

CISplatin and Capecitabine Adjuvant Chemoradiation Therapy

INDICATIONS FOR USE:

INDICATION	ICD10	Regimen Code	Reimbursement Status
Adjuvant treatment of adult patients with resected gastric cancer stage IIA or higher and no distant metastases	C16	00473a	CISplatin Hospital Capecitabine CDS

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 patients individual clinical circumstances.

Chemotherapy is given in 5 cycles as described in Table 1 below:

- cycle 1 and 2 - prior to radiation treatment (21 day cycles),
- cycle 3 - radiation treatment (5 weeks) and
- cycles 4 and 5 - following radiation treatment (21 day cycles).

Cycle 4 to start 2-4 weeks after completion of radiation.

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

Day	Drug	Dose	Route	Diluent & Rate	Cycle
CYCLE 1 and 2					
1	^a CISplatin	60mg/m ²	IV Infusion	1000ml NaCl 0.9% over 2 hours	Every 21 days for 2 cycles
1-14	^{b,c,e} Capecitabine	1000mg/m ² twice daily	PO with food	n/a	Every 21 days for 2 cycles
CYCLE 3					
1-5	^{b,d,e} Capecitabine	825mg/m ² twice daily on each radiotherapy day only	PO with food		Day 1-5 week 1, 2, 3 4, 5 concurrently with radiation.
CYCLE 4 and 5					
1	^a CISplatin	60mg/m ²	IV Infusion	1000ml NaCl 0.9% over 2 hours	Every 21 days for 2 cycles
1-14	^{b,c,e} Capecitabine	1000mg/m ² twice daily	PO with food		Every 21 days for 2 cycles
<p>^a Pre and post hydration therapy required for CISplatin See local hospital policy recommendations. Suggested <u>prehydration</u> for CISplatin therapy: 1. Administer 10mmol magnesium sulphate (MgSO₄) (+/-KCl 20mmol/L if indicated) in 1000 mL sodium chloride 0.9% over 60 minutes. Administer CISplatin as described above <u>Post hydration</u>: Administer 1000 ml 0.9% NaCl over 60mins Mannitol 10% may be used to 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 (2, 3).</p>					
<p>^bThe dose to be administered should consider the available tablet strengths Reference to the NCCP DOSE BANDING TABLES for dosing of capecitabine Here Tablets should be swallowed whole with plenty of water within 30 minutes of eating. Tablets should not be crushed or cut. ^c(Total daily dose = 2000mg/m²) ^d(Total daily dose = 1650mg/m²)</p>					
^e See dose modifications section for patients with identified partial DPD deficiency					

NCCP Regimen: CISplatin and Capecitabine Chemoradiation Therapy	Published: 13/08/2018 Review: 15/07/2025	Version number:4
Tumour Group: Gastrointestinal NCCP Regimen Code: 00473	ISMO Contributor: Prof Maccon Keane	Page 1 of 6
<p>The information contained in this document is a statement of consensus of NCCP and ISMO or IHS professionals regarding their views of currently accepted approaches to treatment. Any clinician seeking to apply or consult these documents is expected to use independent medical judgement in the context of individual clinical circumstances to determine any patient's care or treatment. Use of these documents is the responsibility of the prescribing clinician, and is subject to HSE's terms of use available at http://www.hse.ie/eng/Disclaimer This information is valid only on the day of printing, for any updates please check www.hse.ie/NCCPchemoregimens</p>		

ELIGIBILITY:

- Indications as above
- ECOG 0-1
- Adequate hepatic, renal, and bone marrow function

EXCLUSIONS:

- Hypersensitivity to CISplatin, capecitabine or any of the excipients
- Moderate/severe renal impairment (creatinine clearance < 60 mL/min)
- Significant hearing impairment/tinnitus
- Known complete DPD deficiency
- History of severe and unexpected reactions to fluoropyrimidine therapy
- Pregnancy
- Breast Feeding

PRESCRIPTIVE AUTHORITY:

The treatment plan must be initiated by a Consultant Medical Oncologist

TESTS:

Baseline tests:

- FBC, renal and liver profile
- Audiology and creatinine clearance if clinically indicated
- INR tests if patient is on warfarin as clinically indicated
- DPD testing prior to first treatment with capecitabine using phenotype and/or genotype testing unless patient has been previously tested

Regular tests:

- FBC, renal and liver profile prior to each cycle
- INR tests if patient is on warfarin as clinically indicated

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:

- Consider a reduced starting dose in patients with identified partial DPD deficiency.
 - Initial dose reduction may impact the efficacy of treatment. In the absence of serious toxicity, subsequent doses may be increased with careful monitoring.
- Any dose modification should be discussed with a Consultant.

NCCP Regimen: CISplatin and Capecitabine Chemoradiation Therapy	Published: 13/08/2018 Review: 15/07/2025	Version number:4
Tumour Group: Gastrointestinal NCCP Regimen Code: 00473	ISMO Contributor: Prof Maccon Keane	Page 2 of 6

The information contained in this document is a statement of consensus of NCCP and ISMO or IHS professionals regarding their views of currently accepted approaches to treatment. Any clinician seeking to apply or consult these documents is expected to use independent medical judgement in the context of individual clinical circumstances to determine any patient's care or treatment. Use of these documents is the responsibility of the prescribing clinician, and is subject to HSE's terms of use available at <http://www.hse.ie/eng/Disclaimer>

This information is valid only on the day of printing, for any updates please check www.hse.ie/NCCPchemoregimens

Haematological:

Table 2: Dose modifications in haematological toxicity

ANC (x10 ⁹ /L)		Platelets (x10 ⁹ /L)	Dose
< 1.5	or	<75	Delay chemotherapy for 1 week

After 1 week of delay:

ANC (x10 ⁹ /L)		Platelets (x10 ⁹ /L)	Dose
≥ 1.5	and	≥ 75	100%
1 to <1.5	and	≥ 75	Reduce dose of capecitabine only by 25%
<1	or	<75	Delay for an additional week

After 2nd week of delay:

ANC (x10 ⁹ /L)		Platelets (x10 ⁹ /L)	Dose
≥ 1	and	≥ 75	Reduce dose of capecitabine only by 25%
<1	or	<75	Delay for an additional week

After 3rd week of delay:

ANC (x10 ⁹ /L)		Platelets (x10 ⁹ /L)	Dose
≥ 1	and	≥ 75	Reduce dose of capecitabine only by 50%
<1	or	<75	Omit further chemotherapy

Renal and Hepatic Impairment:

Table 3: Dose modifications of CISplatin and capecitabine in renal and hepatic impairment

Drug	Renal Impairment		Hepatic Impairment
CISplatin	Cr Cl (ml/min)	Dose	No dose modifications for hepatic impairment
	≥60	100%	
	45-59	75%	
	<45	Hold CISplatin or delay with additional IV fluids or go to CARBOplatin	
Capecitabine	≥30	100% dose	*In the absence of safety and efficacy data in patients with hepatic impairment, capecitabine use should be carefully monitored in patients with mild to moderate liver dysfunction, regardless of the presence or absence of liver metastasis.
	<30	Discontinue treatment	

*Reference Table 5 for dose modification of capecitabine in treatment related hepatotoxicity

Management of adverse events:

Table 4: Dose Modification for Adverse Events

Adverse reactions	Recommended dose modification
Nausea grade ≥ 3	Reduce dose of CISplatin by 25%
Non haematological toxicity Grade ≥ 2	Delay chemotherapy until symptoms resolved to Grade 1 or less
Hand –foot syndrome Grade 2	Reduce dose of capecitabine by 25%
Grade 3	Reduce dose of capecitabine by 50%

NCCP Regimen: CISplatin and Capecitabine Chemoradiation Therapy	Published: 13/08/2018 Review: 15/07/2025	Version number:4
Tumour Group: Gastrointestinal NCCP Regimen Code: 00473	ISMO Contributor: Prof Maccon Keane	Page 3 of 6

The information contained in this document is a statement of consensus of NCCP and ISMO or IHS professionals regarding their views of currently accepted approaches to treatment. Any clinician seeking to apply or consult these documents is expected to use independent medical judgement in the context of individual clinical circumstances to determine any patient's care or treatment. Use of these documents is the responsibility of the prescribing clinician, and is subject to HSE's terms of use available at <http://www.hse.ie/eng/Disclaimer>

This information is valid only on the day of printing, for any updates please check www.hse.ie/NCCPchemoregimens

Capecitabine Toxicity

Treatment related hepatotoxicity

Table 5: Dose modification of capecitabine in treatment related hepatotoxicity

Bilirubin		AST,ALT	Dose modification
> 3.0 x ULN	OR	> 2.5 x ULN	Withhold treatment until bilirubin decreases to ≤ 3.0 x ULN or ALT, AST decrease to ≤ 2.5 x ULN

Refer to NCCP regimen 00216 Capecitabine Monotherapy for detailed information on management of capecitabine related adverse events

SUPPORTIVE CARE:

EMETOGENIC POTENTIAL:

CISplatin: High (**refer to local policy**)

Capecitabine: Minimal to low (**refer to local policy**)

PREMEDICATIONS:

Hydration pre and post CISplatin administration (**Reference local policy or see recommendations above**).

OTHER SUPPORTIVE CARE: No specific recommendations

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.

CISplatin

- **Renal toxicity:** Renal toxicity is common with CISplatin. Encourage oral hydration.
- **Ototoxicity and sensory neural damage** should be assessed by history prior to each cycle.

Capecitabine

- **Diarrhoea and dehydration:** This may be dose limiting. Patients with severe diarrhoea should be carefully monitored and given fluid and electrolyte replacement if they become dehydrated.
- **Cardiotoxicity:** Angina-like chest pain, tachycardia, arrhythmias, heart failure, myocardial infarction and cardiac arrest may occur with capecitabine especially in patients with a prior history of coronary artery disease.
- **Dihydropyrimidine dehydrogenase (DPD) deficiency:** DPD is an enzyme encoded by the DPYD gene which is responsible for the breakdown of fluoropyrimidines. Patients with DPD deficiency are therefore at increased risk of fluoropyrimidine-related toxicity, including for example stomatitis, diarrhoea, mucosal inflammation, neutropenia and neurotoxicity. Treatment with 5-Fluorouracil, capecitabine or tegafur-containing medicinal products is contraindicated in patients with known complete DPD deficiency. Consider a reduced starting dose in patients with identified partial DPD deficiency. Initial dose reduction may impact the efficacy of treatment. In the absence of serious toxicity, subsequent doses may be increased with careful monitoring. Therapeutic drug monitoring (TDM) of fluorouracil may improve clinical outcomes in patients receiving continuous 5-fluorouracil infusions.

NCCP Regimen: CISplatin and Capecitabine Chemoradiation Therapy	Published: 13/08/2018 Review: 15/07/2025	Version number:4
Tumour Group: Gastrointestinal NCCP Regimen Code: 00473	ISMO Contributor: Prof Maccon Keane	Page 4 of 6

The information contained in this document is a statement of consensus of NCCP and ISMO or IHS professionals regarding their views of currently accepted approaches to treatment. Any clinician seeking to apply or consult these documents is expected to use independent medical judgement in the context of individual clinical circumstances to determine any patient's care or treatment. Use of these documents is the responsibility of the prescribing clinician, and is subject to HSE's terms of use available at <http://www.hse.ie/eng/Disclaimer>

This information is valid only on the day of printing, for any updates please check www.hse.ie/NCCPchemoregimens

- **Hand-foot syndrome (HFS)**, also known as palmar-plantar erythrodysesthesia (PPE), is a common side effect associated with capecitabine (see Table 4 for dose modification of capecitabine for HFS).

DRUG INTERACTIONS:

- Avoid concurrent use of CISplatin with nephrotoxic drugs (e.g. aminoglycosides, furosemide, NSAIDs) due to additive nephrotoxicity. If necessary, monitor renal function closely.
- Capecitabine enhances the anticoagulant effect of warfarin. Patients taking coumarin derivative anticoagulants should be monitored regularly for alterations in their coagulation parameters and the anti-coagulant dose adjusted accordingly.
- Sorivudine inhibits dihydropyrimidine dehydrogenase thus increasing its toxicity. Therefore, capecitabine must not be administered concomitantly with sorivudine or its chemically related analogues.
- Patients taking phenytoin or fosphenytoin concomitantly with capecitabine should be regularly monitored for increased phenytoin plasma concentrations.
- Current drug interaction databases should be consulted for more information.

ATC CODE:

CISplatin L01XA01
Capecitabine L01BC06

REFERENCES:

1. Lee J, Lim D, Kim S et al. Phase III trial comparing capecitabine plus cisplatin versus capecitabine plus cisplatin with concurrent capecitabine radiotherapy in completely resected gastric cancer with D2 lymph node dissection: The ARTIST Trial. J Clin Oncol 2012;30:268-273.
2. Nephrotoxicity Associated with CISplatin EviQ ID: 184 v.3
<https://www.eviq.org.au/clinical-resources/side-effect-and-toxicity-management/prophylaxis-and-prevention/184-nephrotoxicity-associated-with-CISplatin>
3. Portilla D et al. CISplatin nephrotoxicity. UptoDate Accessed Oct 2017
https://www.uptodate.com/contents/CISplatin-nephrotoxicity?source=search_result&search=CISplatin%20hydration&selectedTitle=1~150
4. Dosage Adjustment for Cytotoxics in Renal Impairment January 2009; North London Cancer Network . Available at <http://londoncancer.org/media/65600/renal-impairment-dosage-adjustment-for-cytotoxics.pdf>
5. NCCP Classification Document for Systemic Anti-Cancer Therapy (SACT) Induced Nausea and Vomiting. V2 2019. Available at: <https://www.hse.ie/eng/services/list/5/cancer/profinfo/chemoprotocols/nccp-classification-document-for-systemic-anti-cancer-therapy-sact-induced-nausea-and-vomiting.pdf>
6. **HPR**A Direct Healthcare Professional Communication. 5-Fluorouracil (i.v.), capecitabine and tegafur containing products: Pre-treatment testing to identify DPD-deficient patients at increased risk of severe toxicity. Accessed Aug 2020 Available at: [https://www.hpra.ie/docs/default-source/default-document-library/important-safety-information-from-marketing-authorisation-holders-of-products-containing-5-fluorouracil-\(i-v-\)-capecitabine-and-tegafur-as-approved-by-the-hpra.pdf?sfvrsn=0](https://www.hpra.ie/docs/default-source/default-document-library/important-safety-information-from-marketing-authorisation-holders-of-products-containing-5-fluorouracil-(i-v-)-capecitabine-and-tegafur-as-approved-by-the-hpra.pdf?sfvrsn=0)
7. Cisplatin 1mg/ml Concentrate for Solution for Infusion. Summary of Product Characteristics Accessed July 2020. Available at <https://www.hpra.ie/img/uploaded/swedocuments/Final%20approved%20SPC%20PA0822.199.001.pdf>
8. Xeloda® Summary of Product Characteristics Accessed July 2020 Available at https://www.ema.europa.eu/en/documents/product-information/xeloda-epar-product-information_en.pdf

NCCP Regimen: CISplatin and Capecitabine Chemoradiation Therapy	Published: 13/08/2018 Review: 15/07/2025	Version number:4
Tumour Group: Gastrointestinal NCCP Regimen Code: 00473	ISMO Contributor: Prof Maccon Keane	Page 5 of 6

The information contained in this document is a statement of consensus of NCCP and ISMO or IHS professionals regarding their views of currently accepted approaches to treatment. Any clinician seeking to apply or consult these documents is expected to use independent medical judgement in the context of individual clinical circumstances to determine any patient's care or treatment. Use of these documents is the responsibility of the prescribing clinician, and is subject to HSE's terms of use available at <http://www.hse.ie/eng/Disclaimer>

This information is valid only on the day of printing, for any updates please check www.hse.ie/NCCPchemoregimens

Version	Date	Amendment	Approved By
1	13/08/2018		Prof Maccon Keane
2	20/03/2020	Updated recommended dose modifications for capecitabine in renal impairment	Prof Maccon Keane
3	15/07/2020	Regimen review Updated emetogenic potential	Prof Maccon Keane
4	2/9/2020	Updated exclusion criteria, baseline testing, dose modifications and adverse events with respect to DPD deficiency as per DHPC from HPRA June 2020 Updated Adverse events regarding palmar-plantar erythrodysesthesia	Prof Maccon Keane

Comments and feedback welcome at oncologydrugs@cancercontrol.ie.

NCCP Regimen: CISplatin and Capecitabine Chemoradiation Therapy	Published: 13/08/2018 Review: 15/07/2025	Version number:4
Tumour Group: Gastrointestinal NCCP Regimen Code: 00473	ISMO Contributor: Prof Maccon Keane	Page 6 of 6

The information contained in this document is a statement of consensus of NCCP and ISMO or IHS professionals regarding their views of currently accepted approaches to treatment. Any clinician seeking to apply or consult these documents is expected to use independent medical judgement in the context of individual clinical circumstances to determine any patient's care or treatment. Use of these documents is the responsibility of the prescribing clinician, and is subject to HSE's terms of use available at <http://www.hse.ie/eng/Disclaimer>

This information is valid only on the day of printing, for any updates please check www.hse.ie/NCCPchemoregimens