

# **DRUG NAME:** Topotecan

SYNONYM(S): Topotecan hydrochloride, NSC-609699

COMMON TRADE NAME(S): HYCAMTIN®

## CLASSIFICATION: Topoisomerase I inhibitor

Special pediatric considerations are noted when applicable, otherwise adult provisions apply.

## **MECHANISM OF ACTION:**

Topotecan is a semisynthetic, water-soluble derivative of camptothecin, which is a cytotoxic alkaloid extracted from plants such as *Camptotheca acuminata*. Topotecan has the same mechanism of action as irinotecan. It inhibits the action of topoisomerase I, an enzyme that produces reversible single-strand breaks in DNA during DNA replication. These single-strand breaks relieve torsional strain and allow DNA replication to proceed. Topotecan binds to the topoisomerase I-DNA complex and prevents re-ligation of the DNA strand, resulting in double strand DNA breakage and cell death.<sup>1</sup> Unlike irinotecan, topotecan is found predominantly in the inactive carboxylate form at neutral pH and it is not a prodrug. As a result, topotecan has different antitumour activities and toxicities from irinotecan.<sup>2</sup> Topotecan is a radiation-sensitizing agent.<sup>3</sup> It is cell cycle phase-specific (S-phase).<sup>4,5</sup>

Interpatient variability	large interpatient and intra	patient variability <sup>5,6</sup>	
Oral absorption		te is being studied in clinical trials <sup>7,8</sup>	
	time to peak plasma concentration	within 1-2 h <sup>7,8</sup>	
Distribution	evenly distributed between blood cells and plasma; extensively distributed into tissues <sup>4</sup>		
	cross blood brain barrier?	CSF to plasma ratio is 29% after a 24-hour infusion and 42% after a 72-hour infusion <sup>4</sup>	
	volume of distribution	130 L (reduced by 25% in patients with CrCl of 20-39 mL/min) <sup>1</sup>	
	plasma protein binding	35% <sup>1,9</sup>	
Metabolism	undergoes reversible, pH-dependent hydrolysis of the active lactone moiety to the inactive hydroxyacid (carboxylate) form (lactone form is present at pH ≤4 and the hydroxyacid form predominates at physiologic pH); relatively small amount of topotecan is metabolized by hepatic microsomal enzymes to an active metabolite, <i>N</i> -demethyltopotecan <sup>1,4,10</sup>		
	active metabolite(s)	lactone moiety and <i>N</i> -demethyltopotecan (clinical significance is unknown <sup>4</sup> )	
	inactive metabolite(s)	hydroxyacid form <sup>1</sup> and glucuronides of topotecan and <i>N</i> - demethyltopotecan <sup>11</sup>	
Excretion	biliary and renal excretion		
	bile	extent of biliary excretion not determined <sup>12</sup>	
	urine	20-60% of dose	
	terminal half life	2-3 h (increased to 5 h in patients with CrCl of 20-40 mL/min) <sup>1</sup>	
	clearance	1030 mL/min (decreased by 33% in patients with CrCl of 40-60 mL/min, by 66% with CrCl 20-40 mL/min); (decreased by 33% with bilirubin of 30-255 $\mu$ mol/L) <sup>1</sup>	
Gender	clearance 24% lower in fer	nales but no dosage adjustment required <sup>1,9</sup>	

## PHARMACOKINETICS:

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Elderly	no clinically significant difference in females; no information found on males
Children	clearance similar to adults when given as a 24-hour continuous infusion
Ethnicity	no information found

Adapted from reference<sup>9</sup> unless specified otherwise. Data pertained to 30 min IV infusion unless specified otherwise.

## USES:

Primary uses:	Other uses:
*Ovarian cancer <sup>13-15</sup>	*Lung cancer, small cell <sup>16-18</sup>
	Gliomas <sup>19</sup>
	Leukemia, acute myelogenous <sup>20,21</sup>
	Leukemia, chronic myelomonocytic <sup>22,23</sup>
	Lung cancer, non-small cell <sup>24</sup>
	Multiple myeloma <sup>25</sup>
	Myelodysplastic syndrome <sup>22,23,26</sup>
	Neuroblastoma <sup>27</sup>
	Pancreatic cancer <sup>28,29</sup>
	Retinoblastoma <sup>27</sup>
	Rhabdomyosarcoma <sup>27,30</sup>
	Sarcoma, Ewing's <sup>27</sup>

\*Health Canada approved indication

### **SPECIAL PRECAUTIONS:**

#### Caution:

 total clearance decreases by 57% with moderate renal impairment; avoid use in patients with severe renal dysfunction (CrCl <20 mL/min)<sup>31</sup>

*Carcinogenicity:* There is some evidence linking therapy with topoisomerase I inhibitors such as topotecan to the development of acute leukemias associated with specific chromosomal translocations. Long-term animal studies have not been done.<sup>1</sup>

*Mutagenicity:* Mutagenic in mammalian *in vitro* and *in vivo* mutation tests, but not mutagenic in bacterial *in vitro* mutation tests.<sup>1,9</sup>

#### Fertility: no information found

**Pregnancy:** Topotecan has been shown in animal studies to cause embryonic and fetal lethality at doses less than human clinical doses. Topotecan caused fetal malformations, primarily of the eye, brain, skull, and vertebrae when administered to pregnant test subjects.<sup>31</sup>

Breastfeeding is not recommended due to the potential secretion into breast milk.1,9

## SIDE EFFECTS:

The table includes adverse events that presented during drug treatment but may not necessarily have a causal relationship with the drug. Because clinical trials are conducted under very specific conditions, the adverse event rates observed may not reflect the rates observed in clinical practice. Adverse events are generally included if they



were reported in more than 1% of patients in the product monograph or pivotal trials, and/or determined to be clinically important.

ORGAN SITE	SIDE EFFECT
	Clinically important side effects are in <i>bold, italics</i>
blood/bone marrow	anemia (89%, severe 37%); nadir 15 days, recovery within 7 days <sup>1,14</sup>
febrile neutropenia	leukopenia (97%, severe 85%) <sup>13,14</sup>
	<i>neutropenia</i> (severe 95-97%) <sup>13,14</sup> ; nadir 12 days, recovery within 7 days <sup>1,14</sup>
	<i>thrombocytopenia</i> (69%, severe 50%) <sup>13,14</sup> ; nadir 15 days, recovery within 5 days <sup>1,14</sup>
	fever or infection with severe neutropenia (25-28%, severe 5%) <sup>13,16</sup>
constitutional symptoms	fatigue (29%, severe 5%)
	fever (28%, severe 1%) <sup>13</sup>
dermatology/skin	extravasation hazard: none4.32
	alopecia (49%)
	rash (16%, severe 1%)
gastrointestinal	emetogenic potential: moderate <sup>13,14,33</sup>
	anorexia (19%, severe 2%)
	constipation (29%, severe 3%)
	diarrhea (32%, severe 4%)
	nausea (64%, severe 8%)
	stomatitis (18%, severe 1%)
	vomiting (45%, severe 5%)
hepatic	bilirubin elevation (severe <2%)
	hepatic enzymes elevation (8%)
neurology	headache (18%, severe 1%)
	neuropathy-sensory (7%)
pain	abdominal pain (22%, severe 4%)
	arthralgia (6%, severe 1%) <sup>13</sup>
	myalgia (4%) <sup>13</sup>
	pain, includes body pain, back pain and skeletal pain (23%, severe 3%)
pulmonary	cough (15%, severe 1%)
	dyspnea (22%, severe 8%)
secondary malignancy	acute leukemias <sup>1</sup>

Adapted from reference<sup>9</sup> unless specified otherwise.



## INTERACTIONS:

AGENT	EFFECT	MECHANISM	MANAGEMENT
docetaxel <sup>34</sup>	administration of docetaxel on day 4 of topotecan therapy decreased docetaxel clearance by 50% and increased docetaxel toxicity	topotecan may alter docetaxel metabolism via CYP3A4 inhibition	administer docetaxel on day 1 of topotecan therapy
phenytoin <sup>10</sup>	increased topotecan clearance	possibly by inducing topotecan hepatic metabolism	may need to increase topotecan dose during concurrent therapy

## SUPPLY AND STORAGE:

### Injection:

Accord Healthcare Inc. supplies topotecan as the hydrochloride salt in 1 mg and 4 mg single dose (preservativefree) vials of ready-to-use solution in a concentration of 1 mg/mL (equivalent to 1 mg and 4 mg topotecan as the free base respectively). Store at room temperature. Protect from light in original packaging.<sup>31</sup>

Pfizer Canada ULC supplies topotecan as the hydrochloride salt in 4 mg single dose (preservative free) vials of ready-to-use solution in a concentration of 1 mg/mL (equivalent to 4 mg topotecan as the free base). Refrigerate. Protect from light in original packaging.<sup>35</sup>

Sandoz Canada Inc. supplies topotecan as the free base in 4 mg single dose (preservative free) vials of ready-touse solution in a concentration of 1 mg/mL. Refrigerate. Protect from light in original packaging.<sup>36</sup>

Teva Canada Limited supplies topotecan as the hydrochloride salt in 1 mg and 4 mg single use (preservative free) vials of lyophilized powder for reconstitution. Store at room temperature. Protect from light in original packaging.<sup>37</sup>

For basic information on the current brand used at BC Cancer, see <u>Chemotherapy Preparation and Stability</u> <u>Chart</u> in Appendix.

## SOLUTION PREPARATION AND COMPATIBILITY:

For basic information on the current brand used at BC Cancer, see <u>Chemotherapy Preparation and Stability</u> <u>Chart</u> in Appendix.

#### Additional information:

Compatibility: consult detailed reference

## PARENTERAL ADMINISTRATION:

BC Cancer administration guideline noted in **bold**, **italics** 

Subcutaneous	no information found
Intramuscular	no information found
Direct intravenous	no information found

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### BC Cancer administration guideline noted in *bold*, *italics*

Intermittent infusion	over 30 min
Continuous infusion	investigational, over 24 h <sup>22,38</sup>
Intraperitoneal	no information found
Intrapleural	no information found
Intrathecal	has been used <sup>39,40</sup>
Intra-arterial	no information found
Intravesical	no information found

## DOSAGE GUIDELINES:

Refer to protocol by which patient is being treated. Numerous dosing schedules exist and depend on disease, response and concomitant therapy. Guidelines for dosing also include consideration of absolute neutrophil count (ANC). Dosage may be reduced, delayed or discontinued in patients with bone marrow depression due to cytotoxic/radiation therapy or in patients with other toxicities.

## <u>Adults</u>:

			BC C	Cancer usual dose noted in <i>bold, italics</i>
	Cycle Length:			
Intravenous:	3 weeks:	<b>1.5 mg/m<sup>2</sup></b> (range 0.75-2 mg/m <sup>2</sup> ) <i>IV once daily for 5</i> <i>consecutive days starting on day 1</i> (total dose per cycle 7.5 mg/m <sup>2</sup> [range 3.75-10 mg/m <sup>2</sup> ]) <sup>9,13</sup>		rting on day 1
	3-4 weeks:	1.25 mg/m²/day (total dose per c		r 24 hours for 5 consecutive days 25 mg/m²) 20
	4-6 weeks:	consecutive day remission, then	vs starti every 4	2 mg/m²/day) IV over 24 hours for 5 ing on day 1 every 4-6 weeks until 4-8 weeks 0 mg/m² [range 5-10 mg/m²]) <sup>22,23</sup>
Dosage in myelosuppression:				tient is being treated; if no guidelines
	available, refer to	o Appendix "Dosa	ige Moo	dification for Myelosuppression"
Dosage in renal failure <sup>9</sup> :		o Appendix "Dosa earance (mL/min		dification for Myelosuppression" Dose
Dosage in renal failure <sup>9</sup> :	Creatinine cl			
Dosage in renal failure <sup>9</sup> :	Creatinine cl	earance (mL/min		Dose
Dosage in renal failure <sup>9</sup> :	Creatinine cl	earance (mL/min 40-60		<b>Dose</b> 1.5 mg/m² (100%)
Dosage in renal failure <sup>9</sup> :	Creatinine cl	earance (mL/min 40-60 20-39		Dose           1.5 mg/m² (100%)           0.75 mg/m² (50%)
Dosage in renal failure <sup>9</sup> :	Creatinine cl	earance (mL/min 40-60 20-39 < 20	)	Dose           1.5 mg/m² (100%)           0.75 mg/m² (50%)           not recommended           N* x (140 - Age) x weight in kg
Dosage in renal failure <sup>9</sup> : Dosage in hepatic failure:	Creatinine cl	earance (mL/min 40-60 20-39 < 20 tinine clearance .23; for females N	)     =  =1.04	Dose         1.5 mg/m² (100%)         0.75 mg/m² (50%)         not recommended         N* x (140 - Age) x weight in kg         serum creatinine in micromol/L

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## Children:

Cycle Length:

3 weeks:

Intravenous:

2 mg/m<sup>2</sup>/day (range 1.5-2 mg/m<sup>2</sup>/day) IV once daily for 5 consecutive days starting on day 1 (total dose per cycle 10 mg/m<sup>2</sup> [range 7.5-10 mg/m<sup>2</sup>])  $^{27}$ 

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