

Medications for Osteoporosis:

An Update

B.C. Provincial Academic Detailing (PAD) Service

Participants will have the opportunity to:

- 1. Apply current evidence to guide prescribing and deprescribing decisions for bisphosphonates, denosumab, raloxifene, teriparatide and romosozumab.
- 2. Incorporate fracture risk reduction and time-to-benefit estimates as part of shared-decision making with patients.
- 3. Compare the principal clinical considerations when choosing between bisphosphonates and denosumab, the most commonly prescribed osteoporosis medications in British Columbia.

Understudied populations in medication clinical trials: Premenopausal females, males, intersex persons, transgender persons, diverse racial or ethnic groups, residents of long term care, multimorbidity and polypharmacy, people taking glucocorticoids, participants defined by FRAX scores.

Brand Name	Generic Name
Fosamax®, Fosavance®	alendronate oral once a day or week
Actonel®, Actonel DR®	risedronate oral once a day or week or month
Aclasta®	zoledronic acid intravenous once a year
Prolia®, Jubbonti®	denosumab subcutaneous every 6 months
Evista®	raloxifene oral once a day
Forteo®, Osnuvo®	teriparatide subcutaneous once a day
Evenity®	romosozumab subcutaneous once a month

BC's Provincial Academic Detailing (PAD) Service is offered free of charge to health care professionals. The service is provided by health authorities and supported by the Ministry of Health. Relevant topics are identified in consultation with various groups. All written materials are externally reviewed by clinicians and experts in critical appraisal.

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Osteoporosis medications: overview

Drug Administration		Contraindications, serious precautions	Important adverse events Duration of therapy		
alendronate risedronate antiresorptives	oral: daily, weekly or monthly	 hypocalcemia alendronate: CrCl < 35 mL/min risedronate: CrCl < 30 mL/min abnormalities of esophagus inability to sit/stand upright for 30 minutes 	 osteonecrosis of the jaw atypical femoral fractures esophageal 	 initial duration of treatment 3-6 	
zoledronic acid antiresorptive	intravenous: annual infusion	 hypocalcemia CrCl < 35 mL/min risk factors for acute kidney injury inability to hydrate pre + post infusion 	 osteonecrosis of the jaw atypical femoral fractures first dose infusion reactions 	years (reassess at 3 years)	
denosumab antiresorptive	subcut: every 6 months	 hypocalcemia: significantly increased risk in renal impairment 	 osteonecrosis of the jaw atypical femoral fractures infections, dermatologic reactions 	 if stopping, transition to a bisphosphonate or seek advice from a consultant 	
raloxifene estrogen receptor modulator	oral: <i>daily</i>	 history or current VTE history of stroke or risk factors for stroke males and premenopausal females 	 thromboembolism vasodilation (hot flushes) leg cramps 	 review for opportunity to deprescribe or transition to another osteoporosis therapy 	
teriparatide anabolic	subcut: <i>daily</i>	 hypercalcemia CrCl < 30 mL/min 	 orthostatic hypotension arthralgia, headache, muscle spasms 	 after 24 months (total exposure), review for continuation or transition to an antiresorptive 	
romosozumab anabolic + antiresorptive	subcut: once a month	 hypocalcemia history of myocardial infarction or stroke 	 possible increased risk of major cardiovascular events † 	 after 12 months, transition to an antiresorptive is recommended 	

antiresorptive: inhibits resorption of bone (osteoclasts); anabolic: stimulates bone formation (osteoblasts); + US FDA 2019 advisory committee: additional data needed to better characterize risk

Osteoporosis medications: indications & basis of approval

Basis of regulatory drug approval for osteoporosis

- 1. Osteoporosis medications generally enter the market with an indication for use in postmenopausal females based on evidence of a reduction in the risk of radiographic vertebral fractures and, in some cases, clinical (symptomatic) fractures.
- 2. Subsequent population indications may be added by demonstrating that the medication increases bone mineral density (BMD) estimates of drug effects (efficacy & safety) for these patient groups are less certain.

Health Canada Indications	Postmenopausal females		Males with	Exposure to medications that increase fracture risk		
	Osteoporosis	Osteopenia	osteoporosis	Glucocorticoid females, males	Aromatase Inhibitor non metastatic breast cancer	Androgen Deprivation non metastatic prostate cancer
bisphosphonates	radiographic vertebral & clinical fractures	BMD ⊕	BMD & radiographic vertebral fractures	BMD †		
denosumab *	radiographic vertebral & clinical fractures		BMD	BMD ††	BMD ⊕	BMD & radiographic vertebral fractures
raloxifene	radiographic vertebral fractures	BMD				
teriparatide *	radiographic vertebral & clinical fractures		BMD	BMD †††		
romosozumab *	radiographic vertebral & clinical fractures					

surrogate outcomes: radiographic vertebral factures and bone mineral density, used in osteoporosis medication clinical trials as a substitute for a direct measure of how a patient feels, functions or survives; radiographic vertebral fractures: detected on scheduled imaging during the clinical trial, may not be symptomatic;

* high risk: indicated for those with a history of osteoporotic fracture or multiple risk factors for fracture; \bigoplus postmarketing trial(s) demonstrate reduction in clinical fractures; † prednisone \geq 7.5 mg per day equivalent fracture or low BMD; †† prednisone \geq 5 mg per day equivalent plus prior fracture or low BMD; †† prednisone \geq 5 mg per day equivalent plus prior fracture or low BMD; †† prednisone \geq 5 mg per day equivalent plus prior fracture or low BMD

Health Canada Drug Product Database; Health Canada Drug Health Product Register; US FDA Approved Drugs; US FDA 2015 public workshop osteoporosis drug development; US FDA table surrogate endpoints drug approval; KEHOE Br J Clin Pharmacol 2019; REID NEJM 2018 osteopenia zoledronic acid; B.C. Ministry of Health 2023 denosumab

B.C. PharmaCare coverage & annual drug cost

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Approximate annual drug cost		PharmaCare coverage	Special Authority Criteria
alendronate Fosamax, Fosavance	\$100 *	Dogular hanafit	
risedronate Actonel	\$100 *	Regular benefit	
zoledronic acid Aclasta	\$390 ◆	Limited Coverage	 osteoporotic fracture <u>plus</u> contraindication to oral bisphosphonate abnormalities of the esophagus such as stricture or achalasia
denosumab Prolia, Jubbonti	\$420 biosimilar \$950 brand	Limited Coverage	 osteoporotic fracture <u>plus</u> contraindication to oral bisphosphonate abnormalities of the esophagus such as stricture or achalasia or immune-mediated hypersensitivity reaction aromatase inhibitor for breast cancer (5-year coverage)
raloxifene Evista	\$200	Limited Coverage	 osteoporotic fracture <u>plus</u> intolerable adverse effects to oral bisphosphonate <u>or</u> unsatisfactory response (postmenopausal females only) – esophageal ulceration, erosion or stricture, lower gastrointestinal symptoms severe enough to cause discontinuation of bisphosphonates after ≥ 1 month trial – new clinically or radiographically documented osteoporotic fracture after 1 year of adherence to alendronate or risedronate
romosozumab Evenity	\$8900	Limited Coverage	 osteoporotic fracture <u>and</u> treatment naïve <u>plus</u> FRAX score ≥ 20% (postmenopausal females only; 1-year coverage)
teriparatide Forteo, Osnuvo	\$8000 biosimilar \$8000 brand	Non-benefit	

* approximate wholesale cost of weekly generic formulation (alendronate 70 mg, risedronate 35 mg); daily (alendronate 5 & 10 mg, risedronate 5 mg), monthly (risedronate 150 mg), delayed release (risedronate 35 mg DR) formulations are more costly; see **BC PAD Osteoporosis Drug Table** for doses, costs, coverage; • excludes infusion costs and potential missed work hours for working patients



Osteoporosis medications: evidence overview

Postmenopausal females§	Hip fracture	Symptomatic vertebral fracture†	Symptomatic fracture††	Radiographic vertebral fracture+++	Serious adverse events	Withdrawals due to adverse events
bisphosphonate vs placebo ; 3 – 4 yrs; meta-analysis baseline VF 0% – 100%	ARR 0.6% RRR 36%	ARR 1.8% RRR 62% *	ARR 2.4% RRR 21% *	ARR 5.6% RRR 51% *	NSS	NSS
denosumab vs placebo; 3 yrs; 1 RCT baseline VF 23%	ARR 0.5% RRR 40%	ARR 1.8% RRR 69%	ARR 1.5% RRR 20%	ARR 4.9% RRR 68%	NSS	NSS
raloxifene vs placebo ; 3 yrs; meta-analysis baseline VF 37% – 56%	NSS	NSS	NSS	ARR 2.8% RRR 41%	NSS	ARI 1.5%
teriparatide vs oral bisphosphonate; 2 yrs; 1 RCT baseline VF 100%	NSS	ARR 2.8% RRR 71%	ARR 4.6% RRR 52%	ARR 6.6% RRR 56%	NSS	NSS
romosozumab vs oral bisphosphonate; 2 – 3 yrs; 1 RCT baseline VF or HF 100%	ARR 1.2% RRR 38%	ARR 1.2% RRR 59%	ARR 3.3% RRR 27%	ARR 3.9% RRR 50%	NSS	NSS

American College of Physicians 2023 Recommendations for Postmenopausal Females with Osteoporosis: bisphosphonates initial pharmacologic therapy (high certainty); denosumab second line (moderate certainty); romosozumab (moderate certainty) or teriparatide (low certainty) followed by a bisphosphonate in females at very high risk of fracture due to age and fracture history; raloxifene not recommended; Males with Osteoporosis: extrapolated from evidence for postmenopausal females: bisphosphonates initial pharmacologic therapy (low certainty); denosumab second line (low certainty);

§ primary osteoporosis: based on BMD or fragility fracture, not secondary to another medical condition or medication; VF HF proportion of participants with a vertebral or hip fracture at baseline; † clinically recognized, symptomatic; †† symptomatic nonvertebral ± vertebral fractures excl. fractures not related to osteoporosis; ††† detected on scheduled imaging, may not be symptomatic, radiographic criteria may vary between trials; ARR absolute risk reduction; ARI absolute risk increase; RRR relative risk reduction; NSS not statistically significantly different; *** bisphosphonates** heterogeneity in baseline fracture risk across RCTs & variability in estimates of drug effect; teriparatide 58% participants previously used a bisphosphonate; romosozumab sequential therapy romosozumab for 1 year followed by alendronate for 1 year; 6% participants previously used a bisphosphonate;

ACP 2023 osteoporosis guideline & systematic review; FREEDOM NEJM 2009 denosumab; VERO Lancet 2018 & Osteo International 2020 teriparatide vs risedronate; ARCH NEJM 2017 romosozumab vs alendronate; US FDA 2019 romosozumab review; Health Canada 2019 romosozumab review; European Medicines Agency 2020 romosozumab review

Bisphosphonates, denosumab: patient population estimates

Secondary versus Primary Prevention: Alendronate, Denosumab								
Postmenopausal females Radiogra			Radiograp	nic vertebral	fracture†	Participant Demographi	Participant Demographics	
- londron - to		ondary prevention vious vertebral fracture		3 years	placebo 15.0% → drug 8.0%	 ages 55 – 81; ambulatory; self rated health good to excellent BMD T score –2.0 or lower; mean age 71; 97% White 		
alendronate	primary prevo without previ		ARR ~2%	4 years	placebo 3.8% → drug 2.1%	 ages 54 – 81; ambulatory; self rated health good t BMD T score –1.6 or lower; mean age 68; 97% W 		
	secondary pr previous vert	revention ebral fracture	ARR ~9%	0	placebo 13.6% → drug 4.6%	 ages 60 – 90; ambulatory; generally in good health BMD T score –2.5 to –4.0; mean age 72; 93% White excluding those who had taken a bisphosphonate for > 3 years or wit the previous year 		
denosumab	primary prevo without previ		ARR ~4%	3 years	placebo 5.2% → drug 1.7%			
Patient Populations: Bisphosphonates								
Population Radiographic vertebral fracture†				Hip fracture				
		NNT 20 (16 baseline risk	•	3 – 4 years	1205 events in 16,902 females	NNT 143 (105 - 333) * baseline risk: ~2%	3 – 4 years	263 events in 16,634 females

males with osteoporosis	NNT 33 (26 − 125) * baseline risk: ~5%	2 years	55 events in 1692 males	not estimable	NNT estimate may include sufficient
people taking glucocorticoids	NNT 30 (20 − 143) * baseline risk: ~8%	1 – 2 years	77 events in 1343 people	not estimable	imprecision to impact clinical or patient decisions

† detected on scheduled imaging, may not be symptomatic; **ARR** absolute risk reduction; **baseline risk + estimates of drug effect** mean difference between drug and placebo for radiographic vertebral fractures varies by baseline risk (e.g., primary vs secondary prevention); this is less apparent for hip fractures where the drug effect varies minimally across trials; **NNT** number of people who need to take a bisphosphonate for one less person to experience a fracture with 95% confidence interval

FIT 1 Lancet 1996 alendronate; FIT 2 JAMA 1998 alendronate; FREEDOM NEJM 2009 & J Bone Min Res 2012 denosumab; ACP 2023 osteoporosis guideline & systematic review; COCHRANE 2016 CD001347





painful osteoporotic fracture: fracture of the vertebrae or another bone considered related to osteoporosis

pad Older adults with frailty & multimorbidity

R	epresentation of people with frailty and multimorbidity
	Mean age of participants across trials: 50 – 85 years People with multimorbidity, polypharmacy and persons in long term
•	care are underrepresented in osteoporosis trials Canadian guideline pharmacotherapy recommendations on the prevention of fractures in long term care are extrapolated from ambulatory, community dwelling females with few or no comorbidities
•	(Osteoporosis Canada 2015) A 2023 systematic review of osteoporosis medications did not find evidence for fracture related mortality, functionality or disability

Consider as part of medication decision making

- 1. Time-to-benefit: onset of symptomatic or hip fracture risk reduction and its relevance to people of advanced age or limited life expectancy
 - bisphosphonates: approximately after 12 months of treatment
 - denosumab: approximately after 6 12 months of treatment
- 2. Administration instructions: that make it difficult to provide medications safely
- **3. Kidney function:** which may preclude medication use or increase the risk of adverse events

Administration				
alendronate	≥ 200 mL plain water	take on empty stomach upon arising for the day; contraindicated =		
risedronate	≥ 120 mL plain water	inability to sit or stand upright for at least 30 minutes, abnormalities of the esophagus		
zoledronic acid	≥ 500 mL of fluids before & after infusion	infusion not less than 15 minutes; check serum calcium before each dose		
denosumab	subcutaneous injection: upper arm, upper thigh, abdomen	check serum calcium before each dose		

Kidney	
alendronate	contraindicated: CrCl < 35 mL/min
risedronate	not recommended: CrCl < 30 mL/min
zoledronic acid	contraindicated: CrCl < 35 mL/min (C-G formula using actual body weight)
denosumab	significantly increased risk of hypocalcemia: CrCl < 30 mL/min

HANDEL BMJ 2023; ACP 2023 osteoporosis guideline & systematic review; Osteoporosis Canada 2015 guideline; GATES Systematic Rev 2023; DEARDORFF JAMA Int Med 2022; FIT 1 Lancet 1996 alendronate; HIP NEJM 2001 risedronate; HORIZON PFT & RFT NEJM 2007 zoledronic acid; REID NEJM 2018 zoledronic acid; FREEDOM NEJM 2009 denosumab; Health Canada Drug Product Database

pad Exit strategies & transitions

Persistence of effect

- Treatment effects of bisphosphonates may persist for years after treatment discontinuation
- Denosumab, teriparatide and romosozumab have a more rapid offset of effect following discontinuation

Bisphosphonates Concern:

longer treatment duration increases the risk of atypical femoral fractures and osteonecrosis of the jaw, although both are rare

- In postmenopausal females, if alendronate is continued for another 5 years after 5 years of initial therapy:
 - symptomatic vertebral factures: ARR 3%
 - atypical femoral fractures: ARI 0.1% 0.4%
- Osteoporosis Canada 2023 suggests (weak recommendations):
 - initial therapy for 3-6 years (low-certainty evidence)
 - BMD measurement 3 years after initiating pharmacotherapy (very low-certainty evidence)
- Health Canada & US FDA: optimal duration of therapy has not been determined

Denosumab Concern:

discontinuing denosumab is associated with an increased risk of multiple vertebral fractures

- Rate of multiple vertebral fractures (starting 7 months after last dose):
 - placebo: 3.6 per 100 patient years
 - denosumab taken for 1-3 years: 3.0 per 100 patient years
 - denosumab taken for > 3 years: 7.5 per 100 patient years
- Consistency with the every 6 months dosing schedule is important
- Osteoporosis Canada suggests: long-term uninterrupted therapy; if discontinuing after ≤ 4 doses, transition to a bisphosphonate; if discontinuing after ≥ 5 doses, seek advice from a consultant with osteoporosis expertise

Teriparatide Concern:

risk of osteosarcoma in animal studies but postmarketing studies in humans do not find a clinical signal of osteosarcoma

- Transition to a bisphosphonate or denosumab typically considered after 24 months of teriparatide
- US FDA 2020 review: evidence no longer supports a warning for osteosarcoma
- Health Canada & US FDA: continue beyond 24 months of lifetime exposure only if high risk for fracture

Romosozumab Concern:

effects on bone mineral density and bone formation markers wane after 12 months of treatment

 Transition to a bisphosphonate or denosumab typically considered after 12 months Health Canada & US FDA: limit treatment duration to 12 months

Health Canada Drug Product Database; US FDA Approved Drugs; AHRQ 2019 osteoporosis long term drug therapy; FLEX extension trial JAMA 2006 alendronate; BLACK NEJM 2020; HORIZON PFT extension trial J Bone Min Res 2012 zoledronic acid; Osteoporosis Canada 2023 guideline; COSMAN J Bone Miner Res 2022 denosumab extension; US FDA 2020 teriparatide review; US FDA 2019 romosozumab review; REID Lancet 2022



Bisphosphonates: clinical considerations

Contraindications	Acute kidney injury	Gastrointestinal	Atypical femoral fractures
oral: abnormalities of esophagus, inability to sit or stand upright for at least 30 minutes, inability to swallow \ge 120 – 200 mL of water	zoledronic acid: ensure adequate hydration (eat & drink normally including at least 500 mL of fluids) prior to and after administration –	oral: may cause or worsen esophagitis, esophageal ulcers, esophageal erosions, stricture or perforation	subtrochanteric or proximal femoral shaft: 1/3 bilateral; may occur in absence of apparent trauma
zoledronic acid: inability to appropriately hydrate pre and	particularly in older adults, those receiving diuretics or nephrotoxic	Musculoskeletal	prodrome: patients should be counselled to report new or
post infusion	medications	bone, joint, muscle pain: possibly	unusual thigh, hip, groin pain
hypocalcemia	monitoring CrCl post dose: recommended in patients at risk	severe; also common infusion reaction symptoms	incidence: $< 0.1\%$ (0 – 5 years)
CrCl < 35 mL/min: alendronate	infusion time: minimum 15		additive risk factors: duration of
CrCl < 30 mL/min: risedronate	minutes	Ophthalmologic	therapy $> 3 - 5$ years, Asian
CrCl < 35 mL/min: zoledronic acid		conjunctivitis, uveitis, episcleritis,	descent, > 1 year of glucocorticoid use
	Infusion reaction	scleritis: incidence ≤ 1%	
Calcium & Vitamin D	zoledronic acid: ~25% of patients		Osteonecrosis of the jaw
zoledronic acid: check serum calcium before each dose –	within 3 days of first infusion, less frequent on subsequent infusions		associated with invasive dental procedures such as tooth
replete calcium and vitamin D if necessary	symptoms: fever, chills, fatigue; musculoskeletal pain; pain &		extraction: consider preventive dentistry/regular dental monitoring
hypocalcemia symptoms: muscle	redness at infusion site		incidence: 0.02 – 0.15%
cramps or twitching, numbness or tingling mouth, fingers or toes	acetaminophen or ibuprofen to prevent or manage symptoms		additive risk factors: higher dose oncology regimens, duration of
See reference list			therapy > 2 – 3 years



Denosumab: clinical considerations

Contraindications	Chronic kidney disease	Multiple vertebral fractures	Atypical femoral fractures	
hypocalcemia	no dose adjustment required	discontinuing denosumab:	subtrochanteric or proximal	
Calcium & Vitamin D	insufficient evidence to evaluate fracture efficacy in patients with	associated with an increased risk of multiple vertebral fractures	femoral shaft: 1/3 bilateral; may occur in absence of apparent	
hypocalcemia: rare if normal	eGFR < 30 mL/min	Osteoporosis Canada suggests	trauma	
kidney function and adequate calcium and vitamin D intake	risk of hypocalcemia increases as eGFR declines:	long-term uninterrupted therapy or an exit strategy that is dependent	prodrome: patients should be counselled to report new or	
check serum calcium: before each	30 – 60 mL/min: < 1%	on duration of therapy	unusual thigh, hip, groin pain	
dose (replete calcium and vitamin D if necessary), and within 2 weeks post dose in patients at	15 – 30 mL/min: 4% < 15 mL/min, dialysis: 24 – 42%	Musculoskeletal	incidence: less well documented for denosumab compared to	
		bone, joint, muscle pain: reported by ~35% in both drug and placebo groups; case reports of severe	bisphosphonates	
risk for hypocalcemia	Infection		Osteonecrosis of the jaw	
risk factors for hypocalcemia:	increase in serious infections	pain	-	
hypoparathyroidism, thyroid or parathyroid surgery, excision of	leading to hospitalization: ENT, GI, cellulitis; absolute risk		associated with invasive dental procedures such as tooth	
small intestine, malabsorption	increase 0.6% over 1-3 years	Biosimilar	extraction: consider preventive	
syndromes, CrCl < 30 mL/min or	caution: patients on	approval based on: comparative	dentistry/regular dental monitoring	
dialysis, previous hypocalcemia hypocalcemia symptoms: muscle	glucocorticoids with active	pharmacology, efficacy, safety	incidence: 0.05 – 0.7% over 7 –10	
	infection or history of recurrent or	efficacy endpoint in confirmatory	years	
cramps or twitching, numbness or tingling mouth, fingers or toes	chronic infection	clinical study: BMD at 1 year in postmenopausal females	additive risk factors: higher dose	
	Dermatologic	approval was extended to all	oncology regimens, longer duration of therapy	
See reference list	rashes, dermatitis, eczema: uncommon; discontinue if severe	denosumab indications		

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The fracture evidence for bisphosphonates and denosumab varies by patient population

- Populations: most osteoporosis medication clinical trials enroll community dwelling, ambulatory, postmenopausal females – evidence for other populations is more limited
- Postmenopausal females: bisphosphonates and denosumab reduce the risk of hip, other clinical, and vertebral fractures
- Comparisons between bisphosphonates and denosumab: indirect comparisons find that denosumab reduces the risk of radiographic vertebral fractures compared to bisphosphonates but not other clinical or hip fractures – however there isn't a large fracture trial comparing the two directly

- Females receiving aromatase inhibitors: denosumab reduces the risk of clinical fractures
- Males receiving androgen deprivation therapy: denosumab reduces the risk of radiographic vertebral fractures
- Males with osteoporosis and people taking glucocorticoids: bisphosphonates may reduce the risk of radiographic vertebral fractures but estimates of effect are imprecise
- Time-to-benefit: incorporate time to benefit estimates when sharing decisions with people of advanced age or limited life expectancy

Develop the exit strategy or transition plan at the time of medication initiation

- Bisphosphonates: Osteoporosis Canada suggests to review after 3 years of treatment
- Denosumab: Osteoporosis Canada suggests long-term uninterrupted therapy; if discontinuing, transition to a bisphosphonate or seek advice from a consultant with expertise in osteoporosis (depending on duration of therapy)
- Teriparatide: approved for a maximum of 24 months of use for most people, guidelines recommend subsequent treatment with a bisphosphonate or denosumab
- Romosozumab: approved for a maximum of 12 months of use, guidelines recommend subsequent treatment with a bisphosphonate or denosumab
- Optimal sequence of osteoporosis medications: there is limited evidence examining the optimal sequence of medications on fracture outcomes

Reference list is available upon request. Materials are designed to be used in conjunction with an academic detailing session provided by a PAD pharmacist. For more information, or to schedule an academic detailing session, please contact: BC Provincial Academic Detailing Service Email: PAD@gov.bc.ca Web: www.bcpad.ca

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