

CARBOplatin (AUC 2) weekly and PACLitaxel 80mg/m² followed by Dose Dense DOXOrubicin cycloPHOSphamide Therapy - Triple Negative Breast Cancer Therapy

INDICATIONS FOR USE:

INDICATION	ICD10	Regimen Code	HSE approved reimbursement status*
Neoadjuvant treatment of triple negative breast carcinoma	C50	00734a	N/A

*This is for post 2012 indications only

TREATMENT:

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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.

CARBOplatin and PACLitaxel are administered on days 1, 8 and 15 of a 21 day cycle for 4 cycles or until disease progression or unacceptable toxicity develops.

This is then followed by DOXOrubicin and cycloPHOSphamide administered once every 14 days for 4 cycles (one cycle = 14 days).

G-CSF support (using standard or pegylated form) is required with all cycles of dose dense chemotherapy.

Facilities to treat anaphylaxis MUST be present when systemic anti-cancer therapy (SACT) is administered.

Admin. Order	Day	Drug	Dose	Route	Diluent & Rate	Cycle
1	1,8,15	PACLitaxel	80mg/m ²	IV infusion	250ml 0.9% NaCl over 60 minutes	Repeat every 21 days for cycles 1 - 4
2	1,8,15	CARBOplatin	AUC 2	IV infusion	250mL glucose 5% over 30 minutes	Repeat every 21 days for cycles 1 - 4
PACLitaxel must be supplied in non-PVC containers and administered using non-PVC giving sets and through an in-line 0.22 µm filter with a microporous membrane.						
PACLitaxe	l should be	diluted to a conce	ntration of 0.	3-1.2mg/mL.		

4 Cycles of PACLitaxel/CARBOplatin (Cycles 1-4 of treatment)

Note: Administration volumes and fluids have been standardised to facilitate electronic prescribing system builds.

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4 Cycles of DOXOrubicin/cycloPHOSphamide (Cycles 5-8 of treatment)

Admin. Order	Day	Drug	Dose	Route	Diluent & Rate	Cycle
1	1	DOXOrubicin ^a	60mg/ m ²	IV push	N/A	Repeat every 14 days for cycles 5 - 8
2	1	cycloPHOSphamide	600mg/m ²	IV infusion ^b	250mL 0.9% NaCl over 30 minutes	Repeat every 14 days for cycles 5 - 8
^a cycloPH0	OSpham	ide may also be administe	ered as an IV bo	lus over 5-10 min	utes.	
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^b 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 belowⁱ and to the age of the patient.

Note: Administration volumes and fluids have been standardised to facilitate electronic prescribing system builds.

CARBOplatin dose:

The dose in mg of CARBOplatin to be administered is calculated as follows:

Dose (mg) = target AUC (mg/mL x min) x (GFR mL/minute +25)

- Measured GFR (e.g. nuclear renogram) is preferred whenever feasible.
- Estimation of GFR (eGFR) can be done by using the Wright formula or using the Cockcroft and Gault formula to measure creatinine clearance.
- The GFR used to calculate the AUC dosing should not exceed 125mL/minute.
- For obese patients and those with a low serum creatinine, for example due to low body weight or post-operative asthenia, the formulae may not give accurate results and measured GFR is recommended.
 - Where obesity (body mass index $[BMI] \ge 30 \text{ kg/m}^2$) or overweight (BMI 25-29.9) is likely to lead to an overestimate of GFR and isotope GFR is not available, the use of the adjusted ideal body weight for Cockcroft and Gault may be considered.
 - Where serum creatinine is less than 63 micromol/L, the use of a creatinine value of 62 micromol/L or a steady pre-operative creatinine value may be considered.
- These comments do not substitute for the clinical judgement of a physician experienced in prescription of CARBOplatin.

WRIGHT FORMULA

There are two versions of the formula depending on how serum creatinine values are obtained, by the kinetic Jaffe method or the enzymatic method. The formula can be further adapted if covariant creatine kinase (CK) values are available (not shown).

1. SCr measured using enzymatic assay.

GFR (mL/minute) = <u>(6230 - 32.8 x Age) x BSA x (1 - 0.23 x Sex)</u> SCr (micromol/min)

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2. SCr measured using Jaffe assay

GFR (mL/minute) = <u>(6580 - 38.8 x Age) x BSA x (1 - 0.168 x Sex)</u> SCr (micromol/min)

Key: Sex = 1 if female, 0 if male; Age in years; BSA= DuBois BSA

COCKCROFT-GAULT FORMULA

GFR (mL/min) = <u>S x (140 - age in years) x wt (kg)</u> serum creatinine (micromol/L)

S = 1.04 for females and 1.23 for males

ELIGIBILITY:

- Indication as above
- ECOG status 0-2
- Adequate organ function; ANC > 1.5 x10⁹/L, platelets 75 x10⁹/L

EXCLUSIONS:

- Hypersensitivity to CARBOplatin*, PACLitaxel, DOXOrubicin, cycloPHOSphamide or any of the excipients
- Congestive heart failure (LVEF < 50%) or other significant heart disease
- Disease progression while receiving platinum based chemotherapy
- Pregnancy or lactation
- Severe hepatic impairment (PACLitaxel)
- Baseline neutrophil count < 1.5 x 10⁹/L

*If it is felt that the patient may have a major clinical benefit from CARBOplatin, it may in exceptional circumstances be feasible to rechallenge a patient with a prior mild hypersensitivity reaction e.g. using a desensitisation protocol, but only with immunology advice, premedication as advised, and a desensitisation protocol under carefully controlled conditions with resuscitation facilities available and medical and/or ITU/ HDU supervision

PRESCRIPTIVE AUTHORITY:

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

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TESTS:

Baseline tests:

- FBC, liver and kidney profile
- Audiometry and creatinine clearance as clinically indicated
- ECG
- MUGA, ECHO (LVEF > 50% to administer DOXOrubicin) if >65 years or dynamic cardiac monitoring (e.g. BNP) if clinically indicated
- Isotope GFR measurement (preferred) or GFR / CrCl estimation

Regular tests:

- FBC weekly during treatment
- Liver and kidney profiles weekly
- Assessment of peripheral neuropathy status as clinically indicated (PACLitaxel only)
- MUGA, ECHO 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:

• Any dose modification should be discussed with a Consultant.

Haematological:

Table 1: Dose modifications of PACLitaxel and CARBOplatin for haematological toxicity

ANC (x 10 ⁹ /L) Pretreatment blood test	
≥1.0	100% dose
0.5 to < 1.0	Delay treatment until recovery
< 0.5	Delay treatment until recovery and consider reducing PACLitaxel and
	CARBOplatin by 25% for subsequent cycles
Febrile neutropenia or previous delay for	Delay treatment until recovery and consider reducing PACLitaxel and
myelosuppression	CARBOplatin by 25% for subsequent cycles
Prolonged recovery greater than two weeks	Delay treatment until recovery, consider reducing PACLitaxel and
delay or 3rd delay for myelosuppression	CARBOplatin by 50% for subsequent cycles or cease
Platelets (x 10 ⁹ /L) Pretreatment blood test	
≥ 100	100% dose
75 to < 100	Clinician's discretion; continue treatment if patient is clinically well.
50 to < 75	Delay treatment until recovery
< 50	Delay treatment until recovery and consider reducing PACLitaxel and
	CARBOplatin by 25% for subsequent cycles

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ANC (x 10 ⁹ /L)		Platelet count (x 10 ⁹ /L)	Dose	
<u>></u> 1.0	and	<u>></u> 100	100%	
< 1.0	and	> 100	Delay for 1 week (or longer if needed), then give	
< 1.0	< 1.0 and		100% dose if ANC > 1.0 and platelets \geq 100.	
			Delay for 1 week (or longer if needed), then give	
<u>></u> 1.0	and	< 100	100% dose if ANC > 1.0 and platelets <u>></u> 100.	
			Dose reduce to 75% after a second delay.	

Table 2: Dose modification of DOXOrubicin and cycloPHOSphamide for haematological toxicity

Renal and Hepatic Impairment:

Table 3: Dose modification in renal and hepatic impairment

Drug	Renal Impairme	nt	Hepatic Impairment				
CARBOplatin ^a	See note below*		No dose mo	dificati	ion required		
PACLitaxel ^b	Renal impairment: no need for dose adjustment is expected.		ALT		Total Bilirubin	Dose	
	Haemodialysis: n	o need for dose	< 10xULN	and	≤ 1.25xULN	80mg/m ²	
	adjustment is ex	pected.	< 10xULN	and	1.26- 2xULN	60mg/m ²	
			< 10xULN	and	2.01- 5xULN	40mg/m ²	
			≥10xULN	and /or	>5xULN	Not recommended	
cycloPHOSphamide ^c	CrCl (mL/min)	Dose	Mild and mo	oderate	e: no need fo	or dose adjustment is	
	≥ 30	100%	expected.				
	10-29	Consider 75% of the original dose.	Severe: not recommended, due to risk of reduced				
	< 10	Not recommended. If unavoidable consider 50% of the original dose.	if				
	Haemodialysis	Not recommended. If unavoidable consider 50% of the original dose.					
DOXOrubicin ^d	CrCl (mL/min)	Dose	Serum Biliru	ıbin (m	nicromol/L)	Dose	
			20-50			50%	
	>10	No dose adjustment is needed.	> 51-86			25%	

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	<10	No need for dose adjustment is expected.	> 86 or Child Pugh C	Not recommended		
	Haemodialysis	75% of the original dose may be considered.				
^a Renal – see note below*, hepatic – Giraud et al 2023						

^{b,c,d} Renal and hepatic dose modifications from Giraud et al 2023

*Renal Dysfunction and CARBOplatin

- Patients with creatinine clearance values of <60mL/min are at greater risk to develop myelosuppression.
- In case of GFR ≤ 20mL/min CARBOplatin should not be administered at all.
- If Cockcroft & Gault or Wright formula are used, the dose should be adjusted as required per cycle based on a serum creatinine obtained within 48 hrs of drug administration.
- If isotope GFR is used, the dose should remain the same provided the serum creatinine is ≤110% of its value at the time of the isotope measurement. If the serum creatinine is higher than this, consideration should be given to re-measuring the GFR or to recalculating using Cockcroft & Gault or Wright formulae.

Management of adverse events:

Table 4: Dose Modification of PACLitaxel for Adverse Events

Adverse reactions	Recommended dose modification
Grade ≥2 motor or sensory neuropathy First Occurrence	Decrease dose of PACLitaxel by 10mg/m ² .
Persistent Grade ≥2 or 2 nd occurrence	Decrease dose of PACLitaxel by a further 10mg/m ²
All other grade 2 non-haematological	Hold treatment until toxicity resolves to ≤ grade 1.
toxicity	Decrease subsequent doses by 10mg/m ² .
≥ Grade 3 reaction	Discontinue

Patients who cannot tolerate treatment after 2 dose reductions or require a treatment delay of greater than 2 weeks, should discontinue treatment

SUPPORTIVE CARE:

EMETOGENIC POTENTIAL:

As outlined in NCCP Classification Document for Systemic Anti Cancer Therapy (SACT) Induced Nausea and Vomiting - <u>Available</u> on the NCCP website

CARBOplatin: Moderate (Refer to local policy) PACLitaxel: Low (Refer to local policy) cycloPHOSphamide/DOXOrubicin : High (Refer to local policy)

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For information:

Within NCIS regimens, antiemetics have been standardised by Medical Oncologists and Haemato-oncologists and information is available in the following documents:

- NCCP Supportive Care Antiemetic Medicines for Inclusion in NCIS (Medical Oncology) Available on the NCCP website
- NCCP Supportive Care Antiemetic Medicines for Inclusion in NCIS (Haemato-oncology) Available on the NCCP website

PREMEDICATIONS:

- All patients must be premedicated with corticosteroids, antihistamines, and H₂ antagonists prior to first dose of PACLitaxel treatment.
- The H₂ antagonist, famotidine, can potentially be omitted from the pre-medication requirements for PACLitaxel but the risk of hypersensitivity with this approach is unknown.
 - Caution is advised particularly for patients receiving PACLitaxel every 3 weeks. It is recommended that if famotidine is omitted that patients are monitored closely for any signs of hypersensitivity. Any hypersensitivity should be managed as per local policy.
 - \circ Where a patient experiences hypersensitivity, consider the use of alternative H₂ antagonists (Refer to local policy).

Table 5 outlines the suggested premedications prior to treatment with PACLitaxel.

Table 5. Suggested premedications prior to treatment with PACEItaker			
Day of treatment	Drug	Dose	Administration prior to PACLitaxel
Day 1	dexAMETHasone ^a	8mg IV	30 minutes
Day 1	Chlorphenamine	10mg IV	30 minutes
Day 1	Famotidine	20mg IV	30 minutes
Day 8 [♭] and thereafter	dexAMETHasone ^a	None	
Day 8 and thereafter	Chlorphenamine	10mg IV	30 minutes
Day 8 and thereafter	Famotidine ^c	20mg IV	30 minutes
^a Dose of dexAMETHasone may be altered, in the event of hypersensitivity reaction, to 20 mg of			
dexAMETHasone orally 12 hr and 6 hr prior to re-challenge with PACLitaxel according to consultant guidance.			
^b Dose of dexAMETHasone may be added from day 8 if increased risk or previous hypersensitivity reaction			
according to consultant guidance.			
Dose of famotidine may be omitted in the absence of hypersensitivity reaction according to consultant			
guidance.			

Table 5: Suggested premedications prior to treatment with PACLitaxel

OTHER SUPPORTIVE CARE:

- Myalgias and arthralgias may occur with PACLitaxel. Analgesic cover should be considered.
- Patients should have an increased fluid intake of 2-3 litres on day 1 to prevent haemorrhagic cystitis associated with cycloPHOSphamide.

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ADVERSE EFFECTS

• Please refer to the relevant Summary of Product Characteristics (SmPC) for details.

DRUG INTERACTIONS:

• Current SmPC and drug interaction databases should be consulted for information.

REFERENCES:

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- cycloPHOSphamide (Endoxana®) Injection 500mg Powder for Solution for Injection. Summary of Product Characteristics. Last updated 21/12/2018. Accessed September 2024. Available at: <u>https://www.hpra.ie/img/uploaded/swedocuments/Licence_PA2299-027-</u>001_21122018112107.pdf

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Version	Date	Amendment	Approved By
1	26/09/2023		Prof Maccon Keane
1a	27/05/2024	Amended title.	Prof Maccon Keane
2	09/12/2024	Reviewed.Updated exclusions section. Updated baseline and regular tests section. Updated emetogenic potential section. Updated other supportive care. Updated regimen in line with NCCP standardisation.	Prof Maccon Keane

Comments and feedback welcome at oncologydrugs@cancercontrol.ie.

ⁱ Cardiotoxicity 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

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.

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