

Regimen Monograph

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A - Regimen Name

CRBPNPAC(W) Regimen

CARBOplatin-Nab-PACLitaxel(Weekly)

Disease Site Lung
Non-Small Cell

Intent Palliative

Regimen Category **Evidence-informed :**

Regimen is considered appropriate as part of the standard care of patients; meaningfully improves outcomes (survival, quality of life), tolerability or costs compared to alternatives (recommended by the Disease Site Team and national consensus body e.g. pan-Canadian Oncology Drug Review, pCODR). Recommendation is based on an appropriately conducted phase III clinical trial relevant to the Canadian context OR (where phase III trials are not feasible) an appropriately sized phase II trial. Regimens where one or more drugs are not approved by Health Canada for any indication will be identified under Rationale and Use.

This **Regimen Abstract** is an **abbreviated** version of a Regimen Monograph and contains only top level information on usage, dosing, schedule, cycle length and special notes (if available). Information in regimen abstracts is accurate to the extent of the ST-QBP regimen master listings, and has not undergone the full review process of a regimen monograph. Full regimen monographs will be published for each ST-QBP regimen as they are developed.

Rationale and Uses For the treatment of advanced or metastatic non-small cell lung cancer, in patients who experienced hypersensitivity reactions to taxanes or have significant contraindications to taxanes and/or their pre-medications.

(Refer to the NDFP eligibility form for detailed funding criteria)

Supplementary [nab-PACLitaxel](#)**Public Funding** New Drug Funding Program (Nab-Paclitaxel - Hypersensitivity Reactions to Taxanes) ([NDFP Website](#))[back to top](#)**B - Drug Regimen****Nab-PACLitaxel is not-interchangeable with other PACLitaxel formulations.**

nab-PACLitaxel	100 mg /m ²	IV	Days 1, 8, 15
CARBOplatin	AUC 5-6*	IV	Day 1

*Adjust Carboplatin dose to AUC target (using Calvert formula) as outlined in "Other Notes" section.

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For a usual total of 4 to 6 cycles unless disease progression or unacceptable toxicity occurs

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D - Premedication and Supportive Measures

Antiemetic Regimen: Moderate + NK1 antagonist (Carboplatin AUC ≥ 5)

- Also refer to [CCO Antiemetic Recommendations](#).

Screen for hepatitis B virus in all cancer patients starting systemic treatment. Refer to the [hepatitis B virus screening and management](#) guideline.

Pre-medications (prophylaxis for infusion reaction):

Nab-paclitaxel:

- No pre-medication to prevent hypersensitivity is required.

Carboplatin:

- There is insufficient evidence that routine prophylaxis with pre-medications reduce infusion reaction (IR) rates.
- Corticosteroids and H1-receptor antagonists $\pm$ H2-receptor antagonists **may** reduce IR rates for some patients (e.g. gynecological patients with a platinum-free interval (PFI) > 12 months or a history of drug allergy who are receiving carboplatin starting from the 7th cycle) but no optimal pre-medication regimen has been established.

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J - Administrative Information

Pharmacy Workload (average time per visit) 36.007 minutes

Nursing Workload (average time per visit) 37.503 minutes

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K - References

CADTH Reimbursement Recommendation: Nab-Paclitaxel (for patients who developed hypersensitivity reactions to taxanes). July 2024.

Carboplatin drug monograph. Ontario Health (Cancer Care Ontario).

Nab-paclitaxel drug monograph. Ontario Health (Cancer Care Ontario).

Prescribing information: ABRAXANE® for Injectable Suspension (paclitaxel protein-bound particles for injectable suspension). Abraxis BioScience, LLC. October 2022.

Socinski MA, Bondarenko I, Karaseva NA, et al. Weekly nab-paclitaxel in combination with carboplatin versus solvent-based paclitaxel plus carboplatin as first-line therapy in patients with advanced non-small-cell lung cancer: final results of a phase III trial. J Clin Oncol. 2012 Jun 10;30(17):2055-62.

January 2026 new ST-QBP regimen

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L - Other Notes

Calvert Formula

DOSE (mg) = target AUC X (GFR + 25)

- AUC = product of serum concentration (mg/mL) and time (min)
- GFR (glomerular filtration rate) expressed as measured Creatinine Clearance or estimated from Serum Creatinine (by Cockcroft and Gault method or Jelliffe method)

(Calvert AH, Newell DR, Gumbrell LA, et al, Carboplatin dosage: Prospective evaluation of a simple formula based on renal function. J Clin Oncol, 1989; 7: 1748-1756)

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M - Disclaimer

Regimen Abstracts

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Regimen Monographs

Refer to the [New Drug Funding Program](#) or [Ontario Public Drug Programs](#) websites for the most up-to-date public funding information.

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