Methotrexate, vinBLAStine, DOXOrubicin, CISplatin (MVAC) -28 Days Therapy

INDICATIONS FOR USE:

<table>
<thead>
<tr>
<th>INDICATION</th>
<th>ICD10</th>
<th>Regimen Code</th>
<th>Reimbursement Status</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neoadjuvant treatment of muscle invasive transitional cell carcinoma (TCC) of the urothelium</td>
<td>C67</td>
<td>00338a</td>
<td>Hospital</td>
</tr>
<tr>
<td>Adjuvant treatment of muscle invasive TCC of the urothelium</td>
<td>C67</td>
<td>00338b</td>
<td>Hospital</td>
</tr>
<tr>
<td>Locally advanced or metastatic TCC of the urothelium</td>
<td>C67</td>
<td>00338c</td>
<td>Hospital</td>
</tr>
</tbody>
</table>

TREATMENT:

The starting dose of the drugs detailed below may be adjusted downward by the prescribing clinician, using their independent medical judgement, to consider each patient's individual clinical circumstances.

Neoadjuvant: Treatment is administered as described in treatment table below every 28 days for 3 cycles unless there is disease progression or unacceptable toxicity develops prior to radical cystectomy.

Adjuvant: Treatment is administered as described in treatment table below every 28 days for 4 cycles unless there is disease progression or unacceptable toxicity develops.

Metastatic: Treatment is administered as described in treatment table below for up to 6 cycles or until disease progression or unacceptable toxicity develops.

Facilities to treat anaphylaxis MUST be present when the chemotherapy is administered.

<table>
<thead>
<tr>
<th>Day</th>
<th>Drug</th>
<th>Dose</th>
<th>Route</th>
<th>Diluent &amp; Rate</th>
<th>Cycle</th>
</tr>
</thead>
<tbody>
<tr>
<td>1, 15, 22</td>
<td>Methotrexate</td>
<td>30mg/m²</td>
<td>IV Bolus</td>
<td>Every 28 days</td>
<td></td>
</tr>
<tr>
<td>2, 15, 22</td>
<td>vinBLAStine</td>
<td>3mg/m²</td>
<td>IV infusion</td>
<td>50ml 0.9% NaCl over 15 min</td>
<td>Every 28 days</td>
</tr>
<tr>
<td>2</td>
<td>DOXOrubicin</td>
<td>30mg/m²</td>
<td>IV Bolus</td>
<td>Every 28 days</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>CISplatin</td>
<td>70mg/m²</td>
<td>IV infusion</td>
<td>500ml 0.9% NaCl over 2 hours</td>
<td>Every 28 days</td>
</tr>
</tbody>
</table>

*VinBLAStine is a neurotoxic chemotherapeutic agent. Refer to NCCP Guidance on the Safe Use of Neurotoxic drugs (including Vinca Alkaloids) in the treatment of cancer. Link*

*Lifetime cumulative dose of DOXOrubicin is 450mg/m²
In establishing the maximal cumulative dose of an anthracycline, consideration should be given to the risk factors outlined below and to the age of the patient.*

*1 Pre and post hydration therapy required for CISplatin
See local hospital policy recommendations.
Suggested prehydration for CISplatin therapy:
1. Administer 10mmol magnesium sulphate (MgSO₄) (±/− KCl 20mmol/L if indicated) in 1000 mL sodium chloride 0.9% over 60 minutes.
Post hydration: Administer 1000 ml 0.9% NaCl over 60mins
Mannitol 10% may be used as per local policy to induce diuresis, although there is no conclusive evidence that this is required. The routine use of furosemide to increase urine flow is not recommended unless there is evidence of fluid overload (6, 7).
ELIGIBILITY:

- Indications as above
- ECOG 0-2

EXCLUSIONS:

- Hypersensitivity to methotrexate, vinBLASTine, DOXOrubicin, CISplatin or any of the excipients
- Congestive heart failure (LVEF < 50%) or other significant heart disease
- Moderate/severe renal impairment (creatinine clearance < 60 mL/min)
- Pregnancy and Lactation
- Pre-existing neuropathies ≥ grade 2
- Significant hearing impairment/tinnitus

PRESCRIPTIVE AUTHORITY:

The treatment plan must be initiated by a Consultant Medical Oncologist.

TESTS:

Baseline tests:

- FBC, Renal and liver profile
- ECG
- MUGA or ECHO (LVEF > 50% to administer doxorubicin) if >65 years or if clinically indicated.
- Audiology and creatinine clearance if clinically indicated

Regular tests:

- FBC, Renal and liver profile prior to each cycle
- If clinically indicated MUGA scan or ECG
- Repeat FBC and renal profile prior to treatment on day 15 and day 22

Disease monitoring:

Disease monitoring should be in line with the patient’s treatment plan and any other test/s as directed by the supervising Consultant.

DOSE MODIFICATIONS:

- Any dose modification should be discussed with a Consultant.
**NCCP Chemotherapy Regimen**

**Haematological:**

Table 1: Dose modification for haematological toxicity on day of chemotherapy

<table>
<thead>
<tr>
<th>ANC (x10^9/L)</th>
<th>Platelets (x10^9/L)</th>
<th>Dose</th>
</tr>
</thead>
<tbody>
<tr>
<td>≥1.5</td>
<td>and</td>
<td>≥100</td>
</tr>
<tr>
<td>&lt;1.5</td>
<td>or</td>
<td>&lt;100</td>
</tr>
</tbody>
</table>

For ANC < 0.5 for 5-7 days or Thrombocytopenic bleeding or platelets < 25, Hold *then 75% of previous dose.

*Do not start a new cycles until ANC ≥1.5 x 10^9/L and platelets ≥100 x10^9/L

**Renal and Hepatic Impairment:**

Table 2: Dose modifications in renal and hepatic impairment

<table>
<thead>
<tr>
<th>Drug</th>
<th>Renal Impairment</th>
<th>Hepatic Impairment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methotrexate</td>
<td>Cr Cl (ml/min)</td>
<td>Bilirubin (micromol/L)</td>
</tr>
<tr>
<td>&gt;80</td>
<td>100%</td>
<td>&lt;50</td>
</tr>
<tr>
<td>60-80</td>
<td>65%</td>
<td>51-85</td>
</tr>
<tr>
<td>45-60</td>
<td>50%</td>
<td>&gt;85</td>
</tr>
<tr>
<td>30-45</td>
<td>Clinical decision</td>
<td></td>
</tr>
<tr>
<td>&lt;30</td>
<td>Contraindicated</td>
<td></td>
</tr>
</tbody>
</table>

Methotrexate: Contraindicated in severe hepatic impairment

<table>
<thead>
<tr>
<th>vinBLASTine</th>
<th>No dose reduction necessary</th>
<th>Bilirubin (micromol/L)</th>
<th>AST/ALT (Units)</th>
<th>Dose</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>26-51</td>
<td>or</td>
<td>60-180</td>
</tr>
<tr>
<td></td>
<td></td>
<td>&gt;51</td>
<td>and</td>
<td>Normal</td>
</tr>
<tr>
<td></td>
<td></td>
<td>&gt;51</td>
<td>and</td>
<td>&gt;180</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>DOXOrubicin</th>
<th>No dose reduction required. Clinical decision in severe impairment.</th>
<th>Bilirubin (micromol/L)</th>
<th>Dose</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>20-51</td>
<td>50%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>51-85</td>
<td>25%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>&gt;85</td>
<td>Omit</td>
</tr>
</tbody>
</table>

If AST 2-3 x normal, give 75% dose.

If AST >3x ULN, give 50% dose

<table>
<thead>
<tr>
<th>CIPlatin</th>
<th>GFR (ml/min)</th>
<th>Dose</th>
<th>No dose reduction necessary</th>
</tr>
</thead>
<tbody>
<tr>
<td>≥60</td>
<td>100%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>45-59</td>
<td>75%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;45</td>
<td>Consider CARBOplatin</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Management of adverse events:**

Table 3: Dose Modification of MVAC Therapy for Adverse Events

<table>
<thead>
<tr>
<th>Adverse reactions</th>
<th>Recommended dose modification</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neurotoxicity</td>
<td></td>
</tr>
<tr>
<td>Grade 2 present at start of next cycle</td>
<td>Reduce dose of CIPlatin and vinBLASTine by 25% dose.</td>
</tr>
<tr>
<td>Grade 3</td>
<td>Discontinue CIPlatin and vinBLASTine</td>
</tr>
</tbody>
</table>

NCCP Regimen: Methotrexate, VinBLASTine, DOXOrubicin and CIPlatin (MVAC)-28day Therapy

Published: 20/06/2016
Review: 08/01/2025
Version number: 3

Tumour Group: Genitourinary
NCCP Regimen Code: 00338

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NCCP Chemotherapy Regimen

SUPPORTIVE CARE:

EMETOGENIC POTENTIAL:

Day 1 Low (Methotrexate)
Day 2 High (Refer to local policy).

<table>
<thead>
<tr>
<th>Drug</th>
<th>Level</th>
<th>Refer to local policy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methotrexate</td>
<td>Low</td>
<td>(Refer to local policy).</td>
</tr>
<tr>
<td>vinBLAStine</td>
<td>Minimal</td>
<td>(Refer to local policy).</td>
</tr>
<tr>
<td>DOXorubicin</td>
<td>Moderate</td>
<td>(Refer to local policy).</td>
</tr>
<tr>
<td>CISplatin</td>
<td>High</td>
<td>(Refer to local policy).</td>
</tr>
</tbody>
</table>

PREMEDICATIONS: None usually required

OTHER SUPPORTIVE CARE:

- Hydration prior and post CISplatin administration (Reference local policy or see recommendations above). Patient should be encouraged to drink large quantities of liquids for 24 hours after the CISplatin infusion to ensure adequate urine secretion.
- Prophylactic laxatives may be required to prevent constipation related to the use of vinca alkaloids.

ADVERSE EFFECTS / REGIMEN SPECIFIC COMPLICATIONS:

The adverse effects listed are not exhaustive. Please refer to the relevant Summary of Product Characteristics for full details.

- **Neutropenia**: Fever or other evidence of infection must be assessed promptly and treated appropriately.
- **Pleural effusion or ascites**: Methotrexate should be used with caution in patients with pleural effusions or ascites, as methotrexate may accumulate in third space fluid compartments.
- **Extravasation**: vinBLAStine And DOXorubicin are vesicants which may cause pain and tissue necrosis if extravasated. (Refer to local extravasation guidelines).
- **Cardiac Toxicity**: DOXorubicin is cardiotoxic and must be used with caution, if at all, in patients with severe hypertension or cardiac dysfunction.
- **Hypersensitivity**: Hypersensitivity reactions have been reported with CISplatin.
- **Renal Toxicity**: Nephrotoxicity is common with CISplatin. Strongly encourage oral hydration. If oral hydration is not possible (e.g. excessive nausea), IV hydration is indicated. Avoid nephrotoxic drugs such as aminoglycoside antibiotics where possible. Where treatment with nephrotoxic drugs must be used, monitor renal function.
- **Ototoxicity and sensory neural damage**: These are associated with CISplatin therapy. They should be assessed by history prior to each cycle

DRUG INTERACTIONS:

- NSAIDs may decrease the clearance of methotrexate by decreasing its renal perfusion and tubular secretion thus increasing its toxicity.
NCCP Chemotherapy Regimen

- Sulphonamides and penicillins may displace bound methotrexate from plasma protein increasing serum methotrexate levels and its toxicity.
- Concomitant administration of drugs that cause folate deficiency may lead to increased methotrexate toxicity.
- Ciprofloxacin may inhibit renal tubular transport of methotrexate, increasing serum methotrexate levels and its toxicity.
- Probenecid may inhibit renal excretion of methotrexate, increasing serum methotrexate levels and its toxicity.
- Co-administration of CISplatin has been reported to cause higher plasma concentrations of vinBLASTine.
- Erythromycin may increase the toxicity of vinBLASTine.
- Concurrent administration of calcium channel blockers with DOXOrubicin should be avoided as they may decrease the clearance of DOXOrubicin.
- CISplatin may potentiate the nephrotoxic and ototoxic effects of loop diuretics and aminoglycosides so concurrent use should be avoided.
- Current drug interaction databases should be consulted for more information.

**ATC CODE:**

- Methotrexate - L01BA01
- vinBLASTine - L01CA01
- DOXOrubicin - L01DB01
- CISplatin - L01XA01

**REFERENCES:**

6. Nephrotoxicity Associated with CISplatin EvIQ ID: 184 v.3

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   https://www.uptodate.com/contents/CISplatin-nephrotoxicity?source=search_result&search=CISplatin%20hydration&selectedTitle=1~150

   Available at: https://www.hpra.ie/img/uploaded/swedocuments/Licence_PA0822-206-006_12112019145755.pdf

   Available at https://www.hpra.ie/img/uploaded/swedocuments/Licence_PA0822-208-001_3111018125148.pdf

    Available at https://www.hpra.ie/img/uploaded/swedocuments/Licence_PA2315-083-001_25022019110336.pdf

    Available at https://www.hpra.ie/img/uploaded/swedocuments/Final%20approved%20SPC%20PA0822.199.001.pdf

12. NCCP Classification Document for Systemic Anti-Cancer Therapy (SACT) Induced Nausea and Vomiting. V1 2018. Available at:

<table>
<thead>
<tr>
<th>Version</th>
<th>Date</th>
<th>Amendment</th>
<th>Approved By</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>20/06/2016</td>
<td></td>
<td>Prof Maccon Keane</td>
</tr>
<tr>
<td>2</td>
<td>11/12/2017</td>
<td>Updated with new NCCP regimen format, updated with revised CISplatin hydration regimen recommendations</td>
<td>Prof Maccon Keane</td>
</tr>
<tr>
<td>3</td>
<td>08/01/2020</td>
<td>Reviewed. Standardisation of treatment table and renal dose modifications. Update of emetogenic potential</td>
<td>Prof Maccon Keane</td>
</tr>
</tbody>
</table>

Comments and feedback welcome at oncologydrugs@cancercontrol.ie.

1Cardiotoxicity is a risk associated with anthracycline therapy that may be manifested by early (acute) or late (delayed) effects.

Risk factors for developing anthracycline-induced cardiotoxicity include:
- high cumulative dose, previous therapy with other anthracyclines or anthracenediones
- prior or concomitant radiotherapy to the mediastinal/pericardial area
- pre-existing heart disease
- concomitant use of other potentially cardiotoxic drugs

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In establishing the maximal cumulative dose of an anthracycline, consideration should be given to the risk factors above and to the age of the patient.