OSTEOPOROSIS MANAGEMENT AMONG CHRONIC GLUCOCORTICOID USERS: A SYSTEMATIC REVIEW

Jordan M. Albaum¹, Soyoung Youn¹, Linda E. Lévesque², Andrea S. Gershon³, Suzanne M. Cadarette¹

¹Leslie Dan Faculty of Pharmacy, University of Toronto, Toronto ON, Canada; ²Queen's University, Kingston ON, Canada; ³Sunnybrook Health Sciences Centre, Toronto ON, Canada

Corresponding Author: Jordan.albaum@mail.utoronto.ca

ABSTRACT

Background

Clinical practice guidelines recommend that all patients starting chronic oral glucocorticoid (GC) therapy receive bone mineral density (BMD) testing and osteoporosis pharmacotherapy.

Objective

We completed a systematic review of observational studies to examine the proportion of patients on chronic oral GC therapy who receive osteoporosis management.

Methods

Two independent reviewers completed a systematic search of Ovid MEDLINE[®] and EMBASE[®] to identify all English language articles that examined the prevalence of osteoporosis management among chronic oral GC users. Clinical trials, abstracts, reviews, commentaries, and letters to the editor were excluded. Study methods and results (use of BMD testing and osteoporosis pharmacotherapy) were abstracted and summarized by year and region.

Results

We identified 29 eligible studies published between 1999 and October 2013: 17 were conducted in North America, 5 in Europe, and 7 in other regions. Heterogeneity between patient populations and methods used to define chronic GC use precluded the direct comparison of results between regions, or over time. Over 80% of studies identified that < 40% of chronic oral GC users received BMD testing or osteoporosis pharmacotherapy. When results of these studies were plotted by year, there was little evidence of improvement in osteoporosis management over time.

Conclusions

Despite consistent recommendations to target osteoporosis prevention at the onset of chronic oral GC therapy, osteoporosis is undermanaged among chronic oral GC users. Targeted interventions are needed to help reduce the burden of fracture-related morbidity associated with GC-induced osteoporosis.

Key Words: Osteoporosis, glucocorticoids, practice patterns, systematic review

Oral glucocorticoids (GC) are commonly prescribed to reduce pain and inflammation in patients with inflammatory arthritis, inflammatory bowel disease, and chronic lung disease.¹ The prevalence of oral GC use is approximately 1.2% among adults aged 20 years or more and it has been estimated that around 7.5% of adults aged 18 years or more have received at least one prescription for an oral GC.²⁻⁴ Due to the severe pain and morbidity associated with chronic inflammatory conditions, the duration of GC therapy can persist for months to years in length.⁴

Chronic oral GC therapy is the leading cause of secondary osteoporosis, a condition labeled GC-induced osteoporosis.⁵ Though there is no single definition of "chronic," it is

commonly accepted that treatment lasting at least 3 months leads to deleterious effects on bone.⁶ Oral GC therapy leads to a rapid reduction in bone formation through inhibition of osteoblast differentiation⁷ and increased osteoclast activity.^{8,9} This results in bone loss which manifests clinically as reduced bone mass and diminished microarchitectural integrity.¹⁰ GC-induced bone loss occurs rapidly at a rate of 6% to 12% within the first year of therapy.¹¹ Fracture risk increases within 3 months of starting therapy with doses as low as 2.5 mg/day prednisone equivalent^{12,13}, and individuals on GC therapy are almost twice as likely to experience a bone fracture compared to non-GC users.³

Fortunately, treatment with osteoporosis medication has shown to improve bone mineral density (BMD) and reduce fracture risk in patients treated with oral GC therapy.¹⁴ As a result, clinical practice guidelines recommend that BMD testing and osteoporosis pharmacotherapy be initiated in patients starting oral GC therapy for \geq 3 months.^{6,15} This recommendation is consistent across osteoporosis guidelines, though the duration (3 to 6 months) and dose (any to 15 mg/day) indicating osteoporosis management differ slightly.¹⁵⁻²²

Since clinical practice guidelines frequently change to reflect the availability of new therapeutic options for osteoporosis, there is little known about how GC-induced osteoporosis management has changed over time, or if regional differences exist. An investigation of studies examining the proportion of chronic GC users that receive osteoporosis management may clarify current practice standards among this high-risk population, and help identify areas for improvement. Thus, we aimed to systematically examine the proportion of chronic oral GC users receiving osteoporosis management, by region and over time. A secondary objective was to identify patient and physician-level predictors of GC-induced osteoporosis management among identified studies.

METHODS

This systematic review follows PRISMA guidelines for the reporting of systematic reviews.²³

Search Strategy

Two reviewers (JMA, SY) independently completed a systematic search of the electronic database Ovid MEDLINE[®] from 1946 (database inception) to October 2013 to identify all English language articles that examined GC-induced osteoporosis management. Ovid EMBASE[®] was also searched through to October 2013. Search terms for "osteoporosis" and "glucocorticoids" were adapted from two separate Cochrane Collaboration reviews^{24,25}, and selected following consultation with a library scientist. A complete list of terms can be found in Appendix 1.

Identification of Relevant Articles

All observational studies that examined the proportion of chronic oral GC users receiving osteoporosis management (BMD testing and osteoporosis pharmacotherapy) were eligible. Studies that included users of inhaled or injectable GCs were included only if management outcomes were reported separately for oral GC users. Eligible osteoporosis pharmacotherapy included bisphosphonates, calcitonin, denosumab, hormone replacement therapy, raloxifene, teriparatide, and testosterone. Clinical trials, commentaries, letters to the editor, reviews, clinical practice guidelines, and abstracts were excluded. Articles that did not state the drug class of pharmacotherapy or that considered calcium/vitamin D as "pharmacotherapy" were excluded since all guidelines recommend drug therapy in addition to calcium/vitamin D to manage osteoporosis. In addition, articles that reported osteoporosis pharmacotherapy only or that did not report use of BMD testing and treatment separately were excluded since all guidelines recommend BMD testing at the onset of chronic oral GC therapy. Titles and abstracts were reviewed independently by two authors (JMA, SY) and inconsistencies settled through consultation with a third author (SMC).

Data Abstraction

Study characteristics including data source, patient demographics, country, GC dose, GC indication, methods, and results were abstracted and summarized by one author (JMA) and verified by a second (SY). Patient characteristics and study methodology were compared between studies. The main osteoporosis clinical practice guidelines, referenced by each included study, were also summarized. Results of eligible studies were plotted by publication and "data year," based on the reported study period. For study periods spanning less than 3 years, data year was defined at the end of the study period. Data year for studies extending 3 or more years was defined as the middle year of the study. For example, studies using data collected from 2001 to 2002 and 2001 to 2003 were both assigned the data year 2002. For studies that reported several data points longitudinally, each reported year was plotted.

RESULTS

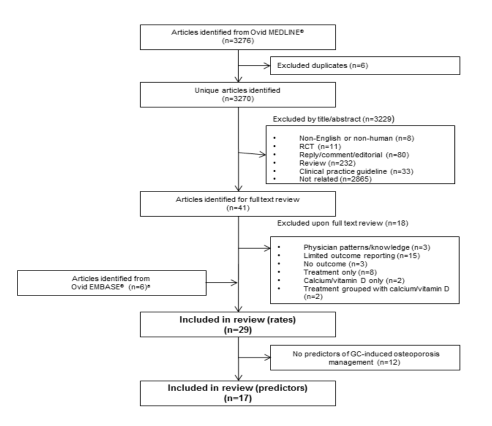
Of 3270 unique articles identified, 41 were retrieved for full text review based on title and abstract screening, Figure 1. After full text review, 18 additional articles were excluded: 3 focused on physician knowledge/prescribing habits²⁶⁻²⁸, 3 reported no outcome²⁹⁻³³, 8 reported osteoporosis pharmacotherapy only³⁴⁻⁴⁴, 2 reported use of calcium/vitamin D only^{45,46}, and 2 included calcium/vitamin D as pharmacotherapy.^{47,48} An additional 6 studies identified using EMBASE® were eligible.

In total, 29 papers published between 1999 and 2013 were eligible: 17 from North America⁴⁹⁻⁶⁵, 5 from Europe⁶⁶⁻⁷⁰, and 7 from other regions (Australia, 2; South Africa, 1; India, 3; Saudi Arabia, 1).⁷¹⁻⁷⁷ These studies referenced several osteoporosis guidelines to inform their methodology including 1996, 2001, and 2010 American College of Rheumatology¹⁵⁻¹⁷, 1998 UK consensus group¹⁸, 1998 and 2002 National Osteoporosis Society^{19,20}, 2003 American Gastroenterological Association²¹, and 2000 South African Medical Association.²² A summary of these guidelines is presented in Table 1.

A variety of data sources were used by these studies to examine GC-induced osteoporosis

management, Table 2. Chart review was the most common source (n=13), followed by administrative databases (n=10) and telephone surveys (n=1). Five studies (17%) used a combination of different data sources. Furthermore, methods used to define chronic GC exposure and report osteoporosis management were not consistent with osteoporosis guidelines, and varied considerably across studies, Table 3. While 83% of studies assessed osteoporosis management in accordance with guideline recommendations, only 45% used guideline criteria to define chronic GC use. When defining osteoporosis pharmacotherapy, all studies included bisphosphonates, while fewer considered the use of hormone replacement therapy (62%), calcitonin (59%), testosterone (28%), raloxifene (24%), and teriparatide (7%). This made it difficult to compare results directly between studies.

FIG. 1 Flow diagram of systematic search results



^aElectronic search of Ovid EMBASE[®] (completed July 2014 in response to reviewer comments) yielded 6077 articles published between 1964 and October 2013. After thorough review, 6 additional articles were included in our systematic review.

	American Co	llege of Rheuma	tology (USA)	American GI Association (USA)	UK Consensus Group (UK)			South Africa
Variable	1996	2001	2010	2003	1998	1998	2002	2000
Treatment Indications								
Prevalent Fracture	Yes	Yes	FRAX	Yes	Yes	Yes	Yes	Yes
GC Dose (mg/day) ^a	NR	5.0	7.5	7.5	15.0	7.5	NR	7.5
GC Duration	≥ 6 months	≥ 3 months	≥ 3 months	> 3 months	≥ 6 months	≥ 6 months	≥ 3 months	≥ 3 months
Screening (BMD Testing	g)							
BMD Testing	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes
Repeat Frequency	0.5 – 1 year	6 months	1 year	1 – 2 years	1 – 3 years	1 – 3 years	1 – 3 years	1 – 2 years
Treatment Options								
Calcium (mg/day) ^b	1500	NR°	1200	1000 – 1500	NR ^c	NR ^c	NR ^c	1000
Vitamin D (IU/day) ^b	800	800	800	800	NR°	NR^{c}	NR°	400
Pharmacotherapy								
Bisphosphonate								х
Alendronate		х	х	х			х	
Etidronate	х				х	х	х	
Pamidronate	х			Х			х	
Risedronate		х	х	Х			х	
Zoledronic Acid			х					
Calcitonin	x	х			х		х	
HRT or T	х	х			х	х	х	
Raloxifene								
Teriparatide			х					

TABLE 1 International guidelines for glucocorticoid-induced osteoporosis management

BMD – bone mineral density, FRAX – WHO fracture risk assessment tool, GC – glucocorticoid, GI – gastroenterological, HRT – hormone replacement therapy, T – testosterone, NR – not reported

^aGlucocorticoid doses given as prednisone equivalent

^bRecommended minimum intake refers to the total intake (diet and supplementation)

^cRecommended, yet dose is not stated in the guideline

TABLE 2	Characteristics o	f studies	identified	through	systematic s	search (n=29)
---------	-------------------	-----------	------------	---------	--------------	---------------

			Main Guideline(s)	Patients, N	Mean age,	GC Indication,	Mean GC dose ^a ,	Pharmacotherapy
Author (year)	Country	Data Source	Referenced	(% female)	years	included (main)	mg/day	Identified ^b
North America (n	=17)							
Buckley (1999)	USA	Telephone survey	1996 ACR	147 (58)	51.0	Any (rheumatic)	10.0	bisphosphonate, calcitonin, HRT
Elliot (2000)	USA	Administrative database & chart review	1996 ACR	72 (0)	57.0	Any (transplant)	12.5	bisphosphonate, calcitonin, testosterone
Osiri (2000)	USA	Chart review	1996 ACR	365 (59)	52.5	Any (rheumatic)	11.4	bisphosphonate, calcitonin, HRT, testosterone
Ettinger (2001)	USA	Administrative database	1996 ACR 1998 UK	8,807 (60)	NR	Any (respiratory)	NR	bisphosphonate, calcitonin
Mudano (2001)	USA	Administrative database	1996 ACR	6,821 (62)	51.9	Any (rheumatic)	13.6	bisphosphonate, calcitonin, HRT, testosterone

			Main Guideline(s)	Patients, N	Mean age,	GC Indication,	Mean GC dose ^a ,	Pharmacotherapy
Author (year)	Country	Data Source	Referenced	(% female)	years	included (main)	mg/day	Identified ^b
Yood (2001)	USA	Administrative	1996 ACR	224 (57)	70.0	Any (respiratory)	8.9	bisphosphonate,
		database &						calcitonin, HRT
		chart review						
Solomon (2002)	USA	Chart review	2001 ACR	236 (80)	60.3	Rheumatic	8.8	bisphosphonate,
								calcitonin, HRT,
								raloxifene
Curtis (2005)	USA	Administrative	1996 ACR	6,281 (66)	50.0	Any, excluding	16.0	bisphosphonate,
		database &	2001 ACR			transplantation		calcitonin, HRT,
		mailed survey				(rheumatic)		raloxifene,
								testosterone
Feldstein (2005)	USA	Administrative	2001 ACR	3,031 (60)	61.4	Any (respiratory)	20.0	bisphosphonate,
		database						calcitonin, raloxifene,
								HRT
Che (2006)	USA	Administrative	2001 ACR	13,862 (58)	NR	Any	NR	bisphosphonate,
		database						calcitonin, raloxifene
Cruse (2006)	USA	Chart review	2001 ACR	370 (0)	64.0	Any (rheumatic)	NR	bisphosphonate,
								calcitonin,
								testosterone
Liu (2006)	USA	Chart review	2001 ACR	35 (60)	54.0	Dermatologic	53.0 ^c	bisphosphonate
, , , , , , , , , , , , , , , , , , ,						C C		

			Main				Mean GC	
Author (year)	Country	Data Source	Guideline(s) Referenced	Patients, N (% female)	Mean age, years	GC Indication, included (main)	dose ^a , mg/day	Pharmacotherapy Identified ^b
Saag (2006)	USA	Administrative database	2001 ACR	3,125 (59-63) ^d	NR	Any (respiratory)	11.0	bisphosphonate, calcitonin, raloxifene, HRT
Guzman-Clark (2007)	USA	Chart review	2001 ACR	100 (6)	73.0	Any (respiratory)	7.5 (median)	bisphosphonate
Ledwich (2009)	USA	Chart review	2001 ACR	73 (82)	60.9	Rheumatic	NR	bisphosphonate, HRT
Majumdar (2012)	Canada	Administrative database	none	15,825 [°] (58)	60.0	Any	NR	bisphosphonate, calcitonin, raloxifene, teriparatide
Thanou (2013)	USA	Administrative database & chart review	2003 AGA	63 (3)	55.0	Inflammatory bowel disease	15.0 (median)	bisphosphonate
Europe (n=5)								
Erb (2002)	UK	Chart review	1998 NOS	235 (71)	NR	Rheumatic	NR	Bisphosphonate, calcitonin, HRT, raloxifene, testosterone

			Main				Mean GC	
			Guideline(s)	Patients, N	Mean age,	GC Indication,	dose ^a ,	Pharmacotherapy
Author (year)	Country	Data Source	Referenced	(% female)	years	included (main)	mg/day	Identified ^b
Gudbjornsson	Iceland	Chart review &	1996 ACR	191 (55)	66.0	Any (rheumatic)	6.0	bisphosphonate,
(2002)		mailed survey	1998 UK					calcitonin, HRT,
								testosterone
Walker-Bone	UK	Chart review	1998 UK	175 (76)	Male: 64.2	Rheumatic	Male: 7,816.5	bisphosphonate, HRT
(2004)					Female: 66.9		Female: 9,465	
							(median	
							cumulative)	
Wall (2008)	UK	Chart review	2002 UK	104 (74)	61.8	Rheumatic	8.6	bisphosphonate,
() ()								HRT, teriparatide
Haroon (2011)	Ireland	Chart review	2002 NOS	81 (64 ⁹)	62.0	Rheumatic	NR	bisphosphonate
Other (n=7)				f				
Hougardy	Australia	Chart review	1998 NOS	212 (58) [†]	69 (median)	Any (respiratory)	10.0 (median)	bisphosphonate, HRT
(2000)								
Rothberg (2000)	South	Administrative	2000 SAMA	1,614 (54)	Male: 51.0	Any (respiratory)	NR	bisphosphonate, HRT
	Africa	database			Female: 53.0			
Smith (2001)	Australia	Chart review	1996 ACR	189 (38)	75.2	Any (respiratory)	NR	bisphosphonate, HRT
. ,				· · ·				

Author (year)	Country	Data Source	Main Guideline(s) Referenced	Patients, N (% female)	Mean age, years	GC Indication, included (main)	Mean GC dose ^a , mg/day	Pharmacotherapy Identified ^b
Gera (2009)	India	Chart review	2002 NOS	105 (64)	42.0	Any (rheumatic)	NR	bisphosphonate, calcitonin, HRT, testosterone
Sadat-Ali (2009)	Saudi Arabia	Administrative database	2001 ACR	165 (39)	Male: 37.0 Female: 40.8	Any (rheumatic)	Male: 15.9 Female: 21.5	bisphosphonate, calcitonin, HRT
Srinivasulu (2010)	India	Administrative database	2001 ACR	151 (58)	52.5	Any (rheumatic)	NR	bisphosphonate
Kohli (2013)	India	Administrative database	2010 ACR	203 (60)	50.5	Any (rheumatic)	NR	bisphosphonate

ACR – American College of Rheumatology, AGA – American Gastroenterological Association, GC – glucocorticoid, HRT – hormone replacement therapy, NOS – National Osteoporosis Society, NR - not reported, SAMA - South American Medical Association.

^a All doses reported in mg prednisone equivalent ^b All studies also reported the proportion of patients who received a BMD test

^c Highest daily dose on file

^d 1996-1997 (62%), 1998-1999 (63%), 2000-2001 (59%)

^e Unique patients identified from 17,736 new long-term systemic GC initiations

^f Total GC users (inhaled + oral); approximately 53.3% were on oral GC therapy only, and 74.1% on combined oral + inhaled GC therapy

⁹% female only reported for 2009 cohort (n=34)

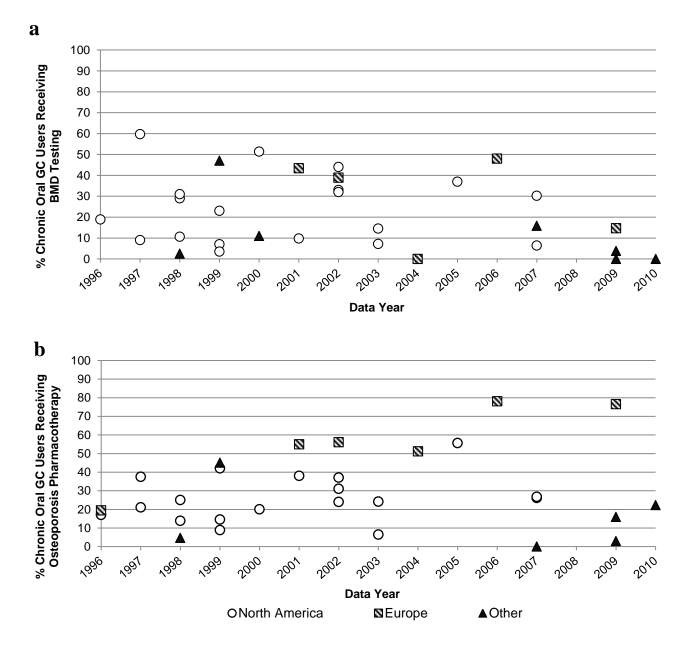
TABLE 3 Summary of study methods (n=29)

		Tota	l (n=29)	North A	merica (n=17)	Eur	ope (n=5)	Other (n=7)	
		Ν	%	Ν	%	Ν	%	Ν	%
Chronic G	lucocorticoid Expo	osure Def	inition ^a						
Duration	Dose								
1 month	10 mg/day	1	3.4	1	5.9	0	0	0	0
2 months	Not specified	1	3.4	1	5.9	0	0	0	0
3 months	5 mg/day	4	13.8	3	17.6	0	0	1	14.3
	7.5 mg/day	4	13.8	1	5.9	0	0	3	42.9
	Not specified	6	20.7	2	11.8	3	60.0	1	14.3
6 months	5 mg/day	1	3.4	1	5.9	0	0	0	0
	7.5 mg/day	6	20.7	2	11.8	2	40.0	2	28.6
Any	2 g	2	6.9	2	11.8	0	0	0	0
-	5 mg/day	1	3.4	1	5.9	0	0	0	0
Any ^b	Any	2	6.9	2	11.8	0	0	0	0
1 prescripti	on/quarter	1	3.4	1	5.9	0	0	0	0
Outcome F	Reporting (Pharma	cotherap	у)						
Drugs incl	uded								
Bisphosp		29	100.0	17	100.0	5	100.0	7	100.0
Calcitonir		17	58.6	13	76.5	2	40.0	2	28.6
HRT		18	62.1	10	58.9	4	80.0	4	57.1
Raloxifen	е	7	24.1	6	35.3	1	20.0	Ō	0
Teriparati		2	6.9	1	5.9	1	20.0	0	0
	Testosterone		27.6	5	29.4	2	40.0	1	14.3
Drug repo	rting								
	macotherapy	8	27.6	7	41.2	1	20.0	0	0
Stratify by		7	24.1	3	17.6	1	20.0	3	42.9
Both	-	14	48.3	7	41.2	3	60.0	4	57.1

HRT – Hormone Replacement Therapy

^aGlucocorticoid doses given as prednisone equivalent ^bResults stratified by dose

FIG. 2 Reported GC-induced osteoporosis management rates among chronic oral GC users, by region over time, (a) BMD testing (n=28), one study⁶⁷ reported no dual x-ray absorptiometry (DXA) machines at the time of study and is not reported here; (b) osteoporosis pharmacotherapy (n=27), two studies^{51,71} did not report total pharmacotherapy and are not reported here. Year was identified by period of time examined in each study and thus does not match publication year.



Prevalence of GC-Induced Osteoporosis Management

In the articles reviewed, the proportion of patients reported to have received BMD testing ranged from 0% to 60%, and pharmacotherapy ranged from 0% to 78%, Figure 2. Over 80% of studies identified that < 40% of chronic oral GC users received BMD testing or osteoporosis treatment.

Only three studies examined trends over time.^{54,56,57} Ettinger et al. (2001) identified an increased prevalence of osteoporosis medication prescribing to a high of 9% over an 18-month time period from January 1999 to June 2000 in California, USA.⁵⁴ However, this increase was attributed to selecting only patients without prior exposure to osteoporosis medication. Saag et al. (2006) similarly described an increase in GCinduced osteoporosis management among older women (aged \geq 65) in the United States from 1996 to 2001. BMD testing increased from 10% to 19% and osteoporosis treatment increased from 24% to 44%. However, little improvement (< 6%) was noted among men and younger women.⁵⁶ More recently, Majumdar et al. (2012) reported an increase in BMD testing or osteoporosis treatment in Manitoba, Canada from 17% to 27% between 1998 and 2008, yet note that the trend appeared to plateau in 2002.⁵⁷

Predictors of GC-Induced Osteoporosis Management

Seventeen studies (59%) identified predictors of GC-induced osteoporosis management, 9 as a primary study objective. Patient sex (88%), patient age (71%), and provider specialty (65%) were the most frequently examined predictors of GC-induced osteoporosis management. Thirteen of the fifteen studies (87%) examining sex identified female sex to be significantly associated with osteoporosis management^{50-60,71,72}, and seven of twelve studies (58%) that investigated age identified older age as a statistically significant predictor.^{50,52-57} Ten of eleven studies (91%) that examined provider specialty reported that provided rheumatologists osteoporosis management more frequently than general practitioners and other specialists.^{50-54,56-58,61,74} Other statistically significant predictors of GCinduced osteoporosis management included: prior fracture^{50,52,54,56,71}, prior BMD test^{54,71,74}, race,^{50-52,65} post-menopausal status^{50,51,59}, greater number of comorbidities,^{52,59} new GC use⁵², higher cumulative GC dose⁵², longer duration of GC use^{50,65,71}, and greater number of healthcare visits.⁵³ Socio-demographic (residence, income, insurance, education)^{51,53,57}, behavioural (GC knowledge, tobacco use)^{51,59}, and provider-related (demographic, training)⁵⁹ factors were also examined by some studies; yet, none of these were statistically significant predictors of GCinduced osteoporosis management.

DISCUSSION

Chronic oral GC therapy is the leading cause of secondary osteoporosis,⁵ and osteoporotic fracture is associated with significant morbidity and mortality.⁷⁸⁻⁸¹ As a result, many organizations have published guidelines outlining the risk of chronic oral GC therapy and consistently testing and osteoporosis recommend BMD pharmacotherapy to minimize bone loss and fracture risk. Despite reduce these recommendations. our systematic review identified low levels of GC-induced osteoporosis management. Furthermore, articles identified in this systematic review provide little evidence that GC-induced osteoporosis management is improving over time, particularly in recent years.

Three studies described an increase in GC-induced osteoporosis management since the late 1990s^{54,56,57}, yet, report that the proportion of chronic GC users who received BMD testing or osteoporosis treatment remains suboptimal (< 30%) in later years. There is also some evidence suggest that GC-induced osteoporosis to management is more common in Europe compared to North America and other regions. All studies completed in Europe after 2002 report rates > 50% for osteoporosis pharmacotherapy. However. these were based entirely in rheumatology clinics where GC-induced osteoporosis management has been observed more commonly than at the general population level.⁵⁰⁻ 54,56-58,61,74 This disparity may indicate a possible disconnect between osteoporosis guidelines and clinical practice.

While 83% of studies assessed osteoporosis management according to guideline recommendations, only 45% used guideline criteria to identify their chronic GC user population. This produced heterogeneity in methods used to define chronic GC use and osteoporosis management that preclude the direct comparison of results between studies. Studies also described highly diverse patient populations that varied substantially in age, sex, and comorbidity, making it difficult to compare the consistency of predictor variables between studies.

Though the majority of studies identify female sex as a predictor of GC-induced osteoporosis management, both studies that exclusively examined men reported treatment prevalence estimates (31% Cruse et al. and 38% Elliot *et al.*)^{49,61} consistent with other studies that included both men and women. In addition, there was no apparent relationship between average patient age (range 37 to 75 years) and rates of osteoporosis management. Several studies noted that patients between the age of 50 and 70 were more likely to receive a BMD test compared to patients under the age of 50, while patients over the age of 70 were more likely to be treated for osteoporosis compared to younger patients.^{52,56,57} This finding may represent an important shift in a physician's priorities from testing to treatment in older patients where age-related bone loss may be more apparent.

Recent reports of adverse events,^{82,83} particularly with bisphosphonates, may contribute to the challenge in managing osteoporosis. In addition, patients requiring chronic oral GC therapy often have disease indicating GC therapy as several additional well as chronic comorbidities. Thus, in the context of chronic oral GC therapy, osteoporosis management may be more difficult to coordinate due to the number of physicians involved in caring for patients with multiple chronic comorbidities. This combination of factors has been noted in similar clinical areas including post-hip fracture osteoporosis management.84

Future work aimed at improving our understanding of patients treated with chronic oral GC therapy may help identify barriers to treatment and provide insight into why GCinduced osteoporosis management is suboptimal. In turn, this may help researchers develop targeted interventions to improve GC-induced osteoporosis management.

There are several limitations that must be noted for our study. First, we acknowledge that some observational studies relating to GC-induced osteoporosis management may have been missed. To mitigate this risk, we used a comprehensive search strategy derived from previous Cochrane systematic reviews, used two search databases, and had two independent reviewers thoroughly conduct the search. Second, due to the heterogeneity between studies and patients, we are unable to comment on overall trends identified and determine the relative importance of predictor variables such as age, sex, and physician specialty for GC-induced osteoporosis management. Nonetheless. all studies consistently report GC-induced suboptimal management of osteoporosis.

CONCLUSION

Despite consistent guideline recommendations to target osteoporosis prevention at the onset of chronic oral GC therapy, results from our systematic review identify that the proportion of chronic GC users receiving osteoporosis management is low, with little evidence of improvement over time, particularly in recent years. Although it is unclear if this represents true mismanagement, it does indicate a missed opportunity for fracture prevention among chronically ill patients requiring long-term GC therapy. Future research should focus on improving our understanding of these patients to identify clinical barriers to treatment.

Acknowledgments

This research was supported by an Ontario Ministry of Research and Innovation Early Research Award and a University of Toronto Connaught New Researcher Award to Dr. Suzanne Cadarette. Dr. Cadarette was supported by a Canadian Institutes of Health Research New Investigator Award in Aging and Osteoporosis (MSH-95364). Jordan Albaum was supported by a Queen Elizabeth II Graduate Scholarship in Science and Technology. Mr. Albaum received funding support from the International Society for Pharmacoepidemiology to present this research at the 29th Annual International Conference on Pharmacoepidemiology and Therapeutic Risk Management in August 2013. Authors thank Joanna Bielecki, Research Librarian at the Leslie Dan Faculty of Pharmacy, University of Toronto for her helpful suggestions regarding systematic literature searches and Lindsay Wong, University of Toronto, Andrea Burden, BSc, MA, University of Toronto and Mina Tadrous, PharmD, MS, University of Toronto for insightful discussions.

APPENDIX 1

Search
1. *Osteoporosis/
2. osteoporos#s.tw.
3. bone loss\$.tw.
4. Bone Density/
5. (bone adj2 (density or fragil\$)).tw.
6. bone mass.tw.
7. bmd.tw.
8. exp Fractures, Bone/
9. fracture\$.tw.
10. Postmenopause/
11. (post menopaus\$ or post-menopaus\$).tw.
12. or/1-11
13. prednisone.tw,sh,rn.
14. glucocorticoid.rn,tw,sh.
15. corticosteroid\$.tw
16. glucocorticoids/
17. or/13-16
18. 12 and 17
19. limit 18 to (English language and humans)
20. limit 19 to clinical trial, all
21. 19 not 20

J Popul Ther Clin Pharmacol Vol 21(3):e486-e504; November 24, 2014 © 2014 Canadian Society of Pharmacology and Therapeutics. All rights reserved.

REFERENCES

- 1. Stahn C, Lowenberg M, Hommes DW, Buttgereit F. Molecular mechanisms of glucocorticoid action and selective glucocorticoid receptor agonists. Mol Cell Endocrinol 2007;275:71-78.
- 2. Diez-Perez A, Hooven FH, Adachi JD, et al. Regional differences in treatment for osteoporosis. The Global Longitudinal Study of Osteoporosis in Women (GLOW). Bone 2011;49:493-498.
- Donnan PT, Libby G, Boyter AC, Thompson P. The population risk of fractures attributable to oral corticosteroids. Pharmacoepidemiol Drug Saf 2005;14:177-186.
- 4. Overman RA, Yeh JY, Deal CL. Prevalence of oral glucocorticoid usage in the United States: a general population perspective. Arthritis Care Res 2013;65:294-298.
- DiPiro JT, Talbert RL, Gary C, Yee PDF. Pharmacotherapy: A Pathophysiologic Approach. (2008) McGraw-Hill Professional Publishing, <u>http://edoqs.com/dipiropharmacotherapy-7th-edition</u>, Accessed 15 August 2013.
- Papaioannou A, Morin S, Cheung AM, et al. 2010 clinical practice guidelines for the diagnosis and management of osteoporosis in Canada: summary. Canadian Medical Association Journal 2010;182:1864-1873.
- 7. Kim HJ, Zhao H, Kitaura H, et al. Glucocorticoids and the osteoclast. Ann NY Acad Sci 2007;335-339.
- 8. Ito S, Suzuki N, Kato S, et al. Glucocorticoids induce the differentiation of a mesenchymal progenitor cell line, ROB-C26 into adipocytes and osteoblasts, but fail to induce terminal osteoblast differentiation. Bone 2007;40:84-92.
- 9. Pereira RC, Delany AM, Canalis E. Effects of cortisol and bone morphogenetic protein-2 on stromal cell differentiation: correlation with CCAAT-enhancer binding protein expression. Bone 2002;30:685-691.
- 10. Weinstein RS. Clinical practice. Glucocorticoid-induced bone disease. N Engl J Med 2011;365:62-70.
- 11. LoCascio V, Bonucci E, Imbimbo B, et al. Bone loss in response to long-term glucocorticoid therapy. Bone and Mineral 1990;8:39-51.

- 12. van Staa TP, Leufkens HG, Abenhaim L, et al. Use of oral corticosteroids in the United Kingdom. QJM 2000;93:105-111.
- Zonana-Nacach A, Barr SG, Magder LS, Petri M. Damage in systemic lupus erythematosus and its association with corticosteroids. Arthritis Rheum 2000;43:1801-1808.
- 14. Lekamwasam S, Adachi JD, Agnusdei D, et al. An appendix to the 2012 IOF-ECTS guidelines for the management of glucocorticoid-induced osteoporosis. Arch Osteoporos 2012;7:25-30.
- 15. Grossman JM, Gordon R, Ranganath VK, et al. American College of Rheumatology 2010 recommendations for the prevention and treatment of glucocorticoid-induced osteoporosis. Arthritis Care Res 2010;62:1515-1526.
- 16. Recommendations for the prevention and treatment of glucocorticoid-induced osteoporosis. American College of Rheumatology Task Force on Osteoporosis Guidelines. Arthritis Rheum 1996;39:1791-1801.
- 17. Recommendations for the prevention and treatment of glucocorticoid-induced osteoporosis: 2001 update. American College of Rheumatology Ad Hoc Committee on Glucocorticoid-Induced Osteoporosis. Arthritis Rheum 2001;44:1496-1503.
- Eastell R, Reid DM, Compston J, et al. A UK Consensus Group on management of glucocorticoid-induced osteoporosis: an update. J Intern Med 1998;244:271-292.
- 19. National Osteoporosis Society. Guidance on the prevention and management of corticosteroid induced osteoporosis. (1998) National Osteoporosis Society, Bath, UK
- 20. Bone and Tooth Society, National Society, Royal College of Osteoporosis Physicians. Glucocorticoid-induced Osteoporosis: Guidelines for Prevention and Treatment. (2002) Royal College of Physicians, London
- 21. Bernstein CN, Leslie WD, Leboff MS AGA technical review on osteoporosis in gastrointestinal diseases. Gastroenterology 2003;124:795-841.
- 22. Hough S. Osteoporosis Clinical Guideline. South African Medical Association--Osteoporosis Working Group. S Afr Med J 2000;90:907-944.
- 23. Moher D, Liberati A, Tetzlaff J, Altman DG. Preferred reporting items for systematic

reviews and meta-analyses: the PRISMA statement. BMJ 2009;339:332-336.

- 24. O'Donnell S, Cranney A, Wells GA, et al. Strontium ranelate for preventing and treating postmenopausal osteoporosis. Cochrane Database of Systematic Reviews 2006; Art. No.: CD005326:
- 25. Cranney A, Welch V, Adachi JD, et al. Calcitonin for the treatment and prevention of corticosteroid-induced osteoporosis. Cochrane Database Syst Rev Art No: CD001983 2000.
- 26. Nielsen BR, Jorgensen NR, Schwarz P. Management of risk of glucocorticoid-induced osteoporosis due to systemic administration in general practice in Denmark. Eur J Gen Pract 2007;13:168-171.
- 27. Duyvendak M, Naunton M, van Roon EN, Brouwers JRBJ. Doctors' beliefs and knowledge on corticosteroid-induced osteoporosis: identifying barriers to improve prevention. J Clin Pharm Ther 2011;36:356-366.
- 28. Soucy E, Bellamy N, Adachi JD, et al. A Canadian survey on the management of corticosteroid induced osteoporosis by rheumatologists. J Rheumatol 2000;27:1506-1512.
- 29. Bell R, Carr A, Thompson P. Managing corticosteroid induced osteoporosis in medical outpatients. J R Coll Physicians Lond 1997;31:158-161.
- Lekamwasam S. A hospital-based study of prophylactic therapy in glucocorticoidinduced osteoporosis. Ceylon Med J 2010;55:44-46.
- 31. Kirigaya D, Nakayama T, Ishizaki T, et al. Management and treatment of osteoporosis in patients receiving long-term glucocorticoid treatment: current status of adherence to clinical guidelines and related factors. Intern Med 2011;50:2793-2800.
- 32. Brask-Lindemann D, Eiken P, Eskildsen P, Abrahamsen B. Time trends for alendronate prescription practices in women with chronic obstructive pulmonary disease and women exposed to systemic glucocorticoids. Osteoporos Int 2013;24:1891-1897.
- 33. Paskins Z, Potter T, Erb N, et al. Audits of the prevention and treatment of corticosteroidinduced osteoporosis in outpatients with rheumatic diseases in the West Midlands. Clin Med 2006;6:183-187.
- 34. Peat ID, Healy S, Reid DM, Ralston SH. Steroid induced osteoporosis: an opportunity

for prevention? Ann Rheum Dis 1995;54:66-68.

- 35. Walsh LJ, Wong CA, Pringle M, Tattersfield AE. Use of oral corticosteroids in the community and the prevention of secondary osteoporosis: a cross sectional study. BMJ 1996;313:344-346.
- 36. Aagaard EM, Lin P, Modin GW, Lane NE. Prevention of glucocorticoid-induced osteoporosis: provider practice at an urban county hospital. Am J Med 1999;107:456-460.
- 37. Hart SR, Green B. Osteoporosis prophylaxis during corticosteroid treatment: failure to prescribe. Postgrad Med J 2002;78:242-243.
- 38. Ryan JG, Morgan RK, Lavin PJ, et al. Current management of corticosteroid-induced osteoporosis: variations in awareness and management. Ir J Med Sci 2004;173:20-22.
- 39. Curtis JR, Westfall AO, Allison J, et al. Agreement and validity of pharmacy data versus self-report for use of osteoporosis medications among chronic glucocorticoid users. Pharmacoepidemiol Drug Saf 2006;15:710-718.
- 40. McKeown E, Bykerk VP, De Leon F, et al. Quality assurance study of the use of preventative therapies in glucocorticoidinduced osteoporosis in early inflammatory arthritis: results from the CATCH cohort. Rheumatology (Oxford) 2012;51:1662-1669.
- 41. Gamez-Nava JI, Zavaleta-Muniz SA, Vazquez-Villegas ML, et al. Prescription for antiresorptive therapy in Mexican patients with rheumatoid arthritis: is it time to reevaluate the strategies for osteoporosis prevention? Rheumatol Int 2013;33:145-150.
- 42. Caplan L, Hines AE, Williams E, et al. An observational study of glucocorticoid-induced osteoporosis prophylaxis in a national cohort of male veterans with rheumatoid arthritis. Osteoporos Int 2011;22:305-315.
- 43. Duyvendak M, Naunton M, Atthobari J, et al. Corticosteroid-induced osteoporosis prevention: Longitudinal practice patterns in The Netherlands 2001-2005. Osteoporos Int 2007;18:1429-1433.
- 44. Mohammad A, Ryan JG, Ralph N, et al. Improving trends in glucocorticoid-induced osteoporosis management: 2002 to 2006. Clin Exp Rheumatol 2007;25:728-733.
- 45. Blalock SJ, Norton LL, Patel RA, Dooley MA. Patient knowledge, beliefs, and behavior concerning the prevention and treatment of

glucocorticoid-induced osteoporosis. Arthritis Rheum 2005;53:732-739.

- 46. Ungprasert S, Wangkaew S, Louthrenoo W. Physician's awareness of the prevention of corticosteroid induced osteoporosis. J Med Assoc Thai 2007;90:59-64.
- 47. Chantler IW, Davie MWJ, Evans SF, Rees JS. Oral corticosteroid prescribing in women over 50, use of fracture prevention therapy, and bone densitometry service. Ann Rheum Dis 2003;62:350-352.
- 48. Shah SK, Gecys GT. Prednisone-induced osteoporosis: an overlooked and undertreated adverse effect. J Am Osteopath Assoc 2006;106:653-657.
- 49. Elliott ME, Farrah RM, Binkley NC, et al. Management of glucocorticoid-induced osteoporosis in male veterans. Ann Pharmacother 2000;34:1380-1384.
- 50. Osiri M, Saag KG, Ford AM, Moreland LW. Practice pattern variation among internal medicine specialists in the prevention of glucocorticoid-induced osteoporosis. J Clin Rheumatol 2000;6:117-122.
- 51. Buckley LM, Marquez M, Feezor R, et al. Prevention of corticosteroid-induced osteoporosis: results of a patient survey. Arthritis Rheum 1999;42:1736-1739
- 52. Curtis JR, Westfall AO, Allison JJ, et al. Longitudinal patterns in the prevention of osteoporosis in glucocorticoid-treated patients. Arthritis Rheum 2005;52:2485-2494.
- 53. Mudano A, Allison J, Hill J, et al. Variations in glucocorticoid induced osteoporosis prevention in a managed care cohort. J Rheumatol 2001;28:1298-1305.
- 54. Ettinger B, Chidambaran P, Pressman A. Prevalence and determinants of osteoporosis drug prescription among patients with high exposure to glucocorticoid drugs. Am J Manag Care 2001;7:597-605.
- 55. Che M, Ettinger B, Nguyen MT, et al. Highdose corticosteroid exposure and osteoporosis intervention in adults. Ann Allergy Asthma Immunol 2006;97:497-501.
- 56. Saag KG, Gehlbach SH, Curtis JR, et al. Trends in prevention of glucocorticoidinduced osteoporosis. J Rheumatol 2006;33:1651-1657.
- 57. Majumdar SR, Lix LM, Yogendran M, et al. Population-based trends in osteoporosis management after new initiations of long-term systemic glucocorticoids (1998-2008). J Clin Endocrinol Metab 2012;97:1236-1242.

- 58. Yood RA, Harrold LR, Fish L, et al. Prevention of glucocorticoid-induced osteoporosis: experience in a managed care setting. Arch Intern Med 2001;161:1322-1327.
- 59. Solomon DH, Katz JN, Jacobs JP, et al. Management of glucocorticoid-induced osteoporosis in patients with rheumatoid arthritis: Rates and predictors of care in an academic rheumatology practice. Arthritis & Rheumatism 2002;46:3136-3142.
- 60. Feldstein AC, Elmer PJ, Nichols GA, Herson M. Practice patterns in patients at risk for glucocorticoid-induced osteoporosis. Osteoporos Int 2005;16:2168-2174.
- 61. Cruse LM, Valeriano J, Vasey FB, Carter JD Prevalence of evaluation and treatment of glucocorticoid-induced osteoporosis in men. J Clin Rheumatol 2006;12:221-225.
- 62. Liu RH, Albrecht J, Werth VP Cross-sectional study of bisphosphonate use in dermatology patients receiving long-term oral corticosteroid therapy. Arch Dermatol 2006;142:37-41
- 63. Guzman-Clark JRS, Fang MA, Sehl ME, et al. Barriers in the management of glucocorticoidinduced osteoporosis. Arthritis Rheum 2007;57:140-146
- 64. Ledwich LJ, Clarke K Screening and treatment of glucocorticoid-induced osteoporosis in rheumatoid arthritis patients in an urban multispecialty practice. J Clin Rheumatol 2009; 15:61-64
- 65. Thanou A, Ali T, Haq O, et al. Utilization of preventive measures for glucocorticoidinduced osteoporosis among veterans with inflammatory bowel disease. ISRN Gastroenterology 2013; 2013: Article ID 862312.
- 66. Erb N, Duncan RC, Raza K, et al. A regional audit of the prevention and treatment of corticosteroid-induced osteoporosis in patients with rheumatic diseases in the West Midlands. Rheumatology 2002;41:1021-1024.
- 67. Gudbjornsson B, Juliusson UI, Gudjonsson FV. Prevalence of long term steroid treatment and the frequency of decision making to prevent steroid induced osteoporosis in daily clinical practice. Ann Rheum Dis 2002;61:32-36.
- 68. Walker-Bone K, Wood A, Hull R, et al. The prevention and treatment of glucocorticoidinduced osteoporosis in clinical practice. Clin Med 2004;4:431-436.

- 69. Wall E, Walker-Bone K. Use of bisphosphonates and dual-energy X-ray absorptiometry scans in the prevention and treatment of glucocorticoid-induced rheumatology. osteoporosis in OJM 2008;101:317-323.
- Haroon M, Rathaille MO, Gradaigh DO. Prevention of glucocorticoid-induced osteoporosis: A re-audit of dual-energy x-ray absorptiometry scan access and management guideline compliance. Int J Clin Rheumtol 2011;6:251-255.
- 71. Hougardy DM, Peterson GM, Bleasel MD, Randall CT. Is enough attention being given to the adverse effects of corticosteroid therapy? J Clin Pharm Ther 2000;25:227-234.
- 72. Sadat-Ali M, Alelq AH, Alshafei BA, et al. Osteoporosis prophylaxis in patients receiving chronic glucocorticoid therapy. Ann Saudi Med 2009;29:215-218.
- 73. Rothberg AD, Matshidze PK. Monitoring and management of bone status in patients on chronic glucocorticoid treatment--the Medscheme experience. S Afr Med J 2000;90:1125-1129.
- 74. Smith MD, Cheah SP, Taylor K, Ahern MJ. Prevention of corticosteroid induced osteoporosis in inpatients recently discharged from a tertiary teaching hospital. J Rheumatol 2001;28:566-570.
- 75. Gera C, Vij AS. Glucocorticoid-induced osteoporosis: unawareness or negligence in India? Int J Rheum Dis 2009;12:230-233.
- 76. Srinivasulu N, Sharma V, Chitnis N, et al. Primary prophylaxis for steroid-induced osteoporosis: Are we doing enough?-An audit from a tertiary care centre. Indian J Rheumatol 2010;5:176-179.

- 77. Kohli B, Kem AK, Bhatia R. A one year study of the adequacy of primary prophylaxis in prevention of glucocorticoid-induced osteoporosis in rural western Uttar Pradesh. Indian J Public Health 2013;4:44-48.
- 78. Hallberg I, Bachrach-Lindstrom M, Hammerby S, et al. Health-related quality of life after vertebral or hip fracture: a sevenyear follow-up study. BMC Musculoskeletal Disorders 2009;10:135.
- 79. Cauley JA, Thompson DE, Ensrud KC, et al. Risk of mortality following clinical fractures. Osteoporos Int 2000;11:556-561.
- 80. Nikitovic M, Wodchis WP, Krahn MD, Cadarette SM. Direct health-care costs attributed to hip fractures among seniors: a matched cohort study. Osteoporos Int 2013;24:659-669.
- Cadarette SM, Burden AM. The burden of osteoporosis in Canada. Can Pharm J (Ott) 2011;144:S3-S3.e1.
- 82. Ruggiero SL, Mehrotra B, Rosenberg TJ, Engroff SL. Osteonecrosis of the jaws associated with the use of bisphosphonates: a review of 63 cases. J Oral Maxillofac Surg 2004;62:527-534.
- Abrahamsen B, Eiken P, Eastell R. Subtrochanteric and diaphyseal femur fractures in patients treated with Alendronate: A register-based national cohort study. J Bone Miner Res 2009;24:1095-1102.
- 84. Solomon DH, Johnston SS, Boytsov NN, et al. Osteoporosis Medication Use after Hip Fracture in U.S. Patients between 2002 and 2011. Journal of Bone and Mineral Research: the official journal of the American Society for Bone and Mineral Research 2014;29(9):1929-1937.