – 5  
 Cyclophosphamide 200mg/m<sup>2</sup> IV infusion Day 1, 2  
 Intrathecal triple chemotherapy Methotrexate + Cytarabine + Hydrocortisone

**Course A (A1, A2 A3)**

Dexamethasone 10mg/m<sup>2</sup>/day PO BD Day 1 – 5  
 Methotrexate 1g/m<sup>2</sup> IV infusion over 24hrs Day 1  
 Folinic acid 15mg/m<sup>2</sup> 6hrly x 3 to 4 doses  
 Intrathecal triple chemotherapy Methotrexate + Cytarabine + Hydrocortisone Day 1  
 Ifosfamide 800mg/m<sup>2</sup> IV infusion Day 1 – 5  
 Cytarabine 150mg/m<sup>2</sup> infusion every 12 hrs Day 4, 5  
 Etoposide 100mg/m<sup>2</sup> IV infusion Day 4, 5

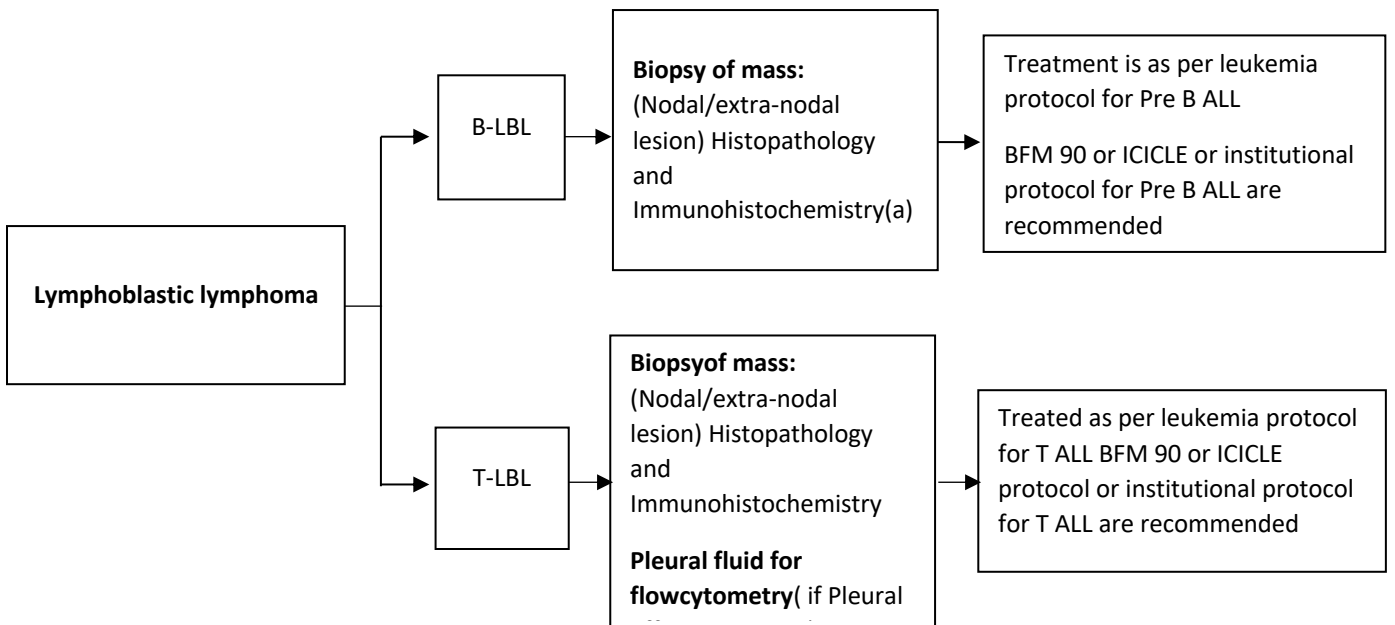
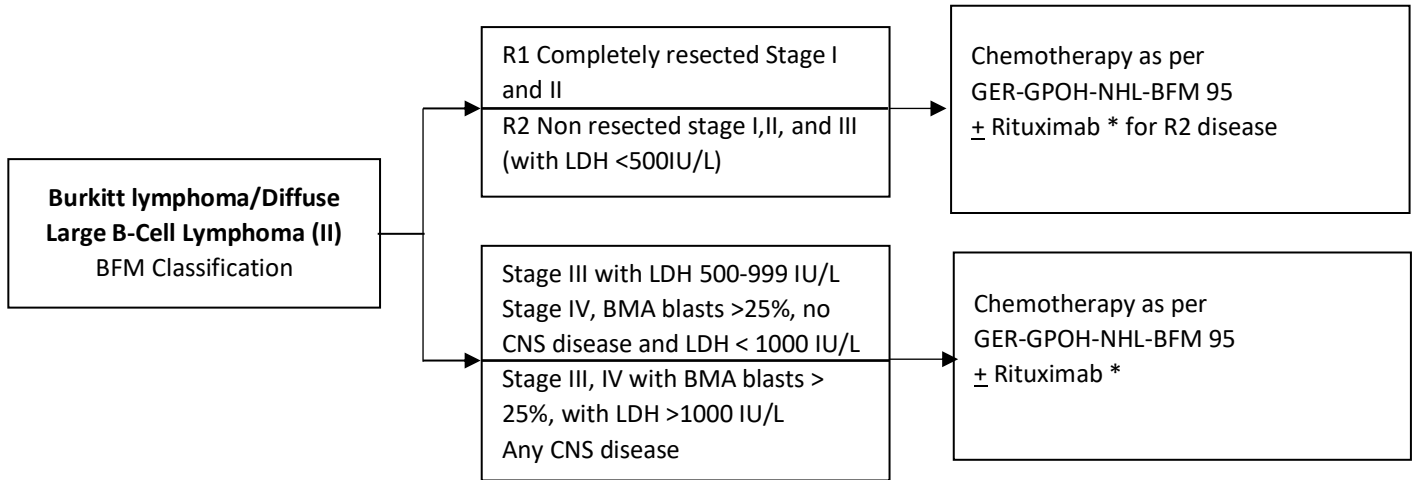
**Course B (B1, B2, B3)**

Dexamethasone 10mg/m<sup>2</sup>/day PO BD Day 1 – 5

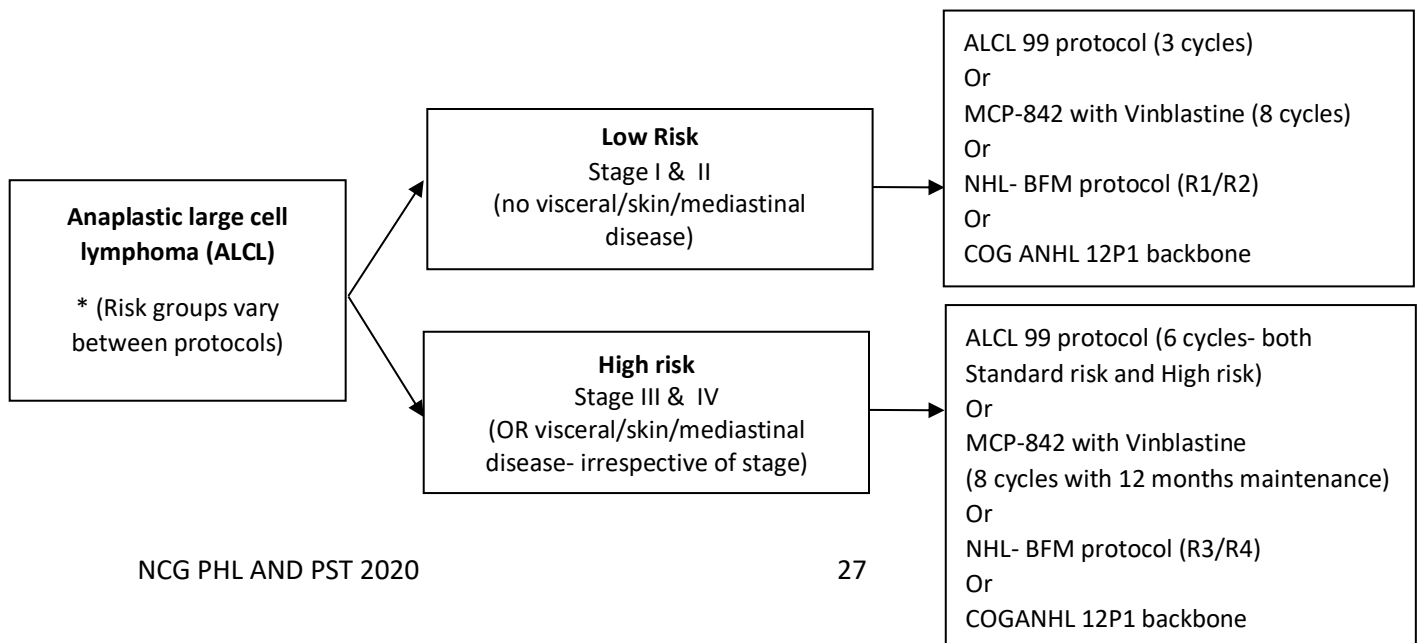
Methotrexate 1g/m<sup>2</sup> IV infusion over 24hrs Day 1  
Folinic acid 15mg/m<sup>2</sup> 6hrly x 3 to 4 doses  
Intrathecal triple chemotherapy Methotrexate + Cytarabine + Hydrocortisone Day 1  
Cyclophosphamide 200mg/m<sup>2</sup> IV infusion Day 1 – 5  
Doxorubicin 25mg/m<sup>2</sup> IV infusion Day 4, 5

**ALCL relapse is salvageable with Vinblastine based chemotherapy followed by autologous or allogeneic bone marrow transplant (b)**

Alternate Regimen option: BFM Classification and treatment



**Primary Mediastinal B cell lymphoma(PMBCL) specific type of B cell lymphoma treated on DA-REPOCH (Dose adjusted R EPOCH) 6 cycles (a) Reassessment scan is done only after the 6 cycle of chemotherapy**



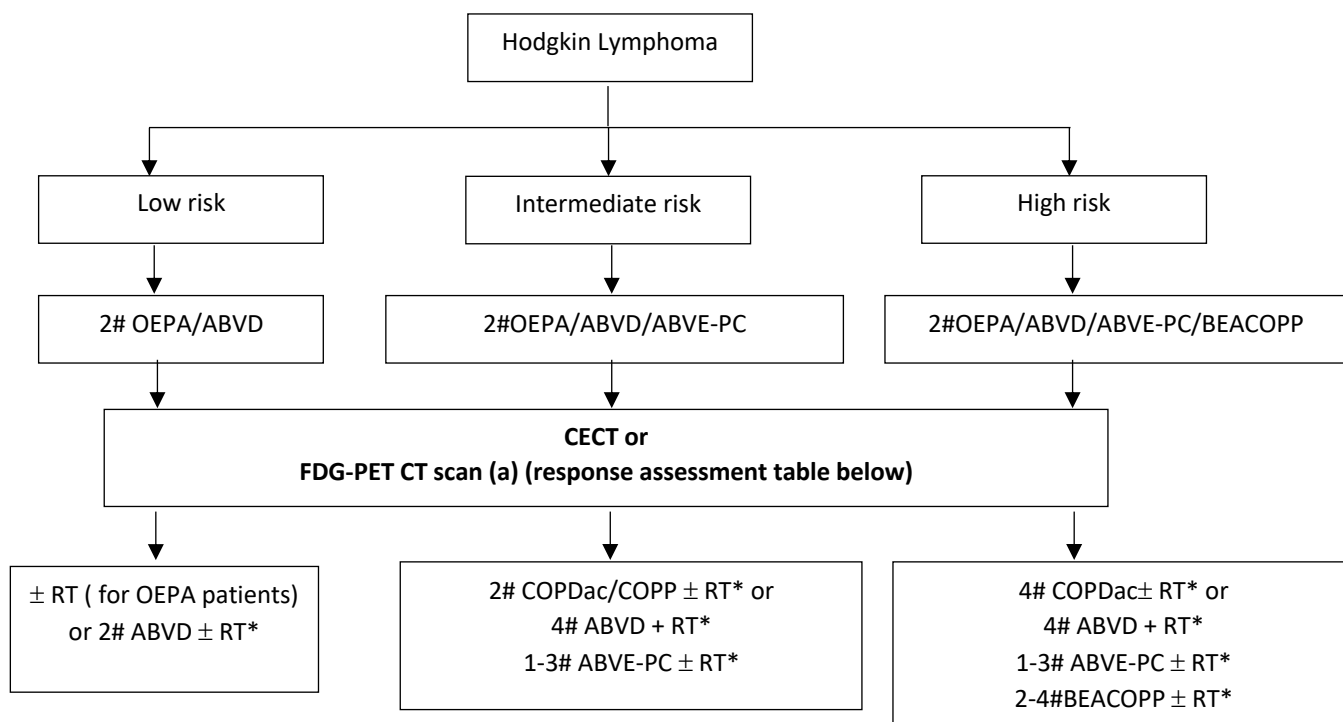


## PEDIATRIC HODGKIN LYMPHOMA

<b>Diagnostic workup</b>		
1	History and physical examination	B symptoms (any one is sufficient) Fever (oral temp >38°C) Night sweats Weight loss > 10% in last 6 months
2	Laboratory	Baseline labs CBC, Renal function test, Liver function test, serum electrolytes LDH Coagulation profile, Viral Markers (b) Bone marrow aspirate and biopsy for stage IV only (b) CSF cytopsin and morphology only if indicated (b)
3	Imaging	CXR CT scan chest abdomen and pelvis (if PET not available) OR PET scan whole body (a) 2D ECHO (a)
4	Histology	Excision biopsy of a peripheral lymph node or Core biopsy of mediastinal or abdominal lymph node if no accessible enlarged peripheral node Immunohistochemistry as in appendix (a)
<b>Staging information (Ann-Arbor staging system)</b>		
Stage I	Involvement of a single lymphatic site (ie. nodal region, Waldeyer's ring, thymus, spleen) (Stage I); or involvement of a single extralymphatic organ or site in the absence of any lymph node involvement (Stage IE)	
Stage II	Involvement of two or more lymph node regions on the same side of the diaphragm (Stage II) or localized involvement of a single extra lymphatic organ or site in association with regional lymph node involvement with out without involvement of other lymph node regions on the same side of the diaphragm (Stage IIE)	
Stage III	Involvement of lymph node regions on both sides of the diaphragm (Stage III) which may also be accompanied by extra lymphatic extension in association with adjacent lymph node involvement (Stage IIIE) or by involvement of the spleen or both (Stage III ES)	
Stage IV	Diffuse or disseminated involvement of one or more extra lymphatic organs, with or without associated lymph node involvement; or isolated extra lymphatic organ involvement in the absence of adjacent regional lymph node involvement, but in conjunction with disease in distant site(s). Stage IV includes any involvement of the liver or bone marrow, lungs (note) or CSF.	
The substage classification A,B,E,S,X amend each stage based on defined features B - B symptoms, A - absence of B symptoms. S - Spleen Involvement E - extra-nodal extension which most commonly involves lung, pleura, pericardium or bone when there is contiguous extension of lymph node disease or the site is proximal to an involved draining lymph node. X - Bulky disease defined differently by various study groups. Bulky peripheral lymphadenopathy is defined as single or conglomerated lymph		

nodal mass >6cm in longest diameter. Bulky mediastinal disease is defined as a mediastinal mass with a horizontal tumor diameter  $\geq 1/3^{\text{rd}}$  the thoracic diameter.

### Treatment algorithm: Pediatric Hodgkin Lymphoma



#### Chemotherapy regimens

ABVD: Doxorubicin, Bleomycin, Vinblastine, and Dacarbazine

DBVE: Doxorubicin, Bleomycin, Vincristine, Etoposide

ABVE-PC: Doxorubicin, Bleomycin, Vincristine, Etoposide, Prednisone, Cyclophosphamide

OEPA: Vincristine, Etoposide, Prednisone, Doxorubicin

COPDac: Cyclophosphamide, Vincristine, Prednisone, Dacrbazine

BEACOPP: Bleomycin, Etoposide, Doxorubicin, Cyclophosphamide, Vincristine, Procarbazine, and

\*RT indicated only for inadequate response (PET)/ bulky disease denoted as (X) in the staging at presentation

RT doses ranging from 19.8 cGy to 25 cGy

## Chemotherapy protocols

### **ABVD**

Adriamycin 25mg/m<sup>2</sup> IV bolus Day 1, 15  
Bleomycin 10 units/m<sup>2</sup> IV bolus Day 1, 15  
Vinblastine 6mg/m<sup>2</sup> IV bolus Day 1, 15  
Dacarbazine 375mg/m<sup>2</sup> IV infusion Day 1, 15

### **OEPA COPDAC**

Vincristine 1.5mg/m<sup>2</sup> IV bolus Day 1, 8, 15  
Etoposide 125mg/m<sup>2</sup> IV infusion day 1 – 4  
Prednisolone 60mg/m<sup>2</sup> PO Day 1 – 15  
Adriamycin 40mg/m<sup>2</sup> IV bolus Day 1, 15

Cyclophosphamide 600mg/m<sup>2</sup> IV infusion Day 1, 8  
Vincristine 1.5mg/m<sup>2</sup> IV bolus Day 1, 8  
Prednisolone 40mg/m<sup>2</sup> PO Day 1 - 15  
Dacarbazine 250mg/m<sup>2</sup> IV infusion Day 1 – 3

### **DBVE**

Adriamycin 25mg/m<sup>2</sup> IV bolus Day 1,15  
Bleomycin 10 units/m<sup>2</sup> IV bolus Day 1,15  
Vincristine 1.5mg/m<sup>2</sup> IV bolus Day 1,15  
Etoposide 100mg/m<sup>2</sup> IV infusion Day 1 – 15

### **ABVE-PC**

Doxorubicin 30mg/m<sup>2</sup> IV bolus Day 1, 2  
Bleomycin 10 units/m<sup>2</sup> IV bolus Day 1, 8  
Vincristine 1.5mg/m<sup>2</sup> IV bolus Day 1, 8  
Etoposide 75mg/m<sup>2</sup> IV infusion Day 1 - 5  
Prednisolone 40mg/m<sup>2</sup> PO Day 1 - 10  
Cyclophosphamide 800mg/m<sup>2</sup> IV infusion Day 1

### **BEACOPP**

Bleomycin 10 units/m<sup>2</sup> IV bolus Day 8  
Etoposide 200mg/m<sup>2</sup> IV infusion Day 1 – 3  
Doxorubicin 35mg/m<sup>2</sup> IV bolus/infusion Day 1  
Cyclophosphamide 1200mg/m<sup>2</sup> IV infusion Day 1, 8  
Vincristine 2mg/m<sup>2</sup> IV bolus Day 7  
Prednisolone 40mg/m<sup>2</sup> PO Day 1 – 14  
Procarbazine 100mg/m<sup>2</sup> PO Day 1 – 7



\*RT as indicated for bulky disease or residual disease between 20 to 35 Gy as per protocol

## APPENDIX A (FLOWCYTOMETRY)

### A. Acute Leukemia- Essential panel (a)

1. Smears stained with a Romanowsky stain and Myeloperoxidase or Sudan Black B
2. NSE, toluidine blue and Iron stain as required.

Note: Morphology is followed by flow cytometric immunophenotyping and other ancillary techniques including cytogenetics and molecular diagnostics. The final diagnosis is based on a combination of all these modalities.

<b>Essential</b>		
Common markers	CD45, CD38, HLADR	
Markers of immaturity	CD34	
	Lineage associated	Lineage Specific
B-cell	CD10, CD19, CD20, surface or cytoplasmic CD22, CyCD79a	
T-cell	CD1a, CD4, CD5, CD7, CD8, TCR $\gamma\delta$	Surface and Cytoplasmic CD3
Myeloid	CD13, CD33, CD117	cyMPO or Cytochemical Myeloperoxidase or Sudan Black B
Monocytic	CD36, CD64	Non Specific Esterase
Megakaryoblastic	X	
NK-cell	CD56	
Plasmacytoid dendritic cells	CD123	

### B. Acute leukemia – Optimal panel (b)

Essential + additional markers as below

<b>Optimal</b>	Lineage associated	Lineage Specific
B - cell	CD73, CD86, CD25, CD304	
T - cell		
Myeloid	CD15	
Monocytic	CD11c, CD14	
Megakaryoblastic	CD41, CD61	
NK – cell		
Plasmacytoid dendritic cells		



### C. Acute leukemia – Optional panel (c)

<b>Optional</b>		
Common markers	CD25, CD45Ra	
Markers of immaturity	CD133, TdT	
	Lineage associated	Lineage Specific
B - cell	CD58, CD81, NG2, CRLF2	IgM, Kappa & Lambda chains
T - cell	CD2, CD99, TCR $\alpha\beta$	
Myeloid	CD15, CD11b, CD16, CD65, CD66c	
Monocytic	CD86, CD300e	
Megakaryoblastic	CD42b	
NK-cell	CD94, CD161	
Plasmacytoid dendritic cells	CD303, CD304	
Mast cells	CD203c	
Erythroid lineage	CD49d, CD71, CD105	CD235a

### D. DNA ploidy by flow cytometry for B-ALL – Optimal (b)

Propidium Iodide  
FxCycle Violet  
DRAQ5  
DAPI (4',6-Diamidino-2-phenyl Indole)

### E. B-ALL Minimal Residual Disease Panel Essential (a) + Optional (c)

<b>Essential</b>	<b>Optimal</b>	<b>Optional</b>
CD10, CD19, CD20, CD34, CD38, CD45, CD73, CD123, CD86, CD304		CD25, CD44, CD66c, CD81, CD200, CD58
Nuclear dye such as Syto13, Syto16, Syto44		

#### Recommendations for processing

- Use Euroflow recommended Bulk-lysis method
- Acquire minimum 10,00,000 CD45-positive events
- Minimum 8-color antibody panel
- Use the template-based analysis
- Should be done in a laboratory with workload of minimum 30 acute leukemia samples per month
- Mention the limit of detection and limit of quantitation of MRD assay
- Mentioned the number of events studied
- Control sample should be evaluated atleast once in month



#### F. T-ALL MRD Optimal (b) + Optional (c)

Optimal	Optional
CD4, CD5, CD7, CD8, CD16, CD34, CD38, CD45, CD56, Surface and cytoplasmic CD3	CD1a, CD2, CD48, CD99, TdT
Nuclear dye such as Syto13, Syto16, Syto44	-

#### G. AML MRD Optimal (b) + Optional (c)

	Optimal	Optional
Deviation from normal	CD13, CD14, CD15, CD33, CD34, CD36, CD38, CD45, CD64, CD117, CD123, HLADR	CD11b, CD65, CD66c, CD71,
Leukemia associated Immunophenotypic markers	CD7, CD19, CD56	CD2, CD4, CD5,

#### H. Lymphoproliferative disorders/Lymphoma Essential (a) + Optimal (b) + Optional (c)

##### B-cell NHL

Essential	Optimal	Optional
CD5, CD10, CD19, CD20, CD23, CD45, CD200, Kappa & Lambda light chains	CD22, CD38, IgM,	CD27, CD43, CD44, CD49d, CD72, CD79b, CD81, CD123, CD148, CD180, CD305, IgD, IgG, FMC7, ROR1, Ki67, BCL2, BCL6, Mum-1

##### T- NHL

Essential	Optimal	Optional
CD3, CD4, CD5, CD7, CD8, CD10, CD16, CD25, CD26, CD45, CD56	CD2, CD30, CD94, CD161, CD185, CD279, ALK-1, Perforin, Granzyme	CD38, CD45RA, CD45RO, TCL1, TCRV $\beta$ -repertoire, KIR, Ki67, TIA-1

***Important:*** The laboratory without expertise in diagnosing hematolymphoid neoplasms and with inadequate IHC/Flow cytometric immunophenotyping panels should refer the sample to any specialized lab dealing with such neoplasms. There cannot be any definite algorithms for diagnosing hematolymphoid neoplasms as each lesion is different and number of reagents used may vary case to case basis.

## APPENDIX B

### Histopathology for lymphomas

#### Tissue /lymph node processing guidelines

##### Essential

- Tissue preservation (avoid frozen processing)
  - Fixative: 10% neutral buffered formalin
- Fixation:
  - Lymph nodes/tissue thicker than 0.8 -1.0cms; should be bisected and large tissue should be serially sliced, perpendicular to the long axis.
  - Tissue  $\leq 4$  cm in greatest dimension should be processed in entirety
  - Should be put for fixation within 30 – 60 minutes of biopsy
  - Fixation volume should be at 3-4 times the volume of the tissue
  - Should not be left in the fixative for more than 48 hrs; and should be processed in 12-24 hrs time (in cases of inevitable delay; should be kept in cold temperature [refrigerator], preferably at 4 degrees centigrade)
- Routine processing and embedding
  - 2-3 micron thick sections with Hematoxylin and eosin stained slides of each paraffin block
  - Immunohistochemistry set up
  - Microscopic evaluation

##### Optional (Immunohistochemistry and Molecular diagnostic laboratories)

*\*For transportation – Either by immersing tissue in the adequate formalin in a sealed container or by paraffin blocks*

#### Lymphoma Tissue Diagnosis

##### Essential: (a)

- Diagnosis:
  - Histological evaluation, i.e. biopsy as a method of investigation with comprehensive IHC panels.
  - Only in instances of inability of get adequate, a fine needle aspiration (FNA) based flow cytometric evaluation should be considered for diagnosis
- Staging
  - Bone marrow biopsy, aspirate and imprint smear

##### Optional/extended work-up: (b)

- Diagnosis:
  - Fine needle aspiration (FNA) based flow cytometricimmunophenotyping along with the biopsy
  - Molecular work-up (c)
- Staging:
  - Flow cytometricimmunophenotypic evaluation (for Non-Hodgkin Lymphoma) (c)

#### A. Hodgkin lymphoma

**(cHL) classical Hodgkin Lymphoma and (NLPHL) Nodular Lymphocyte Predominant Hodgkin Lymphoma-requisites for diagnosis**

### **Classic Hodgkin lymphoma (cHL)**

- Essential: (a)
  - CD3, CD20, CD30, CD15, Pax5, ALK-1\*\*
- Optimal/extended work-up: (b)
  - LCA, CD3, CD20, CD30, CD15, Pax-5, Oct2, Bob1, EBV-LMP1/EBER, Gata 3

### **Nodular lymphocyte predominant Hodgkin lymphoma (NLPHL)**

- Essential: (a)
  - CD20, CD3, CD30, CD15, Pax5
- Optimal/extended work-up: (b)
  - CD3, CD20, CD30, CD15, Pax-5, EBV-LMP1/EBER, PD1, Oct2, Bob1, Gata3, CD4, CD8

\*\* Rule out a possibility of ALK+ve ALCL

## **B. Non-Hodgkin Lymphoma**

### **1. CD20 positive BNHL: large cell morphology**

- Essential: (a)
  - IHC: LCA, CD20, CD3, MIB-1
- Optimal/extended work-up:
  - IHC: CD3, CD20, MIB-1, cyclin D1, CD5, CD10, Bcl6, Mum1, cmyc, Bcl-2, CD30, EBV-LMP1/EBER
  - FISH: CMYC/BCL2/BCL6 gene rearrangement
  - Gene expression/methylation studies – COO subtyping

### **2. CD20 positive BNHL: non-large cell morphology**

- Essential: (a)
  - IHC: LCA, CD20, CD3, MIB-1, CD5, CD23, CD10, bcl6, cyclin D1 (if blastic morphology, please add AMPO, ckit, CD10, CD19/Pax5, Tdt, CD34)
- Optimal/extended work-up: (b)
  - IHC: Mum1, cmyc, Bcl-2, EBV-LMP1/EBER, CD43, CD138, Sox11
  - FISH: CMYC/BCL2/BCL6; IFR4 gene rearrangement
  - Sequencing: MYD88 mutation

### **3. CD3 positive NHL: large cell morphology**

- Essential: (a)
  - IHC: CD20, CD3, CD30, MIB-1, ALK-1, CD4
- Optimal/extended work-up :
  - IHC: CD3, CD20, CD4, CD8, CD2, CD5, CD7, MIB-1, CD56, CD30, ALK-1, CD10, Bcl6, PD1, Mum1, EBV-LMP1/EBER, CD123, Gata3
  - FISH: DUSP22 gene rearrangement

### **4. CD3 positive NHL: non-large cell morphology**

- Essential: (b)
  - IHC: CD20, CD3, CD2, CD5, CD7, CD4, CD8, MIB-1, cyclin D1, Tdt, CD34, CD30, ALK-1
- Optimal/extended work-up :
  - IHC: CD56, CD10, Bcl6, PD1, Mum1, EBV-ISH, CD123, Gata3, CXCL113, CXCR5, ICOS
  - FISH: DUSP22 gene rearrangement



**5. CD3 and CD20 negative NHL- requisites for diagnosis**

- IHC: LCA, CD3, CD20, CD30, CD19, Pax-5, CD138, ALK-1, CD5, CD10, Bcl6, Mum1, EBV-LMP1/EBER, CD56, CD7, CD4, CD8, CD123, MIB-1, c-kit, MPO, CD41, CD61, CD33, CD34, Tdt, CD1a, CD163, S-100 protein, EMA, CD23, CD21, kappa, lambda, MIB-1

***Important:*** *The laboratory without expertise in diagnosing hematolymphoid neoplasms and with inadequate IHC/Flow cytometricimmunophenotyping panels should refer the sample to any specialized lab dealing with such neoplasms. There cannot be any definite algorithms for diagnosing hematolymphoid neoplasms as each lesion is different and number of reagents used may vary case to case basis.*

## APPENDIX C

### Response assessment for Lymphomas

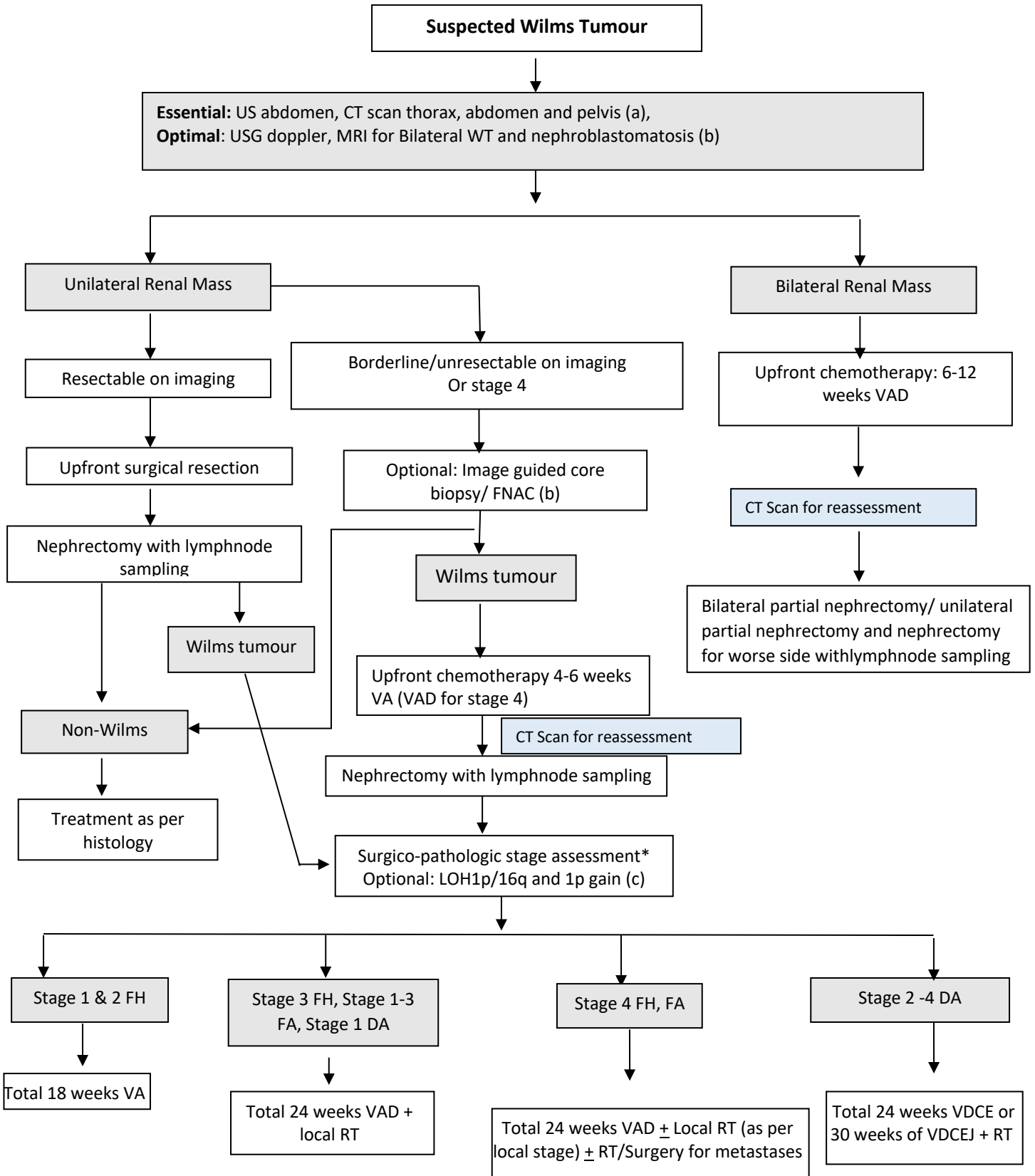
#### Deauville score

1	No FDG uptake
2	FDG uptake $\leq$ mediastinum
3	FDG uptake $>$ mediastinum but $\leq$ liver
4	FDG uptake $>$ liver at any site
5	FDG uptake $>$ liver and new sites of disease
x	New areas of FDG uptake unlikely to be related to lymphoma

## PEDIATRIC SOLID TUMORS

## WILMS TUMOUR

### Treatment Algorithm: Wilms Tumor



- FH=Favourable Histology
  - V= Vincristine, A= Actinomycin D, D= Doxorubicin, C= Cyclophosphamide, E= Etoposide, J= Carboplatin
  - RT= Radiotherapy
  - DA- diffuse anaplasia, FA-focal anaplasia
- 
- Tumours weighing <550g in children aged <24 months and with favorable histology may be observed after surgery
  - In centres adopting the SIOP approach to management of Wilms tumor, post-operative chemotherapy is given as per operative staging and post-operative histology as per table below:

**Chemotherapy :**

Vincristine Inj. Vincristine 0.046mg/kg or 1.5mg/m<sup>2</sup> IV bolus weekly for 24 weeks in all stages

Inj. Actinomycin 45µg/Kg as iv bolus 3-weekly for 24 weeks in all stages

Inj. Adriamycin 2mg/Kg or 60 mg/m<sup>2</sup> as iv bolus 3-weekly for 24 weeks in stages 3 and 4

**Regimen I (VDCE) 24 weeks :**

Vincristine 0.046mg/kg or 1.5mg/m<sup>2</sup> IV bolus weekly

Inj Adriamycin 1.5 mg/kg IV push for <10 kgs or 45 mg/m<sup>2</sup>/day for >10 kgs as infusion over 6 hr, nj. Plus Inj Cyclophosphamide 14.7 mg/kg/day for <10 kgs or 440mg/m<sup>2</sup>/day for >10 kgs as infusion over 2 hrs (D1 to D3)

Inj Etoposide 3.3 mg/kg/day for <10 kgs Or 100mg/m<sup>2</sup>/day for >10 kgs as infusion over 1-2 hrs (D1 to D3)

Plus Inj. Cyclophosphamide 14.7 mg/kg/day for <10 kgs or 440mg/m<sup>2</sup>/day for >10 kgs as infusion over 2 hrs (D1 to D5)

**Regimen UH1 ( VDCEJ) 30 weeks:**

Inj Carboplatin on Day 1 560 mg/m<sup>2</sup>

Inj Cyclophosphamide 1,200 mg/m<sup>2</sup>per day intravenously as infusion over 1-2 hours on day 1

Inj Doxorubicin 37.5 mg/m<sup>2</sup> perday intravenously as infusion over 6 hours (days 1-2)

Inj Etoposide 100 mg/m<sup>2</sup>per day intravenously as infusion over 2 hours (days 1-5)

Inj Irinotecan 20 mg/m<sup>2</sup>per as infusion over 2 hours (days 1-5 and 8-12)

Inj Vincristine 1.5 mg/m<sup>2</sup> IV bolus weekly

### Risk Stratification

Disease	Treatment		
	Stage I	Stage II	Stage III
<b>Low-risk</b>	None	AV (27 weeks)	AV (27 weeks)
<b>Intermediate-risk, all subtypes &lt;500ml</b>	AV (4 weeks)	AV (27 weeks)	AV (27 weeks) + flank RT
<b>Intermediate-risk, stromal or epithelial-type &gt;500 ml</b>	AV (4 weeks)	AV (27 weeks)	AV (27 weeks) + flank RT
<b>Intermediate-risk, nonstromal, nonepithelial</b>	AV (4 weeks)	AVD (27 weeks)	AVD (27 weeks) + flank RT
<b>High-risk blastemal type and diffuse anaplasia Wilms tumour</b>	AVD (27 weeks)	DCEJ (34 weeks) flank RT in DA	DCEJ (34 weeks) + flank RT

### Staging of Wilms Tumour

Stage 1	Tumour limited to the kidney and completely excised.
Stage 2	Tumour extends beyond the kidney but is completely excised The tumour infiltrates the renal sinus and/or adjacent organs or vena cava but is completely resected
Stage 3	Residual nonhematogenous tumor confined to the abdomen; lymph-node involvement, peritoneal spillage, peritoneal implants, either gross or microscopic tumor beyond the surgical margin, or tumor not completely removed.
Stage 4	Hematogenous metastases to lung, liver, bone, brain or other organ.
Stage 5	Bilateral renal involvement at diagnosis.

### RT guidelines in Wilms Tumour

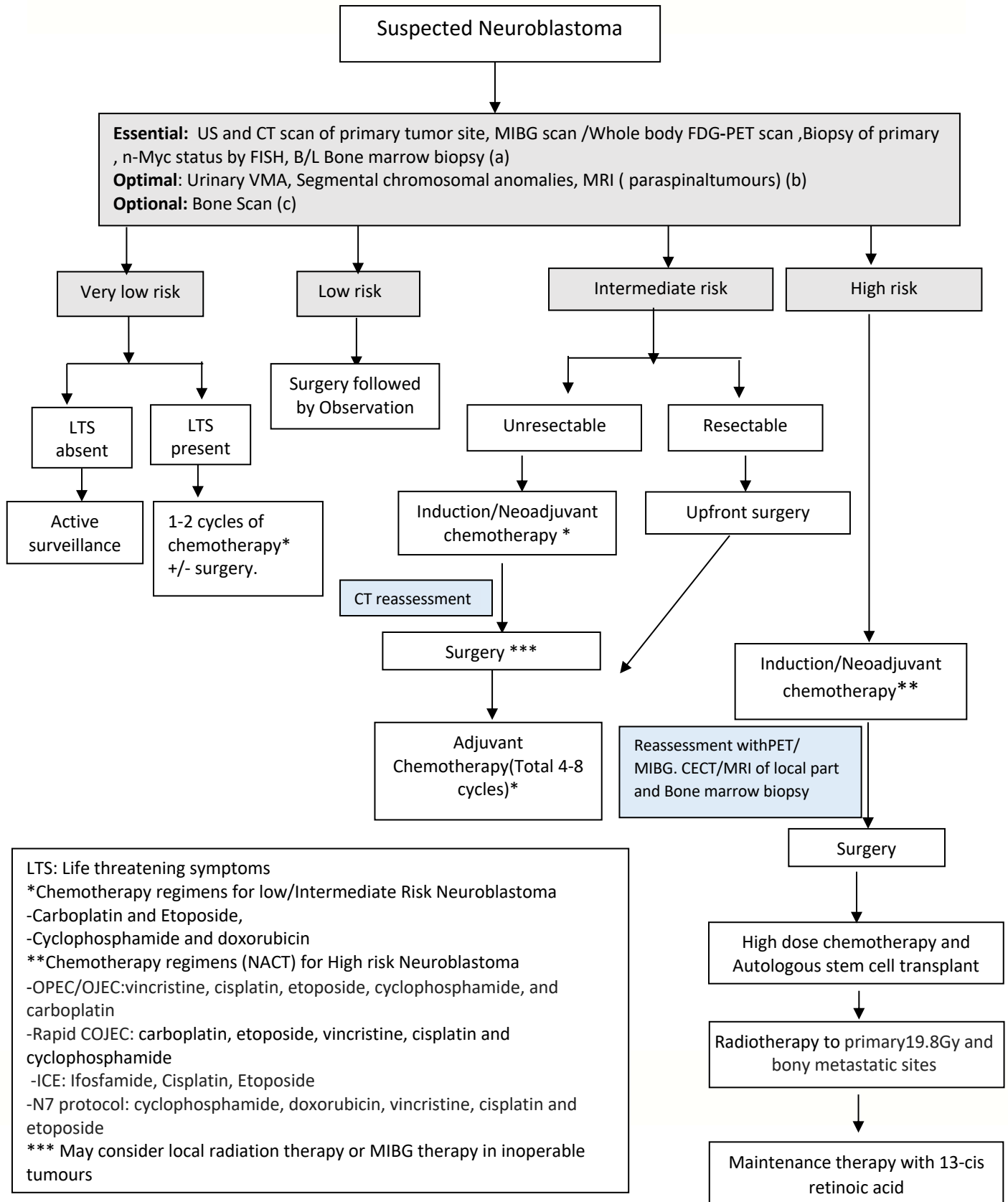
	Abdominal Tumour Stage/ Histology	RT Dose (RT Field)
1.	Stage I & II/ Favorable	No RT
2.	Stage III/ Favorable and Focal Anaplasia	10.8Gy/ 6# @ 1.8Gy/ Fraction (level Ib)
3.	Stage I – II/ Diffuse Anaplasia	10.8Gy/ 6# @ 1.8Gy/ Fraction
4.	Stage III/ Diffuse Anaplasia	19.8Gy/ 11# @ 1.8Gy/ Fraction
5.	Recurrent Abdominal Disease	10.8Gy/ 6# @ 1.8Gy/ Fraction
6.	Lung Mets (Favorable & Unfavorable) Microscopic Disease Gross Disease/ Nodules	12.6Gy/ 7# @ 1.8Gy/ Fraction + 9.0Gy/ 5# @ 1.8Gy/ Fraction (Boost)
7.	Liver Mets (Favorable & Unfavorable Histology)	10.8Gy/ 6# @ 1.8Gy/ Fraction (Whole Liver) + 9.0Gy/ 5# @ 1.8Gy/ Fraction (Boost to Gross residual disease)

8.	Skeletal Mets (Favorable& Unfavorable Histology)	25.2Gy/ 14# @ 1.8Gy/ Fraction (Lesion + 3cm)
9.	Unresected Lymph Nodal Mets (Favorable& Unfavorable Histology)	19.8Gy/ 11# @ 1.8Gy/ Fraction (Nodal Region)

- Whole-lung irradiation may be omitted in cases of FH with complete response (pulmonary) after 6 weeks of VAD (provided there is no extrapulmonary metastases or LOH 1p/16q)
- 3D Conformal RT and Intensity Modulated RT are standard forms of delivery of RT in children with WT

## NEUROBLASTOMA

### Treatment Algorithm: Neuroblastoma



LTS: Life threatening symptoms  
 \*Chemotherapy regimens for low/Intermediate Risk Neuroblastoma  
 -Carboplatin and Etoposide,  
 -Cyclophosphamide and doxorubicin  
 \*\*Chemotherapy regimens (NACT) for High risk Neuroblastoma  
 -OPEC/OJEC: vincristine, cisplatin, etoposide, cyclophosphamide, and carboplatin  
 -Rapid COJEC: carboplatin, etoposide, vincristine, cisplatin and cyclophosphamide  
 -ICE: Ifosfamide, Cisplatin, Etoposide  
 -N7 protocol: cyclophosphamide, doxorubicin, vincristine, cisplatin and etoposide  
 \*\*\* May consider local radiation therapy or MIBG therapy in inoperable tumours



## Risk Stratification

International Neuroblastoma Risk Grouping Staging System (INRGSS)	
Stage	Description
L1	Localized tumor not involving vital structures as defined by the list of Image defined Risk Factors (IDRFs) and confined to one body compartment.
L2	Loco regional tumor with presence of one or more image defined risk factors
M	Distant metastatic disease (except stage MS)
MS	Metastatic disease in children younger than 547 days and metastases confined to skin, liver and/or bone marrow (< 10% of total nucleated cells on smears or biopsy)

### International Neuroblastoma Risk Grouping

<b>Very low Risk</b>	Infant (<18 months) asymptomatic, no high-risk molecular features, Ganglioneuroma any age
<b>Low Risk</b>	Infant (<18 months) NB L2 or stage II/III (non-metastatic)
<b>Intermediate Risk</b>	Stage M infant (<18 months) NB, Stage II/III, L2 non-infantile NB, infant (<18 months) NB L2 or stage II/III with 11q aberration
<b>High Risk</b>	Any NMYC amplified tumour, Metastatic Stage M(non-infant), infant NB MS with 11q aberration

### Chemotherapy doses :

#### RAPID COJEC

Cycle A : Inj Carboplatin 750 mg/m<sup>2</sup> short infusion, Inj Etoposide 175 mg/m<sup>2</sup> as 4-hour infusion, Inj

Vincristine 1.5 mg/m<sup>2</sup> intravenous bolus;

Cycle B : Inj Cisplatin 80 mg/m<sup>2</sup> as a 24-hour continuous infusion, Inj Vincristine 1.5 mg/m<sup>2</sup> intravenous bolus

Cycle C: Inj Cyclophosphamide 1,050 mg/m<sup>2</sup> short infusion. Inj Etoposide 175 mg/m<sup>2</sup> as 4-hour infusion, Inj Vincristine 1.5 mg/m<sup>2</sup> intravenous bolus;

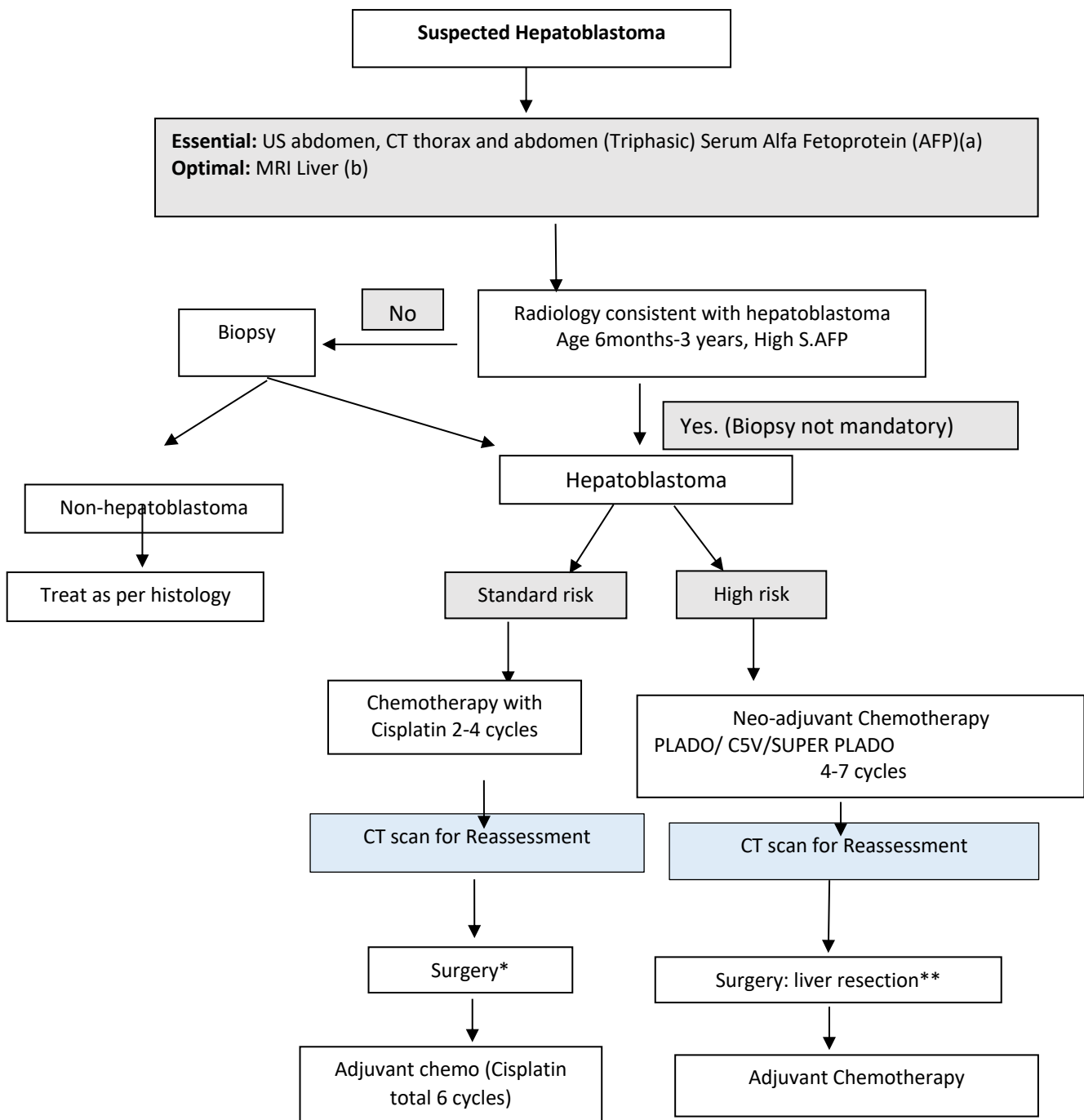
#### Intermediate Risk Neuroblastoma : 4-8 cycles of below drugs

Inj Carboplatin 200 mg/m<sup>2</sup> short infusion (days 1-3) plus Inj Etoposide 150 mg/m<sup>2</sup> as 4-hour infusion ( days 1-3) and /or

Inj. Vincristine 1.5 mg / m<sup>2</sup> IV push (Day 1 & Day 5 ), Inj. Cyclophosphamide 300 mg/m<sup>2</sup> short infusion (Days 1— 5) and Inj. Adriamycin 30 mg / m<sup>2</sup> IV push (Day 4 & Day 5 )

## HEPATOBLASTOMA

### Treatment Algorithm: Hepatoblastoma



- \*Upfront surgical resection may be feasible in selected cases of Standard Risk hepatoblastoma
- \*\*Orthotopic liver transplant indicated in selected cases of high risk hepatoblastoma requiring extensive resection( multifocal/ pretext IV)
- PLADO: Cisplatin + Doxorubicin, (Total 6 cycles)
  - C5V – Cisplatin + 5-fluorouracil+ Vincristine ( Total 6 cycles)
  - SUPERPLADO : Alternating cycles of Carboplatin/Doxorubicin and Cisplatin ( total 10 cycles)

### Risk Stratification

High risk: Patients with any of the following	Standard risk
Serum alpha-fetoprotein <100 µg/l	All other patients
PRETEXT IV	
Small cell undifferentiated subtype	
Additional PRETEXT criteria:	
Extrahepatic intra-abdominal disease (E).	
Distant metastases (M ),	
Nodal metastases (N1, N2)	
Tumor extension into the main and/or both branches of the portal vein (P2, P2a)	
Tumor extension into the vena cava or all three hepatic veins (V3, V3a )	
Intraperitoneal Haemorrhage (H1)	

### Chemotherapy Doses :

#### PLADO: Cisplatin + Doxorubicin, (Total 6 cycles)

Inj. Cisplatin 90mg/m<sup>2</sup> in 500ml NS over 4-6 hr (Day 1) and Inj Adriamycin(30mg/m<sup>2</sup>) in as short infusion (Days 2-3 )

#### PLAVF : 4-6 cycles

Inj. Cisplatin 90mg/m<sup>2</sup> in 500ml NS over 4-6 hr (Day 1) , Inj. VCR (1.4mg/m<sup>2</sup>) IV push Day1 and nj. 5-FU 600 mg/m<sup>2</sup> in 250 ml NS infusion over 24 hr (Day 2)

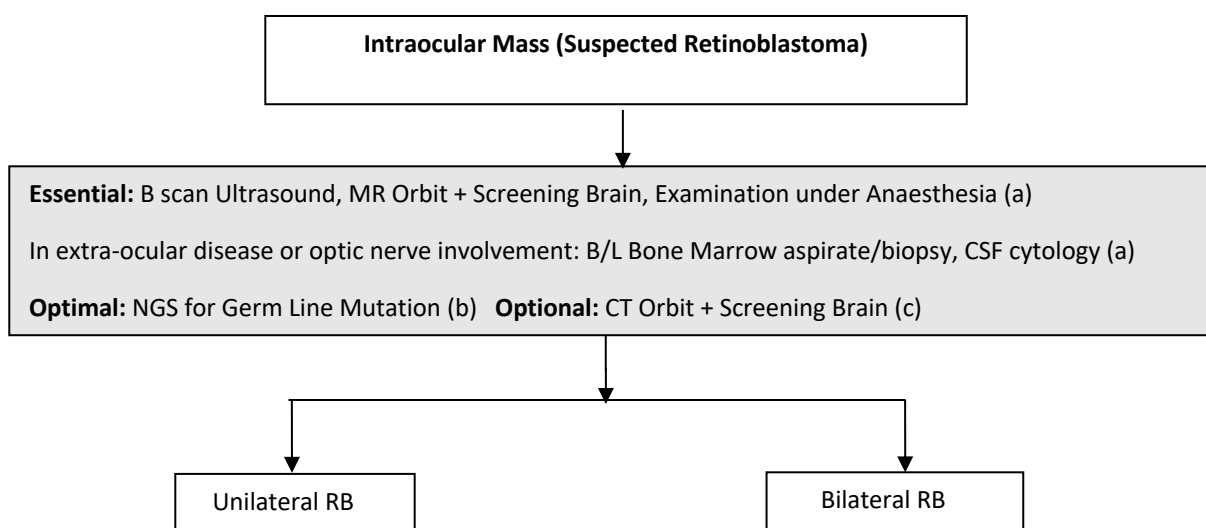
#### SUPERPLADO : Alternating cycles of Carboplatin/Doxorubicin and Cisplatin ( total 10 cycles)

Inj Cisplatin 80 mg/m<sup>2</sup> infusion over 24 hrs

Inj Carboplatin 500mg/m<sup>2</sup> over 1hr and Inj Doxorubicin 60 mg/m<sup>2</sup> infusion over 48hrs

## RETINOBLASTOMA

### Treatment Algorithm: Retinoblastoma



### Grouping for Intraocular Retinoblastoma

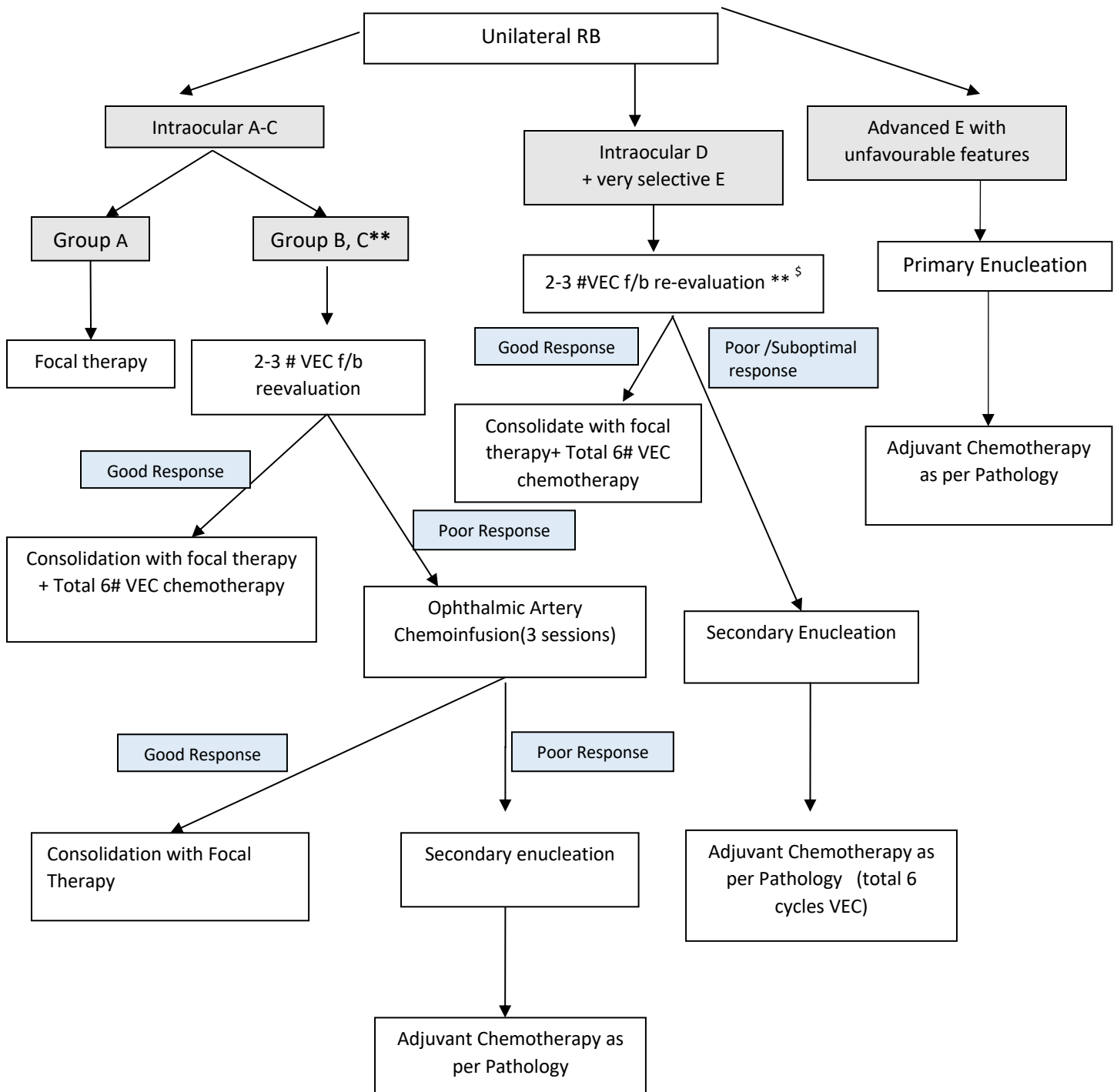
Group A	Small tumors away from foveola and disc <ul style="list-style-type: none"> <li>• Tumors &lt; 3 mm, confined to the retina, and</li> <li>• Located at least 3 mm from the foveola and 1.5 mm from the optic nerve.</li> </ul>
Group B	All remaining tumors confined to the retina <ul style="list-style-type: none"> <li>• all other tumors confined to retina and not in Group A</li> <li>• Subretinal fluid (without subretinal seeding) &lt;3 mm from the base of the tumor.</li> </ul>
Group C	Local vitreous or subretinal seeding <ul style="list-style-type: none"> <li>• Subretinal fluid alone &gt; 3mm and &lt; 6 mm from the tumor</li> <li>• Vitreous or subretinal seeding &lt; 3mm from the tumor</li> </ul>
Group D	Diffuse vitreous or subretinal seeding <ul style="list-style-type: none"> <li>• Subretinal fluid alone &gt; 6 mm from the tumor</li> </ul>

	<ul style="list-style-type: none"> <li>• Vitreous or subretinal seeding &gt; 3mm from the tumor</li> </ul>
Group E	Presence of any one or more of these poor prognosis features <ul style="list-style-type: none"> <li>• More than 2/3 of the globe filled with tumor</li> <li>• Tumor in anterior segment or anterior to vitreous</li> <li>• Tumor in ciliary body</li> <li>• Iris neovascularisation</li> <li>• Neovascular glaucoma</li> <li>• Opaque media from hemorrhage</li> <li>• Tumor necrosis with aseptic orbital cellulitis</li> <li>• Phthisis bulbi</li> </ul>

Genetic counseling needs to be done in all families, with genetic testing indicated in cases of suspected heritable disease

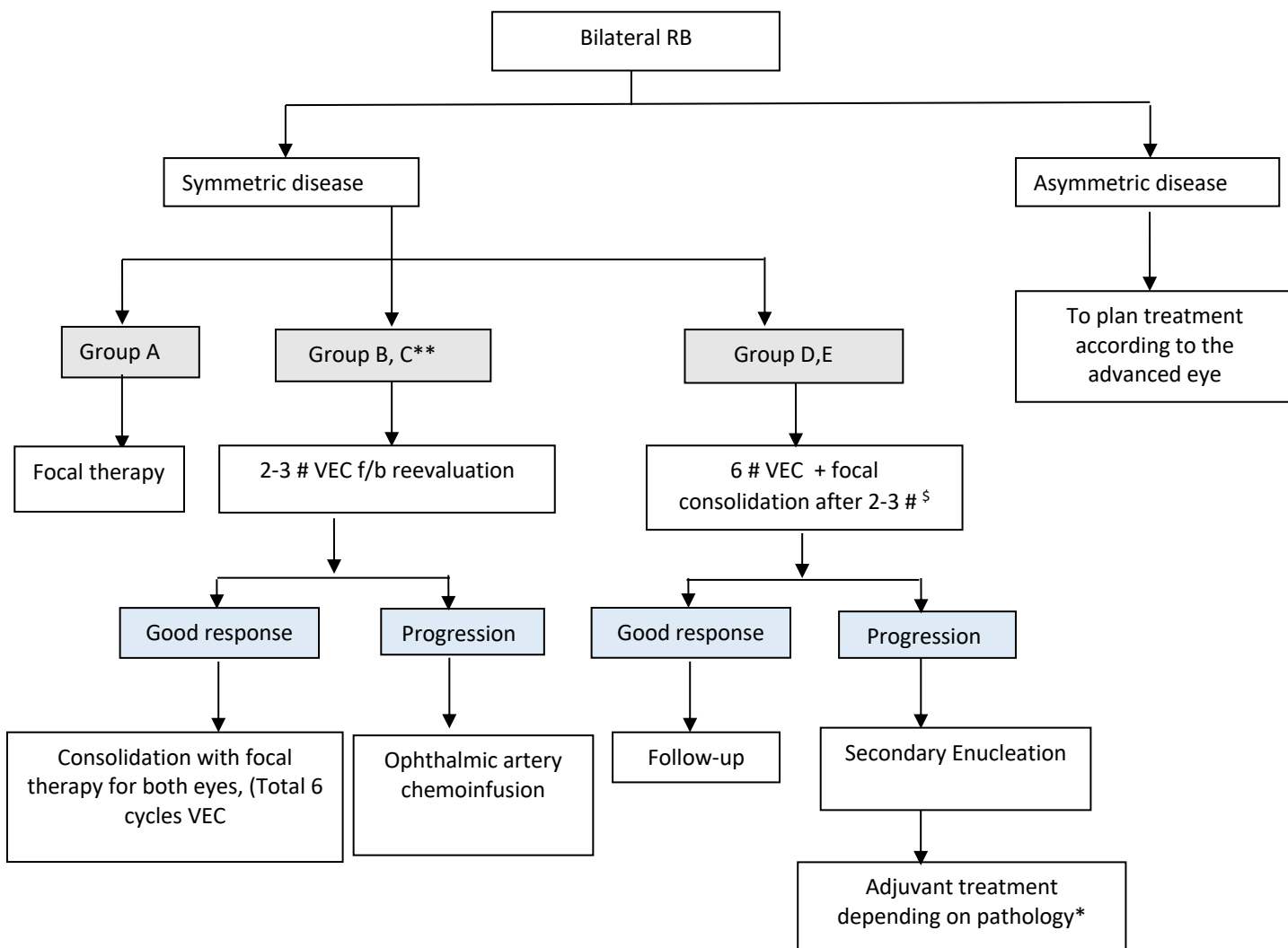
Screening of all siblings less than 7 years of age should be done

Treatment Algorithm: Retinoblastoma- Unilateral



- **Extraocular disease:** 12 cycles of chemotherapy(VEC), Secondary Enucleation ( after 2-4 cycles) and EBRT
- **Response assessment**
  - Intraocular disease : by Examination Under Anaesthesia (EUA) and periodic B-ultrasound (a)
  - Extraocular disease : Additionally, requires MRI (a)

### Treatment Algorithm: Retinoblastoma- Bilateral



- **Extraocular disease:** 12 cycles of chemotherapy(VEC), Secondary Enucleation ( after 2-4 cycles) and EBRT
- **Response assessment**
  - Intraocular disease : by Examination Under Anaesthesia (EUA) and periodic B-ultrasound (a)
  - Extraocular disease : Additionally, requires MRI (a)

**Unfavorable features in group E disease:** Phthisis bulbi, Intraocular haemorrhage, neovascular glaucoma, anterior chamber involvement

**Focal therapies**

- Laser photocoagulation
- TranspupillaryThermo Therapy
- Cryotherapy
- Plaque RT

\*\* To consider **ophthalmic artery chemoinfusion/ intra-arterial chemotherapy** case to case basis and where expertise available

§ To consider **intravitreal chemotherapy** in group D/E

### **Chemotherapy details**

#### **Chemotherapy :VEC (Vincristine, etoposide, Carboplatin)**

- 6 cycles for Intraocular disease
- 12 cycles for extraocular/stage 3 disease (plus EBRT)

#### **Other chemotherapy Options:**

- High dose carboplatin
- Vincristine/doxorubicin/cyclophosphamide (refractory group E tumours/ stage 3 disease)

#### **Stage 4 Bone Marrow disease:**

- Chemotherapy including high dose chemotherapy with Autologous BMT /Stem Cell Transplant

#### **Stage 4 Central nervous system disease.**

- Treatment intent to be decided on a case-to case basis by MDT. To consider palliation

#### **\*Indications for Adjuvant Chemotherapy:**

- Massive choroidal infiltration
- Post-laminar optic nerve involvement (PLONI)
- Scleral or Extra-scleral spread (on radiology or HPR)
- Cut-margin of Optic Nerve positive for tumor,
- Optic Nerve involvement upto the apex at presentation on MRI

#### **\*Indications for Adjuvant Radiotherapy:**

- Extra-scleral spread
- Cut-margin of Optic Nerve positive for tumor on histology
- Extra-ocular mass at presentation
- Optic Nerve involvement upto the apex at presentation on MRI

#### **Radiotherapy doses:**

- Definitive RT 45Gy/25#s over 5 weeks with conformal portals
- Adjuvant RT 39.6Gy/22#s over 5 weeks with conformal portals

### **Chemotherapy Doses**

Inj Vincristine 1.5mg/m<sup>2</sup> IV push over 1 minute on Day 1

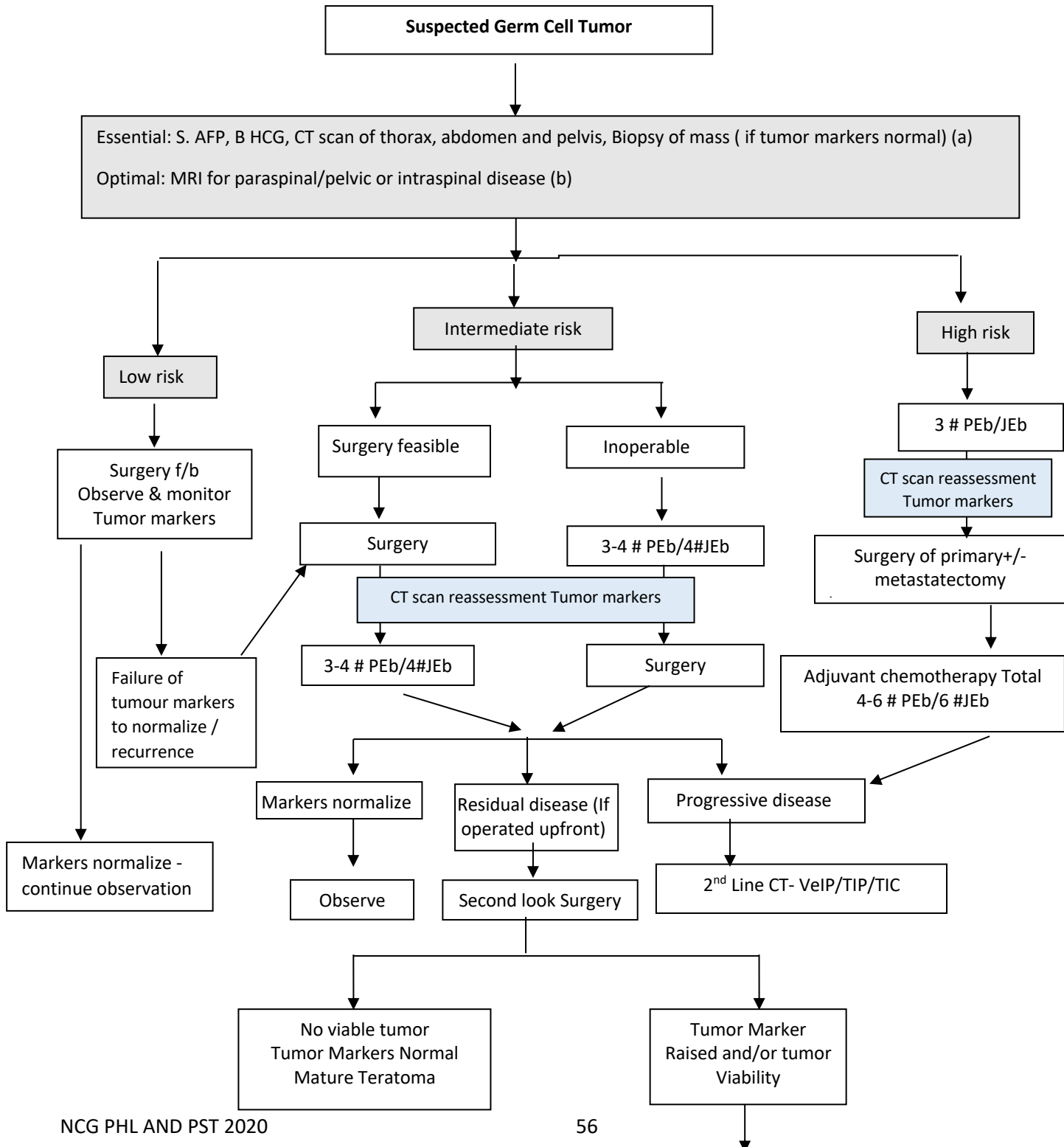
Inj Carboplatin 500-560 mg/m<sup>2</sup> IV over 60 minutes on Day 1

Inj Etoposide 100 mg/m<sup>2</sup> IV over 60 minutes ( Days 1-2)



## EXTRACRANIAL GERM CELL TUMOUR

### Treatment Algorithm: Extra Renal Germ Cell Tumor





**Chemotherapy:**

**PEb:** Cisplatin 20 mg/m<sup>2</sup> as infusion over 4 hours d1-d5, Inj Etoposide 100mg/m<sup>2</sup> as infusion over 2 hours d1-d5, Bleomycin 15mg/m<sup>2</sup> IV push d1 (3-4 cycles in Intermediate Risk and 4-6 cycles in High Risk)

**JEb:** Carboplatin 560mg/m<sup>2</sup> as 1 hour infusion d1, Inj Etoposide 100mg/m<sup>2</sup> as infusion over 2 hours d1-d5, Bleomycin 15mg/m<sup>2</sup> IV push d1 (4 cycles in Intermediate Risk and 6 cycles in High Risk)

**VeIP:** Inj. Velbe 6 mg/m<sup>2</sup> iv bolus (Day 1 only ), Inj. Ifosfamide 1.5 gm/m<sup>2</sup> as infusion over 2-3 hr (Days 1—5) with Inj. Mesna 400mg/m<sup>2</sup> IV at 0,4 and 8 hrs (Days 1—5), Inj. Cisplatin 20mg/m<sup>2</sup> infusion over 1 hr (Days 1-5)

**TIP or TIC:** Inj Paclitaxel 175m/m<sup>2</sup> over 3 hours on d1, Inj. Ifosfamide 1.5 gm/m<sup>2</sup> as infusion over 2-3 hr d2-d5, Inj. Cisplatin 20mg/m<sup>2</sup> infusion over 1 hr d2-d5 (4-6 cycles) OR Inj Carboplatin 560mg/m<sup>2</sup> over 1 hour on Day 1

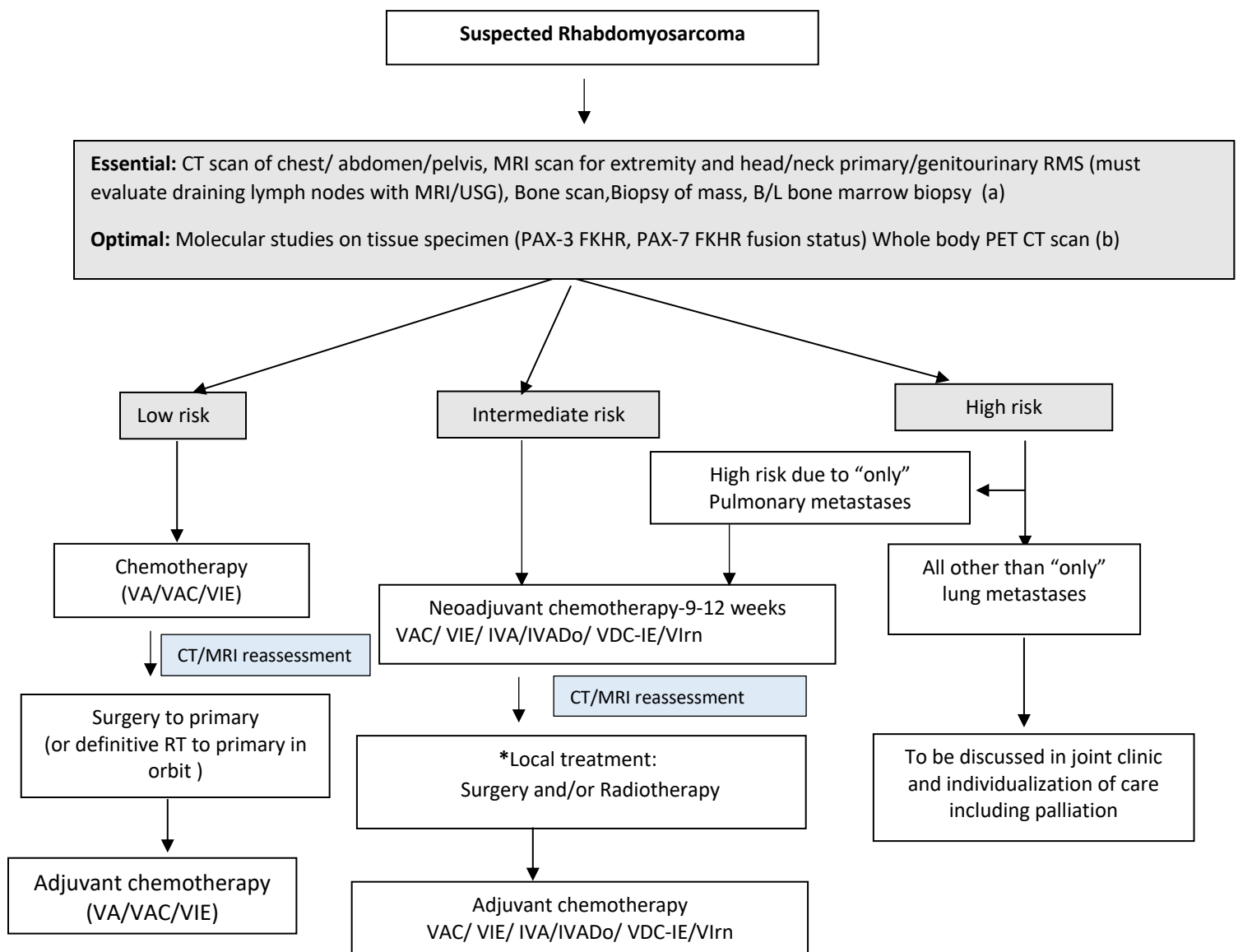
**Risk stratification**

Low risk:	Stage I testes Stage I ovary All immature teratoma (completely excised)
Intermediate risk:	Stage II- IV testes Stage II-III ovary Stage I-II Extragonadal Immature teratoma (incompletely excised)
High risk:	Stage IV ovary Stage III-IV Extragonadal

Tumour markers (Serum Alfa-fetoprotein, AFP and Serum Beta HCG) to be done at diagnosis. If raised, monitoring of disease status may be done by measurement of tumor markers after each cycle of chemotherapy, surgery and at end of treatment. Post treatment surveillance to be done using tumor markers.

## RHABDOMYOSARCOMA

### Treatment Algorithm: Neuroblastoma



### Chemotherapy Options:

- **VA:** Vincristine, Dactinomycin (for low risk only) Inj. Vincristine (1.5mg/m<sup>2</sup>) slow IV push weekly or 3-weekly, Inj Actinomycin slow IVP (> 3yrs/1-3yrs: 0.045mg/kg, <1 yr : 0.025mg/kg) (D1only)
- **VAC:** Inj. Vincristine (1.5mg/m<sup>2</sup>) slow IV push weekly or 3-weekly, Inj Cyclophosphamide over 30 mins (> 3yrs : 2200mg/m<sup>2</sup>, 1-3yrs: 73mg/kg, <1 yr : 6mg/kg) (D1 only) with Inj. Mesna (500 mg/m<sup>2</sup>) at 0,3,6,9 hrs (D1 only) Inj Actinomycin slow IVP (> 3yrs/1-3yrs: 0.045mg/kg, <1 yr : 0.025mg/kg) (D1only)
- **VIE:** Inj. Vincristine (1.5mg/m<sup>2</sup>) slow IV push weekly or 3-weekly, Inj. Ifosfamide (1800 mg/m<sup>2</sup>) as infusion over 3-4 hours (D1 to D5), Inj. Mesna (600 mg/m<sup>2</sup>) at 0,3,6,9 hrs (D1 to D5) and Inj Etoposide (100mg/m<sup>2</sup>) over 1-2 hrs (D1 to D5)
- **IVA:** Inj Ifosfamide 3 g/m<sup>2</sup> as 3-hour intravenous infusion daily ( Days 1 and 2 ), Inj Vincristine 1.5 mg/m<sup>2</sup> IV push on Day 1 Inj actinomycin D at a dose of 1.5 mg/m<sup>2</sup> IV push Day 1
- **IVADo:** Inj Ifosfamide 3 g/m<sup>2</sup> as 3-hour intravenous infusion daily ( Days 1 and 2 ), Inj Vincristine 1.5 mg/m<sup>2</sup> IV push on Day 1 Inj actinomycin D at a dose of 1.5 mg/m<sup>2</sup> IV push Day 1 , Inj Doxorubicin at a dose of 30 mg/m<sup>2</sup> over 6-hour intravenous infusion ( Days 1 and 2 )
- **VDC-IE :** Alternate 14 day cycles of VDC (Inj Vincristine 1.4mg/m<sup>2</sup> IV push Day 1, Inj Doxorubicin 75mg/m<sup>2</sup> over 6 hours and Inj Cyclophosphamide 1200mg/m<sup>2</sup> slow IV infusion Days 1-2 and IE (Inj Ifosfamide 1.8 gm/m<sup>2</sup> as 3-hour intravenous infusion daily ( Days 1-5), Inj Etoposide 100mg/m<sup>2</sup> over 2 hours intravenous infusion daily ( Days 1 -5 )
- **Virn :** Vincristine/Irinotecan

\*Local Therapy options: Complete surgical excision only (upfront surgery)

Complete Surgical excision followed by Radiotherapy(or brachytherapy)

Definitive Radiotherapy (inoperable)

(Indications and doses for adjuvant radiation are given below)

### Radiotherapy Guidelines for Rhabdomyosarcoma

S.No.	Site / Stage / Histology	RT Field	RT Dose
1.	Group I Embryonal Alveolar	No RT Pre - Chemotherapy primary site	36Gy
2.	Group II N0 (microscopic residual disease after surgery) N1 (resected regional lymph node involvement)	Pre - Chemotherapy primary site Pre - Chemotherapy primary site + Nodes	36Gy 41.4Gy
3.	Group III All  Patients undergoing delayed surgical resection with negative margins	Pre - Chemotherapy primary site Pre-chemotherapy primary site	50.4Gy (45 Gy for orbital tumors in complete remission) 36Gy
4.	Group IV	Treat primary site as for other	

groups + all metastatic sites if  
technically feasible & safe

**Staging based on the TNM and site of disease:**

Stage	Sites of Primary Tumor	T Stage	Tumor Size	Regional Lymph Nodes	Distant Metastasis
I	Favorable sites	T1 or T2	Any size	N0 or N1 or NX	M0
II	Unfavorable sites	T1 or T2	a- 5 cm	N0 or NX	M0
III	Unfavorable sites	T1 or T2	a- 5 cm	N1	M0
			b- > 5 cm	N0 or N1 or NX	
IV	Any site	T1 or T2	Any size	N0 or N1 or NX	M1

M0 = absence of metastatic spread; M1 = presence of metastatic spread beyond the primary site; N0 = absence of nodal spread; N1 = presence of nodal spread beyond the primary site; X = unknown N status.

**Grouping based on the surgico-pathological resection**

Group	Definition
I	A localized tumor that is completely removed with pathologically clear margins and no regional lymph node involvement.
II	A localized tumor that is grossly removed with (a) microscopic disease at the margin, (b) involved, grossly removed regional lymph nodes, or (c) both (a) and (b).
III	A localized tumor with gross residual disease after incomplete removal or biopsy only.
IV	Distant metastases are present at diagnosis.

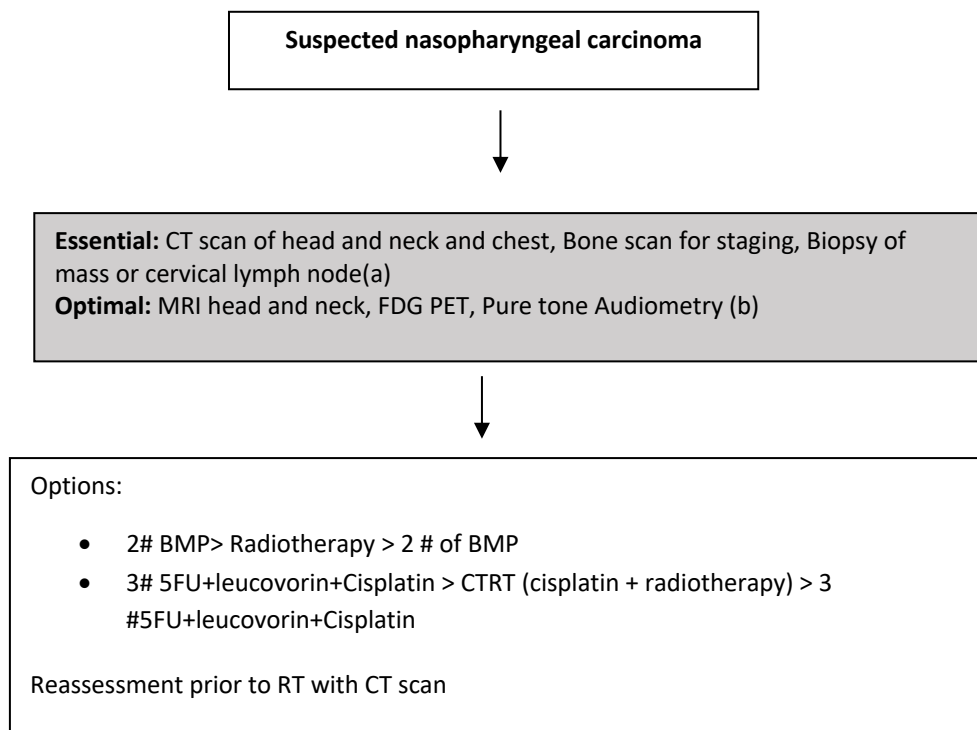
**Risk stratification of RMS**

Risk Group	Histology	Stage	Group
Low risk	Embryonal / (Fusion negative)	1	I, II, III
	Embryonal / (Fusion negative)	2, 3	I, II
Intermediate risk	Embryonal / (Fusion negative)	2, 3	III

	Alveolar / (Fusion positive)	1, 2, 3	I, II, III
High risk	Embryonal or Alveolar (Fusion negative/positive)	4	IV

## NASOPHARYNGEAL CARCINOMA

### Treatment Algorithm: Nasopharyngeal Carcinoma



#### Chemotherapy:

**BMP:** Inj. Bleomycin 10Units/m<sup>2</sup> iv push (D1/D8), Inj. Methotrexate 50mg / m<sup>2</sup> iv push (D1/D8) Inj. Cisplatin 20mg/m<sup>2</sup> infusion over 4 hr d1-d5

**5FU+Cisplatin:** Inj Cisplatin 80 mg/m<sup>2</sup> over 6 hours on day 1 and Inj 5-fluorouracil 1,000 mg/m<sup>2</sup>/d as continuous infusion (days 1 to 4)

#### Radiotherapy :

**Dose:** 55-70 Gy given in fractions of 1.6-2.1 Gy in 33 fractions over 7 weeks

Intensity modulated radiotherapy with/without image guidance

## ANNEXURE -1

### DIAGNOSTIC INVESTIGATIONS FOR COMMON PAEDIATRIC SOLID TUMOURS

Tumor	Local/staging Essential (a)	Local Staging Optimal (b)/Optional (c)
<b>All Paediatric Solid tumours</b>	USG abdomen to be part of basic diagnostic workup	
<b>Neuroblastoma</b>	CT scan of primary tumor site MIBG scan/Whole body FDG-PET scan Biopsy of mass n-Myc status by FISH B/L Bone marrow biopsy	Urinary VMA Segmental chromosomal anomalies MRI ( paraspinaltumours) (b) Bone Scan (c)
<b>Wilms Tumor</b>	CT scan thorax, abdomen and pelvis	USG doppler MRI for Bilateral WT and nephroblastomatosis (b)
<b>Hepatoblastoma</b>	CT thorax and abdomen (Triphasic) Serum Alfa Fetoprotein (AFP)	MRI Liver (b)
<b>Rhabdomyosarcoma</b>	CT scan of chest/ abdomen/pelvis MRI scan for extremity and head/neck primary/genitourinary RMS Bone scan Biopsy of mass B/L Bone marrow biopsy	Molecular studies on tissue specimen (PAX-3 FKHR, PAX-7 FKHR fusion status) Whole body PET CT scan (b)
<b>Germ Cell Tumor</b>	S. AFP, B HCG CT scan of thorax, abdomen and pelvis Biopsy of mass (if tumor markers normal)	MRI for paraspinal/pelvic or intraspinal disease (b) Whole body PET CT scan (c)
<b>Retinoblastoma</b>	B scan Ultrasound MR Orbit + Screening Brain Examination under Anaesthesia In extra-ocular disease or optic nerve involvement: B/L Bone Marrow biopsy, CSF cytology	NGS for Germ Line Mutation (b) CT Orbit + Screening Brain (c)
<b>Nasopharyngeal Carcinoma</b>	CT scan of head and neck and chest Bone scan for staging Biopsy of mass or cervical lymph node	MRI head and neck, FDG PET



Pure tone Audiometry (b)

