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2022 American College of Rheumatology Guideline for the Prevention and Treatment of Glucocorticoid-Induced Osteoporosis

Guideline Summary

A panel of adult and pediatric rheumatologists and endocrinologists updated the systematic literature review and included currently available medications for the prevention and treatment of glucocorticoid (GC)-induced osteoporosis. A patient panel was included in this update.

Similar to the 2017 guideline, we recommend risk stratifying patients as being at low, moderate, or high risk of fracture (Adults ≥40 years, FRAX® 10-year probability of major osteoporotic fracture <10%, 10-19%, or ≥20% respectively). We added a **very high risk category** (prior osteoporotic fracture(s) or bone mineral density (BMD) T score ≤ -3.5 or FRAX (GC-adjusted) 10-year risk of MOF ≥30% or hip ≥4.5% or high GC ≥30 mg/day for >30 days or cumulative doses ≥5 g/year. These cut points were used to stratify PICO questions and weigh potential benefits versus harms, when considering osteoporosis (OP) therapy. For all adults initiating or continuing GC therapy ≥2.5mg/day for >3 months, who have never had fracture risk assessment or been treated with OP therapy, initial clinical fracture risk assessment is strongly recommended over no assessment. Clinical fracture risk factor assessment includes the dose, duration, and pattern of GC use, alcohol use, smoking history, hypogonadism, history of prior fractures, low body weight, significant weight loss, parental history of hip fracture, fall history, thyroid disease, hyperparathyroidism, rheumatoid arthritis, malabsorption, chronic liver disease, inflammatory bowel disease, and height loss. If available, BMD testing with VFA or spinal x-ray is recommended as soon as possible after starting GC therapy for adults and every 1-2 years thereafter while continuing GC therapy.

A strong recommendation was made to use oral bisphosphonates (BPs) over no treatment for adults ≥40 years receiving long-term GCs, at high and very high risk for fracture, based on available fracture data in GIOP populations. Other agents including intravenous BPs, PTH/PTHrP, and denosumab (DEN) are also options and are conditionally recommended given lack of fracture prevention data in GIOP populations. For adults at high risk, we conditionally recommended DEN or PTH/PTHrP over BP. For adults at very high risk, we conditionally recommended PTH/PTHrP over antiresorptives (BP, DEN). Raloxifene (RAL) and romosozumab (ROM) may be used in selected patients, after careful consideration of potential harms including thrombosis, stroke, and cardiovascular events.

Table 1: Definitions of selected terms used in the recommendations and upgraded position statements for GIOP

Term	Adults ≥40 years of age	Adults <40 years of age
Major	Non-traumatic or pathological	Non-traumatic or pathological
osteoporotic	fractures of the spine, hip, wrist,	fractures of the spine, hip, wrist, or
fracture (MOF)	or humerus humerus	
Clinical fracture	History of GC use, evaluation for	History of GC use, evaluation for
risk assessment	falls, fractures, frailty, secondary	falls, fractures, frailty, secondary

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	causes of OP, FRAX® with GC adjustment, BMD with VFA or spinal x-ray	causes of OP, BMD with VFA or spinal x-ray (FRAX® not validated at age <40 years)
Follow up risk	BMD with VFA or spinal x-ray	BMD with VFA or spinal x-ray every
assessment during	every 1-2 years during OP therapy	1-2 years during treatment
GC treatment	BMD with VFA or spinal x-ray	BMD with VFA or spinal x-ray every
GC treatment	every 1-2 years after OP therapy is discontinued	1-2 years after OP therapy is discontinued
FRAX® GC	If GC dose is >7.5 mg/day, multiply	Not applicable as FRAX® is not
correction	the 10-year risk of major	validate in this age group
	osteoporotic fracture by 1.15 and	
	the hip fracture risk by 1.2 *	
Very high fracture risk	Prior OP fracture(s) OR BMD T score ≤ -3.5 OR FRAX® (GC- Adjusted*) 10-year risk of MOF ≥30% or hip ≥4.5% OR	Prior fracture(s) OR GC ≥30mg/day OR cumulative ≥5grams/year
	High GC ≥30 mg/day for >30 days OR cumulative doses ≥5 g/year	
High fracture risk	BMD T score ≤ -2.5 but > -3.5 OR FRAX® (GC-Adjusted*) 10-year risk of MOF ≥20% but <30% or hip ≥3% but <4.5%	
Moderate fracture	FRAX® (GC-Adjusted) 10-year risk	Continuing GC treatment
risk	of MOF ≥10 and <20%, hip >1 and	≥7.5mg/day for ≥6 months AND
	<3% OR BMD T score between -1	BMD Z score < -3 OR significant
	and -2.4	BMD loss (> least significant
		change of DXA)
Low fracture risk	FRAX® (GC-Adjusted) 10-year risk of MOF <10%, hip <1 %, BMD > -1.0	None of the above risk factors other than GC treatment
Daniel III	Ad 11->40 1	A.I. Ita 140
Recommended	Adults ≥40 years at moderate or	Adults <40 years at moderate or
treatment	high risk of fracture	high risk of fracture
strategy		
Calcium and	Optimized intake of dietary and supplemental calcium and vitamin D	
Vitamin D	based on age-appropriate U.S. Recommended Dietary Allowances.	

	We strongly recommend OP	We conditionally recommend
Bisphosphonates	treatment for those at moderate,	treatment for those at moderate or
(BP)	high or very high risk of fracture.	very high risk of fracture with
Alendronate		oral or IV BP ^{\$} , PTH/PTHrP [%] or
(oral)	We strongly recommend oral BP	DEN ^{%&}
Risedronate	over no treatment in high and	
(oral)	very high fracture risk due to	We conditionally recommended
Ibandronate	fracture reduction in GIOP.	against RAL due to harms of VTE
(oral/ IV)		and fatal stroke or ROM due to
Zoledronic acid	We conditionally recommend	uncertain harms including
(IV)	PTH/PTHrP over anti-resorptives	increased myocardial infarction,
PTH & PTHrP	in patients at very high risk of	stroke and death
Agonists	fracture.	
Teriparatide		
(TER)		
Abaloparatide	We conditionally recommend	
(ABL)	DEN ^{\$&} or PTH/PTHrP over oral and	
Anti-RANKL	IV BP in high risk of fracture.	
Denosumab		
(DEN)		
Selective Estrogen	We conditionally recommend IV	
Receptor	BP, ROM, RAL over no treatment	
Modulator	in high and very high risk of	
Raloxifene (RAL)	fracture.	
Anti-sclerostin		
Romosozumab	In moderate risk, we conditionally	
(ROM)	recommend BP, DEN, or	
	PTH/PTHrP in no preferred order	
	among these agents.	
	Except in patients intolerant of	
	other agents, we conditionally	
	recommend <i>against</i> RAL due to	
	harms of VTE and fatal stroke or	
	ROM due to uncertain harms with	

BP = bisphosphonate; PTH = parathyroid hormone; PTHrP = PTH-related protein; RANKL = Receptor activator of NF- $\kappa\beta$ -Ligand; BMD = bone mineral density; GC= glucocorticoid; FRAX® = https://www.shef.ac.uk/FRAX/Tool.jsp; MOF = major osteoporotic fracture; * FRAX® GC correction example: if hip fracture risk is 2.0% multiply by 1.2 for adjusted risk = 2.4%; \$ = Use with caution in women who may become pregnant due higher potency and longer half-life in fetal bones; % Avoid in young adults with open growth plates; & Use with caution in women of child bearing potential due to potential fetal harm. Avoid pregnancy for 5 months after last dose.

increased myocardial infarction,

stroke and death

Osteoporosis Medications for Patients with Chronic Kidney Disease or Following Renal Transplant Bisphosphonates should generally not be used in patients with an eGFR < 35 ml/min. When eGFR is < 35 ml/min, the risk of renal osteodystrophy, including adynamic bone disease, osteomalacia, osteitis fibrosa

cystica and mixed uremic osteodystrophy, is increased. As such, metabolic bone disease expert evaluation for chronic kidney disease-mineral and bone disorder (CKD-MBD) is conditionally recommended to exclude these conditions. Hyperparathyroidism should also be assessed. Once excluded, no dose adjustment is needed when prescribing DEN, PTH/PTHrP, or romosozumab (ROM).

Table 2. Recommendations for initial treatment for prevention of GIOP in adults beginning long-term GC therapy

Recommendations for patients taking prednisone ≥2.5mg/day for >3months	Certainty of evidence	PICO evidence report basis	Page no(s). of evidence tables
For adults and children beginning or continuing chronic GC treatment at low, moderate, high, or very high risk of fracture, we conditionally recommend optimizing dietary and supplemental calcium and vitamin D in addition to lifestyle modifications (CAL/VIT D/LM)	Low or Very Low	1.1a,b,c-1.3a,b,c, 2.1-2.3, 7.16-7.26	6-8, 47-48, 63-65, 141- 144,148- 151
In adults ≥40 years (All additional recomme	endations are in	addition to CAL/VIT I	D/LM.)
For adults ≥40 years with HIGH or VERY HIGH fracture risk, we strongly recommend OP therapy over no treatment. Agents to use include oral BP#, IV BP\$, PTH/PTHrP\$, DEN\$, RAL or ROM.	Low or Very Low	1.4c-1.28c	6-50
For adults ≥40 years with VERY HIGH fracture risk, we conditionally recommend PTH/PTHrP over anti-resorptive (DEN, BP) treatment.	Low	1.13c-1.20c	49-50
For adults ≥40 years with HIGH fracture risk, we conditionally recommend PTH/PTHrP or DEN over BP treatment.	Low	1.13c-1.20c	49-50
For adults ≥40 years with HIGH or VERY HIGH fracture risk, we strongly recommend oral BP over no treatment.	Low	1.4c	8-18
For adults ≥40 years with HIGH or VERY HIGH fracture risk, we conditionally recommend using ROM or RAL in patients intolerant of other agents.	Very low	1.16c, 1.21c, 1.28c	50
For adults ≥40 years with HIGH or VERY HIGH fracture risk, we conditionally recommend <i>against</i> using two different OP medications.	Very Low	1.29-1.35	53-62
For adults ≥40 years with MODERATE fracture risk, we conditionally recommend against ROM except for in patients intolerant of other agents, due to risk of myocardial infarction, stroke or death.	Very Low	1.12b, 1.16b,1.17b,1.21b- 1.25b, 1.28b	40-41, 44- 47

For adults ≥40 years with LOW fracture risk, we strongly recommend <i>against</i> OP medications in addition to CAL/VIT D/LM	Very low	4.4a-4.13a	91-101
due to known risk of harms and no evidence of benefit.			
Adults receiving high-dose GC (initial dose ≥30	 mg/dav for >3	L 30 davs or cumulative	e dose ≥5
gm in 1 year)	,g, uu y 101 - 1	oo days or camalative	. 4000 =0
We conditionally recommend treating with	Low	6.1b6.19a	120-141
PTH/PTHrP over anti-resorptives.			
Oral BP are strongly recommended over no treatment.	Low	6.1b -6.19a	120-141
IV BP and DEN are conditionally	Low	6.1b -6.19a	120-141
recommended over no treatment.			
RAL and ROM are conditionally	Low	6.1b -6.19a	120-141
recommended in those intolerant of other			
agents.			
In adults <40 years (All additional recomme	endations are in	addition to CAL/VIT	D/LM.)
Adults <40 years with MODERATE fracture	Low or Very	2.4-2.22, 3.4-3.17	65-76, 79-
risk, we conditionally recommended oral or	low		84.
IV BP%, DEN%, or PTH/PTHrP therapy.			
Adults <40 years with MODERATE fracture	Very Low	2.9, 3.9	70, 87
risk, we conditionally recommended <i>against</i>			
using ROM due to risk of myocardial			
infarction, stroke or death.			
For adults with solid organ transplants, glome			and no
evidence of chronic kidney disease-mineral ar	nd bone disord	er (CKD-MBD) * or	
hyperparathyroidism	1.	T = 4 = 0.0	100.110
We conditionally recommend expert	Low	5.1-5.26	103-118
evaluation for CKD-MBD in renal transplant			
recipients.		54526	102.110
We conditionally recommend treatment with	Low	5.1-5.26	103-118
oral or IV BP, DEN, PTH/PTHrP, or RAL based			
on individual patient factors.	Vordou	5.9	112
We conditionally recommended <i>against</i> using ROM due to risk of myocardial	Very low	5.9	112
infarction, stroke or death.			
Children ages 4-17 years treated with GCs for	>2 months (lov	u and madarata rick)	
We conditionally recommended optimization	Very low	7.1a-7.4a	141-144
of dietary and supplementation of CAL and	Very low	7.14-7.44	141-144
VIT D as recommended by U.S. RDA			
depending on the age of the child.			
We conditionally recommended against	Very Low	7.5a	144
starting oral or IV BP due to low risk of OP	VCI y LOW	7.50	177
fractures in this age group.			
Children ages 4-17 years with an osteoporotic	fracture who	are continuing treatm	ent with
GCs at a dose of ≥0.1 mg/kg/day for >3 month	=	are communing treatm	CITE WILL

We conditionally recommend treating with	Very low	7.1b-7.2b	148-153
an oral or IV BP.			

GIOP = Glucocorticoid-induced OP, BMP = Bone mineral density, GC = glucocorticoids, BP = bisphosphonate, IV = intravenous, DEN = denosumab, PTH/PTHrP = parathyroid hormone/ parathyroid hormone related protein, RAL = raloxifene, ROM = romosozumab, OP = OP; PICO = Patients, intervention, comparison, outcome; CAL/VIT D/LM = calcium/ vitamin D/ lifestyle modifications; # strong recommendation based on fracture data, \$ conditional due to lack of fracture data, \$ who are not planning on pregnancy during the OP treatment period or are using effective birth control if sexually active; CAL = calcium; CVD = cardiovascular disease; OP = osteoporosis; PICO = Patients, intervention, comparison, outcome; GIOP = Glucocorticoid-induced OP; U.S. RDA = United States Recommended Dietary Allowances; CKD-MBD = chronic kidney disease-mineral and bone disorder; *includes osteomalacia, adynamic bone disease, osteitis fibrosa cystica, mixed uremic osteodystrophy

This updated guideline includes recommendations on abaloparatide (PTHrP) and romosozumab (ROM), which are newly available since the 2017 guideline. It also addresses sequential therapy, which was not addressed in the past. Patients should know that osteoporosis therapy with denosumab (DEN), teriparatide (PTH), PTHrP, or ROM will need sequential osteoporosis therapy to prevent bone loss after these drugs are discontinued. Recommendations for sequential therapies are based in part on initial study designs, long term follow-up studies, and new clinical trials. Patients completing a course of DEN should transition to 1-2 years of a BP. Patients completing a course of PTH, PTHrP, or ROM need to transition to a BP or DEN. Discontinuation of DEN after two or more doses has been associated with rapid bone loss and development of new vertebral compression fractures as soon as 7-9 months after the last dose. As such, BP therapy is recommended beginning at 6-7 months after the last dose of DEN. The precise timing, dose and duration of BP use after DEN cessation is under study, but treatment for at least 1 year seems prudent, until additional research is available. Stopping PTH/PTHrP without transition to another therapy can also result in bone loss, which can be prevented by institution of oral or IV BP or DEN. Stopping ROM without transition to another therapy can result in bone loss, which can be prevented by the institution of oral or IV BP.

Sequential Treatments Recommended When Initial OP Therapy and GC are Discontinued and at Low or Moderate Risk

Initial OP therapy	Subsequent OP therapy options
Oral/IV Bisphosphonate	No subsequent OP therapy needed
RAL	No subsequent OP therapy needed
PTH/PTHrP	Oral or IV Bisphosphonate
Denosumab	Oral or IV Bisphosphonate
Romosozumab	Oral or IV Bisphosphonate

Sequential treatments when a new fracture occurs after ≥12 months of initial OP therapy.

Initial OP therapy	Subsequent OP therapy options
Oral/IV Bisphosphonate	DEN, PTH/PTHrP, ROM
RAL	Oral or intravenous BP, PTH/PTHrP, DEN, ROM
PTH/PTHrP	Oral or intravenous BP, DEN
Denosumab	Oral or IV Bisphosphonate, ROM
Romosozumab	Oral or IV Bisphosphonate, DEN

Sequential Treatments Recommended When Initial OP Therapy and GC are Discontinued and Patient Remains High or Very High Risk

Continue current therapy or switch to IV Bisphosphonate, DEN, PTH/PTHrP, or romosozumab

This summary was initially approved by the ACR Board of Directors on August 13, 2022 and updated on November 9, 2022 and July 9, 2023. These recommendations are included in a full manuscript, which was submitted, peer review pending, for publication in Arthritis & Rheumatology and Arthritis Care and Research.