

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
Leucovorin 50 mg/5 mL 200 mg/20 mL 1000 mg/100 mL (GMP) (F)(PFL) no preservative ¹	N/A	10 mg/mL ¹	50 mg: discard unused portion ^{1,2} 200 mg, 1000 mg: 8 h F ^{1,2}	syringe	8 h RT ^{1,2}	
				0.05-10 mg/mL NS, D5W, Ringer's, LR, D10W, D5-NS ^{1,2} 50-250 mL†	NS , D5W, LR, Ringer's: 24 h RT ¹ D10W, D5-NS: 8 h RT ¹	
Leucovorin 50 mg/5 mL 500 mg/50 mL (Pfizer/Hospira) (F)(PFL) no preservative ³	N/A	10 mg/mL ³	8 h ³	syringe	8 h RT ³	
				0.05–10 mg/mL NS , D5W, LR, Ringer's, D10W, D5NS ³ 50-250 mL†	NS , D5W, LR, Ringer's: 24 h RT ³ D10W, D5NS: 8 h RT ³	

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Leucovorin 50 mg/5 mL 500 mg/50 mL (Teva) (F)(PFL) no preservative ⁴	N/A	10 mg/mL ⁵	discard unused portion ⁵	syringe	8 h ^{6,7}	
				0.4 - 4.8 mg/mL NS , D5W ⁸ 50-250 mL†	72 h F, RT ⁸	
				0.06 - 0.4 mg/mL NS , D5W ⁴ 50-250 mL†	NS: 24 h RT ⁴ D5W: 12 h RT ⁴	
				0.06 - 1 mg/mL Ringer's, Lactated Ringer's, D10W, D10-NS ⁴	Ringer's, LR: 24 h RT ⁴ D10W: 12 h RT ⁴ D10NS: 6 h RT ⁴	

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Lurbinectedin 4 mg (Jazz) (F) no preservative ⁹	8 mL SWI ⁹	0.5 mg/mL ⁹	12 h F , RT ^{9,10}	100-250 mL NS , D5W ⁹	complete administration within 24 h F , RT ⁹	- larger infusion volume is recommended for peripheral line ⁹ - do not use nylon membrane filters for administration if diluted in NS ⁹ ; BD Alaris pumps and syringe sets have polyethersulfone membrane in-line filters ¹¹
Lurbinectedin 4 mg (Pharma Mar) (F) no preservative ¹² (SAP)	8 mL SWI ¹²	0.5 mg/mL ¹²	12 h F , RT ^{10,12}	100–250 mL NS , D5W ¹²	30 h F , RT ¹²	- larger infusion volume is recommended for peripheral line ¹²

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Melphalan 50 mg (Marcan) (RT)(PFL) no preservative ¹³	10 mL supplied diluent ¹³ rapidly add diluent and immediately shake vigorously to dissolve ¹³ record time of reconstitution	5 mg/mL ¹³	2 h RT ¹³ do NOT refrigerate¹³	0.1-0.45 mg/mL NS only¹³	complete administration within 50 min RT from time of initial reconstitution ¹³	- will precipitate if stored in fridge ¹³
Melphalan 50 mg (Taro) (RT)(PFL) no preservative ¹⁴	10 mL supplied diluent ¹⁴ rapidly add diluent and immediately shake vigorously to dissolve ¹⁴ record time of reconstitution	5 mg/mL ¹⁴	2 h RT ¹⁴ do NOT refrigerate¹⁴	0.1-0.45 mg/mL NS only¹⁴	complete administration within 50 min RT from time of initial reconstitution ¹⁴	- will precipitate if stored in fridge ¹⁴

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Mesna 400 mg/4 mL 1000 mg/10 mL (Baxter) (RT) no preservative ¹⁵	N/A	100 mg/mL ¹⁵ (use filter needle to withdraw from ampoule)	discard unused portion ¹⁵	≥1 mg/mL NS , D5W, D5½-NS, LR ¹⁵⁻¹⁷ 100 mL†	24 h RT ¹⁵	
Mesna 1000 mg/10 mL 5000 mg/50 mL (Baxter) (RT) preservative ¹⁵	N/A	100 mg/mL ¹⁵	8 d RT ¹⁵ (vial may be punctured up to 4 times) ¹⁵	≥1 mg/mL NS , D5W, D5½-NS, LR ¹⁵⁻¹⁷ 100 mL†	24 h RT ¹⁵	
Mesna 1000 mg/10 mL (Fresenius Kabi) (RT) preservative ¹⁸	N/A	100 mg/mL ¹⁸	14 d F , RT ^{18,19}	≥1 mg/mL NS , D5W ²⁰ 100 mL†	48 h F, 24 h RT ¹⁸	

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Methotrexate 50 mg/2 mL 500 mg/20 mL 1 g/40 mL (Accord) (RT)(PFL) no preservative ²¹	N/A	25 mg/mL ²¹	50mg: discard unused portion ²¹ 500 mg, 1 g: 8 h RT ²¹	syringe	use within 8 h RT of initial puncture ²¹	- for high-dose regimens (e.g., 1-12 g/m ² as a single dose): use preservative-free methotrexate ²¹ - do not use for IT injection
				0.4–2 mg/mL NS , D5W ²¹ 50-500 mL†	use within 24 h RT of initial puncture ²¹ **(PFL)	
				high dose (e.g., 1-12 g/m ² as a single dose): 1000 mL * NS	use within 24 h RT of initial puncture ²¹ **(PFL)	
Methotrexate <u>intravitreal injection</u> 50 mg/2 mL (Accord) (RT)(PFL) no preservative ²¹	N/A	25 mg/mL ²¹	discard unused portion ²¹	syringe for intravitreal use	use within 4 h of initial puncture ¹⁰	- for intravitreal use preservative-free methotrexate is preferred ²²

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Methotrexate IT Injection Only preservative free methotrexate may be administered by the intrathecal route ²³ 50 mg/2 mL (Accord) (RT)(PFL) no preservative ²¹	N/A	25 mg/mL ²¹	discard unused portion ²¹	IT syringe qs to 6 mL with preservative free NS ^{24,25} diluents containing preservatives should NOT be used for intrathecal administration ²⁶	use within 4 h of initial puncture ¹⁰	- auxiliary info ¹⁰ : IT - label to include route in full (i.e., INTRATHECAL injection) attached to both syringe and outer Ziploc bag ²⁷
Methotrexate 50 mg/2 mL 500 mg/20 mL (Accord) (RT)(PFL) preservative ²¹	N/A	25 mg/mL ²¹	28 d F ^{10,21}	syringe	10 d F ^{10,21}	- contains benzyl alcohol ²¹ - do NOT use for high-dose regimens (e.g., 1-12 g/m ² as a single dose) ²¹ - do NOT use for IT injection ²¹
				0.4–2 mg/mL NS, D5W ²¹ 50-500 mL†	24 h RT ²¹	

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Methotrexate 50 mg/2 mL 500 mg/20 mL 1 g/40 mL 2.5 g/100 mL (Pfizer/Hospira) (RT)(PFL) no preservative ²⁸	N/A	25 mg/mL ²⁸	50mg: discard unused portion ²⁸ 500 mg, 1 g, or 2.5 g: 8 h RT ²⁸	syringe	use within 8 h RT of initial puncture ²⁸	- for high-dose regimens (e.g., 1-12 g/m ² as a single dose): use preservative-free methotrexate ²⁸ - do not use for IT injection
				0.4–2 mg/mL NS , D5W ²⁸ 50-500 mL†	use within 24 h RT of initial puncture ²⁸ **(PFL)	
				high dose (e.g., 1-12 g/m ² as a single dose): 1000 mL * NS	use within 24 h RT of initial puncture ²⁸ **(PFL)	
Methotrexate intravitreal injection 50 mg/2 mL (Pfizer/Hospira) (RT)(PFL) no preservative ²⁸	N/A	25 mg/mL ²⁸	discard unused portion ²⁸	syringe for intravitreal use	use within 4 h of initial puncture ¹⁰	- for intravitreal use preservative-free methotrexate is preferred ²²

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Methotrexate IT Injection Only preservative free methotrexate may be administered by the intrathecal route ²³ 50 mg/2 mL (Pfizer/Hospira) (RT)(PFL) no preservative ²⁸	N/A	25 mg/mL ²⁸	discard unused portion ²⁸	IT syringe qs to 6 mL with preservative free NS ^{24,25} diluents containing preservatives should NOT be used for intrathecal administration ²⁶	use within 4 h of initial puncture ¹⁰	- auxiliary info ¹⁰ : IT - label to include route in full (i.e., INTRATHECAL injection) attached to both syringe and outer Ziploc bag ²⁷
Methotrexate 50 mg/2 mL 500 mg/20 mL (Pfizer/Hospira) (RT)(PFL) preservative ²⁸	N/A	25 mg/mL ²⁸	28 d F ^{10,28}	syringe	10 d F ^{10,28}	- contains benzyl alcohol ²⁸ - do NOT use for high-dose regimens (e.g., 1-12 g/m ² as a single dose) ²⁸ - do NOT use for IT injection ²⁸
				0.4–2 mg/mL NS, D5W ²⁸ 50-500 mL†	24 h RT ²⁸	
Mitomycin 20 mg (Accord) (RT)(PFL) no preservative ²⁹	40 mL SWI ²⁹ shake well ²⁹	0.5 mg/mL ²⁹	12 h F, 6 h RT ^{10,30} **(PFL) ³⁰	syringe	72 h F, 6 h RT ³⁰ **(PFL) ³⁰	

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Mitomycin intravesical 20 mg (Accord) (RT)(PFL) no preservative ²⁹	40 mL SWI ²⁹ shake well ²⁹	0.5 mg/mL ²⁹	12 h F, 6 h RT ^{10,30} **(PFL) ³⁰	syringe	72 h F, 6 h RT ³⁰ **(PFL) ³⁰	
	10 mL SWI ³¹ shake well ²⁹	2 mg/mL ³¹	use immediately after preparation to prevent precipitation ³²	syringe	use immediately after preparation to prevent precipitation ³²	- may precipitate due to low solubility ^{32,33} - do NOT refrigerate ³²
	25 mL SWI shake well	0.8 mg/mL ³⁴	discard unused portion ^{2,34} **(PFL) ^{2,34}	syringe	4 days RT ³⁴ **(PFL) ^{2,34}	- do NOT refrigerate ³⁴
	33.3 mL SWI shake well	0.6 mg/mL ³⁴	discard unused portion ^{2,34} **(PFL) ^{2,34}	syringe	4 days F, RT ³⁴ **(PFL) ^{2,34}	
Mitomycin intraperitoneal 20 mg (Accord) (RT)(PFL) no preservative ²⁹	40 mL SWI ²⁹ shake well ²⁹	0.5 mg/mL ²⁹	12 h F, 6 h RT ^{10,30} **(PFL) ³⁰	0.02-0.04 mg/mL NS , sodium lactate ²⁹	NS: 18 h F, 3 h RT ³⁰ sodium lactate: 6 h F, 3 h RT ³⁰	

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Mitomycin 20 mg (Teva) (RT)(PFL) no preservative ³⁵	40 mL SWI ³⁵ shake well ³⁵	0.5 mg/mL ³⁵	12 h F, 6 h RT ^{10,35} **(PFL) ³⁵	syringe	72 h F, 6 h RT ³⁵ **(PFL) ³⁵	
Mitomycin intravesical 20 mg (Teva) (RT)(PFL) no preservative ³⁵	40 mL SWI ³⁵ shake well ³⁵	0.5 mg/mL ³⁵	12 h F, 6 h RT ^{10,35} **(PFL) ³⁵	syringe	72 h F, 6 h RT ³⁵ **(PFL) ³⁵	
	10 mL SWI ³¹ shake well ³⁵	2 mg/mL ³¹	use immediately after preparation to prevent precipitation ³²	syringe	use immediately after preparation to prevent precipitation ³²	- may precipitate due to low solubility ^{32,33} - do NOT refrigerate ³²
	25 mL SWI shake well	0.8 mg/mL ³⁴	discard unused portion ^{2,34} **(PFL) ^{2,34}	syringe	4 days RT ³⁴ **(PFL) ^{2,34}	- do NOT refrigerate ³⁴
	33.3 mL SWI shake well	0.6 mg/mL ³⁴	discard unused portion ^{2,34} **(PFL) ^{2,34}	syringe	4 days F, RT ³⁴ **(PFL) ^{2,34}	

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Mitomycin intraperitoneal 20 mg (Teva) (RT)(PFL) no preservative ³⁵	40 mL SWI ³⁵ shake well ³⁵	0.5 mg/mL ³⁵	12 h F, 6 h RT ^{10,35} **(PFL) ³⁵	0.02-0.04 mg/mL NS , sodium lactate ³⁵	NS: 18 h F, 6 h RT ³⁵ sodium lactate: 6 h F, RT ³⁵	
mitoXANTRONE 20 mg/10 mL (Fresenius Kabi) (RT) no preservative ³⁶	N/A	2 mg/mL ³⁶	discard unused portion ³⁶	0.2-0.6 mg/mL NS , D5W ³⁶ 50 mL†	24 h RT ³⁶	
mitoXANTRONE 20 mg/10 mL 25 mg/12.5 mL 30 mg/15 mL (Pfizer/Hospira) (RT)(PFL) no preservative ³⁷	N/A	2 mg/mL ³⁷	discard unused portion ³⁷	0.2-0.6 mg/mL NS , D5W ³⁷ 50 mL†	72 h F, 24 h RT ³⁷ **(PFL) ³⁷	
Mogamulizumab 20 mg/5 mL (Kyowa) (F)(PFL) do not shake no preservative ³⁸	N/A	4 mg/mL ³⁸	discard unused portion ³⁸	0.1-3 mg/mL NS 100 mL* mix by gentle inversion; do not shake ³⁸	24 h F ³⁸	- discard if cloudy, discoloured, or visible particulates are present ³⁸ - administer with 0.2 micron in-line filter ³⁸

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Mogamulizumab 20 mg/5 mL (Kyowa) (F)(PFL) do not shake no preservative ³⁹ (SAP)	N/A	4 mg/mL ³⁹	discard unused portion ³⁹	0.1-3 mg/mL NS ³⁹ 100 mL* mix by gentle inversion; do not shake ³⁹	24 h F ³⁹	- discard if cloudy or discoloured ³⁹ - administer with 0.2 micron in-line filter ³⁹
Nelarabine 250 mg/50 mL (Sandoz) (RT) no preservative ⁴⁰	N/A	5 mg/mL ⁴⁰	discard unused portion ⁴⁰	undiluted in empty PVC infusion bag or glass container ⁴⁰	8 h RT ⁴⁰	- discard if discoloured, hazy, or particulates are present ⁴⁰
Nivolumab 40 mg/4 mL 100 mg/10 mL (BMS) (F)(PFL) do not shake no preservative ⁴¹	N/A	10 mg/mL ⁴¹	discard unused portion ⁴¹	1-10 mg/mL NS, D5W ⁴¹ 25-100 mL† mix by gentle inversion; do not shake ⁴¹ OR undiluted in empty infusion bag or glass bottle ⁴¹	complete administration within 7 days F, including max 8 h at RT ⁴¹ **(PFL) ⁴¹ (can be in room light when at RT) ⁴¹	- do not shake ⁴¹ - administer with 0.2 micron in-line filter ⁴¹ - may contain a few amorphous particles ⁴¹ - discard if cloudy, has pronounced colour change (should be clear to pale yellow) ⁴¹

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oBINutuzumab 1000 mg/40 mL (Roche) (F)(PFL)** do not shake no preservative ⁴²	N/A	25 mg/mL ⁴²	discard unused portion ⁴³	NS 100 mg: 100 mL ⁴² 900 mg: 250 mL ⁴² 1000 mg: 250 mL ⁴²	24 h F, 48 h RT ^{42,44}	-once removed from the fridge, diluted product is stable for an additional 48 h RT ^{42,44} - do NOT shake ⁴² - do NOT use dextrose containing solutions ⁴²
Octreotide 50 mcg/1 mL 100 mcg/1 mL 500 mcg/1 mL (Omega) (F)(PFL) no preservative ⁴⁵	N/A	50 mcg/mL ⁴⁵	discard unused portion ⁴⁵	NS⁴⁵ volume adjusted to ensure a continuous infusion of octreotide at 25 mcg/h ⁴⁵	24 h RT ⁴⁵	
		100 mcg/mL ⁴⁵				
		500 mcg/mL ⁴⁵				
Octreotide multidose vial: 1000 mcg/5 mL (Omega) (F)(PFL) preservative ⁴⁵	N/A	200 mcg/mL ⁴⁵	15 d F ⁴⁵	NS⁴⁵ volume adjusted to ensure a continuous infusion of octreotide at 25 mcg/h ⁴⁵	24 h RT ⁴⁵	

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Octreotide (SANDOSTATIN®) 50 mcg/1 mL 100 mcg/1 mL 500 mcg/1 mL (Novartis) (F)(PFL) no preservative ⁴⁶	N/A	50 mcg/mL ⁴⁶	discard unused portion ⁴⁶	NS ⁴⁶ volume adjusted to ensure a continuous infusion of octreotide at 25 mcg/h ⁴⁶	24 h RT ⁴⁶	
		100 mcg/mL ⁴⁶				
		500 mcg/mL ⁴⁶				
Octreotide (SANDOSTATIN®) multi-dose vial: 1000 mcg/5 mL (Novartis) (F)(PFL) preservative ⁴⁶	N/A	200 mcg/mL ⁴⁶	14 d F, RT ⁴⁶	NS ⁴⁶ volume adjusted to ensure a continuous infusion of octreotide at 25 mcg/h ⁴⁶	24 h RT ⁴⁶	

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Octreotide (SANDOSTATIN LAR®) (long acting) 10 mg 20 mg 30 mg (Novartis) (F)(PFL) no preservative ⁴⁶	2 mL supplied diluent ⁴⁶ add diluent: gently run diluent down sides of vial ⁴⁶ do NOT disturb for 2–5 min; then swirl moderately ⁴⁶ record time of reconstitution	10 mg: 5 mg/mL ⁴⁶	discard unused portion ⁴⁶	syringe (for deep intragluteal administration only) ⁴⁶	use within 4 h of initial reconstitution ^{10,46}	- do NOT shake ⁴⁶
		20 mg: 10 mg/mL ⁴⁶				
		30 mg: 15 mg/mL ⁴⁶				

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Octreotide suspension (long acting) 10 mg 20 mg 30 mg (Teva) (F)(PFL) no preservative ⁴⁷	2 mL supplied diluent	10 mg: 5 mg/mL ⁴⁷	discard unused portion ⁴⁷	syringe (for deep intragluteal administration only) ⁴⁷	use within 4 h of initial reconstitution ^{10,47}	- gently shake to resuspend before administration ⁴⁷ - delay in administration may result in sedimentation ⁴⁷
	let stand at RT for 30 min prior to reconstitution ⁴⁷	20 mg: 10 mg/mL ⁴⁷				
	add supplied diluent ⁴⁷ let vial stand for 5 min after adding diluent to saturate powder ⁴⁷ shake moderately in horizontal direction for ≥30 sec to create suspension ⁴⁷ record time of reconstitution	30 mg: 15 mg/mL ⁴⁷				

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Oxaliplatin 50 mg/10 mL 100 mg/20 mL 200 mg/40 mL (Dr. Reddy's) (RT)(PFL) no preservative ⁴⁸	N/A	5 mg/mL ⁴⁸	discard unused portion ⁴⁸	0.2-0.7 mg/mL D5W ⁴⁸ 100-500 mL† do NOT use NS or other chloride- containing solution ⁴⁸ do NOT use aluminum-containing needle and syringe ⁴⁸	0.2-2 mg/mL: 48 h F, 24 h RT ⁴⁸	- do NOT use aluminum- containing needle, syringe, or tubing ⁴⁸
Oxaliplatin 50 mg/10 mL 100 mg/20 mL 200 mg/40 mL (Pfizer/Hospira) (RT) no preservative ⁴⁹	N/A	5 mg/mL ⁴⁹	discard unused portion ⁴⁹	0.2-0.7 mg/mL D5W ⁴⁹ 100-500 mL† do NOT use NS or other chloride- containing solutions ⁴⁹ do NOT use aluminum-containing needle and syringe ⁴⁹	0.2-0.4 mg/mL: 24 h RT ⁴⁹ or 5 d F plus an additional 8 h RT ⁵⁰ 0.5–2 mg/mL: 24 h RT ⁴⁹ or 10 d F, plus an additional 8 h RT ^{10,50} *(PFL) when stored in F ⁵⁰	- do NOT use aluminum- containing needle, syringe, tubing ⁴⁹

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Oxaliplatin 50 mg/10 mL 100 mg/20 mL 150 mg/30 mL 200 mg/40 mL (Sandoz) (RT)(PFL) no preservative ⁵¹	N/A	5 mg/mL ⁵¹	12 h F, RT ^{10,52}	0.2-0.7 mg/mL D5W ⁵¹ 100-500 mL† do NOT use NS or other chloride- containing solution ⁵¹ do NOT use aluminum-containing needle and syringe ⁵¹	0.2-2 mg/mL: 48 h F, 24 h RT ⁵¹	- do NOT use aluminum- containing needle, syringe, tubing ⁵¹
Oxaliplatin 50 mg/10 mL 100 mg/20 mL 200 mg/40 mL (Teva) (RT)(PFL) no preservative ⁵³	N/A	5 mg/mL ⁵³	discard unused portion ⁵³	0.2-0.7 mg/mL D5W ⁵³ 100-500 mL† do NOT use NS or other chloride- containing solution ⁵³ do NOT use aluminum-containing needle and syringe ⁵³	0.2-2 mg/mL: 48 h F, 24 h RT ⁵³	- do NOT use aluminum- containing needle, syringe or tubing ⁵³

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PACLitaxel 30 mg/5 mL 100 mg/16.7 mL 300 mg/50 mL (Accord) (RT)(PFL) preservative ⁵⁴	N/A	6 mg/mL ⁵⁴	30 mg, 100 mg: 28 d RT ^{10,54} 300 mg: 24 h RT ^{10,54}	0.3-1.2 mg/mL NS , D5W, D5NS, D5LR ⁵⁴ 50-500 mL†	complete administration within 27 h RT ⁵⁴	- use non-DEHP bag and tubing ⁵⁴ - administer with 0.2 micron in-line filter ⁵⁴ - avoid excessive shaking ⁵⁴
				0.1 mg/mL NS ⁵⁵	44 h F , RT ⁵⁵	
PACLitaxel 30 mg/5 mL 100 mg/16.7 mL 300 mg/50 mL (Biolyse) (RT) preservative ⁵⁶	N/A	6 mg/mL ⁵⁶	28 d RT ⁵⁷	0.3-1.2 mg/mL NS , D5W ⁵⁶ 50-500 mL†	complete administration within 27 h RT ^{58,59}	- use non-DEHP bag and tubing ⁵⁶ - administer with 0.2 micron in-line filter ⁵⁶
				0.1 mg/mL NS ⁵⁵	44 h F , RT ⁵⁵	
				0.012-0.12 mg/mL NS ⁶⁰	16 h RT ⁵⁸	
				devices with spikes (e.g., chemo dispensing pins) may be used with vials ⁶¹		

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Paclitaxel 30 mg/5 mL 100 mg/16.7 mL 300 mg/50 mL (Sandoz) (RT)(PFL) preservative ⁶²	N/A	6 mg/mL ⁶²	30 mg, 100 mg: 28 d RT ^{10,62} 300 mg: 24 h RT ^{10,62}	0.3-1.2 mg/mL NS , D5W, D5NS ⁶² 50-500 mL†*	complete administration within 27 h RT ⁶²	- use non-DEHP bag and tubing ⁶² - administer with 0.2 micron inline filter ⁶² - avoid excessive shaking
				0.1 mg/mL NS ⁵⁵	44 h F , RT ⁵⁵	
PACLitaxel, nanoparticle, albumin- bound (NAB) 100 mg (Celgene) (RT)(PFL) no preservative ⁶³	20 mL NS ⁶³ slowly direct diluent against side of vial (i.e., ≥1 min) during reconstitution ⁶³ let stand for ≥5 min to wet powder ⁶³ gently swirl or invert for ≥2 min ⁶³	5 mg/mL ⁶³	use immediately (RT) or 8 h F ⁶³ **(PFL) ⁶³	undiluted in empty PVC, non-PVC, or non-DEHP infusion bag ⁶³	48 h F plus an additional 8 h RT ⁶⁴	- each vial contains 900 mg human albumin ⁶³ - to prevent foaming, do NOT inject NS directly onto the powder ⁶³ - some settling may occur; use mild agitation to resuspend ⁶³ - administer with 15 micron filter ONLY ⁶³ (NOTE: filters with pore size less than 15 microns may cause filter blockage) ⁶⁵

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
PACLitaxel, nanoparticle, albumin- bound (NAB) 100 mg (Panacea/Apo) (RT)(PFL) no preservative ⁶⁶	20 mL NS ⁶⁶ slowly direct diluent against side of vial (i.e., ≥1 min) during reconstitution ⁶⁶ let stand for ≥5 min to wet powder ⁶⁶ gently swirl or invert for ≥2 min ⁶⁶ (if foaming occurs, let stand for ≥15 min) ⁶⁶	5 mg/mL ⁶⁶	use immediately (RT) or 8 h F ⁶⁶ **(PFL) ⁶⁶	undiluted in empty PVC, non-PVC, or non-DEHP infusion bag ⁶⁶	56 h F plus an additional 4 h RT ⁶⁷	- each vial contains 900 mg human albumin ⁶⁶ - to prevent foaming, do NOT inject NS directly onto the powder ⁶⁶ - some settling may occur; use gentle inversion to resuspend ⁶⁶ - discard if visible particulates are present ⁶⁶ - administer with 15 micron filter ONLY ⁶⁶
Pamidronate 30 mg/10 mL 60 mg/10 mL 90 mg/10 mL (Fresenius Kabi) (RT) no preservative ⁶⁸	N/A	3 mg/mL ⁶⁸	discard unused portion ⁶⁸	≤0.36 mg/mL ⁶⁸ NS, D5W⁶⁸ 250 mL†	24 h RT ⁶⁸	- do NOT mix with calcium containing solutions (e.g., Lactated Ringer's) ⁶⁸
		6 mg/mL ⁶⁸				
		9 mg/mL ⁶⁸				

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
Pamidronate 30 mg/10 mL 60 mg/10 mL 90 mg/10 mL (Hospira) (RT) no preservative ⁶⁹	N/A	3 mg/mL ⁶⁹	discard unused portion ⁶⁹	0.06–0.36 mg/mL NS , D5W ⁶⁹ 250 mL†	24 h F plus an additional 24 h RT (total 48 h) ⁶⁹ **(PFL) ⁶⁹	- do NOT mix with calcium containing solution (e.g., Lacated Ringer's) ⁶⁹
		6 mg/mL ⁶⁹				
		9 mg/mL ⁶⁹				
Pamidronate 30 mg/10 mL 60 mg/10 mL 90 mg/10 mL (Omega) (RT) no preservative ⁷⁰	N/A ⁷⁰	3 mg/mL ⁷⁰	discard unused portion ⁷⁰	0.06–0.36 mg/mL NS , D5W ⁷⁰ 250 mL†	24 h F plus an additional 24 h RT (total 48 h) ⁷⁰ **(PFL) ⁷⁰	- do NOT mix with calcium containing solution (e.g., Lactated Ringer's) ⁷⁰
		6 mg/mL ⁷⁰				
		9 mg/mL ⁷⁰				
Pamidronate 30 mg/10 mL 60 mg/10 mL 90 mg/10 mL (Pfizer) (RT) no preservative ⁷¹	N/A	3 mg/mL ⁷¹	discard unused portion ⁷¹	0.06-0.36 mg/mL NS , D5W ⁷¹ 250 mL†	24 h F plus an additional 24 h RT (total 48 h) ⁷¹ **(PFL) ⁷¹	- do NOT mix with calcium containing solution (e.g., Lactated Ringer's) ⁷¹
		6 mg/mL ⁷¹				
		9 mg/mL ⁷¹				

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
Pamidronate 30 mg/10 mL 60mg/10 mL 90 mg/10 mL (Sandoz Canada) RT no preservative ⁷²	N/A	3 mg/mL ⁷²	discard unused portion ^{72,73}	NS ; D5W ⁷² 250 mL†	24 h RT ⁷²	- do NOT mix with calcium containing solution (e.g., Lactated Ringer's) ⁷²
		6 mg/mL ⁷²				
		9 mg/mL ⁷²				
PANitumumab 100 mg/5 mL 400 mg/20 mL (Amgen) (F)(PFL) do not shake no preservative ⁷⁴	N/A	20 mg/mL ⁷⁴	discard unused portion ⁷⁴	1-10mg/mL NS ⁷⁴ 100 mL†	24 h F, 6 h RT ⁷⁴⁻⁷⁷	- administer with 0.2 micron in-line filter ⁷⁴ - solution may contain particulates which do not affect product quality ⁷⁴ - do not administer if discoloured ⁷⁴

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
Pegaspargase (pegylated asparaginase <i>E. coli</i>) 3750 units/5 mL (Servier) (F)(PFL) do not shake no preservative ⁷⁸	N/A	750 units/mL ⁷⁸	discard unused portion ⁷⁸	IM ⁷⁸ : max volume: 2 mL in children and adolescents; 3 mL in adults if volume greater than above, use multiple sites ⁷⁸	syringe: use within 4 h of vial puncture ^{2,78}	- do NOT shake ⁷⁸
				IV ⁷⁸ : 100 mL NS , D5W	bag: use within 4 h of vial puncture ^{2,78}	
Pembrolizumab 100 mg/4 mL (Merck) (F)(PFL) do not shake no preservatives ⁷⁹	N/A	25 mg/mL ⁷⁹	discard unused portion ^{2,79}	1-10 mg/mL NS , D5W ⁷⁹ 50 mL * mix by gentle inversion ⁷⁹	complete administration within 96 h F, 6 h RT ⁷⁹	- administer with 0.2 micron in-line filter ⁷⁹ - bring vials and diluted solutions to RT prior to use ⁷⁹ - vials contain 0.25 mL overflow ⁷⁹

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
Pemetrexed 100 mg 500 mg (Accord) (RT) no preservative ⁸⁰	100 mg: 4.2 mL NS ⁸⁰ 500 mg: 20 mL NS ⁸⁰	25 mg/mL ⁸⁰	12 h F, RT ^{10,80}	100 mL NS ⁸⁰	24 h F, RT ⁸⁰	- do NOT mix with calcium containing solution (e.g., Ringer's) ⁸⁰
Pemetrexed 100 mg/4 mL 500 mg/20 mL 850 mg/34 mL 1000 mg/40 mL (Accord) (RT)(PFL) no preservative ⁸¹	N/A	25 mg/mL ⁸¹	discard unused portion ⁸¹	100 mL NS ⁸¹	24 h F ⁸¹	- do NOT mix with calcium containing solution (e.g., Ringer's) ⁸¹
Pemetrexed 100 mg 500 mg (Dr. Reddy's) (RT) no preservative ⁸²	100 mg: 4.2 mL NS ⁸² 500 mg: 20 mL NS ⁸²	25 mg/mL ⁸²	12 h F, RT ^{10,83-85}	100 mL NS ⁸²	24 h F, RT ⁸³⁻⁸⁵	- do NOT mix with calcium containing solution (e.g., Ringer's) ⁸²
Pemetrexed 100 mg 500 mg (Lilly) (RT) no preservative ⁸⁶	100 mg: 4.2 mL NS ⁸⁶ 500 mg: 20 mL NS ⁸⁶	25 mg/mL ⁸⁶	12 h F ^{10,86}	100 mL NS ⁸⁶	24 h F ⁸⁶	- do NOT mix with calcium containing solution (e.g., Ringer's) ⁸⁶

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
Pemetrexed 100 mg 500 mg 1000 mg (Taro) (RT) no preservative ⁸⁷	100 mg: 4.2 mL NS ⁸⁷ 500 mg: 20 mL NS ⁸⁷ 1000 mg: 40 mL NS ⁸⁷	25 mg/mL ⁸⁷	12 h F ^{10,87}	100 mL NS ⁸⁷	24 h F ⁸⁷	- do NOT mix with calcium containing solution (e.g., Ringer's) ⁸⁷
Pentostatin 10 mg (Hospira/Pfizer) (F) no preservative ⁸⁸	5 mL SWI ⁸⁸	2 mg/mL ⁸⁸	8 h RT ⁸⁸	0.18-0.33 mg/mL ⁸⁸ 25-50 mL NS, D5W ⁸⁸	8 h RT ⁸⁸	
PERTuzumab 420 mg/14 mL (Roche) (F)(PFL) no preservative ⁸⁹	N/A	30 mg/mL ⁸⁹ do NOT shake ⁸⁹	discard unused portion ^{43,89}	250 mL NS only ⁸⁹ mix by gentle inversion to avoid foaming ⁸⁹	24 h F, RT ⁸⁹	- do NOT use dextrose containing solutions ⁸⁹

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
<p>PERTuzumab-trastuzumab 1200 mg-600 mg/15 mL 600 mg-600 mg/10 mL (Roche) (F)(PFL) do not shake no preservative⁹⁰</p>	<p>N/A</p>	<p>1200 mg-600 mg⁹⁰: 80 mg/mL pertuzumab and 40 mg/mL trastuzumab</p> <p>600 mg-600 mg⁹⁰: 60 mg/mL pertuzumab and 60 mg/mL trastuzumab</p>	<p>discard unused portion⁹⁰</p>	<p>SC syringe⁹⁰</p>	<p>10 d F, 24 h RT^{10,90}</p>	<p>- do not shake⁹⁰ - contains recombinant human hyaluronidase⁹⁰</p>
<p>Plerixafor 24 mg/1.2 mL (sanofi-aventis) (RT) no preservative⁹¹</p>	<p>N/A</p>	<p>20 mg/mL⁹¹</p>	<p>discard unused portion⁹¹</p>	<p>SC syringe⁹¹</p>	<p>48 h RT^{73,92}</p>	

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
Polatuzumab vedotin 30 mg 140 mg (Hoffman-La Roche) (F)(PFL) do not shake no preservative ⁹³	30 mg: 1.8 mL SWI ⁹³ 140 mg: 7.2 mL SWI ⁹³ direct diluent against side of vial during reconstitution ⁹³ swirl gently to mix ⁹³	20 mg/mL ⁹³ (PFL)	12 h F, RT ^{10,93}	0.72-2.7 mg/mL NS , D5W, ½NS ⁹³ (dilute to a minimum volume of 50 mL) ⁹³ gently invert bag to mix ⁹³	in NS: 72 h F, 4 h RT ⁹³ in D5W or ½NS: 72 h F, 8 h RT ⁹³	- do NOT shake ⁹³ - administer with 0.2 micron in-line filter ⁹³ -discard if discolouration or visible particulates are present ⁹³
Pralatrexate 20 mg/1 mL 40 mg/2 mL (Servier) (F)(PFL) no preservative ⁹⁴	N/A	20 mg/mL ⁹⁴	discard unused portion ²	syringe ⁹⁴	24 h F, RT ⁹⁵ **(PFL) ⁹⁵	- do NOT dilute ⁹⁴
Raltitrexed 2 mg (Pfizer) (F,RT)(PFL) no preservative ⁹⁶	4 mL SWI ⁹⁶	0.5 mg/mL ⁹⁶	12 h F, RT ^{10,96}	50-250 mL NS , D5W ⁹⁶	complete administration within 24 h F, RT ⁹⁶	

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
Ramucirumab 100 mg/10 mL 500 mg/50 mL (Eli Lilly) (F)(PFL) (do not shake) no preservative ⁹⁷	N/A	10 mg/mL ⁹⁷	discard unused portion ⁹⁷	0.4–4 mg/mL NS ^{97,98} 250-500 mL† gently invert to mix ⁹⁷ do NOT shake ⁹⁷	24 h F, 4 h RT ⁹⁷	- administer with 0.2 micron in-line filter ⁹⁷ - do NOT use dextrose containing solutions ⁹⁷
riTUXimab (RITUXAN®) 100 mg/10 mL 500 mg/50 mL (Roche) (F)(PFL) no preservative ⁹⁹	N/A	10 mg/mL ⁹⁹	discard unused portion ⁹⁹	1-4 mg/mL NS, D5W ⁹⁹ 250-500 mL†	NS: 10 d F plus an additional 24 h RT ^{10,99} D5W: 24 h F plus an additional 12 h RT ⁹⁹	
riTUXimab intravitreal injection (RITUXAN®) 100 mg/10 mL (Roche) (F)(PFL) no preservative ⁹⁹	N/A	10 mg/mL ⁹⁹	discard unused portion ⁹⁹	syringe for intravitreal use	use within 4 h of initial puncture ¹⁰	

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
riTUXimab subcutaneous (RITUXAN® SC) 1400 mg/11.7 mL 1600 mg/13.4 mL (Roche) (F)(PFL) no preservative ¹⁰⁰	N/A	120 mg/mL ¹⁰⁰	discard unused portion ¹⁰⁰	SC syringe ¹⁰⁰	48 h F plus 8 h RT ¹⁰⁰	- contains hyaluronidase ¹⁰⁰ - formulations are NOT interchangeable ¹⁰⁰
riTUXimab (RIXIMYO®) 100 mg/10 mL 500 mg/50 mL (Sandoz) (F)(PFL) (do NOT shake) no preservative ¹⁰¹	N/A	10 mg/mL ¹⁰¹	discard unused portion ¹⁰¹	1-4 mg/mL NS, D5W ¹⁰¹ 250-500 mL† gently invert to mix	NS: 10 d F plus an additional 24 h RT ^{10,101} D5W: 24 h F plus an additional 12 h RT ¹⁰¹	
riTUXimab intravitreal injection (RIXIMYO®) 100 mg/10 mL (Sandoz) (F)(PFL) (do NOT shake) no preservative ¹⁰¹	N/A	10 mg/mL ¹⁰¹	discard unused portion ¹⁰¹	syringe for intravitreal use	use within 4 h of initial puncture ¹⁰	

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
riTUXimab (RUXIENCE®) 100 mg/10 mL 500 mg/50 mL (Pfizer) (F)(PFL) no preservative ¹⁰²	N/A	10 mg/mL ¹⁰²	discard unused portion ¹⁰²	1-4 mg/mL NS , D5W ¹⁰² 250-500 mL† gently invert to mix	24 h F plus an additional 24 h RT ¹⁰²	
riTUXimab intravitreal injection (RUXIENCE®) 100 mg/10 mL 500 mg/50 mL (Pfizer) (F)(PFL) no preservative ¹⁰²	N/A	10 mg/mL ¹⁰²	discard unused portion ¹⁰²	syringe for intravitreal use	use within 4 h of initial puncture ¹⁰	
riTUXimab (TRUXIMA®) 100 mg/10 mL 500 mg/50 mL (Teva/Celltrion) (F)(PFL) no preservative ¹⁰³	N/A	10 mg/mL ¹⁰³	discard unused portion ¹⁰³	1-4 mg/mL NS , D5W ¹⁰³ 250-500 mL† gently invert to mix	24 h F plus an additional 12 h RT ¹⁰³	

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
riTUXimab intravitreal injection (TRUXIMA®) 100 mg/10 mL 500 mg/50 mL (Teva/Celltrion) (F)(PFL) no preservative ¹⁰³	N/A	10 mg/mL ¹⁰³	discard unused portion ¹⁰³	syringe for intravitreal use	use within 4 h of initial puncture ¹⁰	
romiDEPsin 10 mg (Celgene Australia) (RT) no preservative ¹⁰⁴	2.2 mL supplied diluent ¹⁰⁴ swirl to mix ¹⁰⁴	5 mg/mL ¹⁰⁴	discard unused portion ¹⁰⁴	500 mL NS ¹⁰⁴	24 h F ¹⁰⁴	- vials contain overfill to allow full drug recovery (drug vial contains 11 mg romidepsin; diluent vial has 2.4 mL diluent) ¹⁰⁴ - solvent contains 80% propylene glycol and 20% anhydrous ethanol ¹⁰⁴

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
Sacituzumab govitecan 180 mg (Gilead) (F)(PFL) no preservative ¹⁰⁵	20 mL NS ¹⁰⁵ bring vials to RT before reconstitution ¹⁰⁵ slowly add diluent to vial and gently swirl; allow to dissolve for up to 15 min ¹⁰⁵ do not shake ¹⁰⁵	10 mg/mL ¹⁰⁵	use immediately after reconstitution to prepare infusion solution ¹⁰⁵ discard unused portion ¹⁰⁵	1.1-3.4 mg/mL NS ¹⁰⁵ 100-1000 mL NS† slowly inject solution to bag to minimize foaming; do not shake ¹⁰⁵	24 h F ¹⁰⁵ , plus an additional 8 h RT including infusion time ¹⁰⁵ ** (PFL) ¹⁰⁵	- do not shake ¹⁰⁵ - protect container from light during administration ¹⁰⁵ - vials contain overflow (~20 mg per vial) ¹⁰⁶
Siltuximab 100 mg 400 mg (Janssen) (F)(PFL) no preservative ¹⁰⁷	100 mg: 5.2 mL SWI ¹⁰⁷ 400 mg: 20 mL SWI ¹⁰⁷ bring vial to RT prior to use (~30 min) ¹⁰⁷ gently swirl, do NOT shake ¹⁰⁷	20 mg/mL ¹⁰⁷	2 h RT ¹⁰⁷	250 mL D5W ¹⁰⁷ dilute to final volume by withdrawing volume from bag equal to volume of drug to be added ¹⁰⁷	complete administration within 6 h RT ¹⁰⁷	- administer with 0.2 micron in-line filter ¹⁰⁷

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
<p>Sirolimus, nanoparticle, albumin- bound (NAB) 100 mg (Aadi) (F)(PFL) no preservative¹⁰⁸ (SAP)</p>	<p>20 mL NS¹⁰⁸ slowly direct diluent against side of vial (over ≥1 min)¹⁰⁸ let stand for ≥5 min to wet powder¹⁰⁸ gently swirl or invert for ≥2 min to avoid foaming¹⁰⁸ if foaming/clumping occurs, let stand until foam subsides (≥15 min)¹⁰⁸</p>	<p>5 mg/mL¹⁰⁸</p>	<p>4 h F^{109,110} **(PFL)¹⁰⁸</p>	<p>undiluted in empty PVC or non-PVC infusion bag¹⁰⁸</p>	<p>9 h F, followed by max 4 h RT¹⁰⁸ **(PFL)¹⁰⁸</p>	<p>- each vial contains ~800-900 mg human albumin^{108,111} - to prevent foaming, do NOT inject NS directly onto the powder¹⁰⁸ - if powder is visible after reconstitution, gently invert to resuspend powder¹⁰⁸ - to prevent administration of proteinaceous strands, administer with 15 micron filter ONLY¹⁰⁸</p>

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
Streptozocin 1g (Keocyt) (F)(PFL) no preservative ¹¹²⁻¹¹⁵ (SAP)	9.5mL NS , SWI, D5W ¹¹²⁻¹¹⁵	100 mg/mL ¹¹²⁻¹¹⁵	12 h F ^{10,113-115}	syringe ¹¹³⁻¹¹⁵	48 h F ^{10,113-115}	
				100-500 mL NS , D5W, SWI ¹¹²⁻¹¹⁵	24 h F ¹¹³⁻¹¹⁵	
Tarlatamab 1 mg (Amgen) (F)(PFL) no preservative ¹¹⁶ (SAP)	1.4 mL SWI ¹¹⁶ do NOT use supplied IV solution stabilizer to reconstitute vials direct diluent against side of vial ¹¹⁶ gently swirl to mix do not shake ¹¹⁶	0.9 mg/mL ¹¹⁶	discard unused portion ¹⁰	250 mL NS ¹¹⁶ add 13 mL supplied IV solution stabilizer to NS bag and gently mix to avoid foaming; do not shake ¹¹⁶ add 1.1 mL reconstituted drug to IV bag following addition of IV solution stabilizer ¹¹⁶ gently mix by inverting bag; do not shake ¹¹⁶	complete administration within 8 h RT ¹¹⁶	- CAUTION: two vial sizes are available; final vial concentrations are different after reconstitution - due to low extractable volume ¹¹⁶ , use filtered venting needle (e.g., Chemo-Vent®) in place of CSTD for compounding - discard if cloudy or has particulates ¹¹⁶

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
<p>Tarlatamab 10 mg (Amgen) (F)(PFL) no preservative¹¹⁶ (SAP)</p>	<p>4.4 mL SWI¹¹⁶</p> <p>do NOT use supplied IV solution stabilizer to reconstitute vials¹¹⁶</p> <p>direct diluent against side of vial¹¹⁶</p> <p>gently swirl to mix do not shake¹¹⁶</p>	<p>2.4 mg/mL¹¹⁶</p>	<p>discard unused portion¹⁰</p>	<p>250 mL NS¹¹⁶</p> <p>add 13 mL supplied IV solution stabilizer to NS bag and gently mix to avoid foaming; do not shake¹¹⁶</p> <p>add 4.2 mL reconstituted drug to IV bag following addition of IV solution stabilizer¹¹⁶</p> <p>gently mix do not shake¹¹⁶</p>	<p>complete administration within 8 h RT¹¹⁶</p>	<p>- CAUTION: two vial sizes are available; final vial concentrations are different after reconstitution - due to low extractable volume¹¹⁶, use filtered venting needle (e.g., Chemo-Vent®) in place of CSTD for compounding - discard if cloudy or has particulates¹¹⁶</p>

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
<p>Tebentafusp 100 mcg/0.5 mL (Immunocore/Medison) (F)(PFL) do not shake no preservative¹¹⁷</p>	<p>N/A</p>	<p>200 mcg/mL¹¹⁷</p>	<p>discard unused portion¹¹⁷</p>	<p>100 mL NS¹¹⁷</p> <p>Step 1: add calculated volume of human albumin 5% to provide 225-275 mcg/mL final concentration¹¹⁷</p> <p>to mix: invert the bag and gently rotate ≥5 times; do NOT shake bag (repeat x3)¹¹⁷</p> <p>Step 2: add calculated volume of drug¹¹⁷</p> <p>to mix: invert the bag and gently rotate ≥5 times; do NOT shake bag (repeat x3)¹¹⁷</p>	<p>complete administration within 24 h F, 4 h RT¹¹⁷</p> <p>bring to RT prior to administration¹¹⁷</p>	<p>- do not use CSTD or filters during preparation¹¹⁷; use filtered venting needle (e.g., Chemo-Vent®) for preparation - CSTD can be used for administration¹¹⁸ - administer using 0.2 micron in-line filter¹¹⁷ - once the bag has been removed from fridge, it must remain at RT¹¹⁷</p>

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
<p>Tebentafusp 100 mcg/0.5 mL (Immunocore/Clinigen) (F)(PFL) do not shake no preservative^{119,120} (SAP)</p>	<p>N/A</p>	<p>200 mcg/mL¹¹⁹</p>	<p>discard unused portion^{2,119,120}</p>	<p>100 mL NS^{119,120}</p> <p>Step 1: add calculated volume of human albumin 5% to provide 225-275 mcg/mL final concentration^{119,120}</p> <p>to mix: invert the bag and gently rotate ≥5 times; do NOT shake bag (repeat x3)^{119,120}</p> <p>Step 2: add calculated volume of drug^{119,120}</p> <p>to mix: invert the bag and gently rotate ≥5 times; do NOT shake bag (repeat x3)^{119,120}</p>	<p>complete administration within 24 h F, 4 h RT^{119,120}</p>	<ul style="list-style-type: none"> - do not use CSTD or filters during preparation¹¹⁹; use filtered venting needle (e.g., Chemo-Vent®) for preparation - CSTD can be used for administration¹¹⁸ - administer using 0.2 micron in-line filter^{119,120} - once the bag has been removed from fridge, it must remain at RT^{119,120}

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
<p>Teclistamab 30 mg/3 mL 153 mg/1.7 mL (Janssen) (F)(PFL) do not shake no preservative¹²¹</p>	<p>N/A</p>	<p>30 mg¹²¹: 10 mg/mL</p> <p>(use for 2.1-52.9 mg doses)*</p>	<p>discard unused portion¹²¹</p>	<p>SC syringe¹²¹</p> <p>if drug volume >2 mL, divide volume into separate syringes for administration¹²¹</p>	<p>20 h F, RT¹²¹</p> <p>if stored in fridge, bring to RT prior to administration¹²¹</p>	<p>- CAUTION: two concentrations are available¹²¹ - do not use CSTD for volumes less than 1 mL¹²²; use filtered venting needle (e.g., Chemo-Vent®) for preparation¹²³</p>
		<p>153 mg¹²¹: 90 mg/mL</p> <p>(use for 53-375 mg doses)*</p>				
		<p>bring to RT before use (~15 min)¹²¹</p> <p>swirl gently for 10 sec to mix; do NOT shake¹²¹</p>				

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DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
Temsirolimus 30 mg/1.2 mL (Pfizer/Wyeth) (F)(PFL) ¹²⁴ no preservative ¹²⁵	1.8 mL supplied diluent ¹²⁴	10 mg/mL ¹²⁴	12 h RT ^{10,124} **(PFL) ¹²⁴	250 mL NS ¹²⁴ record time of dilution ¹²⁴	complete administration within 6 h ¹²⁴ mix by gentle inversion to avoid foaming ¹²⁴	- use non-DEHP bag and tubing - administer with 0.2 micron in-line filter ¹²⁴
Teniposide 50 mg/5 mL (BMS) (RT) preservative ¹²⁶	N/A	10 mg/mL ¹²⁶	discard unused portion	0.1-1 mg/mL NS, D5W ¹²⁶ 50-500 mL *	0.1-0.4 mg/mL: 24 h RT ¹²⁶ 1 mg/mL: complete administration within 4 h RT of preparation ^{126,127}	- do not refrigerate - use non-DEHP bag and tubing ¹²⁶ - do not use if precipitates ^{126,127} - contains DMA*** - excessive agitation may cause precipitation ¹²⁶

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
Thiotepa 15 mg 100 mg (Adienne/Methapharm) (F) no preservative ¹²⁸ (SAP)	15 mg: 1.5 mL SWI ¹²⁸ 100 mg: 10 mL SWI ¹²⁸ to remove haze, filter through 0.22 micron filter after reconstitution ¹²⁹ record time of reconstitution	10 mg/mL ¹²⁸	8 h F ¹²⁸	0.5-1 mg/mL NS ¹²⁸ ≤500 mg: 500 mL ¹²⁸ >500 mg: 1000 mL ¹²⁸ reconstituted solution is hypotonic and must be further diluted with NS prior to use ¹²⁸	24 h F, 4 h RT ¹²⁸	- do not use if precipitates are present ¹²⁸ - reconstituted solution may be used if opalescent ¹²⁸ - administer with 0.2 micron in-line filter ¹²⁸

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
<p>Thiotepa IT injection 15 mg 100mg (Adienne/Methapharm) (F) no preservative¹²⁸ (SAP)</p>	<p>15 mg: 1.5 mL SWI¹²⁸</p> <p>100 mg: 10 mL SWI¹²⁸</p> <p>diluents containing preservatives should NOT be used for intrathecal administration²⁶</p> <p>to remove haze, filter through 0.22 micron filter after reconstitution¹²⁹</p> <p>record time of reconstitution</p>	<p>10 mg/mL¹²⁸</p>	<p>8 h F¹²⁸</p>	<p>IT syringe</p> <p>qs to 6 mL with preservative free NS¹³⁰</p> <p>diluents containing preservatives should NOT be used for intrathecal administration²⁶</p>	<p>use within 4 h of initial reconstitution²</p>	<p>- auxiliary info²⁷: IT - label to include route in full (i.e., INTRATHECAL injection) attached to both syringe and outer ziplock bag²⁷ - do not use if precipitates are present¹²⁸ - reconstituted solution may be used if opalescent¹²⁸</p>
<p>Thyrotropin alfa 1.1 mg (Genzyme) (F)(PFL) no preservative¹³¹</p>	<p>1.2 mL SWI¹³¹</p> <p>swirl gently to mix¹³¹</p> <p>do NOT shake¹³¹</p>	<p>0.9 mg/mL¹³¹</p>	<p>12 h F^{10,131}</p>	<p>syringe¹³¹</p>	<p>24 h F^{10,131}</p>	<p>- do not use if particulates are present¹³¹</p>

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
Tislelizumab 100 mg/10 mL (BeiGene) (F)(PFL) (do not shake) no preservative ¹³² (SAP)	N/A	10 mg/mL ¹³²	discard unused portion ¹³²	2-5 mg/mL NS ¹³² 50 mL* mix by gentle inversion; do not shake ¹³²	complete administration within 20 h F, 4 h RT (max 24 h from preparation) ¹³² bring to RT prior to administration ¹³²	- discard if has visible particulates, or is discoloured or cloudy ¹³² - administer with 0.2 micron in-line filter ¹³²
Tocilizumab 80 mg/4 mL 200 mg/10 mL 400 mg/20 mL (Roche) (F)(PFL) no preservative ¹³³	N/A	20 mg/mL ¹³³	discard unused portion ¹³³	100 mL NS ¹³³ dilute to final volume by withdrawing volume from bag equal to volume of drug to be added ¹³³ gently invert to mix ¹³³	complete administration within 24 h F, RT ¹³³ bring to RT prior to administration ¹³³	- to prevent foaming: slowly add drug to infusion bag and gently invert bag to mix ¹³³
Topotecan 4 mg/4 mL (Accord) (RT)(PFL) no preservative ¹³⁴	N/A	1 mg/mL ¹³⁴	12 h F, RT ^{10,134}	0.025-0.5 mg/mL NS , D5W ¹³⁴ 25-50 mL†	10 d F, 4 d RT ^{10,134}	

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
Topotecan 4 mg/4 mL (Pfizer/Hospira) (F)(PFL) no preservative ¹³⁵	N/A	1 mg/mL ¹³⁵	discard unused portion ¹³⁵	0.02-0.5 mg/mL NS , D5W ¹³⁵ 25-50 mL†	24 h F, RT ¹³⁵	
Topotecan 4 mg/4 mL (Sandoz) (F)(PFL) no preservative ¹³⁶	N/A	1 mg/mL ¹³⁶	discard unused portion ¹³⁶	0.02-0.5 mg/mL NS , D5W ¹³⁶ 25-50 mL†	24 h F ¹³⁶ **(PFL) ¹³⁶	
Trastuzumab (HERCEPTIN®) 440 mg (Roche) (F) no preservative ¹³⁷	20 mL supplied BWI ¹³⁷ swirl vial gently; allow to stand undisturbed for 5 min ¹³⁷	21 mg/mL ¹³⁷	28 d F ¹³⁷	250 mL NS only ¹³⁷ do NOT use dextrose containing solutions ¹³⁷	24 h F, RT ¹³⁷	- do NOT shake ¹³⁷

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
Trastuzumab (HERZUMA®) 150 mg 440 mg (Teva/Celltrion) (F) no preservative ¹³⁸	150 mg: 7.2 mL SWI ¹³⁸	21 mg/mL ¹³⁸	discard unused portion ¹³⁸	250 mL NS only ¹³⁸ do NOT use dextrose containing solutions ¹³⁸	24 h F, RT ¹³⁸	- do NOT shake ¹³⁸ - supplied BWI contains benzyl alcohol ¹³⁸
	440 mg: 20 mL supplied BWI ¹³⁸		28 d F ¹³⁸			
	swirl vial gently; allow to stand undisturbed for 5 min ¹³⁸					
Trastuzumab (OGIVRI®) 150 mg 440 mg (BGP) (F) no preservative ¹³⁹	150 mg: 7.2 mL SWI ¹³⁹	21 mg/mL ¹³⁹	discard unused portion ¹³⁹	250 mL NS only ¹³⁹ do NOT use dextrose containing solutions ¹³⁹	24 h F, RT ¹³⁹	- do NOT shake ¹³⁹ - supplied BWI contains benzyl alcohol ¹³⁹
	440 mg: 20 mL supplied BWI ¹³⁹		28 d F ¹³⁹			
	swirl vial gently; allow to stand undisturbed for 5 min ¹³⁹					

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
Trastuzumab (TRAZIMERA®) 150 mg 440 mg (Pfizer) (F) no preservative ¹⁴⁰	150 mg: 7.2 mL SWI ¹⁴⁰	21 mg/mL ¹⁴⁰	discard unused portion ¹⁴⁰	250 mL NS only ¹⁴⁰ do NOT use dextrose containing solutions ¹⁴⁰	24 h F, RT ¹⁴⁰	- do NOT shake ¹⁴⁰ - supplied BWI contains benzyl alcohol ¹⁴⁰
	440 mg: 20 mL supplied BWI ¹⁴⁰		28 d F ¹⁴⁰			
	swirl vial gently; allow to stand undisturbed for 5 min ¹⁴⁰					

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
<p>Trastuzumab deruxtecan (ENHERTU®) 100 mg (AstraZeneca) (F)(PFL) no preservative¹⁴¹</p>	<p>5 mL SWI¹⁴¹ swirl gently until completely dissolved¹⁴¹ do NOT shake¹⁴¹</p>	<p>20 mg/mL¹⁴¹</p>	<p>12 h F^{10,141} **(PFL)¹⁴¹</p>	<p>100 mL D5W only¹⁴¹ gently invert to mix¹⁴¹ do NOT shake¹⁴¹ do NOT use sodium chloride solution¹⁴¹</p>	<p>complete administration within 24 h F, 4 h RT¹⁴¹ **(PFL)¹⁴¹</p>	<ul style="list-style-type: none"> - do not use if reconstituted solution contains visible particulates or is cloudy or discoloured¹⁴¹ - protect container from light during administration¹⁴² - administer with 0.2 micron in-line filter¹⁴¹ - if stored in fridge, bring bag to RT prior to use¹⁴¹

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
<p>Trastuzumab emtansine (KADCYLA®) 100 mg 160 mg (Roche) (F)(PFL) no preservative¹⁴³</p>	<p>100 mg: 5 mL SWI¹⁴³</p> <p>160 mg: 8 mL SWI¹⁴³</p> <p>swirl gently until completely dissolved</p> <p>do NOT shake¹⁴³</p>	<p>20 mg/mL¹⁴³</p>	<p>12 h F^{10,144}</p>	<p>250 mL NS or ½NS only¹⁴³</p> <p>do NOT shake¹⁴³</p> <p>do NOT use dextrose containing solutions¹⁴³</p>	<p>24 h F¹⁴³</p>	<p>- do not use if reconstituted solution contains visible particulates or is cloudy or discolored¹⁴³ - D5W causes aggregation of the protein¹⁴³ - for infusions prepared in NS: administer with 0.2 micron in-line filter or 0.22 micron polyethersulfane (PES) filter¹⁴³ - for infusions prepared in ½NS: filter is optional for administration¹⁴³</p>

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
Tremelimumab 25 mg/1.25 mL 300 mg/15 mL (AstraZeneca) (F)(PFL) (do not shake) no preservative ¹⁴⁵	N/A	20 mg/mL ¹⁴⁵	discard unused portion ¹⁴⁵	0.1-10 mg/mL NS, D5W ¹⁴⁵ 50 mL* mix by gentle inversion; do NOT shake ¹⁴⁵	24 h F, RT ¹⁴⁵	- administer with 0.2 micron in-line filter ¹⁴⁵ - discard if visible particles are present ¹⁴⁵

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
<p>Treosulfan 1 g 5 g (Medexus) (RT) no preservative¹⁴⁶</p>	<p>1 g¹⁴⁶: 20 mL NS, D5W, SWI, ½NS</p> <p>5 g¹⁴⁶: 100 mL NS, D5W, SWI, ½NS</p> <p>pre-heat diluent to 25-30°C (max)¹⁴⁷</p> <p>shake vial to loosen powder before adding the warmed diluent¹⁴⁸</p> <p>vigorous shaking may be required¹⁴⁸; prolonged standing time may improve solubility¹⁴⁶</p>	<p>50 mg/mL¹⁴⁶</p>	<p>12 h RT^{10,146}</p>	<p>undiluted in empty infusion bag^{146,147}</p>	<p>3 d RT¹⁴⁶</p>	<p>- do NOT refrigerate as may precipitate¹⁴⁶</p>

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
<p>Treosulfan 1 g 5 g (medac) (RT) no preservative^{149,150} (SAP)</p>	<p>1 g^{149,150}: 20 mL SWI, ½NS</p> <p>5 g^{149,150}: 100 mL SWI, ½NS</p> <p>pre-heat diluent to 25-30°C (max)^{149,150}</p> <p>shake vial carefully to loosen powder before adding the warmed diluent^{149,150}</p> <p>gently shake while adding diluent^{149,150}</p> <p>(takes ~2 min to reconstitute)^{149,150}</p>	<p>50 mg/mL^{149,150}</p>	<p>12 h RT^{10,149,151}</p>	<p>undiluted¹⁵²</p> <p>or</p> <p>dilute with NS or D5W in empty infusion bag to final concentration of 20 mg/mL¹⁵¹</p>	<p>4 d RT^{149,151}</p>	<p>- compatible with polytetrafluoroethyl ene filters¹⁵³ - may sometimes require vigorous shaking to reconstitute^{149,150} - do NOT refrigerate as may cause precipitation^{149,150}</p>
<p>vinBLAStine 10 mg/10 mL (Pfizer) (F)(PFL) no preservative¹⁵⁴</p>	<p>N/A</p>	<p>1 mg/mL¹⁵⁴</p>	<p>discard unused portion^{2,154}</p>	<p>25-50 mL NS, D5W¹⁵⁵</p>	<p>use within 4 h of initial vial puncture^{2,154}</p>	<p>- auxiliary info: WARNING: FOR INTRAVENOUS USE ONLY – FATAL IF GIVEN BY OTHER ROUTES^{156,157}</p>

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
vinBLAS tine 10 mg/10 mL (Teva) (F)(PFL) no preservative ¹⁵⁸	N/A	1 mg/mL ¹⁵⁸	discard unused portion ^{2,158}	25-50 mL NS , D5W ¹⁵⁵	use within 4 h of initial vial puncture ^{2,158}	- auxiliary info: WARNING: FOR INTRAVENOUS USE ONLY – FATAL IF GIVEN BY OTHER ROUTES ^{156,157}
vinCRIS tine 2 mg/2 mL 5 mg/5 mL (Pfizer/Hospira) (F)(PFL) no preservative ¹⁵⁹	N/A	1 mg/mL ¹⁵⁹	8 h F, RT ¹⁵⁹	0.01-0.1 mg/mL NS , D5W ¹⁵⁹ 50 mL†	24 h F, RT ¹⁵⁹ **(PFL) ¹⁵⁹	- auxiliary info: WARNING: FOR INTRAVENOUS USE ONLY – FATAL IF GIVEN BY OTHER ROUTES ^{156,157} - for LYEPOCHR protocol, see entry for EPOCHR (3-in-1 solution containing either etoposide or etoposide phosphate AND DOXOrubicin and vinCRIStine)

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
vinCRISTine 1 mg/1 mL 2 mg/2 mL 5 mg/5 mL (Teva) (F)(PFL) no preservative ¹⁶⁰	N/A	1 mg/mL ¹⁶⁰	8 h F, RT ¹⁶⁰	0.01-0.1 mg/mL NS , D5W ¹⁶⁰ 50 mL†	24 h F, RT ¹⁶⁰	- auxiliary info: WARNING: FOR INTRAVENOUS USE ONLY – FATAL IF GIVEN BY OTHER ROUTES^{156,157} - for LYEPOCHR protocol, see entry for EPOCHR (3-in-1 solution containing either etoposide or etoposide phosphate AND DOXOrubicin and vinCRISTine)
Vinorelbine 10 mg/1 mL 50 mg/5mL (Fresenius Kabi) (F)(PFL) no preservative ¹⁶¹	N/A	10 mg/mL ¹⁶¹	discard unused portion ¹⁶¹	0.5-2.0 mg/mL NS , D5W, ½NS, D5-½NS, Ringer's, Ringer's Lactate ¹⁶¹ 50 mL†	24 h F, RT ¹⁶¹	- auxiliary info: WARNING: FOR INTRAVENOUS USE ONLY – FATAL IF GIVEN BY OTHER ROUTES^{156,157}

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
Vinorelbine 10 mg/1 mL 50 mg/5 mL (GMP) (F)(PFL) no preservative ¹⁶²	N/A	10 mg/mL ¹⁶²	discard unused portion ²	0.5-2.0 mg/mL NS , D5W, ½NS, D5-½NS, Ringer's, Ringer's Lactate ¹⁶² 50 mL†	24 h F, RT ¹⁶²	- auxiliary info: WARNING: FOR INTRAVENOUS USE ONLY – FATAL IF GIVEN BY OTHER ROUTES ^{156,157}
Vinorelbine 10 mg/1 mL 50 mg/5 mL (Teva) (F)(PFL) no preservative ¹⁶³	N/A	10 mg/mL ¹⁶³	discard unused portion ¹⁶³	0.5–2.0 mg/mL NS , D5W, ½NS, D5-½NS, Ringer's, Ringer's Lactate ¹⁶³ 50 mL†	24 h F, RT ¹⁶³	- auxiliary info: WARNING: FOR INTRAVENOUS USE ONLY – FATAL IF GIVEN BY OTHER ROUTES ^{156,157}
Zoledronic acid 4 mg/5 mL (Dr Reddy's) (RT) no preservative ¹⁶⁴	N/A	0.8 mg/mL ¹⁶⁴	discard unused portion ¹⁶⁴	100 mL NS , D5W ¹⁶⁴	complete infusion within 24 h of preparation ¹⁶⁴ refrigerate diluted product if not used immediately after preparation; bring to RT prior to use ¹⁶⁴	- do NOT mix with calcium containing solutions (e.g., Lactated Ringer's) ¹⁶⁴

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
Zoledronic acid 4 mg/5 mL (Marcan) (RT) no preservative ¹⁶⁵	N/A	0.8 mg/mL ¹⁶⁵	discard unused portion ¹⁶⁵	100 mL NS , D5W ¹⁶⁵	complete infusion within 24 h of preparation ¹⁶⁵ refrigerate diluted product if not used immediately after preparation; bring to RT prior to use ¹⁶⁵	- do NOT mix with calcium containing solutions (e.g., Lactated Ringer's) ¹⁶⁵
Zoledronic acid 4 mg/5 mL (MDA) (RT) no preservative ¹⁶⁶	N/A	0.8 mg/mL ¹⁶⁶	discard unused portion ¹⁶⁶	100 mL NS , D5W ¹⁶⁶	complete infusion within 24 h of preparation ¹⁶⁶ refrigerate diluted product if not used immediately after preparation; bring to RT prior to use ¹⁶⁶	- do NOT mix with calcium containing solutions (e.g., Lactated Ringer's) ¹⁶⁶

BC CANCER CHEMOTHERAPY PREPARATION AND STABILITY CHART

DRUG & STRENGTH (Storage Prior to Use, Manufacturer, Preservative Status)	Reconstitute With:	To Give:	Vial Stability	Product (for IV bag size selection, see Notes†)	Product Stability	Special Precautions/Notes
Zoledronic acid (ZOMETA) 4 mg/ 5 mL (Novartis) (RT) no preservative ¹⁶⁷	N/A	0.8 mg/mL ¹⁶⁷	discard unused portion ⁴³	100 mL NS, D5W ¹⁶⁷	complete infusion within 24 h of preparation ¹⁶⁷ refrigerate diluted product if not used immediately after preparation; bring to RT prior to use ¹⁶⁷	- do NOT mix with calcium containing solutions (e.g., Lactated Ringer's) ¹⁶⁷
Zoledronic acid 4 mg/5 mL (Sandoz) (RT) no preservative ¹⁶⁸	N/A	0.8 mg/mL ¹⁶⁸	discard unused portion ¹⁶⁸	100 ml NS, D5W ¹⁶⁸	complete infusion within 24 h of preparation ¹⁶⁸ refrigerate diluted product if not used immediately after preparation; bring to RT prior to use ¹⁶⁸	- do NOT mix with calcium containing solutions (e.g., Lactated Ringer's) ¹⁶⁸

* Suggested volume based on usual dose range and any concentration range of stability data

† see [BC Cancer IV Bag Selection table](#): standardized bag sizes are provided for select Benefit Drugs with concentration-dependent stability or large drug volume

** Protect from light means minimizing exposure to direct sunlight over a *storage* period. More specific information on protection from light (eg, protecting container and tubing during *administration*) will be indicated in the Special Precautions/Notes column.

*** Contains DMA (N,N dimethylacetamide). Product may be incompatible with closed system transfer devices (CSTD) such as ChemoLock.

Centres are not to change content locally. All suggestions for change are to be forwarded to the Cancer Drug Manual editor.

Explanatory Notes:

Stability data assumes products prepared using standard aseptic technique in biological safety cabinet at low risk for contamination according to the classification outlined in USP 797.^{169,170}

Vial stability: Stability of solution after first puncture or reconstituted solution.

Storage temperature: If information states same stability with refrigerator and room temperature storage, then fridge stability is bolded as preferred (ie, to minimize growth of micro-organisms).

Discard unused portion: Unused portion from single use vials should be discarded at the end of the day.

“overflow known” is stated if the manufacturer states overflow that is present is within acceptable limits.

“Complete administration within ___” is stated if the manufacturer specifies that the infusion must be completed in a specific time frame following preparation, usually including entire time required for preparation (from first puncture), storage, and administration of infusion.

Nomenclature for **In-line filters** has been standardized to 0.2 micron filter size. For more information, refer to CDM monograph.

Abbreviations:

BWI = bacteriostatic water for injection

CIVI: ambulatory pump = Continuous Intravenous Infusion (e.g., elastomeric infusor)

CSTD = closed system transfer device

D5W = dextrose 5% in water

DMA = N,N dimethylacetamide

F = refrigerate

Non-DEHP = not containing Di(2-ethylhexyl) phthalate (DEHP)

NS = normal saline

PFL = protect from light

RT = room temperature

SAP = drug is approved for use through the Health Canada Special Access Program

SWI = sterile water for injection

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