Risk of delirium associated with antimuscarinics in older adults: a case-time-control study

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Key Points:

1. Antimuscarinics are the backbone of the pharmacological management of overactive bladder. It is increasingly recognised that antimuscarinics are associated with differential effects on cognitive functioning.

2. The safety profiles of antimuscarinics vary because of their dissimilarities to muscarinic receptor-subtype affinities. Individual drug characteristics, including an affinity for the muscarinic receptor subtype M1 and M2 in the brain, ability to cross the blood-brain barrier, drug metabolism and concurrent use of drugs with anticholinergic properties, can increase the risk of delirium in older adults.

3. In this case-time-control study of new users of oxybutynin and solifenacin in older adults (≥ 65 years), the use of oxybutynin is associated with an increased risk of new-onset delirium. The dose-response risk of delirium was significant for oxybutynin but not for solifenacin.

4. Oxybutynin but not solifenacin is associated with a risk of new-onset delirium in older adults. Therefore, prescribers should exercise caution when using oxybutynin in the oldest old, particularly those with pre-existing cognitive impairment.

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ABSTRACT

Background: Older adults are at an increased risk of delirium because of age, polypharmacy, multiple comorbidities and acute illness. Antimuscarinics are the backbone of the pharmacological management of overactive bladder. However, the safety profiles of antimuscarinics vary because of their dissimilarities to muscarinic receptor-subtype affinities and are associated with differential central anticholinergic adverse effects.

Objective: This study aimed to examine delirium risk in new users of oxybutynin and solifenacin in older adults (≥ 65 years). In the secondary analyses, we examined the risk of delirium by type and dose of antimuscarinic.

Method: We applied a case-time-control design to investigate delirium risk in older adults who started taking oxybutynin and solifenacin. We used a nationwide inpatient hospital data (2005-2016), National Minimum Data Set, maintained by the Ministry of Health, New Zealand (NZ), to identify older adults with a new-onset diagnosis of delirium. Eligible patients were older adults aged 65 at entry into the cohort on 1/1/2006. We used dispensing claims data to determine antimuscarinic treatment exposure. The antimuscarinic included in the study were new users of oxybutynin and solifenacin. These two antimuscarinics are subsidised by the Pharmaceutical Management Agency and are the most frequently used antimuscarinic in NZ. A conditional logistic regression model was used to compute matched odds ratios (MORs) and 95% confidence intervals (CIs). In the case-time-control design, we made separate analyses to evaluate the dose-response risk of delirium.

Results: We identified 4818 individuals (mean age 82.14) from 2005 to 2015 with incident delirium and were exposed to at least one of the antimuscarinic of interest. The case-time-control matched odds ratio (MOR) for delirium with oxybutynin was (2.06, 95% confidence interval (CI) 1.07 to 3.96). Solifenacin was not associated with delirium (0.89 95%CI 0.64 to 1.23). In the sensitivity analyses, the case-time-control MOR for delirium using a shorter risk period (0-3 days) did not change the results. The dose-response risk of delirium was significant for oxybutynin ( 0.05, 95%CI 0.02 to 0.08) but not for solifenacin (-0.01, 95%CI -0.03 to 0.00). In addition, in the subgroup analyses, a statistically significant association of delirium
was found for oxybutynin but not for solifenacin in the non-dementia cohort (2.11, 95% CI 1.08-4.13) and the dementia cohort (1.25, 95% CI 0.05-26.9).

**Conclusion:** The study found that oxybutynin but not solifenacin is associated with a risk of new-onset delirium in older adults. The higher blockade of M1 and M2 receptors by oxybutynin is likely to contribute to delirium than solifenacin, which is highly selective for the M3 receptor subtype. Therefore, the treatment choice with an M3 selective agent must be given due consideration, particularly in those with pre-existing cognitive impairment.

**Introduction**

Antimuscarinics are the backbone of the overactive bladder's pharmacological management (OAB) [1-3]. Oxybutynin, darifenacin, propiverine, tolterodine, fesoterodine, solifenacin and trospium are widely used to manage OAB in older adults [1, 4]. Currently, oxybutynin and solifenacin oral formulations are the most frequently used antimuscarinics for the treatment of OAB in New Zealand. The selection of the most appropriate antimuscarinic for treating OAB in older adults depends on their adverse effects profile, as broadly, all of them have similar efficacy [5-7]. Antimuscarinics are associated with peripheral and central anticholinergic adverse effects [8, 9]. One of the most debilitating central anticholinergic adverse effects of antimuscarinics in older adults is drug-induced delirium [10].

Older adults are predisposed to an increased risk of delirium due to polypharmacy, anticholinergic burden, age-related deficits in drug clearance, compromised cholinergic neurotransmission, and impaired blood-brain barrier (BBB) function [11, 12]. It is now increasingly recognised that antimuscarinics are associated with differential central anticholinergic adverse effects [13, 14]. Antimuscarinics that block cholinergic receptors in the brain can contribute to impaired attention, delayed memory, delirium and drowsiness [8]. Individual drug characteristics include an affinity for the muscarinic receptor subtype M1 and M2 in the brain, the ability to cross the blood-brain barrier (BBB), drug metabolism, and concurrent use of anticholinergic drugs increase the risk of delirium in older adults [15]. Data from systematic reviews and network meta-analyses (mostly from healthy study participants) report a wide array of antimuscarinics’ adverse effects in older adults [7, 16]. Still, there is insufficient data from trials on the rate and magnitude of individual CNS adverse effects, including delirium. Instead, they report CNS adverse events as a composite measure, mostly
from the healthy population, to draw any meaningful inferences to a frail older population. In a real-world setting, older adults have higher comorbidity and are frailer and hence have a higher baseline risk of harm from drug exposures than patients recruited in a clinical trial. Hence, the extrapolation of evidence from healthy participants in clinical trials to real-world frail older patients is barely accurate.

To understand and quantify the risk of new-onset delirium posed by antimuscarinics, we need reliable population-level evidence with appropriate control for confounding. A case-control design has been used previously to examine the association of congenital heart defects with antidepressant use in pregnant mothers [17]. Still, to our knowledge, no studies have used a similar design to understand the risk of delirium posed by antimuscarinic drugs in older adults. Therefore, our study chose a case-time-control design to mitigate confounding from unknown time-invariant confounders. In the case-time-control design, a control group can adjust for time trends of antimuscarinic use for OAB. We followed all the recommendations to apply a case-time-control design to our analyses [18-20]. For case-time-control design, the key assumptions are that occurrence of the event must be acute, and the exposure may vary over time [21, 19].

**Method:**

**Ethics**

The Ethical Implications of Research Activity Form (EIRA1-5312) to conduct this study was approved on October 20, 2020, by the University of Bath.

**Data sources**

We used a nationwide inpatient hospital data (2005-2015), National Minimum Data Set (NMDS), maintained by the Ministry of Health, New Zealand, to examine new-onset diagnosis of delirium. We extracted all hospitalisations from the NMDS from 01/01/2005 to 31/12/2015, in which the primary reason was delirium. The NMDS contains clinical (length of hospital stay, diagnosis, procedures) and demographic (age, sex, ethnicity, date of birth, date of event) information for each hospital admission.

**Study population**
Eligible patients were older adults aged 65 at entry into the cohort on 1/1/2006. The cohort attrition table shows the inclusion/exclusion criteria and the final population analysed in the case-control design (Supplementary Table 1). We defined the cohort entry as the date of the first prescription for an antimuscarinic (oxybutynin or solifenacin). We defined incident use as a new prescription for oxybutynin or solifenacin with no previous prescription claims during the 12 months before cohort entry. We censored at new-onset diagnosis of delirium, end of the study period (31/12/2015), discontinuation of oxybutynin and solifenacin (90 days after the end of treatment, crossover to another antimuscarinic (either oxybutynin or solifenacin).

Exposures and effect modifiers

We used dispensing claims data to determine antimuscarinic treatment exposure. We obtained de-identified dispensing claims data for individuals aged 65 years or older for 2005–2015 from the New Zealand (NZ) Ministry of Health (MoH). The PharmS database is a national dispensing claims database maintained by the MoH, which captures subsidised prescriptions dispensed by community pharmacies in NZ. The antimuscarinics included in this study were new users of oxybutynin and solifenacin. These two antimuscarinics are subsidised by the Pharmaceutical Management Agency (PHARMAC) and are the most frequently used antimuscarinic in NZ. In addition, the effect-modifying drugs were sourced from the literature [22-24]. These included antibiotics, antipsychotics (first-generation antipsychotics (FGAs) and second-generation antipsychotics (SGAs), antidepressants (tricyclics and selective serotonin receptor inhibitors), antiepileptics, weak and strong anticholinergics (Appendix 1). We considered these medication classes as separate covariates and examined them individually in the case-time-control model.

Outcomes

The primary outcome was the International Statistical Classification of Diseases and Related Health Problems, Tenth Revision, Australian Modification (ICD-10-AM) code for a new inpatient diagnosis of delirium. We used the ICD-10-AM codes to identify delirium diagnosis (F050, F051, F058 and F059). We excluded patients with a first-time diagnosis of delirium within 180 days after 01/01/2005.

Statistical analyses

Case-time-control cohort
We created a case-time-control cohort for medication exposures, with 5-day observation periods and two 5-week (35 days) washout periods, summed up to an 80-day study period. Case-period is the 5 days before the index date. The reference period is 45-41 days before the index date (Figure 1). We chose two 5-week washout periods based on our previous study and the need to minimise carryover effects and misclassification of medication exposure. First, we calculated the duration of each prescription by dividing the total dose supplied by the daily dose. Next, we determined whether an individual had non-intermittent exposure to the medications of interest within the case and the reference period with the prescription dates.

We compared antimuscarinic use in the case and the control periods, in individuals with delirium (cases) and without a delirium diagnosis (control). First, we calculated the odds ratio in the cases (OR\textsubscript{cases}) by dividing the number of older adults prescribed antimuscarinics in the case period by the number of older adults prescribed antimuscarinics in the reference period. Similarly, we calculated the odds ratio for controls (OR\textsubscript{controls}) to adjust the time trend of exposure. Finally, we derived the MOR\textsubscript{case-time-control} by dividing the ratio of OR\textsubscript{cases} by OR\textsubscript{controls}. Each individual with a delirium diagnosis was matched to 3 individuals without a delirium diagnosis based on age (at cohort entry +/- 180 days), gender, and ethnicity. The index date is the day the individual was diagnosed with delirium for the first time. We used shorter, 3-day case and control periods in the sensitivity analyses and compared antimuscarinic use in the risk and reference periods.

Subgroup analysis:

We were interested to understand the risk of delirium posed by antimuscarinics in the study population with and without dementia. We hypothesised that dementia status might be a potential effect modifier/interaction to explore. We identified dementia cases using ICD-10-AM codes G308, G309, F002, F009, F019, acetylcholinesterase inhibitors, donepezil, and rivastigmine subsidised by PHARMAC for the treatment of dementia.

Dose-response analysis

The dose-response analysis (Table 3) was done using a case-time-control design. The total dosages of drugs exposed within observation periods were computed as continuous variables instead of binary exposure indicators. For this reason, longer, 7-day case and control periods were used.
To calculate the total dosage exposed within the case and the control periods for each prescription, we counted the number of days the prescription overlapped within the case period, multiplied by the daily dose specified, summed all prescriptions, and repeated the calculation for the control period. Then, we computed the increase in total dosage exposed from the control to the case period.

We used the multivariate binary logistic regression model to adjust for covariates (age, gender, ethnicity, any use of effect modifiers, etc.), the change of log(OR) of incident delirium at the end of the case period (i.e. the index date), in response to one unit increase in total dosage exposed from the control to the case period. In Table 3, we report a change in log (odds ratio) with 95% CI with one unit increase in dosage. For effect modifying drugs, these are changes in log(odds ratio) compared to non-exposure.

The pharmaceutical collections (Pharms) and NMDS data were made available as annual, CSV-formatted datasets. The filtering mentioned above and cohort-construction procedures were performed using a computer program written in R (3.4.2, R Core Team, 2016). All analyses were performed using R software, version 3.2.1.5 [25].

Results

Study participants

We identified 4818 individuals (mean age 82.14) from 2005 to 2015 with incident delirium and exposed to at least one of the antimuscarinic of interest, with or without co-exposures to any of the effect-modifying drugs of interest. Of these, 333 had at least one prescription record of the oxybutynin, and 1137 had at least one prescription record of solifenacin within the 80-day study period. The distribution of ages was slightly skewed towards the higher age group, and there were more males than females. However, most of them were NZ Europeans, and only a few belonged to the Māori ethnic group (Table 1).

Primary analyses

The case-time-control matched odds ratio (MOR) for delirium with oxybutynin was (2.06, 95% confidence interval 1.07 to 3.96). Solifenacin was not associated with delirium (0.89, 95% CI 0.64 to 1.23) (Table 2).

Sensitivity analyses
The case-time-control MOR for delirium using a shorter risk period (0-3 days) did not change the results. The MOR for delirium with oxybutynin was (2.24, 95% confidence interval 1.16 to 4.33). Solifenacin was also not associated with delirium (1.00, 95% CI 0.72 to 1.39) (Table 2).

Secondary analyses

The dose-response risk of delirium was significant for oxybutynin (0.05, 95% CI 0.02 to 0.08) but not for solifenacin (-0.01, 95% CI -0.03 to 0.00) (Table 3).

Subgroup analyses

A statistically significant association was found for oxybutynin in the non-dementia cohort (2.11, 95% CI 1.08-4.13) and the dementia cohort (1.25, 95% CI 0.05-26.9). On the other hand, solifenacin was not associated with delirium in non-dementia (0.89, 95% CI 0.64-1.23) as well as dementia cohort (0.64, 95% CI 0.04-8.62) (Table 4).

Discussion

Case-time-control analyses conducted on a population of older adults showed that oxybutynin increased the risk of new-onset delirium.

The safety profiles of antimuscarinics vary because of their dissimilarities to muscarinic receptor-subtype affinities. The five subtypes of muscarinic receptors (M1-M5) are widely distributed within the human body [26]. M3 receptors, mainly located in human detrusor muscle, are primarily responsible for normal micturition contraction. M1 receptors are widely distributed in the neocortex, hippocampus, and neostriatum. M2 receptors are relatively less common than M1 and play a significant role in memory and cognitive function [27]. Therefore, it is postulated that muscarinic receptor selectivity and permeability to the BBB are pivotal to expressing central anticholinergic adverse effects of antimuscarinics. Tertiary-amine antimuscarinics such as solifenacin have relatively fewer cognitive adverse effects than tertiary-amine antimuscarinics such as oxybutynin because the hydrophilic properties are less likely to cross the BBB [28]. Oxybutynin has a relatively higher affinity for M1 and M2 receptors over M3 subtype muscarinic receptors. Due to the higher blockade of M1 and M2 receptors, oxybutynin is more likely to contribute to delirium than solifenacin, as solifenacin is highly selective for the M3 receptor subtype found in the detrusor muscle of the urinary bladder [29, 30]. Therefore, it is biologically plausible that oxybutynin, compared to solifenacin, is more likely to contribute to delirium. Our findings are plausible with this biological mechanism, and the dose-response risk of delirium being greater with oxybutynin
than solifenacin is plausible too. The impact of antimuscarinics on delirium must be evaluated in frail older adults, and the treatment choice with an M3 selective agent must be given due consideration, particularly in those with pre-existing cognitive impairment [31].

Interestingly, our study found differences in dose-response risk of delirium by type of antimuscarinic. A higher risk of delirium is associated with oxybutynin but not with solifenacin.

In both dementia and non-dementia subcohorts, a positive association was found with oxybutynin, and a negative association was found with solifenacin. Oxybutynin is a tertiary amine with a neutral charge, lipophilic, and low molecular weight, and hence can readily cross the blood-brain barrier and induce delirium [32]. In contrast, solifenacin is also a tertiary amine but has a relatively higher molecular weight, is less lipophilic than oxybutynin, and its muscarinic receptor selectivity is higher for the M3 than M1. It is also suggested that comorbid dementia may compromise the BBB function; however, due to the small sample size in our study, the confidence interval for the association of delirium with oxybutynin is wide in the dementia subcohort. Hence, the results must be interpreted with caution.

**Strengths**

This study has several strengths, including its large size, nationwide coverage of older adults in NZ, and a case-time-control design to control for confounding of time-invariant confounders and adjustment for the time-trend bias antimuscarinics use in OAB. The new user design eliminated the bias likely to be introduced by including prevalent users of antimuscarinics. We also demonstrated a dose-response relationship between antimuscarinic exposure and the risk of new-onset delirium. The majority of validity assumptions of the case-control design were fulfilled in this study, including the indication is stable over time [18, 19, 33]. We applied a 5-week washout period considering the differences in half-lives of individual antimuscarinics.

**Limitations**

In our analyses, we extracted the exposures and the outcomes from the administrative datasets. Hence, exposure misclassification due to lack of information on medication consumption, self-medication, and over-the-counter drugs such as NSAIDs linked to delirium may have biased the findings. In addition, we did not validate the ICD-10-AM codes for delirium to confirm a diagnosis, which could have impacted our findings by misclassifying the cases. Previous studies have reported low sensitivity of 9–28% [34, 35] but high 85–99% specificity [36, 37].
of ICD codes for delirium. Literature has also identified the potential for under-reporting and under-recognising delirium in older hospitalised patients [38, 39]. The retrospective nature of our study design does introduce selection bias. The shortcomings of both the case-crossover and case-control designs are carried into the case-time-control design, including bias created by selecting the case and control windows, a control group, and inadequate adjustment for time-varying confounders.

**Conclusion**

The study found that oxybutynin but not solifenacin is associated with a risk of new-onset delirium in older adults. The higher blockade of M1 and M2 receptors by oxybutynin is likely to contribute to delirium than solifenacin, which is highly selective for the M3 receptor subtype. Therefore, the treatment choice with an M3 selective agent must be given due consideration, particularly in those with pre-existing cognitive impairment.

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**Conflict of interest:** The authors declare no conflict of interest.

**References**

1. Natalin R, Lorenzetti F, Dambros M. Management of OAB in Those Over Age 65. Current Urology Reports. 2013;14(5):379-85. doi:10.1007/s11934-013-0338-5.

2. Kachru N, Sura S, Chatterjee S, Aparasu RR. Antimuscarinic Medication Use in Elderly Patients with Overactive Bladder. Drugs & Aging. 2016;33(10):755-63. doi:http://dx.doi.org/10.1007/s40266-016-0399-5.

3. Rai BP, Cody JD, Alhasso A, Stewart L. Anticholinergic drugs versus non-drug active therapies for non-neurogenic overactive bladder syndrome in adults. Cochrane Database Syst Rev. 2012;12(12):Cd003193. doi:10.1002/14651858.CD003193.pub4.
4. Kachru N, Sura S, Chatterjee S, Aparasu RR. Antimuscarinic Medication Use in Elderly Patients with Overactive Bladder. Drugs Aging. 2016;33(10):755-63. doi:10.1007/s40266-016-0399-5.

5. Herbison P, Hay-Smith J, Ellis G, Moore K. Effectiveness of anticholinergic drugs compared with placebo in the treatment of overactive bladder: systematic review. Bmj. 2003;326(7394):841-4. doi:10.1136/bmj.326.7394.841.

6. Hay-Smith J, Herbison P, Ellis G, Morris A. Which anticholinergic drug for overactive bladder symptoms in adults. Cochrane Database Syst Rev. 2005(3):Cd005429. doi:10.1002/14651858.Cd005429.

7. Chapple CR, Khullar V, Gabriel Z, Muston D, Bitoun CE, Weinstein D. The effects of antimuscarinic treatments in overactive bladder: an update of a systematic review and meta-analysis. Eur Urol. 2008;54(3):543-62. doi:10.1016/j.eururo.2008.06.047.

8. Nishtala PS, Salahudeen MS, Hilmer SN. Anticholinergics: theoretical and clinical overview. Expert Opin Drug Saf. 2016;15(6):753-68. doi:10.1517/14740338.2016.1165664.

9. Vouri SM, Kebodeaux CD, Stranges PM, Teshome BF. Adverse events and treatment discontinuations of antimuscarinics for the treatment of overactive bladder in older adults: A systematic review and meta-analysis. Arch Gerontol Geriatr. 2017;69:77-96. doi:10.1016/j.archger.2016.11.006.

10. Inouye SK. Delirium in older persons. N Engl J Med. 2006;354(11):1157-65. doi:10.1056/NEJMra052321.

11. Inouye SK, Westendorp RG, Saczynski JS. Delirium in elderly people. Lancet. 2014;383(9920):911-22. doi:10.1016/s0140-6736(13)60688-1.

12. Hshieh TT, Inouye SK, Oh ES. Delirium in the Elderly. Psychiatr Clin North Am. 2018;41(1):1-17. doi:10.1016/j.psc.2017.10.001.

13. Yamada S, Ito Y, Nishijima S, Kadekawa K, Sugaya K. Basic and clinical aspects of antimuscarinic agents used to treat overactive bladder. Pharmacol Ther. 2018;189:130-48. doi:10.1016/j.pharmthera.2018.04.010.
14. Wagg AS. Antimuscarinic treatment in overactive bladder: special considerations in elderly patients. Drugs Aging. 2012;29(7):539-48. doi:10.1007/bf03262272.

15. Mangoni AA, Jackson SH. Age-related changes in pharmacokinetics and pharmacodynamics: basic principles and practical applications. Br J Clin Pharmacol. 2004;57(1):6-14. doi:10.1046/j.1365-2125.2003.02007.x.

16. Olshansky B, Ebinger U, Brum J, Egermark M, Viegas A, Rekeda L. Differential pharmacological effects of antimuscarinic drugs on heart rate: a randomised, placebo-controlled, double-blind, crossover study with tolterodine and darifenacin in healthy participants ≥ or = 50 years. J Cardiovasc Pharmacol Ther. 2008;13(4):241-51. doi:10.1177/1074248408325404.

17. Sun Y, Pedersen LH, Wu CS, Petersen I, Sørensen HT, Olsen J. Antidepressant use during pregnancy and risk of congenital heart defects: A case-time-control study. Pharmacoepidemiol Drug Saf. 2019;28(9):1180-93. doi:10.1002/pds.4844.

18. Greenland S. Confounding and exposure trends in case-crossover and case-time-control designs. Epidemiology. 1996;7(3):231-9. doi:10.1097/00001648-199605000-00003.

19. Schneeweiss S, Stürmer T, Maclure M. Case-crossover and case-time-control designs as alternatives in pharmacoepidemiologic research. Pharmacoepidemiol Drug Saf. 1997;6 Suppl 3:S51-9. doi:10.1002/(sici)1099-1557(199710)6:3+s51::Aid-pds301>3.0.Co;2-s.

20. Donnan PT, Wang J. The case-crossover and case-time-control designs in pharmacoepidemiology. Pharmacoepidemiol Drug Saf. 2001;10(3):259-62. doi:10.1002/pds.590.

21. Suissa S. The case-time-control design. Epidemiology. 1995;6(3):248-53. doi:10.1097/00001648-199505000-00010.

22. Karlsson I. Drugs that induce delirium. Dementia and Geriatric Cognitive Disorders. 1999;10(5):412-5.

23. Francis J. Drug-Induced Delirium. CNS Drugs. 1996;5(2):103-14. doi:10.2165/00023210-199605020-00003.
24. Clegg A, Young JB. Which medications to avoid in people at risk of delirium: a systematic review. Age Ageing. 2011;40(1):23-9. doi:10.1093/ageing/afq140.

25. Douros A, Bronder E, Klimpel A, Erley C, Garbe E, Kreutz R. Drug-induced kidney injury: A large case series from the Berlin Case-Control Surveillance Study. Clin Nephrol. 2018;89 (2018)(1):18-26. doi:10.5414/cn109212.

26. Glavind K, Chancellor M. Antimuscarinics for the treatment of overactive bladder: understanding the role of muscarinic subtype selectivity. Int Urogynecol J. 2011;22(8):907-17. doi:10.1007/s00192-011-1411-6.

27. Perry EK, Perry RH, Smith CJ, Purohit D, Bonham J, Dick DJ et al. Cholinergic receptors in cognitive disorders. Can J Neurol Sci. 1986;13(4 Suppl):521-7. doi:10.1017/s0317167100037240.

28. Doroshyenko O, Fuhr U. Clinical pharmacokinetics and pharmacodynamics of solifenacin. Clin Pharmacokinet. 2009;48(5):281-302. doi:10.2165/00003088-200948050-00001.

29. Yoshida A, Fujino T, Maruyama S, Ito Y, Taki Y, Yamada S. The forefront for novel therapeutic agents based on the pathophysiology of lower urinary tract dysfunction: bladder selectivity based on in vivo drug-receptor binding characteristics of antimuscarinic agents for treatment of overactive bladder. J Pharmacol Sci. 2010;112(2):142-50. doi:10.1254/jphs.09r14fm.

30. Hampel C, Betz D, Burger M, Nowak C, Vogel M. Solifenacin in the Elderly: Results of an Observational Study Measuring Efficacy, Tolerability and Cognitive Effects. Urol Int. 2017;98(3):350-7. doi:10.1159/000455257.

31. Kay GG, Abou-Donia MB, Messer WS, Jr., Murphy DG, Tsao JW, Ouslander JG. Antimuscarinic drugs for overactive bladder and their potential effects on cognitive function in older patients. J Am Geriatr Soc. 2005;53(12):2195-201. doi:10.1111/j.1532-5415.2005.00537.x.

32. Katz IR, Sands LP, Bilker W, DiFilippo S, Boyce A, D'Angelo K. Identification of medications that cause cognitive impairment in older people: the case of oxybutynin chloride. J Am Geriatr Soc. 1998;46(1):8-13. doi:10.1111/j.1532-5415.1998.tb01006.x.
33. Suissa S. The case-time-control design: further assumptions and conditions. 
Epidemiology. 1998;9(4):441-5.

34. Johnson JC, Kerse NM, Gottlieb G, Wanich C, Sullivan E, Chen K. Prospective versus 
retrospective methods of identifying patients with delirium. J Am Geriatr Soc. 
1992;40(4):316-9. doi:10.1111/j.1532-5415.1992.tb02128.x.

35. Hope C, Estrada N, Weir C, Teng CC, Damal K, Sauer BC. Documentation of delirium in 
the VA electronic health record. BMC Res Notes. 2014;7:208. doi:10.1186/1756-0500-7-208.

36. Katznelson R, Djaiani G, Tait G, Wasowicz M, Sutherland AM, Styra R et al. Hospital 
administrative database underestimates delirium rate after cardiac surgery. Can J Anaesth. 
2010;57(10):898-902. doi:10.1007/s12630-010-9355-8.

37. Bui LN, Pham VP, Shirkey BA, Swan JT. Effect of delirium motoric subtypes on 
administrative documentation of delirium in the surgical intensive care unit. J Clin Monit 
Comput. 2017;31(3):631-40. doi:10.1007/s10877-016-9873-1.

38. Collins N, Blanchard MR, Tookman A, Sampson EL. Detection of delirium in the acute 
hospital. Age Ageing. 2010;39(1):131-5. doi:10.1093/ageing/afp201.

39. Kakuma R, du Fort GG, Arsenault L, Perrault A, Platt RW, Monette J et al. Delirium in 
older emergency department patients discharged home: effect on survival. J Am Geriatr Soc. 
2003;51(4):443-50. doi:10.1046/j.1532-5415.2003.51151.x.
Figure 1: Case-time-control cohort

Reference period

Case period

Oxybutynin
Solifenacin
Drug 1
Drug 2

Panel 5: Case subject:
- At least 1 delirium
- Index date = 1st delirium

Panel 6: Matching:
- Age +/- 100 days
- Sex and ethnicity
- One case to up to 3 controls

Panel 7: Control subject:
- No delirium
- Index date = index date of matching case

Prescription histories (Pharms)
Table 1. Characteristics of the study population according to antimuscarinic type

| Characteristic          | Oxybutynin | Solifenacin |
|-------------------------|------------|-------------|
| Age                     |            |             |
| 65-69                   | 0          | 6           |
| 70-74                   | 11         | 50          |
| 75-79                   | 44         | 172         |
| 80-84                   | 107        | 272         |
| 85-89                   | 100        | 372         |
| 90+                     | 71         | 265         |
| Sex                     |            |             |
| Male                    | 190        | 664         |
| Female                  | 143        | 473         |
| Ethnicity               |            |             |
| NZ European             | 310        | 1062        |
| Māori                   | 6          | 18          |
| Asian                   | 4          | 12          |
| Pacific people          | 0          | 8           |
| MELAA*                  | 0          | 3           |
| Other                   | 13         | 34          |
| Calendar year of the incident event |            |             |
| 2005-2009               | 0          | 395         |
| 2010-2014               | 333        | 742         |

*MELAA- Middle Eastern, Latin American and African*
Table 2: The matched odds ratio (MOR) for delirium diagnosed in older adults (65 years and above) who used antimuscarinics in a case-time-control study

| Drug          | Delirium at index date | Exposures within 5-day observation periods | Exposures within 3-day observation periods |
|---------------|------------------------|------------------------------------------|------------------------------------------|
|               |                        | Neither, Case, Control, Both | Neither, Case, Control, Both | Neither, Case, Control, Both |
| Oxybutynin    | Yes                    | 4485, 85, 57, 191 | 4487, 86, 53, 192 | 2.06 (1.07-3.96) |
|               | No (Reference)         | 13275, 21, 29, 30 | 13276, 21, 29, 29 | 1.00 (0.72-1.39) |
| Solifenacin   | Yes                    | 3681, 212, 230, 695 | 3680, 227, 231, 680 | 0.89 (0.64-1.23) |
|               | No (Reference)         | 13013, 110, 106, 126 | 13020, 105, 107, 123 | 1.00 (0.72-1.39) |
| Antibiotics   | Yes                    | 4607, 79, 45, 87 | 4610, 79, 49, 80 | 1.96 (1.27-3.03) |
|               | No (Reference)         | 12868, 130, 145, 212 | 12869, 132, 146, 208 | 1.78 (1.16-2.73) |
| Anti-epileptics| Yes                   | 4745, 10, 16, 47 | 4744, 12, 17, 45 | 0.62 (0.22-1.81) |
|               | No (Reference)         | 13273, 15, 15, 52 | 13274, 10, 12, 59 | 0.85 (0.28-2.59) |
| Antipsychotics| Yes                   | 4550, 77, 64, 127 | 4564, 63, 64, 127 | 1.09 (0.74-1.62) |
|               | No (Reference)         | 12735, 177, 161, 282 | 12738, 166, 159, 292 | 0.94 (0.63-1.42) |
| Anti-depressants| Yes               | 4380, 105, 74, 259 | 4379, 105, 76, 258 | 1.43 (0.98-2.11) |
|                | No (Reference)         | 12547, 136, 138, 534 | 12545, 132, 128, 550 | 1.44 (0.99-2.11) |
| Weak-ACs      | Yes                    | 3263, 265, 214, 716 | 3613, 258, 224, 723 | 1.24 (1.00-1.53) |
|                | No (Reference)         | 10117, 571, 570, 2097 | 10103, 569, 556, 2127 | 1.13 (0.91-1.39) |
| Strong-ACs    | Yes                    | 4813, 1, 1, 2 | 4814, 1, 1, 2 | 1.00 (0.02-50.4) |
|                | No (Reference)         | 13351, 1, 1, 2 | 13351, 1, 1, 2 | 1.00 (0.02-50.4) |
Table 3 The effect sizes with 95% CI of the dose-response change in the risk of delirium for oxybutynin and solifenacin using case-time-control study design.

| Variable                                      | Oxybutynin       | Solifenacin      |
|-----------------------------------------------|------------------|------------------|
| Age group: 70-74yr                            | 0.047 (0.016 - 0.077) | -0.011 (-0.028 - 0.005) |
| Age group: 75-79yr                            | -0.051 (-0.479 - 0.377) | -0.051 (-0.478 - 0.377) |
| Age group: 80-84yr                            | -0.096 (-0.507 - 0.315) | -0.095 (-0.506 - 0.315) |
| Age group: 85-89yr                            | -0.064 (-0.472 - 0.344) | -0.065 (-0.472 - 0.343) |
| Age group: 90yr +                             | -0.144 (-0.551 - 0.263) | -0.143 (-0.550 - 0.264) |
| Sex: Female                                   | -0.015 (-0.082 - 0.053) | -0.016 (-0.083 - 0.052) |
| Ethnicity: Māori                             | -0.179 (-0.413 - 0.056) | -0.179 (-0.413 - 0.056) |
| Ethnicity: Pacific                           | 0.066 (-0.277 - 0.408) | 0.064 (-0.278 - 0.407) |
| Ethnicity: Asian                             | 0.101 (-0.172 - 0.374) | 0.099 (-0.174 - 0.372) |
| Ethnicity: MELAA                              | 0.469 (-0.219 - 1.158) | 0.463 (-0.226 - 1.152) |
| Ethnicity: Other                             | 0.388 (0.215 - 0.561) | 0.389 (0.216 - 0.562) |
| Antibiotics exposed: case period              | 0.615 (0.336 - 0.894) | 0.627 (0.349 - 0.906) |
| Antibiotics exposed: control period           | 0.008 (-0.295 - -0.312) | -6e-04 (-0.304 - 0.303) |
| Antiepileptics exposed: both periods          | 0.030 (-0.207 - 0.266) | 0.030 (-0.206 - 0.266) |
| Antiepileptics exposed: case period           | 1.474 (0.536 - 2.412) | 1.493 (0.554 - 2.432) |
| Antiepileptics exposed: control period        | 0.675 (-0.325 - 1.675) | 0.654 (-0.347 - 1.656) |
| Antidepressants exposed: both periods         | 0.861 (0.503 - 1.220) | 0.861 (0.503 - 1.219) |
| Antidepressants exposed: case period          | 0.280 (0.041 - 0.519) | 0.278 (0.040 - 0.517) |
| Antidepressants exposed: control period       | 0.075 (-0.180 - 0.330) | 0.075 (-0.180 - 0.330) |
| Antipsychotics exposed: both periods          | 0.155 (-0.046 - 0.356) | 0.152 (-0.049 - 0.353) |
| Antipsychotics exposed: case period           | 0.768 (0.485 - 1.052) | 0.772 (0.489 - 1.055) |
| Antipsychotics exposed: control period        | 0.439 (0.130 - 0.747) | 0.436 (0.128 - 0.744) |
| Antipsychotics exposed: both periods          | 0.302 (0.156 - 0.4483) | 0.302 (0.156 - 0.449) |
| Weak anticholinergics exposed: case period    | 0.028 (-0.140 - 0.196) | 0.039 (-0.129 - 0.207) |
| Weak anticholinergics exposed: control period | 0.021 (-0.153 - 0.195) | 0.010 (-0.164 - 0.185) |
| Weak anticholinergics exposed: both periods   | -0.116 (-0.209 - -0.024) | -0.116 (-0.208 - -0.024) |
| Strong anticholinergics exposed: case period  | 1.237 (-1.537 - 4.011) | 1.142 (-1.632 - 3.916) |
| Strong anticholinergics exposed: control period | 1.065 (-1.709 - 3.838) | 1.064 (-1.710 - 3.837) |
| Strong-AC exposed: both periods               | 1.246 (-0.582 - 3.074) | 1.245 (-0.584 - 3.073) |
| Dementia diagnosis before index-date          | 1.361 (0.798 - 1.925) | 1.369 (0.805 - 1.932) |
Table 4 The matched odds ratio (MOR) for delirium diagnosed in older adults (65 years and above) by dementia status who used antimuscarinics in a case-time-control study

| Drug          | Delirium at index date | Exposures within 5-day observation periods | 5-day observation periods | Exposures within 5-day observation periods | Exposures within 5-day observation periods | Exposures within 5-day observation periods | Exposures within 5-day observation periods |
|---------------|------------------------|---------------------------------------------|---------------------------|---------------------------------------------|---------------------------------------------|---------------------------------------------|---------------------------------------------|
|               |                        | Neither | Case | Control | Both | MOR            | Neither | Case | Control | Both | MOR            | Neither | Case | Control | Both | MOR            |
| Oxybutynin    | Yes                    | 4336    | 80   | 53      | 177  | 2.11 (1.08-4.13) | 149     | 5    | 4       | 14   | 1.25 (0.05-26.9) |
|               | No (Reference)         | 13112   | 20   | 28      | 28   |               | 163     | 1    | 1       | 2    |               |
| Solifenacin   | Yes                    | 3539    | 203  | 223     | 681  | 0.89 (0.64-1.23) | 142     | 9    | 7       | 14   | 0.64 (0.04-8.62) |
|               | No (Reference)         | 12853   | 108  | 105     | 122  |               | 160     | 2    | 1       | 4    |               |
| Antibiotics   | Yes                    | 4442    | 76   | 45      | 83   | 1.96 (1.26-3.05) | 165     | 3    | 0       | 4    | N/A             |
|               | No (Reference)         | 12710   | 124  | 144     | 210  |               | 158     | 6    | 1       | 2    |               |
| Antiepileptics| Yes                    | 4576    | 9    | 15      | 46   | 0.60 (0.20-1.79) | 169     | 1    | 1       | 1    | N/A             |
|               | No (Reference)         | 13107   | 15   | 15      | 51   |               | 166     | 0    | 0       | 1    |               |
| Antipsychotics| Yes                   | 4386    | 73   | 62      | 125  | 1.07 (0.72-1.60) | 164     | 4    | 2       | 2    | 2.00 (0.15-26.7) |
|               | No (Reference)         | 12575   | 175  | 159     | 279  |               | 160     | 2    | 2       | 3    |               |
| Antidepressants| Yes                 | 4228    | 97   | 71      | 250  | 1.43 (0.97-2.11) | 152     | 8    | 3       | 9    | 0.53 (0.04-6.67) |
|               | No (Reference)         | 12404   | 131  | 137     | 516  |               | 143     | 5    | 1       | 18   |               |
| Weak anticholinergics| Yes             | 3135    | 249  | 209     | 693  | 1.19 (0.96-1.48) | 128     | 16   | 5       | 23   | 2.80 (0.67-11.7) |
|               | No (Reference)         | 9992    | 563  | 563     | 2070 |               | 125     | 8    | 7       | 27   |               |
| Strong anticholinergics| Yes             | 4641    | 1    | 1       | 3    | 1.00 (0.02-50.4) | 172     | 0    | 0       | 0    | N/A             |
|               | No (Reference)         | 13184   | 1    | 1       | 2    |               | 167     | 0    | 0       | 0    |               |