BC Cancer Protocol Summary for Palliative Treatment of Metastatic or Locally Advanced Gastric, Gastroesophageal Junction, or Esophageal Carcinoma using Oxaliplatin, Fluorouracil and Leucovorin

Protocol Code:

Tumour Group:

Contact Physician:

ELIGIBILITY:

Patients must have:

- Metastatic or locally advanced (unresectable) gastric, esophagogastric junction, or esophageal carcinoma, and
- No prior palliative chemotherapy, greater than 3 weeks from prior radiation therapy, or at the discretion of clinician, or
- Patients who have received single agent fluorouracil treatment first-line as the result of frailty, but who are now well enough to receive combination chemotherapy, or
- Patients who have progressed on single agent fluorouracil treatment first-line and treatment escalation/combination chemotherapy is desired

Patients should have:

- ECOG performance status 0 to 2
- Adequate marrow reserve, renal and liver function

EXCLUSIONS:

Patients must not have:

- Congenital long QT syndrome
- Severe pre-existing peripheral neuropathy

CAUTIONS:

- Patients with recent myocardial infarction, uncontrolled angina, hypertension, cardiac arrhythmias, congestive heart failure or other serious medical illness
- Patients with baseline greater than 3 loose BM per day (in patients without colostomy or ileostomy)
- Patients with symptomatic peripheral neuropathy

TESTS:

- Baseline CBC & Diff, creatinine, ALT, alkaline phosphatase, total bilirubin, albumin, sodium, potassium, <u>DPYD test</u> (not required if previously tested, or tolerated fluorouracil or capecitabine)
- Baseline if clinically indicated: CEA, CA 19-9, GGT, ECG
- Prior to each cycle: CBC & Diff, creatinine, total bilirubin, ALT
- If clinically indicated: CEA, CA 19-9, alkaline phosphatase, albumin, GGT, sodium, potassium, ECG

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GIGAVFFOX

Gastrointestinal

GI Systemic Therapy

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 For patients on warfarin, weekly INR during fluorouracil therapy until stable warfarin dose established, then INR prior to each cycle

PREMEDICATIONS:

- Antiemetic protocol for moderately emetogenic chemotherapy (see <u>SCNAUSEA</u>)
- If Grade 1 or 2 oxaliplatin hypersensitivity reactions:
 - 45 minutes prior to oxaliplatin:
 - o dexamethasone 20 mg IV in 50 mL NS over 15 minutes
 - 30 minutes prior to oxaliplatin:
 - diphenhydrAMINE 50 mg IV in NS 50 mL over 15 minutes and famotidine 20 mg IV in NS 100 mL over 15 minutes (Y-site compatible)
- Counsel patients to avoid cold drinks and exposure to cold air, especially for 3-5 days following oxaliplatin administration.
- Cryotherapy (ice chips) should NOT be used as may exacerbate oxaliplatininduced pharyngo-laryngeal dysesthesias.

TREATMENT:

A cycle equals:

| Drug | Dose | BC Cancer Administration Guidelines | |
|---------------------------|------------------------|--|--|
| oxaliplatin* | 85 mg/m ² | IV in 250 to 500 mL D5W over 2 hours** | |
| leucovorin [†] | 400 mg/m ² | IV in 250 mL D5W over 2 hours** | |
| fluorouracil [†] | 400 mg/m ² | IV push | |
| fluorouracil | 2400 mg/m ² | 1 | |

Repeat every 14 days until disease progression or unacceptable toxicity.

* Oxaliplatin is not compatible with normal saline. Do not piggyback or flush lines with normal saline.

** Oxaliplatin and leucovorin may be infused over the same two hour period by using a Y- site connector placed immediately before the injection site. Oxaliplatin and leucovorin should not be combined in the same infusion bag.

† fluorouracil IV push is optional in the advanced setting:

| fluorouracil IV push | leucovorin administration options | | |
|------------------------------|--|--|--|
| fluorouracil IV push given | leucovorin given as IV infusion OR leucovorin given as 20 mg/m² IV push | | |
| fluorouracil IV push omitted | leucovorin omitted OR leucovorin given as IV infusion OR leucovorin given as 20 mg/m² IV push | | |

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- *** Alternative administration:
- For 3000 to 5500 mg dose select INFUSOR per dose range below (doses outside dose banding range are prepared as ordered):

| Dose Banding Range | Dose Band INFUSOR (mg) | |
|----------------------|-------------------------------|--|
| Less than 3000 mg | Pharmacy to mix specific | |
| | dose | |
| 3000 to 3400 mg | 3200 mg | |
| 3401 to 3800 mg | 3600 mg | |
| 3801 to 4200 mg | 4000 mg | |
| 4201 to 4600 mg | 4400 mg | |
| 4601 to 5000 mg | 4800 mg | |
| 5001 to 5500 mg | 5250 mg | |
| Greater than 5500 mg | Pharmacy to mix specific dose | |

 Inpatients: 1200 mg/m²/day in 1000 mL D5W by continuous infusion daily over 23 h for 2 days

Patients with PICC lines should have a weekly assessment of the PICC site for evidence of infection or thrombosis.

DOSE MODIFICATIONS (A, B & C):

Fluorouracil Dosing Based on DPYD Activity Score (DPYD-AS)

Refer to "Fluorouracil and Capecitabine Dosing Based on DPYD Activity Score (DPYD-AS)" on www.bccancer.bc.ca/health-professionals/clinical-resources/cancer-drugmanual.

- A. Dose Modifications for NEUROLOGIC Toxicity
- B. Dose Modifications for HEMATOLOGIC Toxicity
- C. Dose Modifications for NON-HEMATOLOGIC, NON-NEUROLOGIC Toxicity

| • | | | | | | |
|-----------------------|--|------------------------|------------------------|--|--|--|
| Agent | Starting Dose | Dose Level -1 | Dose Level -2* | | | |
| oxaliplatin | 85 mg/m² | 65 mg/m² | 50 mg/m ² | | | |
| leucovorin | No dose modifications. If fluorouracil push is omitted, leucovorin may also be omitted or given as 20 mg/m² IV push If oxaliplatin is omitted, leucovorin may be given as 20 mg/m² IV push | | | | | |
| fluorouracil push | 400 mg/m ² | 320 mg/m ² | 200 mg/m ² | | | |
| fluorouracil infusion | 2400 mg/m ² | 2000 mg/m ² | 1600 mg/m ² | | | |

* For any additional dose reductions, use 20% less than previous level or consider discontinuing this regimen.

** The recommended starting doses are based on the modified FOLFOX6 regimen which is widely accepted but has not been studied in comparison to the original FOLFOX6 regimen. Patients may start with oxaliplatin 100 mg/m² as per FOLFOX6 at the discretion of their physician.

Table 2 - Oxaliplatin Neurotoxicity Definitions

| Grade 1 | Paresthesias / dysesthesias of short duration that resolve; do not interfere with function | | |
|---|---|--|--|
| Grade 2 | Paresthesias / dysesthesias interfering with function, but not activities of daily living (ADL) | | |
| Grade 3 | Paresthesias / dysesthesias with pain or with functional impairment which interfere with ADL | | |
| Grade 4 | Persistent paresthesias / dysesthesias that are disabling or life-threatening | | |
| Pharyngo-laryngeal dysesthesias (investigator discretion used for grading): | | | |
| Grade 0 | Grade 0 = none; Grade 1 = mild; Grade 2 = moderate; Grade 3 = severe | | |

Neuropathy may be partially or wholly reversible after discontinuation of therapy; patients with good recovery from Grade 3 (not Grade 4) neuropathy may be considered for re-challenge with oxaliplatin, with starting dose one level below that which they were receiving when neuropathy developed.

| Toxicity Grade | Duration | Persistent (present at start of next cycle) | |
|---|--|--|-----------------------|
| | 1 to 7 days | Greater than 7 days | |
| Grade 1 | Maintain dose level | Maintain dose level | Maintain dose level |
| Grade 2 | Maintain dose level | Maintain dose level | Decrease 1 dose level |
| Grade 3 | 1 st time: ↓ 1 dose level 2 nd time: ↓ 1 dose level | 1 st time: ↓ 1 dose level 2 nd time: ↓ 1 dose level | Discontinue |
| Grade 4 | Discontinue therapy | Discontinue therapy | Discontinue therapy |
| Pharyngo- laryngeal (see precautions) | Maintain dose level | N/A | N/A |

A. Dose Modifications for oxaliplatin NEUROLOGIC Toxicity

B. Dose Modifications for HEMATOLOGIC Toxicity

| | Prior to a Cycle (Day 1) | | Toxicity | Dose Level For Subsequent Cycles | |
|---|--|-------|------------------------------|-------------------------------------|---|
| | | Grade | ANC (x 10 ⁹ /L) | oxaliplatin | fluorouracil |
| • | 1 of cycle, hold treatment. Perform weekly CBC, maximum of 2 times. If ANC is greater than or equal to 1.2 within 2 weeks, proceed with treatment at the dose level noted across from the lowest ANC result of the delayed week(s). | 1 | Greater than or equal to 1.2 | Maintain dose level | Maintain dose level |
| | | 2 | 1.0 to less than 1.2 | Maintain dose level | Maintain dose level |
| • | | 3 | 0.5 to less than 1.0 | ↓ 1 dose level | Maintain dose level |
| • | | 4 | Less than 0.5 | ↓ 1 dose level | omit IV push and ↓ 1 infusion dose level |

| Prior to a Cycle (Day 1) | Toxicity | | Dose Level For Subsequent Cycles | |
|---|----------|-------------------------------------|-------------------------------------|------------------------|
| | Grade | Platelets (x 10 ⁹ /L) | oxaliplatin | fluorouracil |
| If platelets less than 75 on Day 1 of cycle, hold | 1 | Greater than or equal to 75 | Maintain dose level | Maintain dose level |
| treatment. Perform weekly CBC, maximum of 2 times.If platelets greater than or | 2 | 50 to less than 75 | Maintain dose level | Maintain dose level |
| equal to 75 within 2 weeks, proceed with treatment at the dose level noted across from | 3 | 10 to less than 50 | ↓ 1 dose level | Maintain dose level |
| the lowest platelets result of the delayed week(s). If platelets remain less than 75 after 2 weeks, discontinue treatment. | 4 | Less than 10 | ↓ 2 dose levels | Maintain dose level |

| Prior to a Cycle (Day 1) | | | Toxicity | Dose Level For Subsequent Cycles |
|--------------------------|---|-------|--|--|
| | | Grade | Diarrhea | |
| - | If diarrhea greater than or equal to Grade 2 on Day 1 of | 1 | Increase of 2 to 3 stools/day, or mild increase in loose watery colostomy output | Maintain dose level |
| | cycle, hold treatment. Perform weekly checks, maximum 2 times. | 2 | Increase of 4 to 6 stools, or nocturnal stools or mild increase in loose watery colostomy output | Maintain dose level |
| | If diarrhea is less than Grade 2 within 2 weeks, proceed with treatment at the | 3 | Increase of 7 to 9 stools/day or incontinence, malabsorption; or severe increase in loose watery colostomy output | |
| • | dose level noted across from the highest Grade experienced. If diarrhea remains greater than or equal to Grade 2 after 2 weeks, discontinue treatment. | 4 | Increase of 10 or more stools/day or grossly bloody colostomy output or loose watery colostomy output requiring parenteral support; dehydration | ↓ 1 dose level of oxaliplatin, IV push and infusional fluorouracil |
| | | Grade | Stomatitis | |
| • | If stomatitis greater than or equal to | 1 | Painless ulcers, erythema or mild soreness | Maintain dose level |
| | Grade 2 on Day 1 of cycle, hold | 2 | Painful erythema, edema, or ulcers but can eat | Maintain dose level |
| | treatment. Perform weekly checks, maximum 2 times. | 3 | Painful erythema, edema, ulcers, and cannot eat | Ψ 1 dose level of IV push and infusional fluorouracil |
| | If stomatitis is less than Grade 2 within 2 weeks, proceed with treatment at the dose level noted across from the highest Grade experienced. | 4 | As above but mucosal necrosis and/or requires enteral support, dehydration | ↓ 1 dose level of oxaliplatin, IV push and infusional fluorouracil |
| • | If stomatitis remains greater than or equal to Grade 2 after 2 weeks, discontinue treatment. | | | |

C. Dose Modifications for NON-HEMATOLOGIC, NON-NEUROLOGIC Toxicity

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PRECAUTIONS:

1. **Platinum hypersensitivity** can cause dyspnea, bronchospasm, itching and hypoxia. Appropriate treatment includes supplemental oxygen, steroids, epinephrine and bronchodilators. Vasopressors may be required (see table below). For Grade 1 or 2 acute hypersensitivity reactions no dose modification of oxaliplatin is required and the patient can continue treatment with standard hypersensitivity premedication. See Premedications.

Reducing infusion rates (e.g., from the usual 2 hours to 4-6 hours) should also be considered since some patients may develop more severe reactions when rechallenged, despite premedications.

The practice of rechallenging after severe life-threatening reactions is usually discouraged, although desensitization protocols have been successful in some patients. The benefit of continued treatment must be weighed against the risk of severe reactions recurring. The product monograph for oxaliplatin lists rechallenging patients with a history of severe HSR as a contraindication. Various desensitization protocols using different dilutions and premedications have been reported. Refer to SCOXRX: BC Cancer Inpatient Protocol Summary for Oxaliplatin Desensitization for more information.

2. Pharyngo-laryngeal dysesthesia is an unusual dysesthesia characterized by an uncomfortable persistent sensation in the area of the laryngopharynx without any objective evidence of respiratory distress (i.e. absence of hypoxia, laryngospasm or bronchospasm). This may be exacerbated by exposure to cold air or foods/fluids. If this occurs during infusion, stop infusion immediately and observe patient. Rapid resolution is typical, within minutes to a few hours. Check oxygen saturation; if normal, an anxiolytic agent may be given. The infusion can then be restarted at a slower rate at the physician's discretion. In subsequent cycles, the duration of infusion should be prolonged (see Dose Modifications above in the Neurological Toxicity table).

| Clinical Symptoms | Pharyngo-laryngeal Dysesthesia | Platinum Hypersensitivity |
|---------------------------|--|--|
| Dyspnea | Present | Present |
| Bronchospasm | Absent | Present |
| Laryngospasm | Absent | Present |
| Anxiety | Present | Present |
| O ₂ saturation | Normal | Decreased |
| Difficulty swallowing | Present (loss of sensation) | Absent |
| Pruritus | Absent | Present |
| Cold induced symptoms | Yes | No |
| Blood Pressure | Normal or Increased | Normal or Decreased |
| Treatment | Anxiolytics; observation in a controlled clinical setting until symptoms abate or at physician's discretion | Oxygen, steroids, epinephrine, bronchodilators; Fluids and vasopressors if appropriate |

- 3. QT prolongation and torsades de pointes are reported with oxaliplatin. Use caution in patients with history of QT prolongation or cardiac disease and those receiving concurrent therapy with other QT prolonging medications. Correct electrolyte disturbances prior to treatment and monitor periodically. Baseline and periodic ECG monitoring is suggested in patients with cardiac disease, arrhythmias, concurrent drugs known to cause QT prolongation, and electrolyte abnormalities. In case of QT prolongation, oxaliplatin treatment should be discontinued. QT effect of oxaliplatin with single dose ondansetron 8 mg prechemo has not been formally studied. However, single dose ondansetron 8 mg po would be considered a lower risk for QT prolongation than multiple or higher doses of ondansetron, as long as patient does not have other contributing factors as listed above.
- 4. **Neutropenia**: Fever or other evidence of infection must be assessed promptly and treated aggressively.
- 5. Oxaliplatin therapy should be interrupted if symptoms indicative of **pulmonary fibrosis** develop – nonproductive cough, dyspnea, crackles, rales, hypoxia, tachypnea or radiological pulmonary infiltrates. If pulmonary fibrosis is confirmed oxaliplatin should be discontinued.

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- 6. **Extravasation**: Oxaliplatin causes irritation if extravasated. Refer to BC Cancer <u>Extravasation Guidelines</u>.
- 7. Venous Occlusive Disease is a rare but serious complications that has been reported in patients (0.02%) receiving oxaliplatin in combination with fluorouracil. This condition can lead to hepatomegaly, splenomegaly, portal hypertension and/or esophageal varices. Patients should be instructed to report any jaundice, ascites or hematemesis immediately.
- 8. Oxaliplatin therapy should be interrupted if **Hemolytic Uremic Syndrome (HUS)** is suspected: hematocrit is less than 25%, platelets less than 100,000 and creatinine greater than or equal to 135 micromol/L. If HUS is confirmed, oxaliplatin should be permanently discontinued.
- 9. Myocardial ischemia and angina occurs rarely in patients receiving fluorouracil or capecitabine. Development of cardiac symptoms including signs suggestive of ischemia or of cardiac arrhythmia is an indication to discontinue treatment. If there is development of cardiac symptoms patients should have urgent cardiac assessment. Generally re-challenge with either fluorouracil or capecitabine is not recommended as symptoms potentially have a high likelihood of recurrence which can be severe or even fatal. Seeking opinion from cardiologists and oncologists with expert knowledge about fluorouracil / capecitabine toxicity is strongly advised under these circumstances. The toxicity should also be noted in the patient's allergy profile.
- 10. Diarrhea: Patients should report mild diarrhea that persists over 24 hours or moderate diarrhea (4 stools or more per day above normal, or a moderate increase in ostomy output). Mild diarrhea can be treated with loperamide (eg. IMODIUM®) following the manufacturer's directions or per the BC Cancer <u>Guidelines for Management of Chemotherapy-Induced Diarrhea</u>. Note that diarrhea may result in increased INR and the risk of bleeding in patients on warfarin.
- Dihydropyrimidine dehydrogenase (DPD) deficiency may result in severe and unexpected toxicity – stomatitis, diarrhea, neutropenia, neurotoxicity – secondary to reduced drug metabolism. This deficiency is thought to be present in about 3% of the population. Fluorouracil should be permanently discontinued in patients exhibiting exaggerated or prolonged neutropenia, mucositis, and diarrhea.
- 11. **Possible drug interaction with fluorouracil and warfarin** has been reported and may occur at any time. For patients on warfarin, weekly INR during fluorouracil therapy is recommended until a stable warfarin dose is established. Thereafter, INR prior to each cycle. Consultation to cardiology/internal medicine should be considered if difficulty in establishing a stable warfarin dose is encountered. Upon discontinuation of fluorouracil, repeat INR weekly for one month.
- 12. **Possible drug interaction with fluorouracil and phenytoin and fosphenytoin** has been reported and may occur at any time. Close monitoring is recommended. Fluorouracil may increase the serum concentration of these two agents.

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Call the GI Systemic Therapy physician at your regional cancer centre or the GI Systemic Therapy Chair Dr. Theresa Chan at (604) 930-2098 with any problems or questions regarding this treatment program.

References:

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