

Prescribing cascades in community-dwelling adults: a systematic review

Ann Doherty¹, Faiza Shahid², Orla Cotter¹, Frank Moriarty³, Barbara Clyne⁴, Fiona Boland⁵, Tobias Dreischulte², Tom Fahey³, Sean Kennelly⁶, and Emma Wallace¹

¹Royal College of Surgeons Ireland

²University Hospital of Ludwig-Maximilians-University Munich

³Royal College of Surgeons in Ireland

⁴RCSI University of Medicine and Health Sciences

⁵HRB Centre for Primary Care Research

⁶Trinity College

June 23, 2022

Abstract

Abstract Background: The misattribution of an adverse drug reaction (ADR) as a symptom or illness can lead to the prescribing of additional medication, referred to as a prescribing cascade. The aim of this systematic review is to identify published prescribing cascades in community-dwelling adults. **Methods:** Systematic review reported in line with the PRISMA guidelines and pre-registered with PROSPERO. Electronic databases (Medline (Ovid), EMBASE, PsycINFO, CINAHL, Cochrane Library) and grey literature sources were searched. **Inclusion criteria:** Community-dwelling adults; Risk-prescription medication; Outcomes-initiation of new medicine to 'treat' or reduce ADR risk; Study type-cohort, cross-sectional, case-control and case-series studies. **Title/abstract screening, full-text screening, data extraction and methodological quality assessment** was conducted independently in duplicate. **A narrative synthesis was conducted. Results:** A total of 101 studies (reported in 103 publications) were included. Study sample sizes ranged from 126 to 11,593,989 participants and 15 studies examined older adults specifically ([?]60 years). Seventy-eight of 101 studies reported a potential prescribing cascade including calcium channel blockers to loop diuretic (n=5), amiodarone to levothyroxine (n=5), inhaled corticosteroid to topical antifungal (n=4), antipsychotic to anti-Parkinson drug (n=4), and acetylcholinesterase inhibitor to urinary incontinence drugs (n=4). Identified prescribing cascades occurred within three months to one year following initial medication. Methodological quality varied across included studies. **Conclusion and implications:** Prescribing cascades occur for a broad range of medications. ADRs should be included in the differential diagnosis for patients presenting with new symptoms, particularly older adults and those who started a new medication in the preceding 12 months. **Word count:** 245

Prescribing cascades in community-dwelling adults: a systematic review

Ann S Doherty, PhD¹ ; Faiza Shahid²; Orla Cotter¹; Frank Moriarty, PhD³; Fiona Boland^{1, 4}; Barbara Clyne¹; Tobias Dreischulte²; Tom Fahey¹; Seán P Kennelly^{5, 6}; Emma Wallace¹.

¹ Department of General Practice, RCSI University of Medicine and Health Sciences, Dublin 2, Ireland

²Institute of General Practice and Family Medicine, University Hospital of Ludwig-Maximilians-University Munich, Munich, Germany

³School of Pharmacy and Biomolecular Sciences, RCSI University of Medicine and Health Sciences, Dublin 2, Ireland

⁴Data Science Centre, RCSI University of Medicine and Health Sciences, Dublin 2, Ireland

⁵Department of Medical Gerontology, Trinity College Dublin, Dublin 2, Ireland

⁶Department of Age-related Healthcare, Tallaght University Hospital, Dublin 24, Ireland

Corresponding Author:

Professor Emma Wallace

Department of General Practice

RCSI University of Medicine and Health Sciences

123 St Stephen's Green

Dublin 2, Ireland

Phone: +353 (0)1 4023204

Email: emmawallace@rcsi.ie

Word count (excluding title page, abstract, references, tables and figures): 2826

Number of Figures: 3

Number of Tables: 2

Funding: This work was funded by a Health Research Board (HRB) Ireland Emerging Clinician Scientist Award awarded to EW [HRB-ECSA-2020-002]. BC is funded by the HRB Emerging Investigator Award [EIA-2019-09].

Conflict of Interest statement : The authors have no conflicts of interest to declare

Abstract

Background: The misattribution of an adverse drug reaction (ADR) as a symptom or illness can lead to the prescribing of additional medication, referred to as a prescribing cascade. The aim of this systematic review is to identify published prescribing cascades in community-dwelling adults.

Methods: Systematic review reported in line with the PRISMA guidelines and pre-registered with PROSPERO. Electronic databases (Medline (Ovid), EMBASE, PsycINFO, CINAHL, Cochrane Library) and grey literature sources were searched. Inclusion criteria: Community-dwelling adults; Risk-prescription medication; Outcomes-initiation of new medicine to 'treat' or reduce ADR risk; Study type-cohort, cross-sectional, case-control and case-series studies. Title/abstract screening, full-text screening, data extraction and methodological quality assessment was conducted independently in duplicate. A narrative synthesis was conducted.

Results: A total of 101 studies (reported in 103 publications) were included. Study sample sizes ranged from 126 to 11,593,989 participants and 15 studies examined older adults specifically ([?]60 years). Seventy-eight of 101 studies reported a potential prescribing cascade including calcium channel blockers to loop diuretic (n=5), amiodarone to levothyroxine (n=5), inhaled corticosteroid to topical antifungal (n=4), antipsychotic to anti-Parkinson drug (n=4), and acetylcholinesterase inhibitor to urinary incontinence drugs (n=4). Identified prescribing cascades occurred within three months to one year following initial medication. Methodological quality varied across included studies.

Conclusion and implications: Prescribing cascades occur for a broad range of medications. ADRs should be included in the differential diagnosis for patients presenting with new symptoms, particularly older adults and those who started a new medication in the preceding 12 months.

Word count: 245

Keywords: prescribing cascades; systematic review; appropriate prescribing; community-dwelling adults

Background

A prescribing cascade occurs when a medication is used to treat or prevent the adverse effects of another medication.¹⁻³ An unintentional prescribing cascade occurs when the adverse drug reaction (ADR) is misinterpreted as a new medical condition, leading to the prescription of new medication to treat the emerging symptoms.⁴ For example, calcium channel blocker (CCB) induced lower extremity oedema may be misinterpreted as a sign of congestive heart failure and result in the inappropriate prescribing of a loop diuretic to alleviate the oedema instead of simply switching the CCB to an alternative class antihypertensive agent.⁵⁻⁷ Intentional prescribing cascades occur when the ADR is recognised and a subsequent medication is prescribed to combat this ADR either via treatment of the ADR or prevention of it in the first instance.⁴ Prescribing cascades can be further characterised as either appropriate (potential benefits > risks), or inappropriate (risks > potential benefits).⁴ Furthermore, this characterisation of appropriateness is a dynamic entity; an appropriate prescribing cascade can become inappropriate over time, particularly should the clinical circumstances of the patient change.⁴

It is not clear what drives prescribing cascades. Older adults may be more vulnerable due to the nonspecific nature of ADR symptoms in older adults, e.g. falls, fatigue or constipation, all of which have multiple potential causes.⁸ Multimorbidity, which is more common in older adults, may also make the identification of new onset ADRs more challenging.^{9,10} However, the failure to correctly identify an ADR and the resultant prescribing cascade compounds the risk for medication-related harm.

To date prescribing cascades have remained under-researched. A previous scoping review identified only 10 original investigations and seven case reports that examined prescribing cascades.¹¹ In order to optimise prescribing, it is vital that clinically relevant prescribing cascades that commonly occur in practice are identified. The objective of this systematic review was to identify published prescribing cascades in community-dwelling adults.

Methods

Search protocol

The study protocol was previously published¹² and pre-registered with PROSPERO [CRD42021243163].¹³ This study was reported according to the Preferred Reporting Items for Systematic Reviews and Meta-analysis (PRISMA) guidelines.^{14,15} (eTable 1 in Supplementary Material).

Search strategy

Searches were conducted in the following databases: Medline (Ovid), EMBASE, PsycInfo, CINAHL and the Cochrane Library. Searches were initially conducted from inception to March 2021 and updated in February 2022. The search strategy (eBox 1 in Supplementary Material) was developed in consultation with an experienced librarian. No restrictions were placed on language or publication year. Grey literature database searches were conducted in MedNar, Dart Europe, Open Grey, and the Turning Research Into Practice (TRIP) databases using keyword searches. Forwards and backwards citation searching of articles selected for full text review was also conducted. Retrieved results were exported to EndNote X9 prior to screening and study selection using Covidence(r) systematic review management system. Following duplicates removal, titles and abstracts were independently screened by two reviewers (AD and EW, OC or FS) according to inclusion criteria. Disagreements were managed by consensus. Additional information was sought from study authors where necessary.

Studies were included if they met the following criteria:

1. Population: community-dwelling adults ([?]18 years).
2. Risk: prescription of medication that had the potential to cause an ADR that resulted in the prescription of further medication.
3. Outcome: prescribing cascade defined as the initiation of a new medication to ‘treat’ an ADR (unintentional cascade) or to reduce the risk of an ADR (intentional cascade).
4. Study type: prospective or retrospective cohort, cross-sectional, case-crossover, case-control or case-series studies.
5. Setting: primary care and community settings, including ambulatory care.

Exclusion criteria

The following studies were excluded;

1. Population of interest <18 years;
2. Studies conducted solely in nursing homes, residential care, inpatient settings or Emergency Departments (ED);
3. Case reports

Data extraction and quality assessment

Data extraction was conducted by two independent reviewers (AD and EW, OC or FS) using a standardised Microsoft Excel proforma. (see eBox 2, Supplementary Material). The methodological quality of included publications was independently performed in duplicate (AD and EW, OC or FS) using the appropriate JBI- Critical Appraisal checklist (eBox 3, Supplementary Material). Data synthesis was conducted using a narrative synthesis. Alluvial plots of drug pair combinations were created, using R-Studio 2021.09.2 statistical software using the ggalluvial package, to identify the drug-pair combinations examined and to summarise the overall quantitative association reported.

Results

Study identification

The study identification flow diagram is presented in Figure 1. A total of 103 publications relating to 101 studies met the inclusion criteria. Three publications included data from the same study relating to updated data collection time periods (2000-2006; 2000-2010; and 2000-2012).¹⁶⁻¹⁸ Thus, only the final study publication,¹⁸ which contained the entire data collection period, was included in the narrative synthesis.

Study population demographics

Seventy-nine studies presented study participants demographics, of which 15 specifically examined older adults ([?]60 years), with different age-related thresholds (e.g. [?]60 years; [?]65 years; [?]66 years) used across studies.^{5,19-32} Thirteen studies reported analyses stratified by age.^{7,33-44} Total study sample sizes ranged from 126⁴⁵ to 11,593,989⁴⁶ participants. (See eTable 3, Supplementary Material).

Methodological approach to analysis

Most studies (n=88) were retrospective cohort studies^{5,7,21,23-27,29,31-34,36-44,47-115}, three of which incorporated a case-control study within the study design^{49,87,112} and one that conducted a preliminary cross-sectional study¹¹⁴. Five were case-control studies^{19,20}, five cross-sectional studies^{6,46,116-118}, and three case-crossover studies^{45,119,120}. All studies used routine data (health insurance claims, prescription dispensing, clinical databases, national health surveys and pharmacovigilance data). In total, 83 studies examined dispensed prescriptions whereas 18 studies examined prescribed medications (see eTable 3, Supplementary Material).

Of 101 studies, 62 used prescription sequence symmetry analysis (PSSA) to determine the ratio of participants who initiated two medications in both possible sequences (i.e. Drug A- Drug B vs. Drug B- Drug A), with the majority (n=52) adjusting for prescribing trends.

Several studies reported stratified results by dosage,^{5,7,28,29,39} concomitant medication use or polypharmacy,^{7,40,44,87,106} duration,^{32,96} comorbidity,^{36,38,40,44} race³⁴ and nursing home residence.²⁶ For other studies, analyses were adjusted by age,^{20,22,30,52,71,91,121} sex,^{20,22,30,52,71,74,84,121} race,^{22,121} dose,^{52,71} nursing home residence,²² concomitant medication or polypharmacy,^{22,52,71} comorbidity,¹²¹ with some studies conducting adjusted analyses but not reporting the independent association of these covariates.^{23,27,31,90,119,120}

Length of follow up ranged from one month^{55,93,109,120} to seven years¹¹⁵, with the majority over one year (n=33 studies).

Initial medication(s) prescribed to patient

A broad range of medication types were examined as potentially precipitating a prescribing cascade (see Table 1 and column 1, Figure 2a). Ninety-four studies were hypothesis-driven or examined a predefined list of medications (Table 2 and Figure 2a). Seven studies conducting exploratory analyses to identify new signals of potential prescribing cascades are not represented in Figure 2.^{68,76,89,103,104,108,122} Initial medication Anatomical Therapeutic Classification (ATC) codes were not reported for 66 studies and were assigned by our research team.

Suspected adverse reaction(s)

Throughout the included studies, suspected ADRs were presumed to have occurred based on the initiation of the second medication as a treatment. In one study examining the CCB- loop diuretic prescribing cascade, an additional medical chart review was also conducted.¹⁰⁷

The suspected ADRs, symptoms or new diagnoses explored were broad-ranging (see Table 2) most commonly depression (n=13)^{33,37,40,45,52,55,57,73,95,99,100,112,114}; peripheral oedema (n=11)^{5-7,28,36,64,81,98,105-107,118}; urinary incontinence (n=9).^{24,26,41,44,50,53,75,81,119} and parkinsonism (n=9)^{27,29,31,46,57,81,84,123,124}

New medication(s) prescribed

The medication sub-classifications most frequently initiated as a new medication in the 94 studies are summarised in Figure 2a. Seventy-eight studies reported at least one significant positive association, indicating a potential prescribing cascade (Table 1 and Figure 2a-2c).

The most commonly identified prescribing cascades are summarised in Table 2. These include; amiodarone associated with subsequent thyroid hormone prescriptions for hypothyroidism (n=5),^{54,61,87,89,125} CCBs associated with diuretic prescriptions to treat peripheral oedema (n=5),^{5,7,89,106,107} topical antifungals to treat oral candidiasis following inhaled corticosteroids (n=4),^{39,54,56,71} anti-Parkinson medication to treat Parkinsonian symptoms following antipsychotic initiation (n=4),^{19,29,102,124} urinary anticholinergics to treat urinary incontinence following acetylcholinesterase inhibitors (n=4),^{26,44,53,121} and antitussives to treat cough following angiotensin-converting enzyme inhibitors (ACEIs) (n=3).^{18,74,89} Additional prescribing cascades identified included metoclopramide to anti-Parkinson medication (n=3).^{20,31,124} and NSAID to anti-ulcer medication.^{91,93,103}

No association between drug pairs could be determined for several studies, largely due to either a cross-sectional study design examining concurrent drug use, insufficient drug-pair sample size to determine a sequence ratio or reporting of incidence rates with no incidence rate ratio (labelled N/A in Figure 2).^{6,21,25,43,46,53,55,64,81,114,116,117,123,125,126} Several studies reported at least one negative association between drug pairs, indicating a reduced likelihood of the second medication being initiated (see eTable 3 Supplementary materials).^{33,60,68-70,75,83,89,91,95,113}

Modifiers of identified associations

Older people (aged [?]65 years) were more likely to receive; i) anticholinergics for urinary incontinence following SSRI initiation,⁴¹ ii) ulcer drug therapy within 100 days of NSAID initiation,⁹¹ iii) diuretic to treat beta-blocker induced oedema,³⁶ and, iv) thyroxine for hypothyroidism following amiodarone initiation.⁸⁷ Females were more likely to receive an antitussive for cough following ACEI initiation,⁷⁴ anticholinergic medication for urinary incontinence following acetylcholinesterase inhibitor^{24,30} and SSRI initiation,⁴¹ and levothyroxine following amiodarone initiation.⁸⁷

Differential associations were identified for initial medication dosage in nine studies. Those who received higher doses of CCBs^{5,7} and gabapentinoids were more likely to receive a diuretic for oedema;²⁸ higher doses of inhaled corticosteroids were associated with a greater likelihood of treatment for oral candidiasis;³⁹ and higher metoclopramide dosage was found to increase the likelihood for dopaminergic treatment initiation.²⁰ Polypharmacy ([?]5 drugs) was associated with a greater likelihood of receiving thyroid hormones for amiodarone induced hypothyroidism.⁸⁷

Intentional and unintentional cascades

The intentionality of potential prescribing cascades was not reported in any study nor was the intended duration (if any) of the prescription of the second medication. One study provided a breakdown of prescriptions for the initial drug by prescriber type: 23% private cardiologist, 35.5% hospital practitioner, 30.3% General Practitioner, and 11.3% other private specialist, but did not provide details of the prescriber of the second drug.⁹⁵ Another study reported that of the sample who initiated the second drug (irrespective of initiating the first drug), 87.1% of prescriptions were started by family physicians.⁵¹

Clinical importance of prescribing cascade

Two studies reported a number needed to harm (NNTH) for investigated cascades.^{62,104} (See Table 1). One study (n=90) conducted a medical chart validation study of those initiated a loop diuretic after initiating a dihydropyridine CCB (n=64) and determined that 54.7% (n=35) experienced a prescribing cascade.¹⁰⁷

Quality assessment

Overall, the methodological quality varied across included studies (Figure 3 and eTables 4-6, Supplementary Material). Among the retrospective cohort studies (eTable 4) there was a lack of clarity surrounding the similarity of exposed and unexposed groups at baseline and the presence of the outcome at the start of the study. For case-control studies (eTable 5), reporting of baseline comparison of cases and controls was inadequate as well as the appropriateness of matching cases with controls.

Conclusion and implications

Principal findings

This systematic review identified 101 studies across 103 publications that examined potential prescribing cascades across a broad range of pharmacological drug groups. All studies used routine administrative data that included either medication prescribing or dispensed medications data. Of the 101 included studies, 78 (77%) reported at least one significant positive quantitative association that indicates a potential prescribing cascade. The most commonly identified prescribing cascades include; i) CCBs- loop diuretics to treat peripheral oedema (n=5); amiodarone- thyroxine to treat hypothyroidism (n=5); inhaled corticosteroids- topical anti-fungal to treat candidiasis (n=4); antipsychotics- anti-Parkinson medication to treat Parkinsonism (n=4); and acetylcholinesterase inhibitors- drugs for urinary frequency (n=4).

Study methodological quality was variable with a considerable proportion of studies not reporting participant demographics. Almost two-thirds of included studies used PSSA methodology in which all included participants have experienced the outcome at the start of the study. A recent scoping review reported that whilst the PSSA method is a useful tool in detecting prescribing cascades, such cascades need careful clinical review as there is a risk of both false positive and false negative findings.¹²⁷ This is particularly problematic when screening for cascades without predefined hypotheses. In our systematic review, the vast majority of

included studies (n=94, 93%) examined predefined medications as potentially contributing to a prescribing cascade. However, PSSA analyses cannot determine causality and should be interpreted with caution.

Several well-designed cohort and case-control studies examining prescribing cascades were identified. For example, a Canadian population-based study reported that incident CCB users had a higher cumulative incidence of loop diuretic use at one year follow up compared to patients dispensed ACEIs or angiotensin-II-receptor blocker antihypertensives (adjusted hazards ratio 1.4% vs 0.7%, $p < 0.001$).⁵ In a US case-control study, metoclopramide users were three times more likely to begin use of a levodopa-containing medication compared with nonusers (OR = 3.09; 95% CI 2.25 to 4.26).²⁰ Risk increased with increasing daily metoclopramide dose and the effect persisted after adjustment for demographic, health service utilization, and medication use variables.²⁰

Fifteen of 101 studies focused specifically on older populations, with 11 reporting a significant association between increasing age and prescribing cascade occurrence. Older adults are more likely to experience medication-related harm due to increasing prevalence of multimorbidity, polypharmacy and age-related physiological changes that affect drug metabolism.^{9,10,128-130} Furthermore, ADRs are more difficult to diagnose in older adults due to their often non-specific presentation and overlap with pre-existing conditions or conditions likely to develop among older adults.^{1,8,131}

Comparison with existing literature

Two scoping reviews of prescribing cascades have been conducted to date, one that focused on literature surrounding the prevention, detection and reversal of prescribing cascades¹¹ and the second that focused on the use of PSSA as a potential pharmacovigilance tool.¹²⁷ In 2018, Brath et al retrieved 10 original investigations and seven case reports pertaining to prescribing cascades.¹¹ A considerable number of studies have been published since, indicating that this is a rapidly developing field. Morris et al. concluded that PSSA methodology demonstrated only moderate sensitivity and specificity in identifying prescribing cascades and more consistency was required in how these studies were reported.¹²⁷ As described previously, similar issues with methodological quality were identified in this systematic review.

Clinical and research implications

Multi-country studies have shown variation in prescribing cascade likelihood both within and across countries,^{60,97,98} underscoring the need to consider the local prescribing context. Differences in sample demographics, medication availability, approved clinical indications, help-seeking behaviour and prescribing cultures or genetic polymorphisms may influence the incidence of prescribing cascades.

The complexity of optimising prescribing for patients with multimorbidity presents challenges for the prescriber due to the preponderance of single-disease guidelines, resultant polypharmacy, fragmentation and lack of continuity of care and resourcing constraints.¹³² Identification of ADRs remains a clinically challenging area, particularly in relation to older adults. Non-specific presentation of ADR symptoms in older adults, such as delirium, falls, fatigue and constipation, can be challenging to identify as being medication-related as such symptoms have several causes and may overlap with existing multimorbidity.^{8,131} The failure to recognise an ADR may result in a prescribing cascade, furthering the risk for additional medication-related harm.^{1,2} The potential for ADRs should be considered as part of the differential diagnosis for all patients reporting new symptoms, particularly among those who have started a new medication within the previous year.^{1,8,131}

The use of routine administrative data in included studies means that information on the broader clinical context and the rationale for medication prescribing is lacking. The identification of significant negative associations between drug pairs may indicate that prescribers are aware of certain prescribing cascades and proactively avoid their development or that therapeutic alternatives were prescribed. However, no exploration of intentionality of identified cascades could be made based on the data used in included studies.

Overall, it is difficult to determine the clinical importance of prescribing cascades identified as few studies examined clinical endpoints.^{48,62,104} One study examined the association between prescribing cascades that

resulted in prochlorperazine initiation and reported a subsequent 49% increased risk of hip fracture.⁴⁸ Future research is required to determine the relative clinical impact of increased medication exposure and the clinical appropriateness of prescribing cascades.

Strengths and limitations

This systematic review extends the work of previously published scoping reviews^{11,127} by conducting a comprehensive literature search using several databases, including several grey literature searches.

This study also has some limitations. The lack of a MeSH term for prescribing cascades meant broad search terms were used, which led to a high yield of citations to be searched. Additional information was sought from study authors but a small number of studies (n=10) could not be retrieved for eligibility assessment due to the lack of access to the full text or a translated version. The information collated is somewhat limited by the methodological and reporting quality of included studies.

Conclusion

Prescribing cascades are of increasing interest to the research and clinical communities, with a broad range of medications involved. The identification of the most common prescribing cascades can support optimising prescribing as one part of identifying potentially inappropriate prescribing. Few studies have examined the clinical importance or the broader clinical context, including intentionality of prescribing cascades, thereby limiting the inferences that can be drawn about the implications for clinical practice. Challenges remain in differentiating ADR symptoms from that of new onset disease and advancing age and frailty.^{1,8,131} ADRs should be considered as part of the differential diagnosis in patients presenting with new symptoms, particularly for those who have started a new medication in the preceding 12 months.

References

1. Rochon PA, Gurwitz JH. Optimising drug treatment for elderly people: the prescribing cascade. *BMJ-British Medical Journal*.1997;315(7115):1096-1099.
2. Rochon PA, Gurwitz JH. The prescribing cascade revisited.*Lancet (London, England)*. 2017;389(10081):1778-1780.
3. Rochon PA, Gurwitz JH. Drug therapy. *The Lancet*.1995;346(8966):32-36.
4. McCarthy LM, Visentin JD, Rochon PA. Assessing the scope and appropriateness of prescribing cascades. *Journal of the American Geriatrics Society*. 2019;67(5):1023-1026.
5. Savage RD, Visentin JD, Bronskill SE, et al. Evaluation of a Common Prescribing Cascade of Calcium Channel Blockers and Diuretics in Older Adults With Hypertension. *JAMA internal medicine*.2020;180(5):643-651.
6. Vouri SM, van Tuyl JS, Olsen MA, Hong X, Schootman M, Xian H. An evaluation of a potential calcium channel blocker-lower-extremity edema-loop diuretic prescribing cascade. *Journal of the American Pharmacists Association: JAPhA*. 2018;58(5):534-539.
7. Vouri SM, Jiang X, Manini TM, et al. Magnitude of and characteristics associated with the treatment of calcium channel blocker-induced lower-extremity edema with loop diuretics. *JAMA network open*.2019;2(12):e1918425.
8. Lavan AH, Gallagher P. Predicting risk of adverse drug reactions in older adults. *Therapeutic Advances in Drug Safety*.2016;7(1):11-22.
9. Palladino R, Tayu Lee J, Ashworth M, Triassi M, Millett C. Associations between multimorbidity, health-care utilisation and health status: evidence from 16 European countries. *Age and Ageing*.2016;45(3):431-435.

10. Salisbury C, Johnson L, Purdy S, Valderas JM, Montgomery AA. Epidemiology and impact of multi-morbidity in primary care: a retrospective cohort study. *Br J Gen Pract.* 2011;61(582):e12-e21.
11. Brath H, Mehta N, Savage RD, et al. What is known about preventing, detecting, and reversing prescribing cascades: A scoping review. *Journal of the American Geriatrics Society.* 2018;66(11):2079-2085.
12. Moher D, Shamseer L, Clarke M, et al. Preferred reporting items for systematic review and meta-analysis protocols (PRISMA-P) 2015 statement. *Systematic Reviews.* 2015;4.
13. Doherty A, Moriarty F, Boland F, et al. Prescribing cascades in community-dwelling adults: protocol for a systematic review. *HRB open research.* 2021;4:72.
14. Moher D, Liberati A, Tetzlaff J, Altman DG, Grp P. Preferred reporting items for systematic reviews and meta-analyses: The PRISMA statement. *Annals of Internal Medicine.* 2009;151(4):264-W264.
15. Page MJ, McKenzie JE, Bossuyt PM, et al. The PRISMA 2020 statement: an updated guideline for reporting systematic reviews. *BMJ (Clinical research ed).* 2021;372:n71.
16. Vegter S, De Jong-Van Den Berg LTW. Misdiagnosis and mistreatment of a common side-effect - Angiotensin-converting enzyme inhibitor-induced cough. *British Journal of Clinical Pharmacology.* 2010;69(2):200-203.
17. Vegter S, De Boer P, Van Dijk KW, Visser S, De Jong-van Den Berg LTW. Misdiagnosis and mistreatment of ACE-inhibitor-induced cough occurs frequently and decreases therapy compliance. *Pharmaceutisch Weekblad.* 2012;147(42):177-180.
18. Vegter S, de Boer P, van Dijk KW, Visser S, de Jong-van den Berg LTW. The effects of antitussive treatment of ACE inhibitor-induced cough on therapy compliance: a prescription sequence symmetry analysis. *Drug safety.* 2013;36(6):435-439.
19. Avorn J, Bohn RL, Mogun H, et al. Neuroleptic drug exposure and treatment of parkinsonism in the elderly: a case-control study. *Am J Med.* 1995;99(1):48-54.
20. Avorn J, Gurwitz JH, Bohn RL, Mogun H, Monane M, Walker A. Increased incidence of levodopa therapy following metoclopramide use. *Jama.* 1995;274(22):1780-1782.
21. Farkas AH, Winn A, Pezzin LE, Fergestrom N, Laud P, Neuner JM. The Use and Concurrent Use of Side Effect Controlling Medications Among Women on Aromatase Inhibitors. *Journal of Women's Health (15409996).* 2021;30(1):131-136.
22. Gurwitz JH, Avorn J, Bohn RL, Glynn RJ, Monane M, Mogun H. Initiation of antihypertensive treatment during nonsteroidal anti-inflammatory drug therapy. *Jama.* 1994;272(10):781-786.
23. Gurwitz JH, Kalish SC, Bohn RL, et al. Thiazide diuretics and the initiation of anti-gout therapy. *J Clin Epidemiol.* 1997;50(8):953-959.
24. Masurkar PP, Chatterjee S, Sherer JT, Aparasu RR. Antimuscarinic Cascade Across Individual Cholinesterase Inhibitors in Older Adults with Dementia. *Drugs and Aging.* 2021;38(7):593-602.
25. Narayan SW, Pearson SA, Litchfield M, et al. Anticholinergic medicines use among older adults before and after initiating dementia medicines. *Br J Clin Pharmacol.* 2019;85(9):1957-1963.
26. Gill SS, Mamdani M, Naglie G, et al. A prescribing cascade involving cholinesterase inhibitors and anticholinergic drugs. *Archives of Internal Medicine.* 2005;165(7):808-813.
27. Marras C, Herrmann N, Fischer HD, et al. Lithium Use in Older Adults is Associated with Increased Prescribing of Parkinson Medications. *The American journal of geriatric psychiatry : official journal of the American Association for Geriatric Psychiatry.* 2016;24(4):301-309.

28. Read SH, Giannakeas V, Pop P, et al. Evidence of a gabapentinoid and diuretic prescribing cascade among older adults with lower back pain. *Journal of the American Geriatrics Society*. 2021;69(10):2842-2850.
29. Rochon PA, Stukel TA, Sykora K, et al. Atypical antipsychotics and parkinsonism. *Arch Intern Med*. 2005;165(16):1882-1888.
30. Trenaman SC, Bowles SK, Kirkland S, Andrew MK. An examination of three prescribing cascades in a cohort of older adults with dementia. *BMC Geriatrics*. 2021;21(1):1-11.
31. Huh Y, Kim DH, Choi M, et al. Metoclopramide and levosulpiride use and subsequent levodopa prescription in the korean elderly: The prescribing cascade. *Journal of Clinical Medicine*. 2019;8(9).
32. Park SK, Baek YH, Pratt N, Kalisch Ellett L, Shin JY. The Uncertainty of the Association Between Proton Pump Inhibitor Use and the Risk of Dementia: Prescription Sequence Symmetry Analysis Using a Korean Healthcare Database Between 2002 and 2013. *Drug Safety*. 2018;41(6):615-624.
33. Dyson TE, Cantrell MA, Lund BC. Lack of Association between 5 α -Reductase Inhibitors and Depression. *The Journal of urology*. 2020;204(4):793-798.
34. Fox CW, Khaw CL, Gerke AK, Lund BC. Montelukast and neuropsychiatric events—a sequence symmetry analysis. *Journal of Asthma*. 2022.
35. Vouri SM, Jiang X, Brumback B, Winterstein AG. Use of negative controls in a prescription sequence symmetry analysis used to mitigate time-varying bias. *Pharmacoepidemiology and Drug Safety*. 2020;29(SUPPL 3):390-391.
36. Vouri SM, Morris EJ, Jiang X, et al. Evaluation of a Beta-blocker - Edema - Loop Diuretic Prescribing Cascade: A Prescription Sequence Symmetry Analysis. *American journal of hypertension*. 2022.
37. Brandt-Christensen M, Kvist K, Nilsson FM, Andersen PK, Kessing LV. Treatment with antiparkinson and antidepressant drugs: a register-based, pharmaco-epidemiological study. *Movement disorders : official journal of the Movement Disorder Society*. 2007;22(14):2037-2042.
38. Dunvald ACD, Henriksen DP, Hallas J, Christensen MMH, Lund LC. Selective serotonin reuptake inhibitors and the risk of restless legs syndrome: a symmetry analysis. *European Journal of Clinical Pharmacology*. 2020;76(5):719-722.
39. Henriksen DP, Davidsen JR, Christiansen A, Laursen CB, Damkier P, Hallas J. Inhaled Corticosteroids and Systemic or Topical Antifungal Therapy: A Symmetry Analysis. *Annals of the American Thoracic Society*. 2017;14(6):1045-1047.
40. Winkel JS, Damkier P, Hallas J, Henriksen DP. Treatment with montelukast and antidepressive medication—a symmetry analysis. *Pharmacoepidemiology and Drug Safety*. 2018;27(12):1409-1415.
41. Movig KLL, Leufkens HGM, Belitser SV, Lenderink AW, Egberts ACG. Selective serotonin reuptake inhibitor-induced urinary incontinence. *Pharmacoepidemiology and drug safety*. 2002;11(4):271-279.
42. Wang Y, Bos JH, Schuiling-Veninga CCM, et al. Neuropsychiatric safety of varenicline in the general and COPD population with and without psychiatric disorders: a retrospective cohort study in a real-world setting. *BMJ open*. 2021;11(5):e042417.
43. Takeuchi Y, Kajiyama K, Ishiguro C, Uyama Y. Atypical Antipsychotics and the Risk of Hyperlipidemia: A Sequence Symmetry Analysis. *Drug Saf*. 2015;38(7):641-650.
44. Lampela P, Taipale H, Hartikainen S. Use of Cholinesterase Inhibitors Increases Initiation of Urinary Anticholinergics in Persons with Alzheimer's Disease. *J Am Geriatr Soc*. 2016;64(7):1510-1512.
45. Azoulay L, Blais L, Koren G, LeLorier J, Bérard A. Isotretinoin and the risk of depression in patients with acne vulgaris: a case-crossover study. *The Journal of clinical psychiatry*. 2008;69(4):526-532.

46. Onder G, Bonassi S, Abbatecola AM, et al. High Prevalence of Poor Quality Drug Prescribing in Older Individuals: A Nationwide Report From the Italian Medicines Agency (AIFA). *Journals of Gerontology Series A: Biological Sciences & Medical Sciences*. 2014;69(4):430-437.
47. Adimadhyam S, Schumock GT, Calip GS, Smith Marsh DE, Layden BT, Lee TA. Increased risk of mycotic infections associated with sodium-glucose co-transporter 2 inhibitors: a prescription sequence symmetry analysis. *British journal of clinical pharmacology*. 2019;85(1):160-168.
48. Caughey GE, Roughead EE, Pratt N, Shakib S, Vitry AI, Gilbert AL. Increased risk of hip fracture in the elderly associated with prochlorperazine: is a prescribing cascade contributing? *Pharmacoepidemiol Drug Saf*. 2010;19(9):977-982.
49. Corrao G, Botteri E, Bagnardi V, et al. Generating signals of drug-adverse effects from prescription databases and application to the risk of arrhythmia associated with antibacterials. *Pharmacoepidemiology and drug safety*. 2005;14(1):31-40.
50. Fujimoto M, Higuchi T, Hosomi K, Takada M. Association of statin use with storage lower urinary tract symptoms (LUTS): data mining of prescription database. *International journal of clinical pharmacology and therapeutics*. 2014;52(9):762-769.
51. Garrison SR, Dormuth CR, Morrow RL, Carney GA, Khan KM. Nocturnal leg cramps and prescription use that precedes them: A sequence symmetry analysis. *Archives of Internal Medicine*. 2012;172(2):120-126.
52. Hallas J. Evidence of depression provoked by cardiovascular medication: a prescription sequence symmetry analysis. *Epidemiology (Cambridge, Mass)*. 1996;7(5):478-484.
53. Hashimoto M, Hashimoto K, Ando F, Kimura Y, Nagase K, Arai K. Prescription rate of medications potentially contributing to lower urinary tract symptoms and detection of adverse reactions by prescription sequence symmetry analysis. *Journal of Pharmaceutical Health Care and Sciences*. 2015;1(1).
54. Nishtala PS, Chyou TY. Exploring New Zealand prescription data using sequence symmetry analyses for predicting adverse drug reactions. *Journal of clinical pharmacy and therapeutics*. 2017;42(2):189-194.
55. Petri H, de Vet HC, Naus J, Urquhart J. Prescription sequence analysis: a new and fast method for assessing certain adverse reactions of prescription drugs in large populations. *Statistics in medicine*. 1988;7(11):1171-1175.
56. Petri H, Kessels F, Kamakura T. Markers of adverse drug reactions in medication histories. An analysis of inhaled steroid utilization. *Pharmaceutisch weekblad Scientific edition*. 1991;13(2):97-106.
57. Petri H, Leufkens H, Naus J, Silkens R, Van Hessen P, Urquhart J. Rapid method for estimating the risk of acutely controversial side effects of prescription drugs. *Journal of Clinical Epidemiology*. 1990;43(5):433-439.
58. Pouwels K, Visser S, Bos J, Hak E. Angiotensin-converting enzyme inhibitor treatment and the development of urinary tract infection. *Pharmacoepidemiology and Drug Safety*. 2013;22:127-128.
59. Adrian Kym P, Elizabeth Ellen R, Nicole Leanne P. Sequence symmetry analysis graphic adjustment for prescribing trends. *BMC medical research methodology*. 2019;19(1):143.
60. Pratt N, Andersen M, Bergman U, et al. Multi-country rapid adverse drug event assessment: The Asian Pharmacoepidemiology Network (AsPEN) antipsychotic and acute hyperglycaemia study. *Pharmacoepidemiology and Drug Safety*. 2013;22(9):915-924.
61. Pratt N, Chan EW, Choi NK, et al. Prescription sequence symmetry analysis: Assessing risk, temporality, and consistency for adverse drug reactions across datasets in five countries. *Pharmacoepidemiology and Drug Safety*. 2015;24(8):858-864.
62. Rasmussen L, Hallas J, Madsen KG. Cardiovascular drugs and erectile dysfunction - A symmetry analysis. *British Journal of Clinical Pharmacology*. 2015;80(5):1219-1223.

63. Singh S, Cocoros NM, Haynes K, et al. Antidopaminergic-Antiparkinsonian Medication Prescribing Cascade in Persons with Alzheimer’s Disease. *Journal of the American Geriatrics Society*. 2021.
64. Singh S, Cocoros NM, Haynes K, et al. Identifying prescribing cascades in Alzheimer’s disease and related dementias: The calcium channel blocker-diuretic prescribing cascade. *Pharmacoepidemiology and drug safety*. 2021.
65. Sturkenboom MC, Middelbeek A, de Jong van den Berg LT, van den Berg PB, Stricker BH, Wesseling H. Vulvo-vaginal candidiasis associated with acitretin. *Journal of clinical epidemiology*. 1995;48(8):991-997.
66. Takada M, Fujimoto M, Hosomi K. Association between Benzodiazepine Use and Dementia: Data Mining of Different Medical Databases. *International journal of medical sciences*. 2016;13(11):825-834.
67. Takada M, Fujimoto M, Hosomi K. Difference in risk of gastrointestinal complications between users of enteric-coated and buffered low-dose aspirin. *International journal of clinical pharmacology and therapeutics*. 2014;52(3):181-191.
68. Tsiropoulos I, Andersen M, Hallas J. Adverse events with use of antiepileptic drugs: A prescription and event symmetry analysis. *Pharmacoepidemiology and Drug Safety*. 2009;18(6):483-491.
69. Pouwels KB, Widyakusuma NN, Bos JHJ, Hak E. Association between statins and infections among patients with diabetes: a cohort and prescription sequence symmetry analysis. *Pharmacoepidemiology and drug safety*. 2016;25(10):1124-1130.
70. Venalainen O, Bell JS, Kirkpatrick CM, Nishtala PS, Liew D, Ilomaki J. Adverse Drug Reactions Associated With Cholinesterase Inhibitors-Sequence Symmetry Analyses Using Prescription Claims Data. *Journal of the American Medical Directors Association*. 2017;18(2):186-189.
71. van Boven JFM, de Jong-van den Berg LTW, Vegter S. Inhaled corticosteroids and the occurrence of oral candidiasis: a prescription sequence symmetry analysis. *Drug safety*. 2013;36(4):231-236.
72. Vegter S, de Boer P, van Dijk KW, Visser S, de Jong-van den Berg LTW. The effects of antitussive treatment of ACE inhibitor-induced cough on therapy compliance: a prescription sequence symmetry analysis. *Drug Safety*. 2013;36(6):435-439.
73. Hersom K, Neary MP, Levaux HP, Klaskala W, Strauss JS. Isotretinoin and antidepressant pharmacotherapy: a prescription sequence symmetry analysis. *Journal of the American Academy of Dermatology*. 2003;49(3):424-432.
74. Bowman L, Carlstedt BC, Miller ME, McDonald CJ. Evaluation of ACE-inhibitor (ACE-I) associated cough using modified prescription sequence analysis (PSA). *Pharmacoepidemiology and Drug Safety*. 1995;4(1):17-22.
75. Kalisch Ellett LM, Pratt NL, Barratt JD, Rowett D, Roughead EE. Risk of medication-associated initiation of oxybutynin in elderly men and women. *Journal of the American Geriatrics Society*. 2014;62(4):690-695.
76. King CE, Pratt NL, Craig N, et al. Detecting Medicine Safety Signals Using Prescription Sequence Symmetry Analysis of a National Prescribing Data Set. *Drug Safety*. 2020;43(8):787-795.
77. Ko HHT, Lareu RR, Dix BR, Hughes JD, Parsons RW. A sequence symmetry analysis of the interrelationships between statins, diabetes and skin infections. *British Journal of Clinical Pharmacology*. 2019.
78. Knowledge and confidence in medication management. *Nursing management (Harrow, London, England : 1994)*. 2017;24(5):14.
79. Lai EC-C, Hsieh C-Y, Yang Y-HK, Lin S-J. Detecting potential adverse reactions of sulpiride in schizophrenic patients by prescription sequence symmetry analysis. *PLoS ONE*. 2014;9(2).
80. Lai EC-C, Yang Y-HK, Lin S-J, Hsieh C-Y. Use of antiepileptic drugs and risk of hypothyroidism. *Pharmacoepidemiology and drug safety*. 2013;22(10):1071-1079.

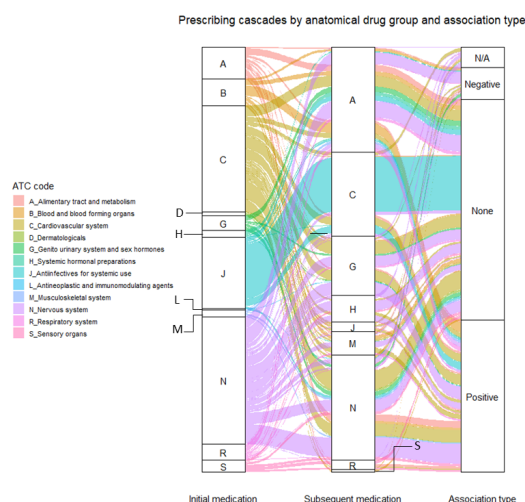
81. Trenaman SC, Bowles SK, Kirkland S, Andrew MK. An examination of three prescribing cascades in a cohort of older adults with dementia. *BMC geriatrics*. 2021;21(1):297.
82. Wang Y, van Boven JFM, Bos JHJ, et al. Risk of neuropsychiatric adverse events associated with varenicline treatment for smoking cessation among Dutch population: A sequence symmetry analysis. *Pharmacoepidemiology and Drug Safety*. 2021.
83. Bytzer P, Hallas J. Drug-induced symptoms of functional dyspepsia and nausea. A symmetry analysis of one million prescriptions. *Alimentary pharmacology & therapeutics*. 2000;14(11):1479-1484.
84. Brandt-Christensen M, Kvist K, Nilsson FM, Andersen PK, Kessing LV. Treatment with antidepressants and lithium is associated with increased risk of treatment with antiparkinson drugs: a pharmacoepidemiological study. *Journal of neurology, neurosurgery, and psychiatry*. 2006;77(6):781-783.
85. Iwasawa M, Sagami K, Yokoyama S, Hosomi K, Takada M. Adherence to guidelines for antiulcer drug prescription in patients receiving low-dose aspirin therapy in Japan. *International journal of clinical pharmacology and therapeutics*. 2019;57(4):197-206.
86. Yokoyama S, Ieda S, Nagano M, et al. Association between oral anticoagulants and osteoporosis: Real-world data mining using a multi-methodological approach. *Int J Med Sci*. 2020;17(4):471-479.
87. Yokoyama S, Tanaka Y, Hosomi K, Takada M. Polypharmacy is associated with amiodarone-induced hypothyroidism. *International Journal of Medical Sciences*. 2021;18(15):3574-3580.
88. Yokoyama S, Wakamoto S, Tanaka Y, Nakagawa C, Hosomi K, Takada M. Association Between Antipsychotics and Osteoporosis Based on Real-World Data. *The Annals of pharmacotherapy*. 2020;54(10):988-995.
89. Chen Y, Huang ST, Hsu TC, Peng LN, Hsiao FY, Chen LK. Detecting Suspected Prescribing Cascades by Prescription Sequence Symmetry Analysis of Nationwide Real-World Data. *Journal of the American Medical Directors Association*. 2021.
90. Gadzhanova S, Pratt N, Roughead E. Use of SGLT2 inhibitors for diabetes and risk of infection: Analysis using general practice records from the NPS MedicineWise MedicineInsight program. *Diabetes research and clinical practice*. 2017;130:180-185.
91. Hallas J, Bytzer P. Screening for drug related dyspepsia: an analysis of prescription symmetry. *European journal of gastroenterology & hepatology*. 1998;10(1):27-32.
92. Genia English, August 10, 1996. *National Library of Medicine*.
93. Pouwels K, Kalkman A, Schagen D, Visser S, Hak E. Do SSRIs increase the risk of gastrointestinal adverse effects? *Pharmacoepidemiology and Drug Safety*. 2013;22:127.
94. Silwer L, Petzold M, Hallas J, Lundborg CS. Statins and nonsteroidal anti-inflammatory drugs-an analysis of prescription symmetry. *Pharmacoepidemiol Drug Saf*. 2006;15(7):510-511.
95. Maura G, Billionnet C, Coste J, Weill A, Neumann A, Pariente A. Non-bleeding Adverse Events with the Use of Direct Oral Anticoagulants: A Sequence Symmetry Analysis. *Drug Saf*. 2018;41(9):881-897.
96. Park KR, Kim KB, Baek YH, et al. Signal detection of benzodiazepine use and risk of dementia: sequence symmetry analysis using South Korean national healthcare database. *International journal of clinical pharmacy*. 2018;40(6):1568-1576.
97. Roughead EE, Chan EW, Choi NK, et al. Proton pump inhibitors and risk of *Clostridium difficile* infection: a multi-country study using sequence symmetry analysis. *Expert opinion on drug safety*. 2016;15(12):1589-1595.
98. Roughead EE, Chan EW, Choi NK, et al. Variation in Association Between Thiazolidinediones and Heart Failure Across Ethnic Groups: Retrospective analysis of Large Healthcare Claims Databases in Six Countries. *Drug Saf*. 2015;38(9):823-831.

99. Roughead EE, Kalisch LM, Pratt NL, Killer G, Barnard A, Gilbert AL. Managing glaucoma in those with co-morbidity: not as easy as it seems. *Ophthalmic epidemiology*. 2012;19(2):74-82.
100. Lindberg G, Hallas J. Cholesterol-lowering drugs and antidepressants—a study of prescription symmetry. *Pharmacoepidemiol Drug Saf*. 1998;7(6):399-402.
101. Janetzki JL, Sykes MJ, Ward MB, Pratt NL. Proton pump inhibitors may contribute to progression or development of chronic obstructive pulmonary disease—A sequence symmetry analysis approach. *Journal of Clinical Pharmacy and Therapeutics*. 2021;46(6):1687-1694.
102. Hirano Y. Risk of Extrapyramidal Syndromes Associated With Psychotropic Polypharmacy: A Study Based on Large-Scale Japanese Claims Data. *Therapeutic innovation & regulatory science*. 2020;54(2):259-268.
103. Hallas J, Wang SV, Gagne JJ, Schneeweiss S, Pratt N, Pottegård A. Hypothesis-free screening of large administrative databases for unsuspected drug-outcome associations. *European journal of epidemiology*. 2018;33(6):545-555.
104. Hellfritsch M, Rasmussen L, Hallas J, Pottegård A. Using the Symmetry Analysis Design to Screen for Adverse Effects of Non-vitamin K Antagonist Oral Anticoagulants. *Drug Saf*. 2018;41(7):685-695.
105. Alaskar MA, Brown JD, Voils SA, Vouri SM. Loop diuretic use following fluid resuscitation in the critically ill. *American journal of health-system pharmacy : AJHP : official journal of the American Society of Health-System Pharmacists*. 2022;79(3):165-172.
106. Vouri SM, Jiang X, Morris EJ, Brumback BA, Winterstein AG. Use of negative controls in a prescription sequence symmetry analysis to reduce time-varying bias. *Pharmacoepidemiology and Drug Safety*. 2021;30(9):1192-1199.
107. Vouri SM, Morris EJ, Usmani SA, et al. Evaluation of the key prescription sequence symmetry analysis assumption using the calcium channel blocker: Loop diuretic prescribing cascade. *Pharmacoepidemiology and Drug Safety*. 2022;31(1):72-81.
108. Wahab IA, Pratt NL, Ellett LK, Roughead EE. Sequence Symmetry Analysis as a Signal Detection Tool for Potential Heart Failure Adverse Events in an Administrative Claims Database. *Drug Saf*. 2016;39(4):347-354.
109. Thacker EL, Schneeweiss S. Initiation of acetylcholinesterase inhibitors and complications of chronic airways disorders in elderly patients. *Drug Saf*. 2006;29(11):1077-1085.
110. de Jong JC, van den Berg PB, Tobi H, de Jong-van den Berg LT. Combined use of SSRIs and NSAIDs increases the risk of gastrointestinal adverse effects. *Br J Clin Pharmacol*. 2003;55(6):591-595.
111. Hachiken H, Murai A, Wada K, Kuwahara T, Hosomi K, Takada M. Difference between the frequencies of antisecretory drug prescriptions in users of buffered vs. enteric-coated low-dose aspirin therapies. *International journal of clinical pharmacology and therapeutics*. 2013;51(10):807-815.
112. Hagberg KW, Divan HA, Nickel JC, Jick SS. Risk of Incident Antidepressant-Treated Depression Associated with Use of 5 α -Reductase Inhibitors Compared with Use of α -Blockers in Men with Benign Prostatic Hyperplasia: A Population-Based Study Using the Clinical Practice Research Datalink. *Pharmacotherapy*. 2017;37(5):517-527.
113. Lund LC, Højlund M, Henriksen DP, Hallas J, Kristensen KB. Sodium-glucose cotransporter-2 inhibitors and the risk of gout: A Danish population based cohort study and symmetry analysis. *Pharmacoepidemiol Drug Saf*. 2021;30(10):1391-1395.
114. Thiessen BQ, Wallace SM, Blackburn JL, Wilson TW, Bergman U. Increased prescribing of antidepressants subsequent to beta-blocker therapy. *Arch Intern Med*. 1990;150(11):2286-2290.

115. Wahab IA, Pratt NL, Kalisch LM, Roughead EE. Comparing time to adverse drug reaction signals in a spontaneous reporting database and a claims database: a case study of rofecoxib-induced myocardial infarction and rosiglitazone-induced heart failure signals in Australia. *Drug Saf.* 2014;37(1):53-64.
116. Kalisch Ellett LM, Pratt NL, Kerr M, Roughead EE. Antipsychotic polypharmacy in older Australians. *International psychogeriatrics.* 2018;30(4):539-546.
117. Vouri SM. Rhinorrhea as a Result of Alzheimer's Disease Treatment. *The Senior care pharmacist.* 2020;35(4):148-149.
118. Morris EJ, Brown JD, Manini TM, Vouri SM. Differences in Health-Related Quality of Life Among Adults with a Potential Dihydropyridine Calcium Channel Blocker–Loop Diuretic Prescribing Cascade. *Drugs and Aging.* 2021;38(7):625-632.
119. Kröger E, Van Marum R, Souverein P, Carmichael PH, Egberts T. Treatment with rivastigmine or galantamine and risk of urinary incontinence: results from a Dutch database study. *Pharmacoepidemiol Drug Saf.* 2015;24(3):276-285.
120. Pouwels KB, Bos JH, Hak E. ACE inhibitors and urinary tract infections. *Epidemiology.* 2014;25(3):466-467.
121. Masurkar PP, Chatterjee S, Sherer JT, Aparasu RR. Antimuscarinic Cascade Across Individual Cholinesterase Inhibitors in Older Adults with Dementia. *Drugs & Aging.* 2021;38(7):593-602.
122. Lai EC-C, Hsieh C-Y, Kao Yang Y-H, Lin S-J. Detecting potential adverse reactions of sulpiride in schizophrenic patients by prescription sequence symmetry analysis. *PloS one.* 2014;9(2):e89795.
123. Singh S, Cocoros NM, Haynes K, et al. Antidopaminergic-Antiparkinsonian Medication Prescribing Cascade in Persons with Alzheimer's Disease. *Journal of the American Geriatrics Society.* 2021;69(5):1328-1333.
124. Kim S, Cheon SM, Suh HS. Association Between Drug Exposure and Occurrence of Parkinsonism in Korea: A Population-Based Case-Control Study. *The Annals of pharmacotherapy.* 2019;53(11):1102-1110.
125. Lai ECC, Yang YHK, Lin SJ, Hsieh CY. Use of antiepileptic drugs and risk of hypothyroidism. *Pharmacoepidemiology and Drug Safety.* 2013;22(10):1071-1079.
126. Lai ECC, Hsieh CY, Yang YHK, Lin SJ. Detecting potential adverse reactions of sulpiride in schizophrenic patients by prescription sequence symmetry analysis. *PLoS ONE.* 2014;9(2).
127. Morris EJ, Hollmann J, Hofer AK, et al. Evaluating the use of prescription sequence symmetry analysis as a pharmacovigilance tool: A scoping review. *Research in social & administrative pharmacy : RSAP.* 2021.
128. Moriarty F, Hardy C, Bennett K, Smith SM, Fahey T. Trends and interaction of polypharmacy and potentially inappropriate prescribing in primary care over 15 years in Ireland: a repeated cross-sectional study. *BMJ open.* 2015;5(9).
129. Mangoni AA, Jackson SHD. Age-related changes in pharmacokinetics and pharmacodynamics: basic principles and practical applications. *British Journal Of Clinical Pharmacology.* 2004;57(1):6-14.
130. Guthrie B, Makubate B, Hernandez-Santiago V, Dreischulte T. The rising tide of polypharmacy and drug-drug interactions: Population database analysis 1995-2010. *Bmc Medicine.* 2015;13.
131. Petrovic M, van der Cammen T, Onder G. Adverse Drug Reactions in Older People. *Drugs & Aging.* 2012;29(6):453-462.
132. Wallace E, Salisbury C, Guthrie B, Lewis C, Fahey T, Smith SM. Managing patients with multimorbidity in primary care. *BMJ-British Medical Journal.* 2015;350.

133. Pouwels KB, Visser ST, Bos HJ, Hak E. Angiotensin-converting enzyme inhibitor treatment and the development of urinary tract infections: a prescription sequence symmetry analysis. *Drug safety*.2013;36(11):1079-1086.
134. Takada M, Fujimoto M, Yamazaki K, Takamoto M, Hosomi K. Association of statin use with sleep disturbances: data mining of a spontaneous reporting database and a prescription database. *Drug safety*.2014;37(6):421-431.
135. Gau CS, Chang CJ, Tsai FJ, Chao PF, Gau SS. Association between mood stabilizers and hypothyroidism in patients with bipolar disorders: a nested, matched case-control study. *Bipolar disorders*.2010;12(3):253-263.
136. Vouri SM, Possinger MC, Usmani S, Solberg LM, Manini T. Evaluation of the Potential Acetylcholinesterase Inhibitor-Induced Rhinorrhea Prescribing Cascade. *Journal of the American Geriatrics Society*.2020;68(2):440-441.
137. Pouwels KB, Kalkman GA, Schagen D, Visser ST, Hak E. Is combined use of SSRIs and NSAIDs associated with an increased risk of starting peptic ulcer treatment? *Br J Clin Pharmacol*. 2014;78(1):192-193.

Figure 1: PRISMA flow diagram of included studies



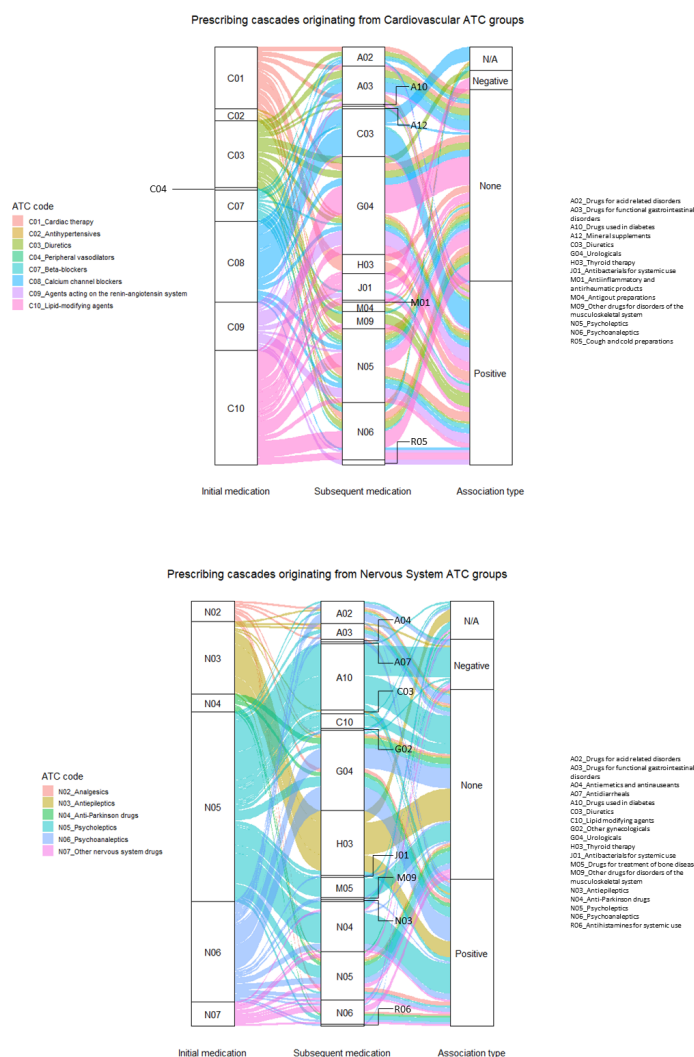


Figure 2: Prescribing cascades examined in non-exploratory studies (n=94) stratified by ATC classification. These alluvial plots represent initial (column 1) and subsequent (column 2) medication pairs examined and the primary quantitative association identified (column 3). The height of the strata in columns 1 and 2 are proportional to the number of instances that the relevant medication has been examined across included studies. The height of the strata in column 3 is proportional to the number of identified quantitative associations that belong to each association type. The width of the linkage between column 1 and column 2 is proportional to the number of instances that the unique medication pair has been examined across included studies. The width of the linkage between column 2 and column 3 is proportional to the number of tested medication pairs that result in a prescribing cascade (positive association), do not result in a prescribing cascade (none), indicate a lower likelihood of a prescribing cascade (negative association), or where no association could be examined due to study reporting (N/A: non-applicable); Figure 2a: ATC1 level; Figure 2b: Cardiovascular medications (ATC3 level); Figure 2c: Nervous system medications (ATC3 level)

[CHART][CHART][CHART]

Figure 3: Quality appraisal summary of included studies (n=98): Figure 3a: cohort studies; Figure 2b: case-control studies; Figure 3c: cross-sectional studies

Table 1: Primary results of included studies by ATC pharmacological classification (n=101)

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Alimentary tract and metabolism Adimadhyam (2019) ⁴⁷	Alimentary tract and metabolism Sodium/Glucose cotransporter-2 inhibitors (SGLT2-I)	Alimentary tract and metabolism Genital mycotic infections	Alimentary tract and metabolism Antifungal	Alimentary tract and metabolism <i>PSSA SGLT2-I- Antifungal</i> ± 365 days aSR 1.24 (95%CI 1.20–1.28)
Avorn (1995b) ^{a, 20}	Metoclopramide	Extrapyramidal symptoms (EPS)	Anti-Parkinson drug (APD)	<i>Metoclopramide- APD</i> (< 90 days) aOR 3.04 (95%CI 2.22-4.17)
Gadzhanova (2017) ⁹⁰	SGLT2-I Dipeptidyl peptidase 4 inhibitor (DPP4-I)	Urinary or genital infections	Trimethoprim Nitrofurantoin Norfloxacin	<i>Risk of UTI</i> (< 6 months) SGLT2-I users (3.6%) compared to DPP4-I users (4.9%), aHR 0.90 (95%CI 0.66-1.24) <i>Risk of genital infections</i> (< 6 months) SGLT2-I users (2.9%) compared with DPP4-I users (0.9%), aHR 3.50 (95%CI 1.95-5.89)
Janetzki (2021) ¹⁰¹	PPI	Development or exacerbation of chronic obstructive pulmonary disease (COPD)	Long-acting muscarinic antagonist (LAMA) or long-acting beta-2 agonist (LABA) listed for the treatment of COPD	<i>PSSA: PPI- LAMA/LABA</i> ± 1 year Omeprazole: aSR=1.29 (95%CI 1.22-1.36) Esomeprazole: aSR=1.25 (95%CI 1.22-1.29) Rabeprazole: aSR=1.15 (95%CI 1.08-1.21) Pantoprazole: aSR=1.08 (95%CI 1.05-1.12) Lansoprazole: aSR=1.08 (95%CI 0.96-1.22)

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Lund (2021) ¹¹³	SGLT2-I Glucagon-like peptide-1 receptor agonists (GLP1-RA)	Gout	Any uric acid lowering therapy, colchicine or first hospital diagnosis of gout (composite)	<i>Risk of gout <3 years: intention to treat analysis</i> HR: 0.58 (0.44 to 0.75) [GLP1-RA as referent] <i>Risk of gout <3 years: per-protocol analysis</i> HR: 0.48 (0.33 to 0.70) [GLP1-RA as referent] <i>PSSA: SGLT2-I- Gout ±365 days</i> aSR 0.63 (95%CI 0.47-0.84) <i>PSSA: GLP1-RA- Outcome ±-365 days</i> aSR 0.94 (95%CI 0.78-1.13) <i>PSSA: PPI- Anti-dementia medication ±3 years</i> aSR 1.38 (95%CI 1.28-1.48); n=3025 <i>PSSA: H2RA- Anti-dementia medication ±-3 years</i> aSR 2.35 (2.13-2.59); n=2308 <i>PSSA: Rosiglitazone- Furosemide ±1 year</i> Pooled (Australia and Canada): aSR 1.65 (95%CI 1.58-1.72) Pooled (Asia): aSR 1.21 (95%CI 1.01-1.45) <i>PSSA: Pioglitazone- Furosemide ±-1 year</i> Pooled (Australia and Canada): aSR 1.47 (95%CI 1.41-1.91) Pooled (Asia): aSR 1.11 (95%CI 0.86-1.32)
Park (2018b) ³²	PPI Histamine 2 receptor antagonist (H2RA)	Dementia	Anti-dementia medication (secondary outcome)	
Roughead (2015) ⁹⁸	Pioglitazone Rosiglitazone	Oedema	Furosemide	

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Roughead (2016) ⁹⁷	PPI	Clostridium difficile infection	Oral vancomycin	<i>PSSA: PPI- Oral vancomycin ± 1 year Pooled estimate: aSR 2.40 (95%CI 1.88-3.05) Pooled estimate (Asia only): aSR 3.16 (95%CI 1.95-5.10)</i>
Wahab (2014) ¹¹⁵	Rosiglitazone	Heart failure	Furosemide	<i>PSSA: Rosiglitazone- Furosemide (Jul 2000 to Dec 2007) aSR=1.73 (99%CI 1.34-2.24)</i>
Blood and blood forming organs Hachiken (2013) ¹¹¹	Blood and blood forming organs Low dose aspirin (LDA)	Blood and blood forming organs Gastrointestinal (GI) complications	Blood and blood forming organs H2RAs PPIs	<i>Blood and blood forming organs PSSA: LDA- PPIs ± 365 days Enteric coated LDA: aSR 1.87 (95% CI 1.26-2.83)</i>

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Maura (2018) ⁹⁵	Direct oral anticoagulants (DOACs; excluding edoxaban)	GI events (composite) Nausea Constipation Depression Glaucoma	Gastrointestinal medications (composite) Gastrointestinal medications without acid disorder drugs Antiemetics Drugs for constipation	<i>PSSA: DOAC- Gastrointestinal medications (composite) ± 360 days aSR 0.95 (95%CI 0.92-0.97); n=24,916 Apixaban- Gastrointestinal medications ± 360 days aSR 1.18 (95%CI 1.10-1.26); n=3440 PSSA: DOAC- Gastrointestinal medications (without acid disorder drugs ± 360 days) aSR 1.26 (95%CI 1.24-1.29); n=37,764 PSSA: DOAC- Antiemetic ± 360 days aSR 1.25 (95%CI 1.22-1.28); n=27,080 PSSA: DOAC- Drugs for constipation ± 360 days aSR 1.25 (95%CI 1.22-1.27); n=43,112 DOAC- Antidepressant medication ± 360 days aSR 1.26 (95%CI 1.23-1.30); n=20,613 DOAC- Glaucoma medication ± 360 days aSR 1.01 (95%CI 0.97-1.05); n=9,473</i>

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Takada (2014) ⁶⁷	Low dose aspirin (LDA) Enteric coated Buffered	GI complications	H2RAs PPIs	<i>PSSA: LDA- PPIs ±12 months</i> Enteric-coated LDA: aSR 1.20 (95%CI 0.97-1.49) Buffered LDA: aSR 0.59 (95%CI 0.33-1.05) <i>PSSA: LDA- H2RAs +12 months</i> Enteric-coated LDA: aSR 0.83 (95%CI 0.67-1.02) Buffered LDA: aSR 0.78 (95%CI 0.350-1.21)
Yokoyama (2020b) ⁸⁶	Oral anticoagulants	Osteoporosis	Bisphosphonate	<i>PSSA: Warfarin- Bisphosphonate ±12 months</i> aSR 1.43 (95%CI 1.02-2.03); n=148
Cardiovascular system Bowman (1995) ⁷⁴	Cardiovascular system Angiotensin converting enzyme inhibitor (ACEI)	Cardiovascular system Cough	Cardiovascular system Antitussive	Cardiovascular system <i>ACEI- Antitussive (<1 year; adjusted)</i> aOR 1.53 (95%CI 1.17-2.01)
Fujimoto (2014) ⁵⁰	Statins	Lower urinary tract symptoms (LUTS)	Drugs for storage LUTS	<i>PSSA: Statins- Drugs for storage LUTS ±365 days</i> All statins: aSR 1.17 (95% CI 1.05-1.30) Pravastatin: aSR 1.27 (95%CI 1.05-1.54) Statins- Solifenacin: aSR 1.47 (95% CI 1.25-1.73) Statins- Oxybutynin: aSR 1.71 (95% CI 1.09-2.72)

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Gurwitz (1997) ²³	Antihypertensive medication (see Supplementary Material)	Gout	Anti-gout medication (see Supplementary Material)	<i>Antihypertensive- Anti-gout medication < 365 days</i> Non-thiazide antihypertensive alone: aRR 1.00 (95%CI 0.65-1.53) Thiazide diuretic alone: aRR 1.99 (95%CI 1.21-3.26) Thiazide diuretic plus non-thiazide antihypertensive: aRR 2.29 (95%CI 1.55-3.37)
Hallas (1996) ⁵²	Beta blockers Cardiovascular drugs (see Supplementary Material)	Depression	Antidepressants	<i>Beta-blocker- Antidepressant (study period)</i> aRR 1.09 (95% CI 0.95, 1.26) <i>ACEIs- Antidepressant</i> aRR 1.29 (95% CI 1.08, 1.56) <i>Calcium channel blockers- Antidepressant</i> aRR 1.31 (95% CI 1.14, 1.51)
Lindberg & Hallas (1998) ¹⁰⁰	Cholesterol-lowering medication	Depression	Antidepressants	<i>PSSA: Cholesterol-lowering drug- Antidepressant (study period)</i> All drugs: aSR 0.90 (95%CI 0.68-1.22); n=184 Simvastatin: aSR 1.59 (1.08-2.45); n=91 Among 5,458,467 DH CCB users (weighted), 185,130 individuals (3.4% weighted) were identified with new loop diuretic use.
Morris (2021) ^{c, 118}	Dihydropyridine calcium channel blockers (DH-CCBs)	Oedema	Loop diuretic	

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Pouwels (2013) ¹³³	ACEI	Urinary tract infection (UTI)	Nitrofurantoin	<i>PSSA: ACEI- Nitrofurantoin ± 4 weeks</i> aSR 1.68 (95% CI 1.21-2.36); n=161
Pouwels (2014) ^{b, 120}	ACEI	UTI	Nitrofurantoin	<i>ACEI- Nitrofurantoin (<30 days vs <60-90 days)</i> Crude OR=1.84 (95%CI 1.51-2.25)
Pouwels (2016) ⁶⁹	Statin	Infection	Antibiotic	<i>PSSA: Statin- Antibiotic ± 13 months</i> Any antibiotic: aSR 0.86 (95%CI 0.81-0.91)
Pratt (2015) ⁶¹	Amiodarone	Hypothyroidism	Thyroxine	<i>PSSA: Amiodarone- Thyroxine ± 12 months</i> Pooled aSR 2.63 (95%CI 1.47-4.72)

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Savage (2020) ⁵	Calcium channel blockers (CCBs) ACEIs or Angiotensin receptor blockers (ARBs) (comparator)	Oedema	Loop diuretic	<i>CCB- Loop diuretic</i> <i><90 days</i> Incident CCB users had a higher cumulative incidence of loop diuretic than the comparators (1.4% vs. 0.7% [other antihypertensive comparator] and 0.5% [general comparator], p<0.001). <i>CCB</i> <i>versus other</i> <i>antihypertensive</i> <i>(ACEI or ARB)</i> 1-30 days: aHR 1.68 (95%CI 1.38-2.05) 31-60 days: aHR 2.26 (95%CI 1.76-2.92) 61-90 days: aHR 2.40 (95%CI 1.84-3.13) 91-180 days: aHR 2.24 (95%CI 1.86-2.71) 181-365 days: aHR 1.64 (95%CI 1.38-1.94) <i>PSSA: Statin-</i> <i>NSAID ±365 days</i> aSR 0.94 (95%CI 0.85-1.05)
Silwer (2006) ⁹⁴	Statin	Muscle pain	NSAID	

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Singh (2021b) ⁶⁴	CCBs	Lower extremity oedema	Diuretics	<i>CCB- Diuretic day 8-day 365 Cohort 1:</i> 161 incident diuretic users among 3,304 incident CCB users (4.9%, 95%CI 4.2-5.7). <i>Cohort 2:</i> 1,586 incident diuretic users among 36,462 prevalent CCB users (1.3%, 95%CI 4.1-4.6). <i>Cohort 3:</i> 130 incident diuretic use among 2,525 participants with polypharmacy at the day of incident CCB dispensing (5.1, 95%CI 4.3-6.0). <i>PSS: Statin- Hypnotic drugs ±365 days aSR 1.18 (95%CI 1.11-1.25) Beta-blocker- Antidepressant <34 days (concurrent use) Beta-blocker: RR 2.6 (95%CI 2.3-3.0) PSSA: ACEI- Cough medication ±6 months 2000-2012: SR 2.0 (95%CI 1.8-2.2)</i>
Takada (2014) ¹³⁴	Statins	Sleep disturbance	Hypnotic drugs	
Thiessen (1990) ^{e, 114}	Beta-blocker	Depression	Antidepressants	
Vegter (2013) ¹⁸	ACEI	Cough	Cough medication	

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Vouri (2018) ^{c, 6}	DH-CCBs	Lower extremity oedema	Loop diuretic	<i>DH-CCB- Loop diuretic (2014)</i> The potential prescribing cascade was identified in 2.2 million visits (4.6%) using the primary definition of prescribing cascade.
Vouri (2019) ⁷	DH-CCBs	Lower extremity oedema	Loop diuretic	<i>PSSA: DH-CCB- Loop diuretic (2014)</i> ± 360 days aSR 1.87 (95%CI 1.84-1.90)
Vouri (2021a) ¹⁰⁷	DH-CCBs	DH-CCB induced oedema	Loop diuretic	<i>PSSA: DH-CCB- Loop diuretic ± 360 days</i> aSR 2.27 (95% CI 1.44-3.58)
Vouri (2021b) ¹⁰⁶	DH-CCB	DH-CCB induced oedema	Loop diuretic	<i>PSSA: DH-CCB- Loop diuretic ± 360 days</i> Relative to levothyroxine initiators: aSR 1.72 (95%CI 1.66-1.78) Relative to ACEI/ARBs initiators: aSR 1.45 (1.41-1.49)
Vouri (2022) ³⁶	Beta-blocker	Oedema	Loop diuretic	<i>PSSA: Beta-blocker- Loop diuretic ± 90 days</i> aSR 1.78 (99%CI 1.72-1.84)
Yokoyama (2021) ^{d, 87}	Amiodarone	Hypothyroidism	Thyroid preparations	<i>PSSA: Amiodarone- Thyroid preparations ± 12 months</i> aSR 12.8 (95%CI 8.44-20.28)
Dermatologicals Azoulay (2007) ^{b, 45}	Dermatologicals Isotretinoin	Dermatologicals Depression	Dermatologicals Antidepressants	Dermatologicals <i>Isotretinoin- Antidepressant (5 month risk and control windows)</i> aRR 2.68 (95%CI 1.10-6.48)

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Hersom (2003) ⁷³	Isotretinoin Minocycline	Depression	Antidepressants (MAOIs excluded)	<i>Isotretinoin- Antidepressant (study period)</i> aRR 0.97 (95%CI 0.92–1.02) <i>Minocycline- Antidepressant (study period)</i> aRR 0.98 (95%CI 0.95–1.02)
Sturkenboom (1995) ⁶⁵	Acitretin	Vulvo-vaginal infection	Vulvo-vaginal anti-infective drug	<i>Acitretin- Vulvo-vaginal anti-infective (study period)</i> Pooled Mantel-Haenszel IRR: 3.3 (95%CI 1.1–9.6)
Genito urinary system and sex hormones Dyson (2020) ³³	Genito urinary system and sex hormones 5- α reductase inhibitors (5-ARI)	Genito urinary system and sex hormones Depression	Genito urinary system and sex hormones Antidepressant	Genito urinary system and sex hormones <i>PSSA: 5-ARI- Antidepressant ± 365 days</i> Crude SR 0.84 (95% CI 0.80–0.89)
Hagberg (2017) ^{d, 112}	5-ARI Alpha blocker (AB)	Depression	Antidepressant (<90 days of depression diagnosis)	<i>5ARI- Antidepressant (compared with AB only users)</i> 5-ARIs only: aIRR=0.94 (95%CI 0.85–1.04) 5-ARIs + ABs: aIRR= 1.04 (94%CI 0.89–1.21) <i>Nested case-control analysis (compared with AB only users)</i> 5-ARIs only: aOR 0.88 (95%CI 0.78–1.01) 5-ARIs+ABs: aOR 0.90 (95%CI 0.73–1.10).
Anti-infectives for systemic use	Anti-infectives for systemic use	Anti-infectives for systemic use	Anti-infectives for systemic use	Anti-infectives for systemic use

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Corrao (2005) ^{d, 49}	Antibacterial drugs for systemic use	Arrhythmia triggered by prolonged QT interval	Antiarrhythmic	<p><i>PSSA:</i></p> <p><i>Antibacterial-</i> <i>Antiarrhythmic</i> <i>(study period)</i></p> <p>Erythromycin aSR 1.78 (95%CI 1.09, 2.89); n=73</p> <p>Ciprofloxacin aSR 1.17 (95%CI 1.02, 1.33); n=870</p> <p><i>Cohort</i> <i>analysis</i> <i>(standardised</i> <i>incidence ratios)</i></p> <p>Erythromycin: 1.96 (95%CI 1.45-2.59); n=8,956</p> <p>Clarithromycin: 1.18 (95%CI 1.08-1.29); n=97,900</p> <p>Rokitamycin: 1.27 (95%CI 1.00-1.66); n=15,247</p> <p>Ciprofloxacin: 1.25 (95%CI 1.14-1.37); n=58,070</p> <p>Norfloxacin: 1.17 (95%CI 1.00-1.36); n=22,421</p> <p>Levofloxacin: 1.33 (95%CI 1.03-1.38); n=14,159</p> <p><i>Case-control</i> <i>analysis</i></p> <p>Erythromycin: OR 1.89 (95%CI 1.33-2.68)</p> <p>Clarithromycin: OR 1.18 (95%CI 1.04-1.34)</p> <p>Ciprofloxacin: OR 1.21 (95%CI 1.05-1.39)</p> <p>Levofloxacin: OR 1.33 (95%CI 1.04-1.70)</p>

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Antineoplastic and immunomod- ulating agents Farkas (2021) ²¹	Antineoplastic and immunomod- ulating agents Aromatase inhibitors (AI)	Antineoplastic and immunomod- ulating agents For the treatment of menopausal symptoms Vasomotor symptoms, vaginal dryness, arthralgias, pain	Antineoplastic and immunomod- ulating agents See Supplementary Material	Antineoplastic and immunomod- ulating agents <i>Medication use in 12 months before AI:</i> Any new side effect medication: 7,436 (40.2%) Opiates 31.5% ; SSRIs 16.1%; Gabapentin 7.0% <i>Medication use in the 24 months after AI:</i> Any new side effect medication: 13,179 (71.2%) Opiates 55.1%; SSRIs 22.6%; Benzodiazepines 18.4%; Tramadol 17.7%; Gabapentin 14.6%
Musculo-skeletal system Gurwitz (1994) ^{a, 22}	Musculo-skeletal system NSAID	Musculo-skeletal system Hypertension	Musculo-skeletal system Antihypertensive	Musculo-skeletal system <i>NSAID- Antihypertensive (<365 days)</i> OR=2.01 (95%CI 1.89-2.14)
Nervous system Avorn (1995a) ^{a, 19}	Nervous system Neuroleptics	Nervous system Extrapyramidal symptoms	Nervous system APD (excluding amantadine monotherapy)	Nervous system <i>Any Anti-Parkinson drug (<90 days)</i> Any neuroleptic: aOR 5.4 (95%CI 4.8-6.1) <i>Anticholinergic anti-Parkinson drug (<90 days)</i> Any neuroleptic: aOR 8.5 (95%CI 4.8-6.1) <i>Dopaminergic agent (<90 days)</i> Any neuroleptic: aOR 2.2 (95%CI 1.9-2.7)

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Brandt-Christensen (2007) ³⁷	APD Control 1: Antidiabetics Control 2: unexposed	Depression	Antidepressants	<i>Anti-Parkinson drug- Antidepressant (versus unexposed)</i> APD cohort: RR 2.10 (95%CI 2.04-2.16) Antidiabetic cohort: RR 1.34 (95%CI 1.32-1.36) <i>PSSA: SSRI - RLS drug ±365 days</i> Any drug: aSR 0.99 (95%CI 0.95-1.02) Dopamine agonist only: aSR 1.21 (95%CI 1.12-1.32); n=2,267 <i>Likelihood for incident hypothyroidism (study period)</i> Lithium: OR 1.41 (95%CI 1.14-1.74) Carbamazepine: OR 1.37 (95%CI 1.13-1.65) Valproate: OR 1.72 (95%CI 1.40-2.11) <i>AChEI- Anticholinergic</i> Patients dispensed cholinesterase inhibitors were more likely to receive an anticholinergic medication in follow-up (4.5% vs 3.1%; p<0.001).
Dalgard Dunvald (2020) ³⁸	Selective serotonin reuptake inhibitors (SSRI)	Restless leg syndrome (RLS)	Dopamine agonist Quinine	
Gau (2010) ^{a, 135}	Lithium Carbamazepine Valproate	Hypothyroidism	Thyroxine, liothyronine or thyroid hormone and hypothyroidism diagnosis (composite)	
Gill (2005) ²⁶	Acetylcholinesterase inhibitors (AChEI)	Urge urinary incontinence	Urinary anticholinergics	

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Hirano (2020) ¹⁰²	Anxiolytic Hypnotic Antidepressants Antipsychotics	EPS	Diagnosis of EPS and APD prescription in same month (composite)	<i>PSSA:</i> <i>Psychotropic</i> <i>medication- EPS</i> <i>and APD ±12</i> <i>months</i> Anxiolytic: aSR 2.48 (95%CI 2.16-2.85); n=992 Hypnotic: aSR 2.28 (95%CI 1.97-2.64); n=872 Antidepressant: aSR 2.26 (95%CI 1.93-2.66); n=728 Antipsychotic: aSR 9.24 (95%CI 7.35-11.8); n=817
Kalisch Ellett (2018) ^{c, 116}	Antipsychotics	EPS Hyperprolacti- naemia Diabetes mellitus	Anticholinergic Hy- perprolactinaemia medications Oral diabetes medications	<i>Concomitant</i> <i>medication use</i> Anticholinergic: n=51 (0.7%) Hyper- prolactinaemia medications: n=8 (0.1%) Oral diabetes medicines: n=874 (11.8%)
Kroger (2015) ^{b, 119}	AChEI	Urinary incontinence	Drugs for urinary frequency and incontinence	<i>AChEI- Drugs for</i> <i>urinary frequency</i> <i><90 days</i> All patients (n=2,700): aHR 1.13 (95%CI 0.97-1.32) Rivastigmine patients (n=1,853): aHR 1.13 (95%CI 0.95-1.34) Galantamine patients (n=1,043): aHR 1.10 (95%CI 0.81-1.50)

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Lai (2013) ⁸⁰	Antiepileptic drugs (AEDs)	Hypothyroidism	Levothyroxine	<i>PSSA: AEDs- Levothyroxine ±12 months Any AED: aSR 1.13 (99%CI 1.09-1.18) Carbamazepine: aSR 1.21 (99%CI 1.08-1.34) Phenobarbital: aSR 1.25 (99%CI 1.15-1.36) Phenytoin: aSR 1.75 (99%CI 1.58-1.94) Valproate: aSR 1.34 (99%CI 1.20-1.49) Oxcarbazepine: aSR 1.22 (99%CI 1.03-1.46)</i>
Lampela (2016) ⁴⁴	AChEI or Memantine	Urinary incontinence	Urinary anticholinergics	<i>AChEI- Urinary anticholinergics (versus memantine users) <6 months: aHR 1.47 (95%CI 1.17-1.86) <12 months: aHR 1.41 (95%CI 1.17-1.69)</i>
Marras (2016) ²⁷	Lithium Valproic acid Antidepressant	Drug induced tremor diagnosed as Parkinson's Disease (PD)	Anti-Parkinson drug or PD diagnosis (<i>see Supplementary Material</i>)	<i>Start of dopaminergic drug (no previous antipsychotic use) Lithium (versus antidepressant): aHR (95%CI 1.06-3.30) Start of anti-Parkinson drug or PD diagnosis (no previous antipsychotic use) Lithium (versus antidepressant): aHR 1.68 (95%CI 1.13-2.48)</i>

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Masurkar (2021) ²⁴	AChEI	Overactive bladder	Urinary anticholinergic	<i>AChEI- Anticholinergic cascade <6 months</i> Rivastigmine: aHR=1.0 Donepezil: aHR=1.55 (95%CI 1.31-1.83) Galantamine: aHR=1.17 (95%CI 0.87-1.58)
Movig (2002) ⁴¹	SSRI	Urinary incontinence	Spasmolytic agent or 30 or more units of incontinence wear	<i>SSRI- Spasmolytic agent/incontinence wear <3 month</i> During SSRI (versus before SSRI): IDR 1.57 (95%CI 1.38-1.79) During SSRI (versus after SSRI): IDR 2.03 (95%CI 1.76-2.34) During SSRI (versus before and after SSRI): IDR 1.75 (95%CI 1.56-1.97) <i>Risk for incontinence during exposed period (versus non-exposed)</i> aRR 1.61, 95%CI 1.42-1.82
Narayan (2019) ²⁵	AChEI or Memantine	Several ADRs examined relating to anticholinergic medication use	Anticholinergics (see Supplementary Material)	<i>Anti-dementia drug- Marker medication ±180 days</i> Exposed to at least one anticholinergic ±180 days: n=1439 Exposed to at least one anticholinergic after anti-dementia drug: n=416

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Onder (2014) ^{c, 46}	Anti-Parkinson drugs and antipsychotics (concomitant use)	Parkinsonism (side effect of antipsychotics); Behavioural disorders (side effect of anti-Parkinson drugs)	Anti-Parkinson drugs and antipsychotics (concomitant use)	<i>Prevalence of concomitant use of anti-Parkinson and antipsychotic medication (2011)</i> Total population: n=25,949 (0.2%) 65-74 years: n=10,200 (0.2%) 75-84 years: n=10,625 (0.2%) [?]85 years: n=5,124 (0.3%) <i>PSSA:</i> <i>Benzodiazepines- Anti-dementia drugs ± 3 years aSR 2.19 (95%CI 1.92-2.49); n=1,285</i> <i>Flunarizine- Antidepressant <30 days</i> Number of antidepressant starts during or within 30 days after flunarizine use was 5 out of a total of 34 histories <i>Flunarizine- Antidepressant (study period)</i> Incidence Rate=1.342 (95%CI 1.00-1.80) <i>Flunarizine- Anti-Parkinson drug</i> In a subset of 777 flunarizine recipients there were 10 participants who received anti-Parkinson drugs
Park (2018a) ⁹⁶	Benzodiazepines	Dementia	Anti-dementia drugs	
Petri (1988) ⁵⁵	Flunarizine	Depression	Antidepressant	
Petri (1990) ⁵⁷	Flunarizine	Depression or Parkinsonism	Antidepressant or Anti-Parkinson drug	

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Pratt (2013) ⁶⁰	Antipsychotics	Acute hyperglycaemia	Insulin	<i>PSSA: Olanzapine- Insulin ±12 months</i> USA Public: aSR 1.14 (95%CI 1.1-1.17) Sweden: aSR 1.53 (95%CI 1.13-2.06) <i>Risperidone- Insulin +-12 months</i> USA Public: aSR 1.09 (95%CI 1.07-1.12) <i>Gabapentinoid- Diuretic <90 days (versus non-users)</i> aHR 1.44 (95%CI 1.23-1.70).
Read (2021) ²⁸	Gabapentinoid	Oedema	Diuretic	<i>Antipsychotic- Anti-Parkinson drug/diagnosis <1 year (versus atypical antipsychotic)</i> Typical antipsychotics: adjusted HR 1.30 (95%CI 1.04-1.58) No therapy: aHR 0.40 (95%CI 0.29-0.43)
Rochon (2005) ²⁹	Antipsychotic	Parkinsonism	Anti-Parkinson drug or Parkinson diagnosis (composite)	<i>PSSA: Benzodiazepine- Anti-dementia drug ±12 months</i> 12 months: aSR 1.23 (95%CI 1.11-1.37)
Takada (2016) ⁶⁶	Benzodiazepine	Dementia	Anti-dementia drug	<i>PSSA: Atypical antipsychotics- Anti- hyperlipidemic drugs</i> Olanzapine ±360 days: aSR 2.19 (95%CI 1.55-3.12)
Takeuchi (2015) ⁴³	Atypical antipsychotics	Hyperlipidemia	Anti- hyperlipidemic drugs	

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Thacker (2006) ¹⁰⁹	AChEI	Drug-induced airways complications	Antibacterial and oral corticosteroid	<i>AChEI- Antibacterial and oral corticosteroid <1 month</i> Fully-adjusted RR=1.19 (95%CI 0.52-2.74)
Venäläinen (2017) ⁷⁰	AChEI	Nausea Dyspepsia Diarrhoea Urinary incontinence Seizures Anxiety Insomnia Depression	Antiemetics PPIs/H2RAs Loperamide/ Oral rehydration sachets Oxybutynin Anxiolytics Anticonvulsants Hypnotics and sedatives Antidepressants	<i>AChEI- Marker drug ±1 year</i> Loperamide/Oral rehydration: aSR 1.42 (95%CI 1.14-1.77); n=348 Anxiolytics: aSR 1.16 (95%CI 1.01-1.34); n=807 Hypnotics and sedatives: aSR 1.19 (95%CI 1.05-1.36); n=963 Antiemetics: aSR 1.18 (95%CI 1.05-1.32); n=1,202 Anticonvulsants: (aSR 1.26 (95%CI 1.03-1.55); n=389 PPI/H2RAs: aSR 0.87 (95%CI 0.77-0.98), n=1,079 Antidepressant: aSR 0.77 (95%CI 0.70-0.85), n=1,698 Oxybutynin: aSR 1.04 (95%CI 0.81-1.34), n=261

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Vouri (2020) ^{c, 136}	AChEI or Memantine	Rhinorrhea	Rhinorrhea medications (see Supplementary Material)	<i>AChEI/Memantine- Rhinorrhea medications (concomitant use)</i> AChEI users were more likely to use a rhinorrhea medication compared to non-AChEI users, OR 7.16 (95%CI 2.25-22.73); adjusted OR=4.7 (95%CI 1.53-14.43)
Wang (2021a) ⁸²	Varenicline	Neuropsychiatric adverse events: Depression Anxiety Sleep disorders	Antidepressant Anxiolytics Hypnotics and sedatives (composite outcome)	<i>PSSA: Varenicline- Any NPAE drug ±365 days</i> aSR 1.00 (95%CI 0.89-1.13) <i>PSSA: Varenicline- Hypnotics and sedatives ±-365 days</i> Sleep disorder drug: aSR=1.25 (95% CI 1.05-1.48)

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Wang (2021b) ⁴²	Varenicline (Nicotine replacement therapy [NRT] as comparator)	Neuropsychiatric adverse events: Depression Anxiety Insomnia	Antidepressants Antidepressants in combination with psycholeptics Anxiolytics Hypnotics and sedatives (composite outcome)	<i>General population with psychiatric disorders <24 weeks</i> Any NPAE medication: adjusted OR 0.82 (95% CI 0.68 to 0.99) <i>General population without psychiatric disorders <24 weeks</i> Any NPAE medication: adjusted OR 0.85, 95% CI 0.72 to 1.00) <i>COPD population with psychiatric disorders <24 weeks</i> Any NPAE medication: adjusted OR 0.97 (95% CI 0.66 to 1.44) <i>COPD population without psychiatric disorders <24 weeks</i> Any NPAE medication: adjusted OR 0.81 (95% CI 0.54 to 1.20)
Yokoyama (2020a) ⁸⁸	Antipsychotics	Osteoporosis	Bisphosphonate	<i>PSSA: Antipsychotic- Bisphosphonate</i> No association identified.
Respiratory system Fox (2022) ³⁴	Respiratory system Montelukast	Respiratory system Neuropsychiatric adverse events (NPAE)	Respiratory system Antidepressants Benzodiazepines Hypnotics Antipsychotics Mood stabilisers Buspirone (composite outcome)	Respiratory system <i>PSSA: Montelukast- Any NPAE medication ±14-365 days</i> SR 0.84 (95%CI 0.80-0.89)

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Henriksen (2017) ³⁹	Inhaled corticosteroids	Oral candidiasis	Systemic or topical antifungal	<i>PSSA: Inhaled corticosteroid- Topical antifungal ±12 months</i> Crude SR 2.89 (95%CI 2.80-2.97) <i>PSSA: Inhaled corticosteroid- Systemic antifungal +/-12 months</i> Crude SR 1.50 (95%CI 1.46-1.54)
Petri (1991) ⁵⁶	Inhaled corticosteroids	Oral candidiasis	Topical antifungal	<i>Inhaled corticosteroids- Topical antifungal <90 days</i> Crude OR=1.66 (n=21)
Van Boven (2013) ⁷¹	Inhaled corticosteroids	Oral candidiasis	Topical antifungal	<i>PSSA: Inhaled corticosteroids- Topical antifungal ±12 months</i> Crude SR 1.94 (95%CI 1.71-2.21)
Winkel (2018) ⁴⁰	Montelukast	Depression	Antidepressant (excluding bupropion)	<i>PSSA: Montelukast- Antidepressant ±1 year</i> Crude SR 1.19 (95%CI 1.11-1.28)
Sensory organs	Sensory organs	Sensory organs	Sensory organs	Sensory organs

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Roughead (2012) ⁹⁹	Timolol Latanoprost Bimatoprost Pilocarpine Brimonidine	Exacerbation of airways disease Exacerbation of depression	Inhaled beta-agonists Inhaled corticosteroids Oral corticosteroids SSRI	<i>PSSA: Glaucoma- marker medications ±1 year</i> Timolol- Inhaled beta agonist: aSR 1.48 (95%CI 1.22-1.78); n=786 Timolol- Inhaled corticosteroid: aSR 1.43 (95%CI 1.13-1.81); n=494 Latanoprost- Inhaled beta agonist: aSR 1.24 (95%CI 1.11-1.38); n=2,251 Latanoprost- Oral corticosteroid: aSR 1.14 (95%CI 1.00-1.29); n=1,671 Timolol- Antidepressant: aSR 1.24 (95%CI 1.07-1.43); n=1,253 Timolol- SSRI: aSR 1.30 (95%CI 1.08-1.56); n=791 Latanoprost- Antidepressant: aSR 1.16 (95%CI 1.03-1.31); n=1,871 Latanoprost- SSRI: aSR 1.20 (95%CI 1.03-1.39); n=1,155 Multiple medication groups examined
Multiple medication groups examined	Multiple medication groups examined	Multiple medication groups examined	Multiple medication groups examined	Multiple medication groups examined

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Brandt-Christensen (2006) ⁸⁴	Antidepressant Lithium Antidiabetic	Parkinsonism	APD (see Supplementary Material for exclusions)	<i>Index drug- Anti-Parkinson drug (versus unexposed)</i> Antidepressant: RR 1.79 (95%CI 1.72-1.86) Lithium: RR 1.88 (95%CI 1.60-2.20) Antidiabetic: RR 0.80 (95%CI 0.74-0.86)

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Bytzer & Hallas (2000) ⁸³	Predefined list of 32 index medications (see Supplementary Material)	Dyspepsia or nausea	Cisapride or Metoclopramide	<i>PSSA: Index medication- Cisapride <100 days NSAIDs:</i> aSR=1.33 (95%CI 1.02-1.77); n=211 Methylxanthines: aSR=2.36 (1.00-8.44); n=18 <i>PSSA: Index medication- Metoclopramide <100 days Insulin</i> aSR 2.91 (95%CI 1.40-8.11); n=28 Opioids: aSR 2.84 (95%CI 2.48-3.28); n=1017 Potassium supplement: 1.42 (95%CI 1.15-1.79); n=324 Digoxin: 2.87 (95%CI 2.01-4.35); n=138 Nitrates: 1.74 (95%CI 1.16-2.77); n=88 Loop diuretics: 1.50 (95%CI 1.23-1.85); n=383 ACEIs: 2.27 (95%CI 1.46-3.85); n=77 Oral corticosteroids: 1.33 (95%CI 1.11-1.60); n=458 Antibiotics: 1.40 (95%CI 1.24-1.60); n=974 Penicillins: 1.38 (95%CI 1.21-1.59); n=868 Macrolides: 1.58 (95%CI 1.31-1.94); n=414 NSAIDs: 1.48 (95%CI 1.28-1.74); n=676 Asthma drugs: 1.42 (95%CI 1.14-1.79); n=307 Methylxanthines: 2.03 (95%CI 1.25-3.65); n=63

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Caughey (2010) ⁴⁸	Medicines commonly associated with dizziness identified (see Supplementary Material)	Dizziness	Prochlorperazine	<i>PSSA: Index medication- Prochlorperazine ±12 months</i> Cardiac therapy: aSR=1.14 (95%CI 1.06-1.22); n=3,017 Nitrates: aSR=1.11 (95%CI 1.03-1.21); n=2,224 Isosorbide mononitrate: aSR=1.21 (95%CI 1.07-1.38); n=918 Diuretic: aSR=1.07 (95%CI 1.01-1.14); n=3,845 Beta-blocker: aSR=1.13 (95%CI 1.05-1.21); n=3,156 CCBs: aSR=1.22 (95%CI 1.16-1.36); n=2,696 ACE inhibitors: aSR=1.22 (95%CI 1.14-1.31); n=3,162 AR2B: aSR=1.20 (95%CI 1.11-1.30); n=2,577 Statins: aSR=1.50 (95%CI 1.40-1.61); n=3,411 NSAIDs: aSR=1.37 (95%CI 1.27-1.47); n=3,079 Opioids: aSR=1.24 (95%CI 1.17-1.31); n=5,266 Sedatives: aSR=1.18 (95%CI 1.11-1.26); n=3,470

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
de Jong (2003) ¹¹⁰	Antidepressant with or without NSAID	GI adverse effects	H2RAs PPIs Prostaglandins	<i>Antidepressant- Ulcer drugs (compared with TCA only)</i> SSRI: IRR 1.2 (95%CI 0.5-2.8); n=1181 SSRI + NSAID: IRR 12.4 (95%CI 3.2-48.0); n=86
Garrison (2012) ⁵¹	Statin Diuretic Inhaled long-acting beta-agonists (LABA)	Nocturnal leg cramps	Quinine	<i>PSSA: Index drug- Quinine \pm 1 year</i> All statins: aSR 1.16 (95%CI 1.04-1.29); n=1,326 All LABAs: aSR 2.42 (95%CI 2.02-2.89); n=576 LABA alone: aSR 2.17 (95%CI 1.56-3.02); n=137 LABA- corticosteroid: aSR 2.55 (95%CI 2.06-3.12); n=439 All diuretics: aSR 1.47 (95%CI 1.33-1.63); n=1,590 Loop diuretic: aSR 1.20 (95%CI 1.00-1.44); n=447 Thiazide diuretic: aSR 1.48 (95%CI 1.29-1.68); n=977 Potassium-sparing diuretic: aSR 2.12 (95%CI 1.61-2.78); n=206

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Hallas & Bytzer (1998) ⁹¹	Predefined list of 33 medications (see Supplementary Material)	Dyspepsia	Ulcer drug prescription	<i>PSSA: Index drug- Ulcer drug prescription ± 100 days</i> NSAIDs: aSR 1.80 (95%CI 1.64-1.99) CCBs: aSR 1.40 (95%CI 1.18-1.67) Oral corticosteroids: aSR 1.15 (95%CI 1.02-1.30) ACEIs: aSR 1.38 (1.12-1.73) Methylxanthines: aSR 1.49 (1.05-2.19)

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Hashimoto (2015) ⁵³	Medicines that cause storage symptoms; Medicines that cause voiding symptoms	LUTS	Medications for (LUTS)	<i>PSSA: Index drug- Medications for LUTs ±12 months</i> Oxycodone: aSR 1.20 (95%CI 1.03–1.41) Morphine: aSR: 1.29 (95%CI 1.14– 1.45) Donepezil: aSR: 1.98 (95%CI 1.57–2.50) Intestinal lavage solution: aSR 1.86 (95%CI 1.65–2.10) Cyclophosphamide: aSR 1.52 (95%CI 1.14–2.04) Levodo- pa/benserazide: aSR 1.82 (95%CI 1.18–2.81) Amantadine: aSR 1.53 (95%CI 1.12–2.09) Paroxetine: aSR 1.77 (95%CI 1.33–2.36) Milnacipran: aSR 2.10 (95%CI 1.28–3.45) Diazepam: aSR 1.73 (95%CI 1.46–2.06) Risperidone: aSR 1.55 (95%CI 1.34–1.79) Levomepromazine: aSR 2.20 (95%CI 1.34–1.79) Sulpiride: aSR 1.32 (95%CI 1.01–1.72) Cimetidine: aSR 1.99 (95%CI 1.24–3.20) Scopolamine butylbromide: aSR 1.72 (95%CI 1.55–1.92) Tiotropium bromide: aSR 1.75 (95%CI 1.42–2.16) Cibenzoline: sSR 2.97 (95%CI 1.92–4.59) Amezinium metilsufate: aSR 1.89 (95%CI

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Huh (2019) ³¹	Metoclopramide or levosulpiride	Drug induced Parkinsonism	Levodopa	<i>PSSA:</i> <i>Metoclopramide -</i> <i>Levodopa <90 days</i> aOR 2.94 (95%CI 2.35, 3.67) <i>PSSA:</i> <i>Levosulpiride -</i> <i>Levodopa <90 days</i> aOR 3.30 (95%CI 2.52, 4.32)
Kalisch Ellett (2014) ⁷⁵	See Supplementary Material	Urinary incontinence	Oxybutynin	<i>PSSA: Index</i> <i>medication-</i> <i>Oxybutynin ±12</i> <i>months</i> Prazosin (women only): aSR 1.84 (95%CI 1.29-2.63); n=135 Low-ceiling diuretics, excluding thiazides: aSR 1.22 (95%CI 1.06-1.41); n=750 CCBs: aSR 1.45 (95%CI 1.33-1.57); n=2,230 ACEIs: aSR 1.28 (95%CI 1.19-1.39); n=2,616 ACEIs + diuretic: aSR 1.35 (1.15-1.58); n=620 ARBs: aSR 1.42 (1.30-1.55); n=2,040 ARB+ diuretic: aSR 1.32 (1.16-1.49); n=999 HRT: aSR 1.54 (95%CI 1.42-1.67); n=2,446 Antipsychotics: aSR 0.83 (95%CI 0.78-0.89); n=2,121 Hypnotic sedatives: aSR 1.10 (95%CI 1.03-1.18); n=3,326

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Kim (2019) ^{a, 124}	Propulsives Antipsychotics Antivertigo agent (see Supplementary Material)	Drug induced Parkinsonism	APD or Parkinson diagnosis (composite) (see Supplementary Material)	<i>Index medication- Anti-Parkinson drug/ diagnosis <1 year</i> Levosulpiride: OR 4.3 (95%CI 3.5-5.3); n=595 Mosapride: OR 2.1 (95%CI 1.7-2.6); n=430 Domperidone: OR 2.1 (95%CI 1.6-2.8); n=247 Metoclopramide: OR 2.7 (95%CI 1.8-4.1); n=121 Itopride: OR 1.6 (95%CI 1.2-2.2); n=232 Clebopride: OR 12.8 (95%CI 2.8-57.0); n=19 Combined propulsive use: OR 3.9 (95%CI-2.8-5.5); n=219 Typical antipsychotic: OR 6.4 (95%CI 1.4-28.2); n=17 Atypical antipsychotic: OR 2.4 (95%CI 1.2-4.9); n=56 Risperidone: OR 13.5 (95%CI 1.8-102.1); n=23 Flunarizine: OR 5.0 (95%CI 2.7-9.0); n=86

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Ko (2019) ⁷⁷	Statins	Skin and soft tissue infection New onset diabetes mellitus	Dicloxacillin/ Flucloxacillin Antidiabetic	<i>PSSA: Statin- Antibiotic ±365 days</i> aSR 1.40 (95%CI 1.34-1.47); n=7,726 <i>PSSA: Statin- Antidiabetic ±-365 days</i> aSR 1.09 (95%CI 1.04-1.15); n=6,794 <i>PSSA: Antidiabetic- Antibiotic ±-365 days</i> aSR 1.24 (95%CI 1.15-1.33); n=2,828
Nishtala & Chyou (2017) ⁵⁴	Amiodarone Lithium Frusemide Fluticasone Simvastatin	Hypothyroidism Hyperthyroidism Hypokalaemia Oral candidiasis Muscle cramps	Thyroxine Carbimazole Potassium Nystatin Quinine sulphate	<i>PSSA: Amiodarone- Thyroxine ±360 days</i> aSR 3.57 (95% CI 3.17-4.02) <i>Lithium- Thyroxine ±-360 days</i> aSR 3.43 (95% CI 2.55-4.70) <i>Amiodarone- Carbimazole ±-360 days</i> aSR 8.81 (95% CI 5.86-13.77) <i>Simvastatin- Quinine sulphate ±-360 days</i> aSR 1.69 (95% CI 1.61-1.77) <i>Fluticasone- Nystatin ±-360 days</i> aSR 2.34 (95% CI 2.19-2.50) <i>Frusemide- Potassium ±-360 days</i> aSR 2.94 (95% CI 2.83-3.05)

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Pouwels (2013) ¹³⁷	SSRI with or without NSAID	Peptic ulcer	Peptic ulcer drug treatment	<i>PSSA: SSRI +/- NSAID- Peptic ulcer treatment ±4 weeks</i> SSRI: aSR 0.83 (95%CI 0.65-1.06) NSAID: aSR 2.50 (95%CI 2.27-2.76) SSRI + NSAID: aSR 1.48 (95%CI 0.90-2.49)
Rasmussen (2015) ⁶²	Antithrombotic drugs Cardiovascular drugs (see Supplementary Material)	Erectile dysfunction	5-phosphodiesterase inhibitor	<i>PSSA: Cardiovascular drugs- 5-phosphodiesterase inhibitor ±6 months</i> Thiazides: aSR 1.28 (95%CI 1.20, 1.38); NNTH 370 (95%CI 300, 500); n=3,118 β-blockers: aSR 1.18 (95%CI 1.09, 1.28); NNTH 680 (95%CI 480, 1200); n=2,511 CCBs: aSR 1.29 (95%CI 1.21, 1.38); NNTH 330 (95%CI 270, 440); n=3,379 ACEIs: aSR 1.29 (95%CI 1.21, 1.37); NNTH 350 (95%CI 290, 440); n=4,182 ARBs: aSR 1.16 (95%CI 1.06, 1.26); NNTH 540 (95%CI 360, 1200); n=2,082

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Singh (2021a) ⁶³	Antipsychotic or Metoclopramide	Parkinsonism	Anti-Parkinson drug	<i>Antipsychotic/metoclopramide- Anti-Parkinson drug < day 8-365 Cohort 1: 36 (0.8%) incident anti-Parkinson drug users among 4534 incident antipsy- chotic/metoclopramide users Cohort 2: 20 (0.5%) incident users of anti-Parkinsonian drugs among 3485 antipsy- chotic/metoclopramide users</i>
Trenaman (2021) ⁸¹	AChEIs Metoclopramide CCBs	Urinary incontinence Parkinsonism Pedal oedema	Urinary medications Anti-Parkinson drug Diuretic	<i>AChEI- Urinary medications < 6 months 60 cases of prescribing cascade were identified. Extending to 365 days resulted in 52 additional cases. Metoclopramide- Anti-Parkinson drug < 6 months 11 cases of the prescribing cascade were identified. Extending to 365 days resulted in 5 additional cases. CCB- Diuretic < 6 months 289 cases of prescribing cascade were identified. Extending to 365 days resulted in 369 cases.</i>
Exploratory studies	Exploratory studies	Exploratory studies	Exploratory studies	Exploratory studies

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Tsiropoulos (2009) ⁶⁸	AEDs	Exploratory analysis	Any other medication presented in the same period	<p><i>PSSA: All AEDs-Marker medication</i> Propulsives ±183 days: aSR 1.31 (95%CI 1.11-1.56); n=571 Laxatives ±183 days: aSR 1.57 (95%CI 1.29-1.92); n=432 Topical corticosteroids ±183 days: aSR 1.32 (95%CI 1.16-1.52); n=900 <i>PSSA:</i> <i>Carbamazepine-Marker medication</i> Propulsives +-183 days: aSR 1.57 (95%CI 1.14-2.19); n=163 Laxatives +-183 days: aSR 1.61 (95%CI 1.01-2.59); n=82 Topical corticosteroids +-183 days: aSR 1.48 (95%CI 1.17-1.87); n=305 Anti-acne preparations +-183 days: aSR 3.66 (95%CI 1.31-2.62); n=23 Bone disease treatment +-548 days: aSR 1.98 (95%CI 1.03-3.92); n=43 <i>PSSA:</i> <i>Oxcarbazepine-Marker medication</i> Propulsives +-183 days: aSR 2.54 (95%CI 1.71-3.85); n=119 Laxatives +-183 days: aSR 3.74 (95%CI 2.31-6.29); n=103 Topical corticosteroids +-183 days: aSR 1.40 (95%CI 1.08-1.83); n=245 <i>Phenobarbital-Marker medication</i> Bone disease treatment +-548</p>

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
King (2020) ⁷⁶	654 different medications examined	New onset heart failure	Furosemide	<p><i>PSSA: Index drug-Furosemide ± 12 months</i></p> <p>Fosaprepitant: aSR 2.60 (95%CI 2.42-2.81); n=3,394</p> <p>Granisetron: aSR 2.24 (95%CI 2.42-2.81); n=2,299</p> <p>Tropisetron: aSR 1.43 (95%CI 1.08-1.79); n=340</p> <p>Degarelix: aSR 1.66 (95%CI 1.29-2.06); n=293</p> <p>Brinzolamide: aSR 1.18 (95%CI 1.06-1.32); n=1,304</p> <p>Travoprost: aSR 1.18 (95%CI 1.01-1.35); n=788</p> <p>Latanoprost: aSR 1.11 (95%CI 1.04-1.19); n=3,619</p> <p>Brimonidine: aSR 1.10 (95%CI (1.00-1.20); n=2,012</p> <p>Pizotifen: aSR 1.27 (95%CI 1.11-1.44); n=978</p> <p>Rizatriptan: aSR 1.16 (95%CI 1.03-1.31); n=1,036</p> <p>Sumatriptan: aSR 1.16 (95%CI 1.03-1.29); n=1,250</p> <p>Benzhexol: aSR 1.65 (95%CI 1.12-2.24); n=142</p> <p>Mesalazine: aSR 1.33 (95%CI 1.13-1.54); n=646</p> <p>Levetiracetam: aSR 1.13 (95%CI 1.03-1.23); n=2,005</p> <p>Fluorometholone: aSR 1.11 (95%CI 1.07-1.15); n=9,410</p> <p>Ranitidine: aSR 1.08 (95%CI 1.04-1.12); n=10,875</p> <p>Denosumab: aSR 1.07 (95%CI 1.03-1.10); n=16,714</p>

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Wahab (2016) ¹⁰⁸	691 different medications examined	Heart failure	Furosemide	<i>PSSA: Index medication- Furosemide ± 1 year</i> Teriparatide: aSR 5.02 (95% CI 1.07–23.7); n=10 Lodoxamide: aSR 2.50 (95% C; 1.06–5.91); n=27 Famotidine: aSR 1.69 (95% CI 1.38–2.08); n=423 Latanoprost: aSR 1.48 (95% CI 1.38–1.59); n=3,107 Pilocarpine: aSR 1.43 (95% CI 1.16–1.77); n=632 Brinzolamide: aSR 1.37 (95% CI 1.16–1.62); n=564 Betahistine: aSR 1.31 (95% CI 1.07–1.62); n=359 Ranitidine: aSR 1.24 (95% CI 1.17–1.31); n=5,554 Paracetamol: aSR 1.06 (95% CI 1.04–1.09;; n=24,210

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Chen (2021) ⁸⁹	<i>Confirmatory analysis</i> Amiodarone <i>Exploratory analysis</i> ACEIs Statins Buffered LDA Enteric-coated LDA DH-CCBs	Hypothyroidism Gout Cough UTI Storage LUTS Depression Sleep disturbances Hepatotoxicity Muscle pain Skin and soft tissue infection Infection in those with type-2 diabetes GI complications Oedema	<i>Confirmatory analysis</i> Thyroxine Allopurinol <i>Exploratory analysis</i> (see Supplementary Material)	<i>Confirmatory PSSA</i> ± 1 year Amiodarone- Thyroxine: aSR 3.77 (95%CI 3.43-4.14); n=2,667 Amiodarone- Allopurinol: aSR 0.83 (95%CI 0.76-0.90); n=2,071 <i>Exploratory PSSA</i> ± 1 year ACEIs- Antitussive: aSR 1.33 (95% CI 1.31-1.34); n=141,924 Statins- Drugs for urinary frequency: aSR 1.17 (95% CI 1.16-1.19); n=107,422 Statins- Antidepressants: aSR 1.19 (95% CI 1.18-1.21); n=117,443 Statins- Hypnotics: aSR 1.10 (95% CI 1.09-1.12); n=124,061 Statins- Ursodeoxycholic acid: aSR 1.26 (95% CI 1.21-1.31); n=11,231 Statins- NSAIDs: aSR 1.02 (95% CI 1.02-1.03); n=430,774 Statins- Di- cloxacillin/Flucloxacillin: aSR 1.18 (95% CI 1.15-1.22); n=23,068 Statins- Antibiotic treatment (those with type 2 diabetes): aSR 1.38 (95% CI 1.36-1.39); n=150,016 DH-CCBs- Loop diuretic: aSR 1.46 (95% CI 1.45-1.48); n=139,375

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Lai (2014) ¹²²	Sulpiride Non-sulpiride antipsychotics	EPS Diabetes Hy- perprolactinaemia Cardiac arrhythmias	<i>Confirmatory analyses:</i> Anticholinergics Oral hyperglycaemics Prolactine inhibitors Class 1B antiarrhythmics [2] <i>Exploratory analyses:</i> all medications prescribed after the index date	<i>Confirmatory PSSA analyses ±12 months</i> Sulpiride- Anticholinergics: aSR 1.73 (95%CI 1.46-2.06); n=568 Haloperidol- Anticholinergics: aSR 1.99 (95%CI 1.68-2.35); n=611 Risperidone- Anticholinergics: aSR 1.21 (95%CI 1.04-1.41); n=702 Olanzapine- Anticholinergics: aSR 0.73 (95%CI 0.58-0.93); n=281 Amisulpiride- Anticholinergics: aSR 0.54 (95%CI 0.40-0.73); n=188 Sulpiride- Prolactine inhibitors: aSR 12.0 (95%CI 1.59-91.2); n=16 Amisulpiride- Prolactine inhibitors: aSR 8.05 (95%CI 1.00-65.4); n=8 Haloperidol- Class 1b antiarrhythmics: sSR 2.81 (95%CI 1.03-7.66); n=21 <i>Exploratory PSSA analyses: Sulpiride- Marker medication ±-12 months</i> Stomatological preparations: aSR 1.86 (95%CI 1.13-3.07); n=71 Corticosteroids for local oral treatment: aSR 1.71 (95%CI 1.00-2.91); n=59 Beta blockers, any: aSR 1.42 (95%CI 1.12-1.71); n=371 Beta blockers, non-selective: aSR 1.61 (95%CI 1.28-2.03); n=304 Dermatological

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Hallas (2018) ¹⁰³	186,758 associations tested in the main analysis; 30 best signals reported	Exploratory analysis	30 strongest signals reported	<i>PSSA: Index- Marker medication ±12 months</i> Opioids- Drugs for constipation (crude SR 2.34, 95%CI 2.31–2.38); n=84,020 High ceiling diuretics- Potassium SR 3.31 (95%CI 3.24-3.38); n=48539 Thiazide- Potassium SR 3.46 (95%CI 3.39-3.54); n=45,175 Opioids- Propulsives SR 2.14 (95%CI 2.10-2.17); n=62,139 NSAIDS- Anti-ulcer drugs SR 1.71 (95%CI 1.67-1.74); n=49646 Antithrombotic- Anti-ulcer drugs SR 1.41 (95%CI 1.39-1.44); n=54,841 Cough suppressants- Drugs for constipation SR 1.95 (95%CI 1.90-2.00); n=26,0015 Corticosteroids, systemic use- Drugs affecting bone structure and mineralisation SR 3.40 (95%CI 3.27-3.54); n=13,023

Primary author (Year)	Initial medication(s)	Suspected ADR	New medication(s)	Quantitative association (primary analysis or association at 1 year)
Hellfritzsche (2018) ¹⁰⁴	Non-vitamin K oral anticoagulants (NOAC)	Exploratory analysis	20 strongest signals reported	<i>PSSA: NOAC- Marker drug ±6 months</i> Benzodiazepines, hypnotic: cSR 8.28 (95%CI 6.01-12.05); NNTH 193 Osmotic laxatives: cSR 1.35 (95%CI 1.25-1.46); NNTH 133 Benzodiazepines, sedative: cSR 1.99 (95%CI 1.74-2.30); NNTH 174 Corticosteroids, anal use: cSR 2.03 (95%CI 1.76-2.35); NNTH 176 SSRI: cSR 1.57 (95%CI 1.37-1.77); NNTH 202 Other antidepressant: cSR 1.59 (95%CI 1.41-1.80); NNTH 207 PPI: cSR 1.19 (95%CI 1.11-1.28); NNTH 209 Phenylpiridine opioids: cSR 2.12 (95%CI 1.81-2.51); NNTH 215 Propulsives: cSR 1.51 (95%CI 1.35-1.71); NNTH 216 Iron bivalent, oral: cSR 1.62 (95%CI 1.42-1.86); NNTH 238 Contact laxatives: cSR 1.29 (95%CI 1.17-1.43); NNTH 253

Note: aSR: adjusted sequence ratio; cSR: crude sequence ratio; aHR: adjusted hazard ratio; HR: hazard ratio; aOR: adjusted odds ratio; IDR: incidence density ratio; IRR: incidence rate ratio; aIRR: adjusted incidence rate ratio; PSSA: prescription sequence symmetry analysis; NNTH: number needed to harm; ^a:

case-control study;^b: case-crossover study; ^c: cross-sectional study; ^d: includes case-control study; e: includes cross-sectional study

Table 2: Summary of findings for the most commonly identified prescribing cascades

Initial medication	Suspected ADR	Second medication	Main findings
DH-CCB	Oedema	Loop diuretic	<p><1 year: aSR 1.46 (95% CI 1.45-1.48); n=139,375⁸⁹ <360 days: aSR 1.87 (95%CI 1.84-1.90); 55,818⁷ <360 days: aSR 2.27 (95% CI 1.44–3.58); n=90¹⁰⁷ <360 days: aSR 1.72 (95%CI 1.66-1.78) relative to levothyroxine negative control; aSR 1.45 (1.41-1.49) relative to ACEI/ARB negative control³⁵ Rate of being dispensed a loop diuretic versus general comparator group⁵ 1-30 days: aHR 2.51 (95%CI 2.13-2.96) 31-60 days: aHR 2.99 (95%CI 2.43-3.69) 61-90 days: aHR 3.89 (95%CI 3.11-4.87) 91-180 days: aHR 3.20 (95%CI 2.72-3.76) 181-365 days: aHR 2.22 (95%CI 1.90-2.60)</p>

Initial medication	Suspected ADR	Second medication	Main findings
Amiodarone	Hypothyroidism	Thyroxine	<p><1 year: aSR 3.77 (95% CI 3.43-4.14); n=2,667⁸⁹</p> <p><360 days: aSR 3.57 (95%CI 3.17-4.02)⁵⁴</p> <p><1 year: aSR 2.14 (99%CI 1.92-2.39); n=2,613¹²⁵</p> <p><1 year Australia: aSR 5.30 (95%CI 4.69-5.96); n=1,979⁶¹</p> <p><1 year Hong Kong: aSR 2.33 (95%CI 1.99-2.72); n=754</p> <p><1 year Japan: aSR 1.77 (95%CI 0.61-5.08); n=6</p> <p><1 year Korea: aSR 1.52 (95%CI 1.29-1.80); n=657</p> <p><1 year Taiwan: aSR 3.26 (95%CI 2.26-4.70); n=153</p> <p><1 year: Pooled aSR 2.63 (95%CI 1.47-4.72)</p> <p><6 months: aSR 13.6 (95%CI 7.73-25.96)⁸⁷</p> <p><12 months: aSR 12.8 (95%CI 8.44-20.28)</p> <p><18 months: aSR 11.4 (95%CI 7.98-16.80)</p> <p><24 months: aSR 11.7 (95%CI 8.32-16.94)</p> <p><30 months: aSR 10.8 (95%CI 7.86-15.29)</p> <p><36 months: aSR 10.8 (95%CI 7.89-15.00)</p>
Inhaled corticosteroids	Oral candidiasis	Topical antifungals	<p><90 days OR 1.66; n=21⁵⁶</p> <p><1 year: SR 2.89 (95%CI 2.80-2.97)³⁹</p> <p><1 year: SR SR 1.94 (95%CI 1.71-2.21)⁷¹</p> <p><360 days: aSR 2.34 (95% CI 2.19-2.50)⁵⁴</p>

Initial medication	Suspected ADR	Second medication	Main findings
Neuroleptics/Antipsychotic	Parkinsonian symptoms/ extrapyramidal symptoms	Anti-parkinson medication or Parkinson diagnosis	<p><90 days: aOR 5.4 (95%CI 4.8-6.1)¹⁹ <1 year (1 antipsychotic): aSR 9.24 (7.35-11.8); n=817¹⁰² <1 year (2 antipsychotics): aSR 22.2 (9.94-61.7); n=137 <1 year ([?]3 antipsychotics): aSR 34.8 (5.87-1413.8); n=37 Never use: aOR 1.0 (referent); n=10,714¹²⁴ Very-late use ([?]181 days): aOR 1.1 (95%CI 0.6-1.8); n=61 Late use (31-180 days): aOR 2.0 (95%CI 1.2-3.3); n=94 Early use (8-30 days): aOR 6.0 (95%CI 2.3-15.9); n=43 Current use ([?]7 days): aOR 3.0 (95%CI 1.7-5.4); n=80 Typical: aOR 6.4 (95%CI 1.4-28.2); n=17 Haloperidol: aOR 4.3 (95%CI 0.9-20.1); n=12 Atypical: aOR 2.4 (95%CI 1.2-4.9); n=56 Quetiapine: aOR 0.9 (95%CI 0.4-2.2); n=26 Risperidone: aOR 13.5 (95%CI 1.8-102.1); n=23 Combined use: aOR 3.2 (95%CI 0.6-17.9); n=7 Typical antipsychotics: aHR 1.30 (95%CI 1.04-1.58) versus atypical antipsychotic use²⁹ No therapy: aHR 0.40 (95%CI 0.29-0.43)</p>

Initial medication	Suspected ADR	Second medication	Main findings
Acetylcholinesterase inhibitors	Urinary incontinence	Drugs for urinary frequency and incontinence	During follow-up (1 st June 1999-31 st March 2003): older adults dispensed acetylcholinesterase inhibitors had a higher risk of subsequently receiving an anticholinergic medication to treat urge urinary incontinence (aHR, 1.55,95% CI, 1.39-1.72) ²⁶ Donepezil-Medication for managing Lower Urinary Tract Symptoms (LUTS) ⁵³ <3 months: 1.32 (95%CI 1.00-3.50); n=243 <12 months: aSR: 1.98 (95%CI 1.57-2.50); n=319 <6 months: aHR 1.47 (95%CI 1.17-1.86) versus memantine users ⁴⁴ <12 months: aHR 1.41 (95%CI 1.17-1.69) versus memantine users Donepezil: aHR 1.55 (95%CI 1.31-1.83) versus rivastigmine use ²⁴ Galantamine: aHR 1.17 (95%CI 0.87-1.58) versus rivastigmine use <90 days aOR 3.04 (95%CI 2.22-4.17) ²⁰ <90 days aOR 2.94 (95%CI 2.35-3.67) ³¹ Anti-Parkinson medication or diagnosis <1 year: aOR 2.7 (95%CI 1.8-4.1); n=121 ¹²⁴
Metoclopramide	Parkinsonian symptoms	Levodopa	<1 year OR=1.58 (95%CI 1.21-2.07) ⁷⁴ <6 months: SR 2.0 (95%CI 1.8-2.2); n=1,898; estimated 13.4% mistreated cough ¹⁸ <1 year: aSR 1.33 (95% CI 1.31-1.34); n=141,924 ⁸⁹
ACE inhibitors	Cough	Antitussive	

Initial medication	Suspected ADR	Second medication	Main findings
NSAID	GI symptoms	Anti-ulcer medication	<4 weeks: aSR 2.50 (95%CI 2.27-2.76); n=2,016 ¹³⁷ <100 days: aSR 1.80 (95%CI 1.64-1.99); n=1,814 ⁹¹ <1 year: SR 1.71 (95%CI 1.67-1.74); n=49,646 ¹⁰³
Ranitidine	Heart failure	Furosemide	<1 year: aSR 1.08 (95%CI 1.04-1.12); n=10,875 ⁷⁶ <1 year: aSR 1.24 (95% CI 1.17-1.31); n=5,554 ¹⁰⁸
Rosiglitazone	failure	Furosemide	<1 year Australia-1: aSR 1.70 (95%CI 1.34-2.15) ⁹⁸ <1 year Australia-2: aSR 1.63 (95%CI 1.51-1.76) <1 year Canada: aSR 1.65 (95%CI 1.57-1.73) <1 year Pooled estimate (Australia & Canada): aSR 1.65 (95%CI 1.58-1.72) <1 year Hong Kong: aSR 3.37 (95%CI 1.69-6.72) <1 year Korea: aSR 1.14 (95%CI 1.08-1.21) <1 year Taiwan: aSR 1.12 (95%CI 0.99-1.25) <1 year Pooled estimate (Asia): aSR 1.21 (95%CI 1.01-1.45) July 2000-December 2007: aSR 1.73 (99%CI 1.34-2.24) ¹¹⁵
SGLT2-I	Genital infections	Antifungal	<30 days: aSR 1.35 (95%CI 1.26-1.44) ⁴⁷ <60 days: aSR 1.48 (95%CI 1.40-1.56) <90 days: aSR 1.53 (95% CI 1.43-1.60) <180 days: aSR 1.42 (95%CI 1.37-1.47) <365 days: aSR 1.24 (95%CI 1.20-1.28) Genital infection occurred more frequently among SGLT2-I users than DPP-4 users (2.9% vs, 0.9%, aHR 3.50, 95%CI 1.95-5.89) ⁹⁰

Initial medication	Suspected ADR	Second medication	Main findings
DOAC	Depression	Antidepressant	<3 months: aSR 1.29 (95%CI 1.23-1.35); n=7,253 ⁹⁵ <6 months: aSR 1.28 (95%CI 1.24-1.33); n=12,530 <12 months: aSR 1.26 (95%CI 1.23-1.30); n=20,613 SSRI <6 month: SR 1.57 (1.37-1.77); n=1,137; NNTH 202 ¹⁰⁴ Other antidepressant <6 month: SR 1.59; 1,076; (1.41-1.80); NNTH 207 Furosemide <360 days: aSR 2.94 (95% CI 2.83-3.05) ⁵⁴ High ceiling diuretic <1 year: SR 3.31 (95%CI 3.24-3.38); n=48,539 ¹⁰³
High ceiling diuretics	Hypokalaemia	Potassium	
Statins	Lower urinary tract symptoms (LUTS)	Drugs for urinary frequency and incontinence	<91 days: aSR 1.21 (95% CI 1.00, 1.46); n=446 ⁵⁰ <182 days: aSR 1.19 (95% CI 1.04, 1.38); n=785 <365 days: aSR 1.17 (95% CI 1.05, 1.30); n=1,373 <1 year: aSR 1.17 (95% CI 1.16-1.19); n=107,422 ⁸⁹
Statins	Skin soft tissue infection	Antibiotic (Dicloxacillin or Flucloxacillin)	<1 year: aSR 1.18 (95% CI 1.15-1.22); n=23,068 ⁸⁹ <91 days: aSR 1.40 (95%CI 1.29-1.52); n=2,498 ⁷⁷ <182 days: aSR 1.41 (95%CI 1.33-1.50); n=4,277 <365 days: aSR 1.40 (95%CI 1.34-1.47); n=7,726
Statins	Depression	Antidepressant	<1 year: aSR 1.19 (95% CI 1.18-1.21); n=117,443 ⁸⁹ Simvastatin-Antidepressant (April 1991-December 1995): aSR 1.59 (1.08-2.45); n=91 ¹⁰⁰
Statins	Muscle cramps	Quinine	<360 days: aSR 1.69 (95% CI 1.61-1.77) ⁷⁰ <1 year: aSR=1.16 (95%CI 1.04-1.29); n=1,326 ⁵¹

Initial medication	Suspected ADR	Second medication	Main findings
Brinzolamide	Heart failure	Furosemide	<1 year Brinzolamide: aSR 1.18 (95%CI 1.06-1.32); n=1,304 ⁷⁶ <1 year Brinzolamide: aSR 1.37 (95% CI 1.16–1.62); n=564 ¹⁰⁸
Latanoprost	Heart failure	Furosemide	<1 year Latanoprost: aSR 1.11 (95%CI 1.04-1.19); n=3,619 ⁷⁶ <1 year Latanoprost: aSR 1.48 (95% CI 1.38–1.59); n=3,107 ¹⁰⁸
Carbamazepine	Hypothyroidism	Levothyroxine	1998-2004: aOR 1.37 (95%CI 1.13-1.65) ¹³⁵ <1 year: aSR 1.21 (99%CI 1.08-1.34) ⁸⁰
Valproate	Hypothyroidism	Levothyroxine	1998-2004: aOR 1.72 (95%CI 1.40-2.11) ¹³⁵ <1 year: aSR 1.34 (99%CI 1.20-1.49) ⁸⁰
Lithium	Drug induced tremor Parkinson	Anti-parkinson drug	Jan 1995-December 1999: RR 1.88 (95%CI 1.60-2.20) ⁸⁴ Up to 2 year follow-up (referent valproic acid): aHR 1.50 (95%CI 0.68-3.36) ²⁷ Up to 2 year follow-up (referent antidepressant): aHR 1.56 (95%CI 0.98-2.48)
Lithium	Hypothyroidism	Thyroxine	1998-2004: aOR 1.41 (95%CI 1.14-1.74) ¹³⁵ <360 days: aSR 3.43 (95% CI 2.55-4.70) ⁵⁴

Initial medication	Suspected ADR	Second medication	Main findings
Benzodiazepine	Dementia	Anti-dementia drug	<3 months: aSR 1.24 (95%CI 1.05-1.45); n=625 ⁶⁶ <6 months: aSR 1.20 (95%CI 1.06-1.37); n=973 <12 months: aSR 1.23 (95%CI 1.11-1.37); n=1,450 <24 months: aSR 1.34 (95%CI 1.23-1.47); n=2,049 <36 months: aSR 1.41 (95%CI 1.29-1.53); n=2,408 <48 months: aSR 1.44 (95%CI 1.33-1.56); n=2,653 <3 years: aSR 2.19 (95%CI 1.92-2.49); n=1,285 ⁹⁶ <2 years: aSR 2.00 (95%CI 1.71-2.34); n=780 <1 year: aSR 1.77 (95%CI 1.39-2.27); n=286
SSRI	Urinary incontinence	Drugs for urinary frequency and incontinence (or incontinence products ⁴¹)	Paroxetine <1 year: aSR 1.77 (95%CI 1.33–2.36) ⁵³ During SSRI (before SSRI as referent): IDR 1.57 (95%CI 1.38-1.79) ⁴¹ During SSRI (after SSRI as referent): IDR 2.03 (95%CI 1.76-2.34) During SSRI (before and after SSRI as referent): IDR 1.75 (95%CI 1.56-1.97) Patients had a 61% higher risk for incontinence (aRR 1.61, 95%CI 1.42-1.82)

Note: aSR: adjusted sequence ratio; SR: crude sequence ratio; aHR: adjusted hazard ratio; aOR: adjusted odds ratio; IDR: incidence density ratio; NNTH: number needed to har