

## IE/VC

### INDICATION (ICD10) C40, C41, C49

1. Ewing sarcoma

### REGIMEN

#### IE cycles 1, 3 and 5

Days 1, 2, 3, 4 and 5

Mesna 1000mg/m<sup>2</sup> IV bolus one hour prior to ifosfamide

IFOSFAMIDE 1800mg/m<sup>2</sup> in #ml sodium chloride 0.9% IV infusion over 60 minutes

Mesna 1800mg/m<sup>2</sup> in 1000ml sodium chloride 0.9% IV infusion over 60 minutes  
concurrently with ifosfamide

ETOPOSIDE 100mg/m<sup>2</sup> in #ml sodium chloride 0.9% IV infusion over 2 hours

Mesna 1200mg/m<sup>2</sup> in 1000ml sodium chloride 0.9% IV infusion over 16 hours

#### VC cycles 2 and 4

Day 1 VINCRISTINE 2mg/m<sup>2</sup> (maximum 2mg) in 50ml sodium chloride 0.9% IV  
infusion over 10 minutes

Mesna 1000mg/m<sup>2</sup> IV bolus one hour prior to cyclophosphamide

CYCLOPHOSPHAMIDE 1200mg/m<sup>2</sup> in #ml sodium chloride 0.9% IV infusion over 60  
minutes

Mesna 1200mg/m<sup>2</sup> in 1000ml sodium chloride 0.9% IV infusion over 60 minutes  
concurrently with ifosfamide

Mesna 800mg/m<sup>2</sup> in 1000ml sodium chloride 0.9% IV infusion over 23 hours

# diluent and diluent volume for dose prescribed as per national standardised product specification

### CYCLE FREQUENCY AND NUMBER OF CYCLES

Every 14 days or on haematological recovery to absolute neutrophil count  $\geq 0.75 \times 10^9/L$ , platelets  $\geq 75 \times 10^9/L$ .

Equivalent to cycles 10 to 14 following VDC/IE

### ANTI-EMETICS

High emetic risk days 1, 2, 3, 4 and 5 cycles 1, 3 and 5 (consider aprepitant)

High emetic risk day 1 cycles 2 and 4 (consider aprepitant)

### CONCURRENT MEDICATION REQUIRED

Cyclophosphamide	Ensure mesna administered, using separate lumen from cyclophosphamide. Ensure adequate oral fluid intake. Cotrimoxazole 480mg bd M/W/F for duration of chemotherapy. Benzydamine mouthwash
Ifosfamide	Ensure mesna administered. Ensure adequate oral fluid intake. Cotrimoxazole 480mg bd M/W/F for duration of chemotherapy.
Vincristine	Laxatives should be prescribed
GCSF	Starting at least 24 hours after chemotherapy and stop at least 24 hours before commencing chemotherapy

## EXTRAVASATION AND TYPE OF LINE / FILTERS

Cyclophosphamide - neutral  
Etoposide - irritant  
Ifosfamide – neutral  
Vincristine – vesicant

Double lumen central line

## INVESTIGATIONS

Blood results required before SACT administration  
FBC, U&E and LFTs every week  
Neutrophils x 10<sup>9</sup>/L ≥0.75  
Platelets x 10<sup>9</sup>/L ≥75  
DTPA baseline  
Creatinine clearance >55ml/min  
Serum creatinine every cycle  
Haematuria monitoring every specimen IE cycles, pre treatment only VC cycles  
Vitamin D baseline  
Hepatitis B status baseline  
ECG (possible ECHO) required if patient has preexisting cardiac disease  
Baseline weight and every cycle

## MAIN TOXICITIES AND ADVERSE REACTIONS

Cyclophosphamide	May irritate bladder, drink copious volumes of water. Microscopic Haemorrhagic cystitis: additional bolus dose 600mg/m <sup>2</sup> then continue infusion at double dose. Grade ≥2 macroscopic haemorrhagic cystitis: discontinue chemotherapy and continue double dose MESNA and hydration x 24 hours post-chemotherapy
Ifosfamide	Ifosfamide encephalopathy. Nephrotoxicity: Irreversible renal failure and tubular damage can occur, and this is more frequent with cumulative doses over 25–50g/m <sup>2</sup> of Ifosfamide. Haematuria.
Vincristine	Neuropathy

## INTERACTIONS WHICH MAY REQUIRE DOSE MODIFICATIONS

(not exhaustive list check SPC/BNF/Stockleys)

Cyclophosphamide	Cytochrome P450 enzyme inducers (e.g. rifampicin, carbamazepine, phenytoin, St Johns Wort, corticosteroids): may increase active cyclophosphamide metabolites. Allopurinol, Cimetidine and protease inhibitors: may increase active metabolites. Aprepitant, Ciprofloxacin, Fluconazole, Itraconazole: may reduce activation of cyclophosphamide and alter the effectiveness of treatment. Grapefruit juice: decreased or delayed activation of cyclophosphamide. Patients should be advised to avoid grapefruit juice.
Ifosfamide	Aprepitant and fosaprepitant are predicted to increase the exposure to ifosfamide. Caution.

## DOSE MODIFICATIONS

Doxorubicin maximum lifetime dose

= 400mg/m<sup>2</sup> (in patients with cardiac dysfunction or exposed to mediastinal irradiation)

= 450-550mg/m<sup>2</sup> (with normal cardiac function).

## Haematological

If platelets and ANC not recovering by day 22 give 80% VC/IE doses in subsequent cycles.

## Non-haematological

Cardiac Toxicity

Fractional shortening (FS) <29% or left ventricular (LVEF) <40% or decrease by an absolute value of ≥10 percentile points from previous tests then delay chemotherapy course for 7 days and repeat cardiac tests. If FS has recovered to ≥29% then proceed to the next course. If FS remains <29% then omit doxorubicin and substitute dactinomycin 1.5mg/m<sup>2</sup> on day 1 only (max 1.5mg) or use liposomal doxorubicin when meet funding criteria.

Gastrointestinal toxicity

Grade 3/4 mucositis beyond day 22 after IE give 80% IE.

Ifosfamide

Neural and nephrotoxicity grade

Toxicity Grade	GFR (ml/min/1.73m <sup>2</sup> )	Tp/C <sub>crea</sub> (T <sub>mp</sub> /GFR) (mmol/l)	HCO <sub>3</sub> <sup>*</sup> (mmol/l)	Action (apply worst grade)
Grade 0/1	≥60	≥1.00	≥17.0	give 100% dose
Grade 2	40-59	0.8-0.99	14.0-16.9	give 70% dose
Grade 3/4	≤40	≤0.8	≤14.0	**Switch to cyclophosphamide

\*Low values of HCO<sub>3</sub> should be re-checked when the patient is clinically stable (to rule out infection as a cause, etc) before modifying treatment.

\*\*Discuss with consultant before and to confirm substitution of ifosfamide with cyclophosphamide 2100mg/m<sup>2</sup>/day day 1 only.

Fractional phosphate clearance calculated

$$Tp/C_{crea} [\text{mmol/ml}] = \text{Phosphate}_{\text{serum}} - \frac{\text{Phosphate}_{\text{urine}} \times \text{creatinine}_{\text{serum}}}{\text{Creatinine}_{\text{urine}}}$$

## Hepatic impairment

Etoposide

Bilirubin ≥50micromol/L or decreased albumin	give 50% dose
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Ifosfamide

Bilirubin >17micromol/L or AST and ALP >2.5xULN	discuss
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Vincristine

Bilirubin 25-51 or AST 60-180u/L	give 50%
Bilirubin >51micromol/L and normal AST	give 50%
Bilirubin >51micromol/L and AST >180u/L	not recommended

## Renal impairment

### Cyclophosphamide

CrCl 10-29ml/min	give 75% dose
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### Etoposide

GFR<60ml/min/1.73m<sup>2</sup> then give 70% etoposide dose

Defer therapy and monitor renal function and discuss with consultant if there is a significant rise in serum creatinine, even if CrCl >60mls/min as ifosfamide may cause delayed renal impairment.

### Ifosfamide

CrCl >50ml/min	give 100% dose
CrCl <50ml/min	Clinical decision

## REFERENCES

1. EuroEwing 2012