

# BEERS LIST- WHAT IS IT & WHY IS IT IMPORTANT?

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# OBJECTIVES

- ◉ Describe the Beers List, how to access and how it's used
- ◉ Identify common OTC medications to be avoided or used with caution in older adults and in LTC practice
- ◉ Demonstrate how to access and utilize the Beers List through case study

# THE BEERS LIST DEFINED

- ⦿ NO it's not a list of adult beverages to drink while you're here at Niagara Falls.



# THE BEERS LIST DEFINED

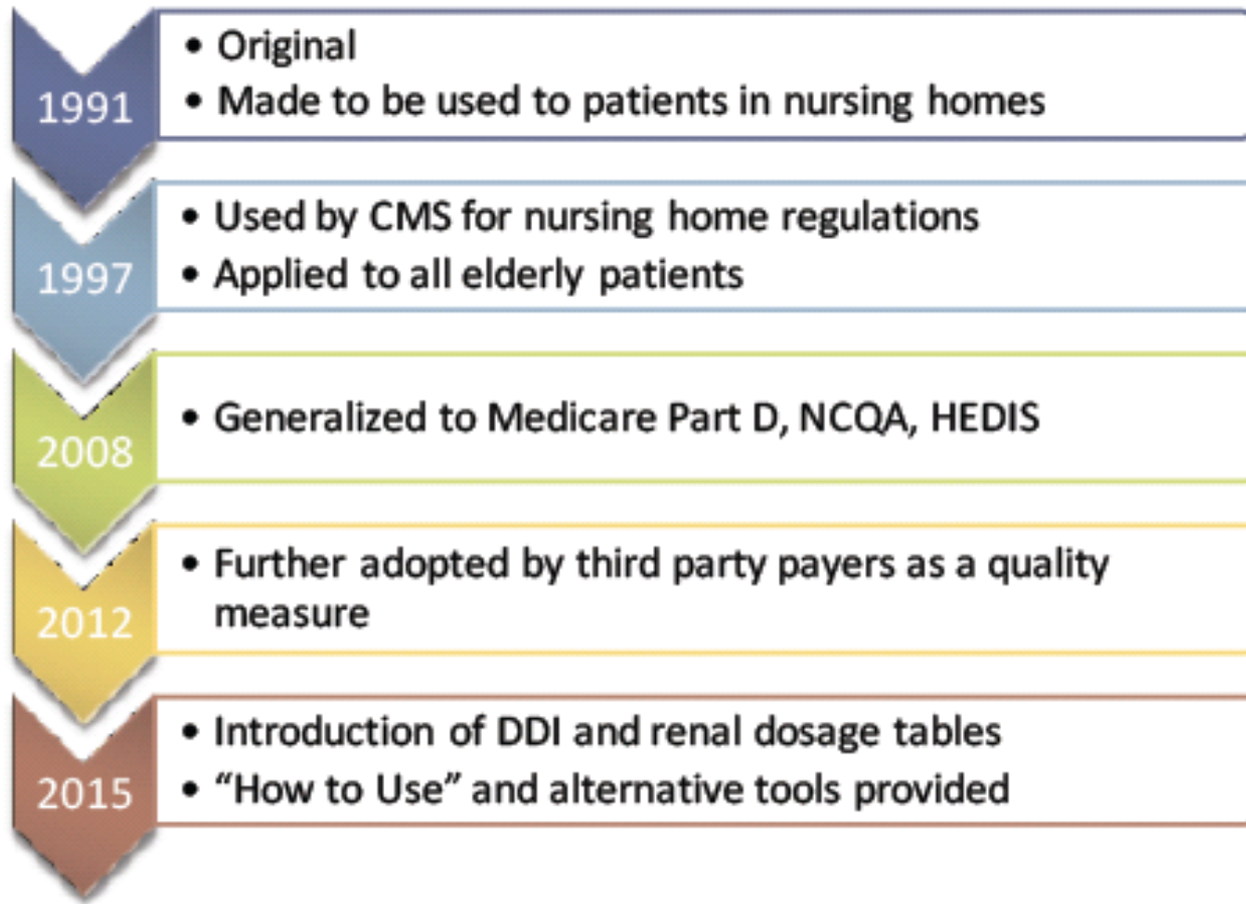
- Recourse to identify potentially inappropriate medication for older adults because of ineffectiveness or a high risk of adverse events; and to improve medication selection and overall medication safety in the elderly population.

# BACKGROUND

- The Beers Criteria is named after a lead researcher, Dr. Mark H Beers
- It was developed in 1991 to target nursing home residents
- It has been revised over the years
  - Some of the medications on the list you may not be familiar with because they have fallen out of favor in practice



# HISTORY OF THE BEERS CRITERIA



# PURPOSE OF THE BEERS CRITERIA

- To identify potentially inappropriate medications that should be avoided in many older adults
- To reduce adverse drug events and drug related problems, and to improve medication selection and medication use in older adults
- Designed for use in any clinical setting

# WHY SHOULD PHARMACY PERSONAL PROVIDING CARE FOR OLDER PATIENTS FAMILIARIZE THEMSELVES WITH THE BEERS CRITERIA?

- Increased risk of hospitalization in patients on PIM
- Higher healthcare cost

# SELF-ASSESSMENT

- **What is the purpose of the Beers Criteria?**
  - A. To identify potentially inappropriate medications to be avoided in older adults
  - B. To reduce adverse drug events and drug related problems
  - C. To improve medication selection and medication use in older adults
  - D. All of the above

# SELF-ASSESSMENT

- What is the purpose of the Beers Criteria?
  - A. To identify potentially inappropriate medications to be avoided in older adults
  - B. To reduce adverse drug events and drug related problems
  - C. To improve medication selection and medication use in older adults
  - D. All of the above**

# HOW TO USE THE BEERS LIST

- The creators of the Beers List put ratings on how important the recommendations are.
- QE: Quality of Evidence
  - Low, moderate , or strong
- SR: Strength of Warning
  - Strong or Weak

**Table 1. Designations of Quality of Evidence and Strength of Recommendations**

Quality of Evidence	
High	Evidence includes consistent results from well-designed, well-conducted studies in representative populations that directly assess effects on health outcomes ( $\geq 2$ consistent, higher-quality randomized controlled trials or multiple, consistent observational studies with no significant methodological flaws showing large effects)
Moderate	Evidence is sufficient to determine risks of adverse outcomes, but the number, quality, size, or consistency of included studies; generalizability to routine practice; or indirect nature of the evidence on health outcomes ( $\geq 1$ higher-quality trial with $>100$ participants; $\geq 2$ higher-quality trials with some inconsistency; $\geq 2$ consistent, lower-quality trials; or multiple, consistent observational studies with no significant methodological flaws showing at least moderate effects) limits the strength of the evidence
Low	Evidence is insufficient to assess harms or risks in health outcomes because of limited number or power of studies, large and unexplained inconsistency between higher-quality studies, important flaws in study design or conduct, gaps in the chain of evidence, or lack of information on important health outcomes
Strength of Recommendation	
Strong	Benefits clearly outweigh harms, adverse events, and risks, or harms, adverse events, and risks clearly outweigh benefits
Weak	Benefits may not outweigh harms, adverse events, and risks
Insufficient	Evidence inadequate to determine net harms, adverse events, and risks

**Table 2. 2015 American Geriatrics Society Beers Criteria for Potentially Inappropriate Medication Use in Older Adults**

Organ System, Therapeutic Category, Drugs	Rationale	Recommendation	Quality of Evidence	Strength of Recommendation
<b>Anticholinergics</b>				
First-generation antihistamines Brompheniramine Carbinoxamine Chlorpheniramine Clemastine Cyproheptadine Dexbrompheniramine Dexchlorpheniramine Dimenhydrinate Diphenhydramine (oral) Doxylamine Hydroxyzine Meclizine Promethazine Triprolidine	Highly anticholinergic; clearance reduced with advanced age, and tolerance develops when used as hypnotic; risk of confusion, dry mouth, constipation, and other anticholinergic effects or toxicity  Use of diphenhydramine in situations such as acute treatment of severe allergic reaction may be appropriate	Avoid	Moderate	Strong
Antiparkinsonian agents Benztropine (oral) Trihexyphenidyl	Not recommended for prevention of extrapyramidal symptoms with antipsychotics; more-effective agents available for treatment of Parkinson disease	Avoid	Moderate	Strong
Antispasmodics Atropine (excludes ophthalmic) Belladonna alkaloids Clidinium-Chlordiazepoxide Dicyclomine Hyoscyamine Propantheline Scopolamine	Highly anticholinergic, uncertain effectiveness	Avoid	Moderate	Strong
<b>Antithrombotics</b>				
Dipyridamole, oral short-acting (does not apply to the extended-release combination with aspirin)	May cause orthostatic hypotension; more effective alternatives available; intravenous form acceptable for use in cardiac stress testing	Avoid	Moderate	Strong
Ticlopidine	Safer, effective alternatives available	Avoid	Moderate	Strong
<b>Anti-infective</b>				
Nitrofurantoin	Potential for pulmonary toxicity, hepatotoxicity, and peripheral neuropathy, especially with long-term use; safer alternatives available	Avoid in individuals with creatinine clearance <30 mL/min or for long-term suppression of bacteria	Low	Strong
<b>Cardiovascular</b>				
Peripheral alpha-1 blockers Doxazosin Prazosin Terazosin	High risk of orthostatic hypotension; not recommended as routine treatment for hypertension; alternative agents have superior risk–benefit profile	Avoid use as an antihypertensive	Moderate	Strong

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Organ System, Therapeutic Category, Drug	Rationale	Recommendation	Quality of Evidence	Strength of Recommendation
<b>Anticholinergics</b>				
First-generation antihistamines	Highly anticholinergic; clearance reduced with advanced age, and tolerance develops when used as hypnotic; risk of confusion, dry mouth, constipation, and other anticholinergic effects or toxicity	Avoid	Moderate	Strong
Brompheniramine				
Carbinoxamine				
Chlorpheniramine				
Clemastine				
Cyproheptadine	Use of diphenhydramine in situations such as acute treatment of severe allergic reaction may be appropriate			
Dexbrompheniramine				
Dexchlorpheniramine				
Dimenhydrinate				
Diphenhydramine (oral)				
Doxylamine				
Hydroxyzine				
Meclizine				
Promethazine				
Triprolidine				
<b>Antiparkinsonian agents</b>				
Benztropine (oral)	Not recommended for prevention of extrapyramidal symptoms with antipsychotics; more-effective agents available for treatment of Parkinson disease	Avoid	Moderate	Strong
Trihexyphenidyl				
<b>Antispasmodics</b>				
Atropine (excludes ophthalmic)	Highly anticholinergic, uncertain effectiveness	Avoid	Moderate	Strong
Belladonna alkaloids				
Clidinium-Chlordiazepoxide				
Dicyclomine				
Hyoscyamine				
Propantheline				
Scopolamine				
<b>Antithrombotics</b>				
Dipyridamole, oral short-acting (does not apply to the extended-release combination with aspirin)	May cause orthostatic hypotension; more effective alternatives available; intravenous form acceptable for use in cardiac stress testing	Avoid	Moderate	Strong
Ticlopidine	Safer, effective alternatives available	Avoid	Moderate	Strong
<b>Anti-infective</b>				
Nitrofurantoin	Potential for pulmonary toxicity, hepatotoxicity, and peripheral neuropathy, especially with long-term use; safer alternatives available	Avoid in individuals with creatinine clearance <30 mL/min or for long-term suppression of bacteria	Low	Strong
<b>Cardiovascular</b>				
Peripheral alpha-1 blockers				
Doxazosin	High risk of orthostatic hypotension; not recommended as routine treatment for hypertension; alternative agents have superior risk-benefit profile	Avoid use as an antihypertensive	Moderate	Strong
Prazosin				
Terazosin				

Table 2 (Contd.)

Organ System, Therapeutic Category, Drugs	Rationale	Recommendation	Quality of Evidence	Strength of Recommendation
<p>Antidepressants, alone or in combination</p> <p>Amitriptyline Amoxapine Clomipramine Desipramine Doxepin &gt;6 mg/d Imipramine Nortriptyline Paroxetine Protriptyline Trimipramine</p>	<p>Highly anticholinergic, sedating, and cause orthostatic hypotension; safety profile of low-dose doxepin (<math>\leq 6</math> mg/d) comparable with that of placebo</p>	Avoid	High	Strong
<p>Antipsychotics, first- (conventional) and second- (atypical) generation</p>	<p>Increased risk of cerebrovascular accident (stroke) and greater rate of cognitive decline and mortality in persons with dementia Avoid antipsychotics for behavioral problems of dementia or delirium unless nonpharmacological options (e.g., behavioral interventions) have failed or are not possible <b>and</b> the older adult is threatening substantial harm to self or others</p>	Avoid, except for schizophrenia, bipolar disorder, or short-term use as antiemetic during chemotherapy	Moderate	Strong
<p>Barbiturates</p> <p>Amobarbital Butabarbital Butalbital Mephobarbital Pentobarbital Phenobarbital Secobarbital</p>	<p>High rate of physical dependence, tolerance to sleep benefits, greater risk of overdose at low dosages</p>	Avoid	High	Strong
<p>Benzodiazepines</p> <p><i>Short- and intermediate- acting</i></p> <p>Alprazolam Estazolam Lorazepam Oxazepam Temazepam Triazolam</p>	<p>Older adults have increased sensitivity to benzodiazepines and decreased metabolism of long-acting agents; in general, all benzodiazepines increase risk of cognitive impairment, delirium, falls, fractures, and motor vehicle crashes in older adults</p>	Avoid	Moderate	Strong

(Continued)

Table 2 (Contd.)

Organ System, Therapeutic Category, Drugs	Rationale	Recommendation	Quality of Evidence	Strength of Recommendation
Insulin, sliding scale	Higher risk of hypoglycemia without improvement in hyperglycemia management regardless of care setting; refers to sole use of short- or rapid-acting insulins to manage or avoid hyperglycemia in absence of basal or long-acting insulin; does not apply to titration of basal insulin or use of additional short- or rapid-acting insulin in conjunction with scheduled insulin (i.e., correction insulin)	Avoid	Moderate	Strong
Megestrol	Minimal effect on weight; increases risk of thrombotic events and possibly death in older adults	Avoid	Moderate	Strong
Sulfonylureas, long-duration Chlorpropamide	Chlorpropamide: prolonged half-life in older adults; can cause prolonged hypoglycemia; causes syndrome of inappropriate antidiuretic hormone secretion	Avoid	High	Strong
Glyburide	Glyburide: higher risk of severe prolonged hypoglycemia in older adults			
<b>Gastrointestinal</b>				
Metoclopramide	Can cause extrapyramidal effects, including tardive dyskinesia; risk may be greater in frail older adults	Avoid, unless for gastroparesis	Moderate	Strong
Mineral oil, given orally	Potential for aspiration and adverse effects; safer alternatives available	Avoid	Moderate	Strong
Proton-pump inhibitors	Risk of <i>Clostridium difficile</i> infection and bone loss and fractures	Avoid scheduled use for >8 weeks unless for high-risk patients (e.g., oral corticosteroids or chronic NSAID use), erosive esophagitis, Barrett's esophagitis, pathological hypersecretory condition, or demonstrated need for maintenance treatment (e.g., due to failure of drug discontinuation trial or H <sub>2</sub> blockers)	High	Strong
<b>Pain medications</b>				
Meperidine	Not effective oral analgesic in dosages commonly used; may have higher risk of neurotoxicity, including delirium, than other opioids; safer alternatives available	Avoid, especially in individuals with chronic kidney disease	Moderate	Strong

(Continued)

Table 2 (Contd.)

Organ System, Therapeutic Category, Drugs	Rationale	Recommendation	Quality of Evidence	Strength of Recommendation
<b>Non-cyclooxygenase-selective NSAIDs, oral:</b> Aspirin >325 mg/d Diclofenac Diflunisal Etodolac Fenoprofen Ibuprofen Ketoprofen Meclofenamate Mefenamic acid Meloxicam Nabumetone Naproxen Oxaprozin Piroxicam Sulindac Tolmetin	Increased risk of gastrointestinal bleeding or peptic ulcer disease in high-risk groups, including those aged >75 or taking oral or parenteral corticosteroids, anticoagulants, or antiplatelet agents; use of proton-pump inhibitor or misoprostol reduces but does not eliminate risk. Upper gastrointestinal ulcers, gross bleeding, or perforation caused by NSAIDs occur in approximately 1% of patients treated for 3–6 months and in ~2–4% of patients treated for 1 year; these trends continue with longer duration of use	Avoid chronic use, unless other alternatives are not effective and patient can take gastroprotective agent (proton-pump inhibitor or misoprostol)	Moderate	Strong
Indomethacin	Indomethacin is more likely than other NSAIDs to have adverse CNS effects. Of all the NSAIDs, indomethacin has the most adverse effects.	Avoid	Moderate	Strong
Ketorolac, includes parenteral	Increased risk of gastrointestinal bleeding, peptic ulcer disease, and acute kidney injury in older adults			
Pentazocine	Opioid analgesic that causes CNS adverse effects, including confusion and hallucinations, more commonly than other opioid analgesic drugs; is also a mixed agonist and antagonist; safer alternatives available	Avoid	Low	Strong
Skeletal muscle relaxants Carisoprodol Chlorzoxazone Cyclobenzaprine Metaxalone Methocarbamol Orphenadrine	Most muscle relaxants poorly tolerated by older adults because some have anticholinergic adverse effects, sedation, increased risk of fractures; effectiveness at dosages tolerated by older adults questionable	Avoid	Moderate	Strong
Genitourinary Desmopressin	High risk of hyponatremia; safer alternative treatments	Avoid for treatment of nocturia or nocturnal polyuria	Moderate	Strong

The primary target audience is practicing clinicians. The intentions of the criteria are to improve the selection of prescription drugs by clinicians and patients; evaluate patterns of drug use within populations; educate clinicians and patients on proper drug usage; and evaluate health-outcome, quality-of-care, cost, and utilization data.

CNS = central nervous system; NSAIDs = nonsteroidal anti-inflammatory drugs.

# NSAIDS

- Very common and easily accessible
  - Since NSAIDS are so easily obtained, patients may be unaware of the potentially hazardous side effects
    - HTN
    - GI Bleeds
    - Kidney Injury

# CASE STUDY

Your friend's grandma Bernice comes into the pharmacy and asks you what aisle the Ibuprofen is in. How would you respond?

- A. Take Bernice to the IBU because it's good customer service
- B. Show Bernice where the Tylenol is and recommend it instead of the IBU due to it being on the Beers List
- C. Get your pharmacist to consult with Bernice on the need to purchase IBU

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Table 3. 2015 American Geriatrics Society Beers Criteria for Potentially Inappropriate Medication Use in Older Adults Due to Drug–Disease or Drug–Syndrome Interactions That May Exacerbate the Disease or Syndrome

Disease or Syndrome	Drug(s)	Rationale	Recommendation	Quality of Evidence	Strength of Recommendation
<b>Cardiovascular</b>					
Heart failure	NSAIDs and COX-2 inhibitors Nondihydropyridine CCBs (diltiazem, verapamil) —avoid only for heart failure with reduced ejection fraction Thiazolidinediones (pioglitazone, rosiglitazone) Cilostazol Dronedarone (severe or recently decompensated heart failure)	Potential to promote fluid retention and exacerbate heart failure	Avoid	NSAIDs: moderate CCBs: moderate Thiazolidinediones: high Cilostazol: low Dronedarone: high	Strong
Syncope	AChEIs Peripheral alpha-1 blockers Doxazosin Prazosin Terazosin Tertiary TCAs Chlorpromazine Thioridazine Olanzapine	Increases risk of orthostatic hypotension or bradycardia	Avoid	Peripheral alpha-1 blockers: high TCAs, AChEIs, antipsychotics: moderate	AChEIs, TCAs: strong Peripheral alpha-1 blockers, antipsychotics: weak
<b>Central nervous system</b>					
Chronic seizures or epilepsy	Bupropion Chlorpromazine Clozapine Maprotiline Olanzapine Thioridazine Thiothixene Tramadol	Lowers seizure threshold; may be acceptable in individuals with well-controlled seizures in whom alternative agents have not been effective	Avoid	Low	Strong
Delirium	Anticholinergics (see Table 7 for full list) Antipsychotics Benzodiazepines Chlorpromazine Corticosteroids <sup>a</sup> H <sub>2</sub> -receptor antagonists Cimetidine Famotidine Nizatidine Ranitidine Meperidine Sedative hypnotics	Avoid in older adults with or at high risk of delirium because of the potential of inducing or worsening delirium Avoid antipsychotics for behavioral problems of dementia or delirium unless nonpharmacological options (e.g., behavioral interventions) have failed or are not possible <b>and</b> the older adult is threatening substantial harm to self or others Antipsychotics are associated with greater risk of cerebrovascular accident (stroke) and mortality in persons with dementia	Avoid	Moderate	Strong

Table 3. 2015 American Geriatrics Society Beers Criteria for Potentially Inappropriate Medication Use in Older Adults Due to Drug–Disease or Drug–Syndrome Interactions That May Exacerbate the Disease or Syndrome

Disease or Syndrome	Drug(s)	Rationale	Recommendation	Quality of Evidence	Strength of Recommendation
Cardiovascular					
Heart failure	NSAIDs and COX-2 inhibitors Nondihydropyridine CCBs (diltiazem, verapamil) —avoid only for heart failure with reduced ejection fraction Thiazolidinediones (pioglitazone, rosiglitazone) Cilostazol Dronedarone (severe or recently decompensated heart failure)	Potential to promote fluid retention and exacerbate heart failure	Avoid	NSAIDs: moderate CCBs: moderate Thiazolidinediones: high Cilostazol: low Dronedarone: high	Strong
Syncope	AChEIs Peripheral alpha-1 blockers Doxazosin Prazosin Terazosin Tertiary TCAs Chlorpromazine Thioridazine Olanzapine	Increases risk of orthostatic hypotension or bradycardia	Avoid	Peripheral alpha-1 blockers: high TCAs, AChEIs, antipsychotics: moderate	AChEIs, TCAs: strong Peripheral alpha-1 blockers, antipsychotics: weak
Central nervous system					
Chronic seizures or epilepsy	Bupropion Chlorpromazine Clozapine Maprotiline Olanzapine Thioridazine Thiothixene Tramadol	Lowers seizure threshold; may be acceptable in individuals with well-controlled seizures in whom alternative agents have not been effective	Avoid	Low	Strong
Delirium	Anticholinergics (see Table 7 for full list) Antipsychotics Benzodiazepines Chlorpromazine Corticosteroids <sup>a</sup> H <sub>2</sub> -receptor antagonists Cimetidine Famotidine Nizatidine Ranitidine Meperidine Sedative hypnotics	Avoid in older adults with or at high risk of delirium because of the potential of inducing or worsening delirium Avoid antipsychotics for behavioral problems of dementia or delirium unless nonpharmacological options (e.g., behavioral interventions) have failed or are not possible <b>and</b> the older adult is threatening substantial harm to self or others Antipsychotics are associated with greater risk of cerebrovascular accident (stroke) and mortality in persons with dementia	Avoid	Moderate	Strong

Table 3 (Contd.)

Disease or Syndrome	Drug(s)	Rationale	Recommendation	Quality of Evidence	Strength of Recommendation
Dementia or cognitive impairment	Anticholinergics (see Table 7 for full list) Benzodiazepines H <sub>2</sub> -receptor antagonists Nonbenzodiazepine, benzodiazepine receptor agonist hypnotics Eszopiclone Zolpidem Zaleplon Antipsychotics, chronic and as-needed use	Avoid because of adverse CNS effects  Avoid antipsychotics for behavioral problems of dementia or delirium unless nonpharmacological options (e.g., behavioral interventions) have failed or are not possible <b>and</b> the older adult is threatening substantial harm to self or others. Antipsychotics are associated with greater risk of cerebrovascular accident (stroke) and mortality in persons with dementia	Avoid	Moderate	Strong
History of falls or fractures	Anticonvulsants Antipsychotics Benzodiazepines Nonbenzodiazepine, benzodiazepine receptor agonist hypnotics Eszopiclone Zaleplon Zolpidem TCAs SSRIs Opioids	May cause ataxia, impaired psychomotor function, syncope, additional falls; shorter-acting benzodiazepines are not safer than long-acting ones  If one of the drugs must be used, consider reducing use of other CNS-active medications that increase risk of falls and fractures (i.e., anticonvulsants, opioid-receptor agonists, antipsychotics, antidepressants, benzodiazepine-receptor agonists, other sedatives and hypnotics) and implement other strategies to reduce fall risk	Avoid unless safer alternatives are not available; avoid anticonvulsants except for seizure and mood disorders	High  Opioids: moderate	Strong  Opioids: strong
Insomnia	Oral decongestants Pseudoephedrine Phenylephrine Stimulants Amphetamine Armodafinil Methylphenidate Modafinil Theobromines Theophylline Caffeine	CNS stimulant effects	Avoid	Moderate	Strong

(Continued)

Table 3 (Contd.)

Disease or Syndrome	Drug(s)	Rationale	Recommendation	Quality of Evidence	Strength of Recommendation
Dementia or cognitive impairment	Anticholinergics (see Table 7 for full list) Benzodiazepines H <sub>2</sub> -receptor antagonists Nonbenzodiazepine, benzodiazepine receptor agonist hypnotics Eszopiclone Zolpidem Zaleplon Antipsychotics, chronic and as-needed use	Avoid because of adverse CNS effects  Avoid antipsychotics for behavioral problems of dementia or delirium unless nonpharmacological options (e.g., behavioral interventions) have failed or are not possible <b>and</b> the older adult is threatening substantial harm to self or others. Antipsychotics are associated with greater risk of cerebrovascular accident (stroke) and mortality in persons with dementia	Avoid	Moderate	Strong
History of falls or fractures	Anticonvulsants Antipsychotics Benzodiazepines Nonbenzodiazepine, benzodiazepine receptor agonist hypnotics Eszopiclone Zaleplon Zolpidem TCAs SSRIs Opioids	May cause ataxia, impaired psychomotor function, syncope, additional falls; shorter-acting benzodiazepines are not safer than long-acting ones  If one of the drugs must be used, consider reducing use of other CNS-active medications that increase risk of falls and fractures (i.e., anticonvulsants, opioid-receptor agonists, antipsychotics, antidepressants, benzodiazepine-receptor agonists, other sedatives and hypnotics) and implement other strategies to reduce fall risk	Avoid unless safer alternatives are not available; avoid anticonvulsants except for seizure and mood disorders	High  Opioids: moderate	Strong  Opioids: strong
Insomnia	Oral decongestants Pseudoephedrine Phenylephrine Stimulants Amphetamine Armodafinil Methylphenidate Modafinil Theobromines Theophylline Caffeine	CNS stimulant effects	Avoid	Moderate	Strong

(Continued)

- \*Video
- <http://elearning.pharmacist.com/products/4721/steady-the-pharmacists-role-in-older-adult-fall-prevention>

**Table 4. 2015 American Geriatrics Society Beers Criteria for Potentially Inappropriate Medications to Be Used with Caution in Older Adults**

Drug(s)	Rationale	Recommendation	Quality of Evidence	Strength of Recommendation
Aspirin for primary prevention of cardiac events	Lack of evidence of benefit versus risk in adults aged $\geq 80$	Use with caution in adults aged $\geq 80$	Low	Strong
Dabigatran	Increased risk of gastrointestinal bleeding compared with warfarin and reported rates with other target-specific oral anticoagulants in adults aged $\geq 75$ ; lack of evidence of efficacy and safety in individuals with CrCl $< 30$ mL/min	Use with caution in adults aged $\geq 75$ and in patients with CrCl $< 30$ mL/min	Moderate	Strong
Prasugrel	Increased risk of bleeding in older adults; benefit in highest-risk older adults (e.g., those with prior myocardial infarction or diabetes mellitus) may offset risk	Use with caution in adults aged $\geq 75$	Moderate	Weak
Antipsychotics Diuretics Carbamazepine Carboplatin Cyclophosphamide Cisplatin Mirtazapine Oxcarbazepine SNRIs SSRIs TCAs Vincristine	May exacerbate or cause syndrome of inappropriate antidiuretic hormone secretion or hyponatremia; monitor sodium level closely when starting or changing dosages in older adults	Use with caution	Moderate	Strong
Vasodilators	May exacerbate episodes of syncope in individuals with history of syncope	Use with caution	Moderate	Weak

The primary target audience is the practicing clinician. The intentions of the criteria are to improve selection of prescription drugs by clinicians and patients; evaluate patterns of drug use within populations; educate clinicians and patients on proper drug usage; and evaluate health-outcome, quality-of-care, cost, and utilization data.

CrCl = creatinine clearance; SNRIs = serotonin-norepinephrine reuptake inhibitors; SSRIs = selective serotonin reuptake inhibitors; TCAs = tricyclic antidepressants.

**Table 5. 2015 American Geriatrics Society Beers Criteria for Potentially Clinically Important Non-Anti-infective Drug-Drug Interactions That Should Be Avoided in Older Adults**

Object Drug and Class	Interacting Drug and Class	Risk Rationale	Recommendation	Quality of Evidence	Strength of Recommendation
ACEIs	Amiloride or triamterene	Increased risk of Hyperkalemia	Avoid routine use; reserve for patients with demonstrated hypokalemia while taking an ACEI	Moderate	Strong
Anticholinergic	Anticholinergic	Increased risk of Cognitive decline	Avoid, minimize number of anticholinergic drugs (Table 7)	Moderate	Strong
Antidepressants (i.e., TCAs and SSRIs)	≥2 other CNS-active drugs <sup>a</sup>	Increased risk of Falls	Avoid total of ≥3 CNS-active drugs <sup>a</sup> ; minimize number of CNS-active drugs	Moderate	Strong
Antipsychotics	≥2 other CNS-active drugs <sup>a</sup>	Increased risk of Falls	Avoid total of ≥3 CNS-active drugs <sup>a</sup> ; minimize number of CNS-active drugs	Moderate	Strong
Benzodiazepines and nonbenzodiazepine, benzodiazepine receptor agonist hypnotics	≥2 other CNS-active drugs <sup>a</sup>	Increased risk of Falls and fractures	Avoid total of ≥3 CNS-active drugs <sup>a</sup> ; minimize number of CNS-active drugs	High	Strong
Corticosteroids, oral or parenteral	NSAIDs	Increased risk of Peptic ulcer disease or gastrointestinal bleeding	Avoid; if not possible, provide gastrointestinal protection	Moderate	Strong
Lithium	ACEIs	Increased risk of Lithium toxicity	Avoid, monitor lithium concentrations	Moderate	Strong
Lithium	Loop diuretics	Increased risk of Lithium toxicity	Avoid, monitor lithium concentrations	Moderate	Strong
Opioid receptor agonist analgesics	≥2 other CNS-active drugs <sup>a</sup>	Increased risk of Falls	Avoid total of ≥3 CNS-active drugs <sup>a</sup> ; minimize number of CNS drugs	High	Strong
Peripheral Alpha-1 blockers	Loop diuretics	Increased risk of Urinary incontinence in older women	Avoid in older women, unless conditions warrant both drugs	Moderate	Strong
Theophylline	Cimetidine	Increased risk of Theophylline toxicity	Avoid	Moderate	Strong
Warfarin	Amiodarone	Increased risk of Bleeding	Avoid when possible; monitor international normalized ratio closely	Moderate	Strong
Warfarin	NSAIDs	Increased risk of Bleeding	Avoid when possible; if used together, monitor for bleeding closely	High	Strong

<sup>a</sup>Central nervous system (CNS)-active drugs: antipsychotics; benzodiazepines; nonbenzodiazepine, benzodiazepine receptor agonist hypnotics; tricyclic antidepressants (TCAs); selective serotonin reuptake inhibitors (SSRIs); and opioids.

ACEI = angiotensin-converting enzyme inhibitor; NSAID = nonsteroidal anti-inflammatory drug.

**Table 6. 2015 American Geriatrics Society Beers Criteria for Non-Anti-Infective Medications That Should Be Avoided or Have Their Dosage Reduced with Varying Levels of Kidney Function in Older Adults**

Medication Class and Medication	Creatinine Clearance, mL/min, at Which Action Required	Rationale	Recommendation	Quality of Evidence	Strength of Recommendation
<b>Cardiovascular or hemostasis</b>					
Amiloride	<30	Increased potassium, and decreased sodium	Avoid	Moderate	Strong
Apixaban	<25	Increased risk of bleeding	Avoid	Moderate	Strong
Dabigatran	<30	Increased risk of bleeding	Avoid	Moderate	Strong
Edoxaban	30–50 <30 or >95	Increased risk of bleeding	Reduce dose Avoid	Moderate	Strong
Enoxaparin	<30	Increased risk of bleeding	Reduce dose	Moderate	Strong
Fondaparinux	<30	Increased risk of bleeding	Avoid	Moderate	Strong
Rivaroxaban	30–50 <30	Increased risk of bleeding	Reduce dose Avoid	Moderate	Strong
Spirolactone	<30	Increased potassium	Avoid	Moderate	Strong
Triamterene	<30	Increased potassium, and decreased sodium	Avoid	Moderate	Strong
<b>Central nervous system and analgesics</b>					
Duloxetine	<30	Increased Gastrointestinal adverse effects (nausea, diarrhea)	Avoid	Moderate	Weak
Gabapentin	<60	CNS adverse effects	Reduce dose	Moderate	Strong
Levetiracetam	≤80	CNS adverse effects	Reduce dose	Moderate	Strong
Pregabalin	<60	CNS adverse effects	Reduce dose	Moderate	Strong
Tramadol	<30	CNS adverse effects	Immediate release: reduce dose Extended release: avoid	Low	Weak
<b>Gastrointestinal</b>					
Cimetidine	<50	Mental status changes	Reduce dose	Moderate	Strong
Famotidine	<50	Mental status changes	Reduce dose	Moderate	Strong
Nizatidine	<50	Mental status changes	Reduce dose	Moderate	Strong
Ranitidine	<50	Mental status changes	Reduce dose	Moderate	Strong
<b>Hyperuricemia</b>					
Colchicine	<30	Gastrointestinal, neuromuscular, bone marrow toxicity	Reduce dose; monitor for adverse effects	Moderate	Strong
Probenecid	<30	Loss of effectiveness	Avoid	Moderate	Strong

CNS = central nervous system.

**Table 7. Drugs with Strong Anticholinergic Properties**

<b>Antihistamines</b> Brompheniramine Carbinoxamine Chlorpheniramine Clemastine Cyproheptadine Dexbrompheniramine Dexchlorpheniramine Dimenhydrinate Diphenhydramine (oral) Doxylamine Hydroxyzine Meclizine Triprolidine	<b>Antiparkinsonian agents</b> Benztropine Trihexyphenidyl	<b>Skeletal muscle relaxants</b> Cyclobenzaprine Orphenadrine
<b>Antidepressants</b> Amitriptyline Amoxapine Clomipramine Desipramine Doxepin (>6 mg) Imipramine Nortriptyline Paroxetine Protriptyline Trimipramine	<b>Antipsychotics</b> Chlorpromazine Clozapine Loxapine Olanzapine Perphenazine Thioridazine Trifluoperazine	<b>Antiarrhythmic</b> Disopyramide
<b>Antimuscarinics (urinary incontinence)</b> Darifenacin Fesoterodine Flavoxate Oxybutynin Solifenacin Tolterodine Tropium	<b>Antispasmodics</b> Atropine (excludes ophthalmic) Belladonna alkaloids Clidinium- chlordiazepoxide Dicyclomine Homatropine (excludes ophthalmic) Hyoscyamine Propantheline Scopolamine (excludes ophthalmic)	<b>Antiemetic</b> Prochlorperazine Promethazine

**Table 8. Medications Moved to Another Category or Modified Since 2012 Beers Criteria**

Independent of Diagnoses or Condition (Table 2)	Considering Disease or Syndrome Interactions (Table 3)
Nitrofurantoin—recommendation and rationale modified	Heart failure—rationale and quality of evidence modified
Dronedarone—recommendation and rationale modified	Chronic seizures or epilepsy—quality of evidence modified
Digoxin—recommendation and rationale modified	Delirium—recommendation and rationale modified
Benzodiazepines—recommendation modified	Dementia or cognitive impairment—recommendation and rationale modified; new drugs added
Nonbenzodiazepine, benzodiazepine receptor agonist hypnotics—recommendation modified	History of falls or fractures—recommendation and rationale modified; new drugs added
Meperidine—recommendation modified	Parkinson disease—recommendation and rationale modified
Indomethacin and ketorolac, includes parenteral—rationale modified	Chronic kidney disease Stage IV or less (creatinine clearance <30 mL/min)—triamterene moved to Tables 5 and 6
Antipsychotics—recommendation and rationale modified	Insomnia—new drugs added
Estrogen—recommendation modified	
Insulin, sliding scale—rationale modified	

**Table 9. Medications Removed Since 2012 Beers Criteria**

Independent of Diagnoses or Condition (Table 2)	Considering Disease and Syndrome Interactions (Table 3)
Antiarrhythmic drugs (Class 1a, 1c, III except amiodarone) as first-line treatment for atrial fibrillation	Chronic constipation—entire criterion
Trimethobenzamide	Lower urinary tract—inhaled anticholinergic drugs
Mesoridazine—no longer marketed in United States	
Chloral hydrate—no longer marketed in United States	

**Table 10. Medications Added Since 2012 Beers Criteria**

Independent of Diagnoses or Condition (Table 2)	Considering Disease and Syndrome Interactions (Table 3)
Proton-pump inhibitors	Falls and fractures—opioids
Desmopressin	Insomnia—armodafinil and modafinil
Anticholinergics, first-generation antihistamines—meclizine	Dementia or cognitive impairment—eszopiclone and zaleplon Delirium—antipsychotics

# WARNING

- Think of the Beers Criteria as a warning light
  - Why is the patient taking the drug? Is it truly needed?
  - Are there safer more effective alternatives?
  - Characteristics that increase risk with this medication



# ACCESSING BEERS CRITERIA

## ○ American Geriatrics Society

- Free Access to PDF of Updated Beers Criteria
  - <https://www.americangeriatrics.org/>
    - Search Updated Beers Criteria
    - Access Content
- Minimal charge for digital pocket cards

# KEY PRINCIPLES

- Medications in the Beers Criteria are potentially inappropriate, not definitely inappropriate
- Reading the rationale and recommendations is important for guidance
- Understanding why medications are included on the list is important in medication selection
- Optimal application of the Beers Criteria involves offering safer non-pharmacologic and pharmacologic therapies
- The Beers Criteria should be a starting point for a comprehensive process of identifying and improving medication appropriateness and safety

# SELF-ASSESSMENT

## The Beers List

- A. is the end all be all.
- B. is utilized in all countries.
- C. should be used as a warning light.

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## The Beers List

- A. is the end all be all.
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- C. should be used as a warning light.

- The Beers Criteria continues to be improving the care of older adults by reducing their exposure to potentially inappropriate medications.

# ACKNOWLEDGEMENTS

- ◉ Why Pharmacists Should Check Out the Beers List  
<https://www.pharmacytimes.com/contributor/gunda-siska-pharmd/2018/01/why-pharmacists-should-check-out-the-beers-list>
- ◉ American Geriatrics Society 2015 Updated Beers Criteria for Potentially Inappropriate Medication Use in Older Adults  
<https://onlinelibrary.wiley.com/doi/full/10.1111/jgs.13702>
- ◉ American Geriatrics Society 2015 Beers Criteria Update: Implications for Practice  
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