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Potential Risk Factors for, and Clinical Implications of, Delirium

during Inpatient Rehabilitation: A Matched Case-Control Study

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ABSTRACT

Objectives: To investigate the association between a wide set of baseline characteristics (age, sex, rehabilitation discipline), functional scores [Functional Independence Measure (FIM), cumulative Illness Rating Scale (CIRS)], diseases, and administered drugs and incident delirium in rehabilitation inpatients and, furthermore, to assess clinical implications of developing delirium during rehabilitation. *Design:* Matched case-control study based on electronic health record data.

Setting and participants: We studied rehabilitation stays of inpatients admitted between January 1, 2015, and December 31, 2018, to ZURZACH Care, Rehaklinik Bad Zurzach, an inpatient rehabilitation clinic in Switzerland.

Methods: We conducted unconditional logistic regression analyses to estimate adjusted odds ratios (AORs) with 95% CIs of exposures that were recorded in \geq 5 cases and controls.

Results: Among a total of 10,503 rehabilitation stays, we identified 125 validated cases. Older age, undergoing neurologic rehabilitation, a low FIM, and a high CIRS were associated with an increased risk of incident delirium. Being diagnosed with a bacterial infection (AOR 2.62, 95% CI 1.06-6.49), a disorder of fluid, electrolyte, or acid-base balance (AOR 2.76, 95% CI 1.19-6.38), Parkinson's disease (AOR 5.68, 95% CI 2.54-12.68), and administration of antipsychotic drugs (AOR 8.06, 95% CI 4.26-15.22), antiparkinson drugs (AOR 2.86, 95% CI 1.42-5.77), drugs for constipation (AOR 2.11, 95% CI 1.25-3.58), heparins (AOR 2.04, 95% CI 1.29-3.24), or antidepressant drugs (AOR 1.88, 95% CI 1.14-3.10) during rehabilitation, or an increased anticholinergic burden (ACB \geq 3) (AOR 2.59, 95% CI 1.41-4.73) were also associated with an increased risk of incident delirium.

Conclusions and Implications: We identified a set of factors associated with an increased risk of incident delirium during inpatient rehabilitation. Our findings contribute to detect patients at risk of delirium during inpatient rehabilitation.

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Delirium is an etiologically nonspecific organic cerebral syndrome characterized by concurrent impairment of consciousness, attention, perception, thinking, memory, psychomotor behavior, emotions, and the sleep-wake cycle and can vary in duration and severity.^{1,2} The

underlying pathomechanisms are likely multifactorial, and identified risk factors in a hospital setting are older age, male sex, decreased functional ability, high burden of disease, comorbidities such as degenerative neurologic disorders or infections, dehydration, malnutrition, immobility, prolonged hospital stay, and polypharmacy.³⁻¹⁰ Several studies have suggested that acetylcholine deficiency may be involved in the pathophysiology of delirium, and that the use of anticholinergic medications may increase the risk of delirium.¹¹⁻¹⁷

In the inpatient rehabilitation setting, as in the acute setting, delirium has been associated with a longer duration of stay and higher mortality.^{18–23} Because of the inability of delirious patients to follow

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the challenging interdisciplinary therapeutic schedule, delirium has also been associated with poor functional rehabilitation outcome.^{24,25} Two studies assessing the Functional Independence Measure (FIM) of patients undergoing rehabilitation showed that patients who developed delirium during the stay had a more severe impairment at the beginning, and a more limited FIM improvement during rehabilitation than patients who did not.^{26,27}

Older age is a common risk factor for delirium among rehabilitation inpatients.²⁶⁻²⁹ Also, a retrospective study identified traumatic brain injury, depression, diabetes mellitus and musculoskeletal disorders, as well as several out-of-range laboratory parameters as risk factors for delirium among rehabilitation inpatients.²⁹

Identifying risk factors for incident delirium during rehabilitation, including specific conditions and administered drugs, is useful to detect patients who are susceptible to develop delirium.

The aim of this study was to explore the association between incident delirium during inpatient rehabilitation and a wide range of factors such as patient characteristics, rehabilitation discipline, functional scores at admission, diagnoses, and administered drugs. Furthermore, this study aimed to describe functional rehabilitation outcome and length of rehabilitation stay in patients who developed delirium and in patients who did not.

Methods

Data Source and Study Design

We conducted a retrospective matched case-control study using data from the electronic health records of ZURZACH Care, Rehaklinik Bad Zurzach, an inpatient rehabilitation clinic in Switzerland. Electronic health records comprise medical notes (suggestive of incident delirium, as validated in a previous study),³⁰ patient- and rehabilitation-specific characteristics such as age, sex, rehabilitation discipline and length of stay, as well as clinical data such as diagnoses [recorded as *International Classification of Diseases, Tenth Revision (ICD-10)* codes],² administered drugs [recorded as Anatomical Therapeutic Chemical (ATC) codes],³¹ FIM,³² and the Cumulative Illness Rating Scale (CIRS).³³ This study was approved by the Ethics Committee Northwest/Central Switzerland (Project-ID 2018-01351).

Study Population

We included all rehabilitation stays of patients who were admitted for inpatient rehabilitation between January 1, 2015, and December 31, 2018. Single patients may have contributed to more than 1 rehabilitation stay, if they were referred for rehabilitation several times during the study period. We excluded all stays with missing patient characteristics such as age, sex, or rehabilitation discipline.

Cases and Controls

Cases were patients who developed delirium at some point after the admission date. The definition and validation of delirium in the data set has been described in detail previously.³⁰ Briefly, we defined 15 keywords commonly used to describe delirious patients in medical notes. Profiles of patients with at least 2 recorded keywords and no diagnosis of delirium at admission were reviewed by at least 2 independent physicians, based on predefined evaluation criteria to confirm or refute the diagnosis of delirium. In confirmed cases, the first recorded keyword was defined as the date of onset of delirium (index date). Eligible controls were patients who did not have any record of delirium predictive keywords in their electronic health records and no diagnosis of delirium at admission. For each validated case, we matched 4 controls on calendar time [by assigning the index date $(\pm 1 \text{ month})$ of the cases to their controls] and time span between admission date and index date.

Exposure

For cases and controls, we assessed age and length of stay as continuous variables, and sex (male; female), age groups (<65; 65-74; 75-84; \geq 85 years), rehabilitation discipline (neurology; non-neurology), and primary diagnosis for rehabilitation as categorical variables. Furthermore, we assessed FIM, including cognitive FIM (C-FIM) and motoric FIM (M-FIM) in categories of severity, adapted from the German Modification of the *ICD-10*,³⁴ and evaluated its change between admission and discharge. We assessed disease burden at admission, by categorizing the CIRS into quartiles. We assessed the prevalence of comorbidities recorded in \geq 5 cases and controls (see Supplementary Table 1 for the complete *ICD-10* codes list).

Additionally, we assessed the number of administered drugs at admission as continuous variable, and the administered drug classes that were recorded in \geq 5 cases and controls at any time between admission and index date (see Supplementary Table 2 for the complete ATC codes list). We defined "users" of the above drugs as patients with at least 1 administration between admission and index date, and "nonusers" as those with no recorded administration in the same interval. Lastly, we calculated the Anticholinergic Cognitive Burden (ACB) at admission and assessed whether cases and controls were exposed to an increased ACB (\geq 3 or <3).³⁵

Statistical Analysis

We summarized continuous variables providing means and SDs, and categorical variables as absolute and relative frequencies.

We conducted unconditional logistic regression analyses to calculate odds ratios with 95% CIs for each exposure variable. We adjusted all analyses for sex, age, and rehabilitation discipline to calculate adjusted odds ratios (AORs) with 95% CIs. Given the unconditional analysis of matched sets, we also adjusted all analyses for the 2 matching factors (index date and time span between admission and index date).³⁶

All statistical analyses were conducted using SAS 9.4 (SAS Institute). Graphics were composed using Prism GraphPad 9.4 (GraphPad Software).

Results

Of 9406 patients who underwent a total of 10,503 rehabilitation stays during the study period, we identified 125 validated incident delirium episodes and 500 matched controls (Supplementary Figure 1). Patients and rehabilitation characteristics of cases and controls are reported in Table 1. Diseases of the nervous system (53.6%), among these cerebral infarction (26.4%), were the most frequent primary diagnoses for rehabilitation among cases. Diseases of the musculoskeletal system (48.0%), among these spondylopathies (7.4%) and other dorsopathies (12.8%), were the most frequent primary diagnoses for rehabilitation among controls (Supplementary Table 3).

Older age and undergoing neurologic rehabilitation were associated with increased risks of incident delirium (Table 1).

Severe functional impairment (FIM \leq 65) and severe burden of disease (CIRS \geq 14) were also associated with increased risks of incident delirium (Table 2).

Several comorbidities were associated with an increased risk of incident delirium during inpatient rehabilitation (Figure 1; see Supplementary Table 4 for exact numbers). Being diagnosed with bacterial infections or disorders of fluid, electrolyte, and acid-base balance was associated with a moderately increased risk of incident delirium (AORs 2.62, 95% CI 1.06-6.49, and 2.76, 95% CI 1.19-6.38,

Table 1	l
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Characteristic	Cases (n = 125)	Controls ($n = 500$)	OR (95% CI)	AOR (95% CI)*
Sex				
Female	55 (44.0)	275 (55.0)	1 ref.	1 ref.
Male	70 (55.0)	225 (45.0)	1.56 (1.05-2.31)	1.39 (0.89-2.17)
Age, y				
<65	13 (10.4)	227 (45.4)	1 ref.	1 ref.
65-74	23 (18.4)	110 (22.0)	3.67 (1.79-7.53)	3.54 (1.69-7.45)
75-84	62 (49.6)	128 (25.6)	8.48 (4.49-16.02)	9.06 (4.68-17.56
≥85	27 (21.2)	35 (7.0)	13.64 (6.42-28.99)	12.99 (5.89-28.67
Age, y, mean (SD)	77.2 (9.9)	64.6 (15.7)	n/a	n/a
Rehabilitation discipline				
Neurology	89 (71.2)	167 (33.4)	4.97 (3.23-7.65)	4.89 (3.07-7.79)
Nonneurology	36 (28.8)	333 (66.6)	1 ref.	1 ref.
Days between admission date and index date, mean (SD)	10.3 (10.3)	10.3 (10.3)	n/a	n/a

n/a, not applicable; ref., referent.

Values are n (%) unless otherwise noted. Controls were matched to cases on index date (±1 month) and time between the admission date and the index date (days between admission date and index date). All ORs were calculated with unconditional logistic regression and adjusted for matching factors (index date and exposure time).

*Sex adjusted on age, rehabilitation discipline (neurology/nonneurology); age adjusted on sex, rehabilitation discipline (neurology/nonneurology); rehabilitation discipline adjusted on age, sex.

[†]Frequencies (%) within nonneurology disciplines (cases/controls): angiology (4.0/7.4), cardiology (4.8/9.6), rheumatology (1.6/9.8), orthopedics (15.2/26.2), headache (0/4.0), or pain (0.8/7.4) programs.

respectively), compared to not having these diagnoses. Parkinson's disease, and more generally extrapyramidal and movement disorders, were strongly associated with the risk of incident delirium compared to not having these conditions (AOR 5.68, 95% CI 2.54-12.68, and 3.51, 95% CI 1.89-6.52, respectively). Other comorbidities were not associated with incident delirium after adjusting for sex, age, and rehabilitation discipline.

Cases had a higher number of administered drugs at admission compared to controls [mean (SD), 9.0 (3.4) vs 6.7 (3.8)]. The administration of different drug classes was associated with an increased risk of incident delirium (Figure 2; see Supplementary Table 5 for exact numbers). The use of drugs for constipation (AOR 2.11, 95% CI 1.25-3.58), heparins (AOR 2.04, 95% CI 1.29-3.24), and antidepressants (AOR 1.88, 95% CI 1.14-3.10) was associated with a moderately increased risk of incident delirium, whereas the use of dopaminergic agents and antipsychotic drugs was associated with a markedly increased risk of incident delirium compared to non-use of these drug classes (AOR 2.86, 95% CI 1.42-5.77, and 8.06, 95% CI 4.26-15.22, respectively). Several drug classes were not associated with

incident delirium after adjusting for sex, age, and rehabilitation discipline.

The ACB was higher within cases than controls [mean (SD), 0.9 (1.3) vs 0.6 (1.1)], and having a high ACB (\geq 3) was associated with an increased risk of delirium compared to having a low ACB (<3) (AOR 2.59, 95% CI 1.41-4.73).

Cases had a longer mean rehabilitation stay than controls [mean days (SD), 33.1 (18.7) vs 27.8 (16.5)], and the FIM of cases improved less between admission and discharge [Δ FIM (SD), 7.4 (17.1) vs 17.9 (12.6)] than that of controls (Figure 3).

Discussion

In this retrospective matched case-control study based on inpatient clinical data, we identified older age, neurologic rehabilitation, reduced FIM, and high disease or anticholinergic burden at admission as factors associated with a considerably increased risk of incident delirium during rehabilitation.

Table 2

Odds Ratios of FIM and CIRS Scores at Admission Among Cases With Incident Delirium and Matched Controls

Measure	Cases $(n = 125)$	Controls $(n = 499)^*$	OR (95% CI)	AOR (95% CI) [†]
FIM score at admission				
FIM, mean (SD)	45.6 (18.4)	78.7 (19.3)	n/a	n/a
Cognitive FIM, mean (SD)	13.2 (5.7)	22.3 (5.4)	n/a	n/a
Motor FIM, mean (SD)	32.5 (15.0)	56.4 (15.5)	n/a	n/a
FIM low to medium impairment (66-126)	16 (12.8)	382 (76.6)	1 ref.	1 ref.
FIM high impairment (18-65)	109 (87.2)	117 (23.4)	25.88 (14.42-46.46)	13.19(7.03-24.72)
Cognitive FIM low to medium impairment (11-35)	73 (58.4)	488 (97.8)	1 ref.	1 ref.
Cognitive FIM high impairment (5-10)	52 (41.6)	11 (2.2)	32.37 (16.08-65.16)	19.11(8.64-42.27)
Motor FIM low to medium impairment (27-91)	76 (60.8)	471 (94.4)	1 ref.	1 ref.
Motor FIM high impairment (13-26)	49 (39.2)	28 (5.6)	11.50 (6.73-19.64)	6.75(3.65-12.51)
CIRS score at admission				
CIRS, mean (SD)	18.8 (8.2)	15.1 (9.6)	n/a	n/a
CIRS low severity (0-8)	10 (8.0)	141 (28.3)	1 ref.	1 ref.
CIRS medium severity (9-13)	23 (18. 4)	131 (26.3)	2.70 (1.23-5.92)	1.63 (0.69-3.83)
CIRS high severity (14-20)	45 (36.0)	102 (20.4)	6.98 (3.31-14.70)	2.95 (1.29-6.74)
CIRS very high severity (21-56)	47 (37.6)	125 (25.1)	6.12 (2.90-12.90)	2.65 (1.16-6.07)

n/a, not applicable; ref., referent.

Values are n (%) unless otherwise noted. Controls were matched to cases on index date (±1 month) and time between the admission date and the index date (days between admission date and index date). All ORs were calculated with unconditional logistic regression and adjusted for matching factors (days between admission date and index date).

*Missing database entries (FIM and CIRS) for 1 control.

[†]Adjusted on age, sex, rehabilitation discipline (neurology/nonneurology).

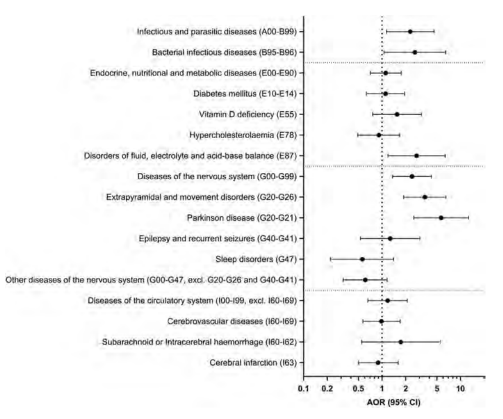


Fig. 1. Forest plot of adjusted odds ratios (95% CIs) among cases with incident delirium and matched controls for exposure to different comorbidities defined as a record of the *International Classification of Diseases, Tenth Revision,* code. Controls were matched to cases on index date (±1 month) and days between the admission date and the index date. Odds ratios were calculated with unconditional logistic regression and adjusted for matching factors, age, sex, and rehabilitation discipline (neurology/nonneurology).

Patients with infectious diseases, disorders of fluid, electrolyte, and acid-base balance, and Parkinson's disease at admission, and patients treated with laxatives, heparins, antidepressants, dopaminergic agents, and antipsychotics during rehabilitation, were at an increased risk of developing delirium.

Furthermore, patients who developed incident delirium had a longer mean rehabilitation stay and a poorer functional rehabilitation outcome, quantified by the FIM change between admission and discharge, than patients without delirium.

Patient and Rehabilitation Characteristics

Our results suggest that patients who have become delirious during rehabilitation were more frequently men and older than patients who have not. Compared to patients aged <65 years, patients between 65 and 74 years of age had a 3.5-fold increased risk, patients aged between 75 and 84 years a 9.1-fold increased risk, and patients >85 years a 13.0-fold increased risk of delirium. The results are consistent with previous studies, which reported that patients who developed delirium during rehabilitation were older²⁶⁻²⁹ and more often men^{26,28,29} than patients who did not develop delirium. In our study, most cases underwent neurologic rehabilitation, and patients among this rehabilitation discipline had a 4.9-fold increased risk of incident delirium compared to patients among other rehabilitation disciplines. This observation could be explained by neurologic imbalance caused by degenerative neurologic conditions that may trigger the pathophysiology of delirium.³⁷ The cognitive and motoric FIM at admission was lower among cases than controls [mean (SD), 13.2 (5.7) vs 22.3 (5.4) and 32.5 (15.0) vs 56.4 (15.5), respectively], and patients with an FIM lower than 65 points at admission had a 13.2-fold increased risk of incident delirium as compared to patients with an FIM higher than 65 points. These results suggest that patients with an impaired functional degree are more likely to develop delirium during rehabilitation, which is consistent with 2 previously published studies that assessed the FIM among patients with and without delirium.^{26,27} Bushi et al²⁶ found that patients with delirium had a significantly lower cognitive and motor FIM on admission than patients without delirium [mean (SD), 15.2 (5.8) vs 24.2 (6.0) and 24.3 (9.6) vs 31.3 (9.1), respectively] and that patients with delirium more often had a primary neurologic diagnosis for rehabilitation than patients without delirium.²⁶

Burden of Disease and Comorbidities

We observed a 2.6- to 2.9-fold increased risk of delirium among patients with an increased burden of disease (CIRS) compared to patients with low burden of disease. This is comparable with the observations of Stelmokas et al,³⁸ who reported a 4.5-fold increased risk of delirium among patients with an elevated Age-Adjusted Charlson Index.

Patients with prevalent infectious diseases had a 2.3-fold increased risk of delirium, patients with disorders of fluid, electrolyte, and acidbase balance had a 2.7-fold increased risk of delirium, and patients with extrapyramidal and movement disorders even had a 3.5-fold (among them, patients with Parkinson's disease a 5.7-fold) increased risk of delirium compared with patients who did not have a diagnosis of these conditions. These results are only partially comparable to those of a previous study, which assessed comorbidities and laboratory parameters as potential risk factors for delirium in the rehabilitation setting.²⁹ Jang et al²⁹ observed an increased risk of delirium among patients with traumatic brain injuries, depression, diabetes mellitus, and musculoskeletal disorders, as well as among patients with increased white blood cells, erythrocyte sedimentation

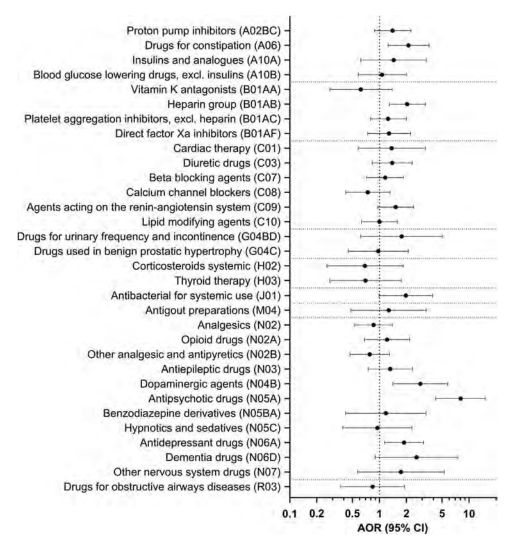


Fig. 2. Forest plot of adjusted odds ratios (95% CIs) among cases with incident delirium and matched controls for exposure to selected drug groups defined as at least 1 record of an administered code of the respective Anatomical Therapeutic Chemical (ATC) class at any time from the admission date until the index date. Controls were matched to cases on index date (±1 month) and days between the admission date and the index date. Odds ratios were calculated with unconditional logistic regression and adjusted for matching factors, age, sex, and rehabilitation discipline (neurology/nonneurology).

rate, C-reactive protein and decreased potassium and phosphorus levels.²⁹ In our study we could not assess brain injuries, depression, and musculoskeletal disorders (<5 observations for cases and/or controls), and we did not observe an increased risk of delirium among patients with diabetes mellitus. Nevertheless, the increased inflammatory or infectious parameters (ie, white blood cells, erythrocyte sedimentation rate, and C-reactive protein) observed by Jang et al²⁹ are consistent with the increased risk of delirium we observed among patients with infectious diseases, and the decreased potassium and phosphorus levels are consistent with the increased risk we observed among patients with disorders of fluid, electrolyte, and acid-base balance. These findings are consistent with the current state of research suggesting that neurodegenerative diseases affecting dopamine levels and conditions of inflammation or electrolyte imbalance are favorable conditions for the development of delirium.³⁷

Anticholinergic Burden and Comedications

Among our study population, cases on average used more drugs than controls. The resulting anticholinergic burden was higher among cases than controls, and patients with an ACB of \geq 3 points had a

2.6-fold increased risk of incident delirium compared to patients with an ACB <3. These observations support the hypothesis of several studies that polypharmacy, particularly involving drugs with anticholinergic potential, may cause neurotransmitter imbalance and thus promote the pathophysiology of delirium.^{10-17,37}

Patients who used laxatives, heparins or antidepressants had an approximately 2-fold increased risk of developing delirium, patients who used dopaminergic agents had a 2.9-fold increased risk, and those who used antipsychotics had an approximately 8-fold increased risk compared with nonuse of these drug classes. From a pharmacologic point of view, only some of these results are attributable to the direct effect of these drug classes on the onset of delirium, whereas others may be indirectly but not causally associated with delirium. For instance, in inpatient setting, heparins are often used to prevent thromboembolic conditions,³⁹ and laxatives to prevent constipation among patients with reduced mobility. It is reasonable to assume that the observed association is rather due to the prolonged immobility than to a direct pathogenic effect of these classes of drugs on delirium. We also observed a statistically significant association of both Parkinson's disease and dopaminergic drugs with an increased risk of delirium. Although this may be plausible from a pharmacologic point

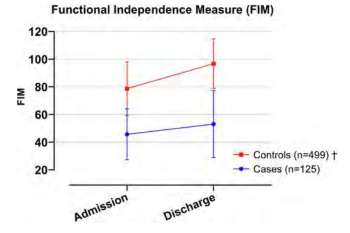


Fig. 3. Functional Independence Measure (FIM) scores at rehabilitation admission and at discharge for cases with incident delirium and matched controls, mean (SD). FIM improvement between admission and discharge, mean (SD): 7.4 (17.1) for cases; 17.9 (12.6) for controls. †Missing database entries for 1 control.

of view, the association between dopaminergic drugs and increased risk of delirium could reflect that almost all Parkinson's patients receive this drug class as a standard treatment.

Clinical Implications

We observed that patients who experienced incident delirium during rehabilitation on average had a 5 days longer rehabilitation stay and a poorer functional rehabilitation outcome at discharge [Δ FIM (SD), 7.4 (17.1) vs 17.9 (12.6)] than patients who did not. These observations are consistent with previous studies,²⁶⁻²⁹ particularly one study reported a significantly lower change in FIM between admission and discharge for patients with delirium compared with patients without delirium [Δ FIM (SD), 10.5 (13.1) vs 19.4 (15.4)].²⁷

Strengths and Limitations

The following limitations of our study have to be considered. First, our analyses were based on clinical routine data, which were not primarily collected for research purposes. However, the consistency of our results with previous studies corroborates the validity of our data. Second, although we rigorously assessed medication use prior to the index date and time, potential protopathic bias must be considered. For example, the substantially increased risk of delirium observed in association with antipsychotic drugs may be explained by the administration of this drug class to patients presenting with early symptoms of delirium, rather than by a direct association between antipsychotic drug use and delirium. Because of the nonspecific and off-label use of antipsychotic drugs in clinical practice and the short follow-up time, we were not able to detect and limit this type of bias by shifting the index date. Third, because the aim of our study was not to test formal hypotheses, we assessed a wide range of potential risk factors simultaneously. Therefore, the results should be considered as a set of factors associated with, rather than causing delirium. Fourth, because of the low prevalence of certain drug classes and also the short observation time of our study, we were not able to differentiate between occasional, prolonged, or cumulative use of medication. This would have helped us to understand whether the increased risk of delirium is associated with chronic use of certain drugs, or whether even occasional use is associated with delirium. However, given the pathophysiology of delirium, which typically develops within hours or days, we believe that our approach was appropriate for the assessed drug classes.

An important strength of our study is the high quality of the data set, which comprised accurate and structured entries of each single drug administration and diagnosis record. This allowed us to precisely define exposures without the use of proxy parameters.

Considering the above-mentioned limitations, our study offers a broad overview of the main risk factors for incident delirium during inpatient rehabilitation. Especially, our study adds knowledge to the existing literature regarding associations between administered drug classes and incident delirium during rehabilitation.

Conclusions and Implications

Our study suggests that among inpatients undergoing rehabilitation, older age, neurologic rehabilitation, reduced FIM, and high disease or anticholinergic burden, as well as a number of prevalent comorbidities and coadministered drug classes, are potential risk factors for incident delirium. Moreover, incident delirium during rehabilitation seems to be associated with worse functional rehabilitation outcome and longer rehabilitation stay.

These findings may be relevant for health care providers working in the rehabilitation setting. Identifying patients potentially at risk of delirium during rehabilitation by considering a set of risk factors at rehabilitation admission, such as age, functional scores, comorbidities, and preexisting drug prescriptions could represent an innovative method compared to the more conventional delirium assessment tools, which are based on the observation of patients over time and are therefore time consuming and require staff training.⁴⁰ Furthermore, modifiable risk factors such as new drug prescriptions or the anticholinergic drug burden should be proactively considered to reduce the risk of incident delirium.

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Supplementary Material

Supplementary Table 1

List of Assessed Comorbidities, Inclusive ICD-10 Codes, and Subcodes

Comorbidities	ICD-10 Codes
Infectious and parasitic diseases	A00-B99
Bacterial infectious diseases	B95-B96
Endocrine, nutritional, and metabolic diseases	E00-E90
Diabetes mellitus	E10-E14
Vitamin D deficiency	E55
Hypercholesterolemia	E78
Disorders of fluid, electrolyte and acid-base balance	E87
Diseases of the nervous system	G00-G99
Extrapyramidal and movement disorders	G20-G26
Parkinson disease	G20-G21
Epilepsy and recurrent seizures	G40-G41
Sleep disorders	G47
Other diseases of the nervous system	G00-G47, excl. G20-G26; G40-G41
Diseases of the circulatory system	100-199, excl. 160-169
Cerebrovascular diseases	160-169
Subarachnoid or Intracerebral hemorrhage	160-162
Cerebral infarction	I63

ICD-10, International Classification of Diseases, Tenth Revision.

Supplementary Table 2

List of Assessed Co-administered Drug Classes Inclusive ATC Codes and Subcodes

Administered Drug Classes	ATC Codes
Proton pump inhibitors	A02BC
Drugs for constipation	A06
Insulins and analogues	A10A
Blood glucose lowering drugs, excl. insulins	A10B
Vitamin K antagonists	B01AA
Heparin group	B01AB
Platelet aggregation inhibitors, excl. heparin	B01AC
Direct factor Xa inhibitors	B01AF
Cardiac therapy	C01
Diuretic drugs	C03
Beta blocking agents	C07
Calcium channel blockers	C08
Agents acting on the renin-angiotensin system	C09
Lipid modifying agents	C10
Drugs for urinary frequency and incontinence	G04BD
Drugs used in benign prostatic hypertrophy	G04C
Corticosteroids systemic	H02
Thyroid therapy	H03
Antibacterial for systemic use	J01
Antigout preparations	M04
Analgesics	N02
Opioid drugs	N02A
Other analgesic and antipyretics	N02B
Antiepileptic drugs	N03
Dopaminergic agents	N04B
Antipsychotic drugs	N05A
Benzodiazepine derivatives	N05BA
Hypnotics and sedatives	N05C
Antidepressant drugs	N06A
Dementia drugs	N06D
Other nervous system drugs	N07
Drugs for obstructive airways diseases	R03

ATC, Anatomical Therapeutic Chemical.

Supplementary Table 3

Primary Diagnosis for Rehabilitation of Cases With Incident Delirium and Matched Controls

Primary Diagnoses for Rehabilitation (ICD-10)	Cases $(n = 125) n (\%)$	Controls (n = 500) n (%)
Neoplasms (C00-D48)	6 (4.8)	10 (2.0)
Diseases of the nervous system (G00-G99, I60-I63)	67 (53.6)	138 (27.6)
Morbus Parkinson or other extrapyramidal disorders (G20-G26)	14 (11.2)	5 (1.0)
Multiple sclerosis or other demyelinating diseases (G35-G37)	1 (0.8)	5 (1.0)
Migraine or other headache syndromes (G43-G44)	0	25 (5.0)
Guillain—Barré syndrome and other polyneuropathies (G61-G62)	1 (0.8)	3 (0.6)
Cerebral palsy and other paralytic syndromes (G80-G83)	1 (0.8)	9 (1.8)
Cerebral haemorrhage (I60-I62)	7 (5.6)	7 (1.4)
Cerebral infarction (I63)	33 (26.4)	60 (12.0)
Other diseases of the nervous system*	10 (8.0)	24 (4.8)
Diseases of the circulatory system (100-199, excl. 160-163)	14 (11.2)	79 (15.8)
Ischemic heart diseases (I20-I25)	4 (3.2)	21 (4.2)
Valvular heart diseases (105-108, 134-136)	1 (0.8)	11 (2.2)
Other forms of heart disease [†]	1 (0.8)	22 (4.4)
Peripheral artery disease (I73)	4 (3.2)	6 (1.2)
Lymphoedema or other noninfective disorders of lymphatic vessels (189)	1 (0.8)	18 (3.6)
Other diseases of the circulatory system [‡]	3 (2.4)	1 (0.2)
Diseases or injuries of the musculoskeletal system (M00-M99, S00-T98)	34 (27.2)	240 (48.0)
Coxarthrosis (M16)	0	23 (4.6)
Gonarthrosis (M17)	2 (1.6)	28 (5.6)
Arthrosis or other arthropathies (M18-M19)	0	5 (1.0)
Spondylopathies (M45-M49)	4 (3.2)	37 (7.4)
Other dorsopathies [§]	1 (0.8)	64 (12.8)
Myalgia or rheumatism (M79)	1 (0.8)	8 (1.6)
Osteopathies and chondropathies (M80-M94)	2 (1.6)	2 (0.4)
Intracranial injury (SO6)	10 (8.0)	4 (0.8)
Fracture of femur (S72)	4 (3.2)	15 (3.0)
Fracture of lower leg (S82)	0	8 (1.6)
Other fractures or injuries (S00-S99, excl. S06, S72, S82)	6 (4.8)	29 (5.8)
Complication of internal joint prosthesis (T84)	4 (3.2)	17 (3.4)
Other diseases	4 (3.2)	33 (6.6)

ICD-10, International Classification of Diseases, Tenth Revision.

*Meningitis and other neurologic inflammatory diseases (G00-G09); Atrophies primarily affecting the central nervous system (G10-G14); Nerve and plexus disorders (G50-G59); Myopathies (G72); Hydrocephalus (G91); or Cerebral cysts (G93).

[†]Endocarditis (I33, I38-I39); Dilated cardiomyopathy (I42); Arrhythmias (I49).

[‡]Aortic aneurysm or dissection (I71); Venous thromboembolism (I82).

⁸Cervicalgia (M50); Intervertebral disc disorders (M51); Sciatica (M54.3); Lumbago (M54.5). ^{II}Infections (A00-B99); Endocrine, nutritional and metabolic diseases (E00-E90); Diseases of the digestive system (K00-K93); or Diseases of the respiratory system (J00-J99).

Supplementary Table 4 Odds Ratios of Comorbidities Among cases With Incident Delirium and Matched Controls

Comorbidities (ICD-10 Codes)	Cases, n (%) ($n = 125$)	Controls, n (%) ($n = 500$)	OR (95% CI)	AOR (95% CI)*
Infectious and parasitic diseases (A0	D-B99)			
No [†]	105	470	1 ref.	1 ref.
Yes [‡]	20	30	3.06 (1.66-5.64)	2.29 (1.14-4.61)
Bacterial infectious diseases (B95-B9	6)			
No [†]	112	486	1 ref.	1 ref.
Yes‡	13	14	4.16 (1.88-9.21)	2.62 (1.06-6.49)
Endocrine, nutritional and metabolic	diseases (E00-E90)			
No [†]	58	274	1 ref.	1 ref.
Yes [‡]	67	226	1.41 (0.95-2.09)	1.11 (0.71-1.75)
Diabetes mellitus (E10-E14)			· · · ·	
No [†]	100	414	1 ref.	1 ref.
Yes‡	25	86	1.20 (0.73-1.98)	1.11 (0.63-1.94)
Vitamin D deficiency (E55)				
No [†]	109	461	1 ref.	1 ref.
Yes [‡]	16	39	1.75 (0.94-3.27)	1.55 (0.76-3.17)
Hypercholesterolemia (E78)				1.55 (0.70 5.17)
No [†]	106	436	1 ref.	1 ref.
Yes‡	19	64	1.23 (0.70-2.14)	0.91 (0.49-1.68)
Disorders of fluid, electrolyte and ac		5	1.25 (0.70 2.14)	0.51 (0.45-1.00)
No [†]	110	484	1 ref.	1 ref.
Yes [‡]	15	16	4.15 (1.99-8.67)	2.76 (1.19-6.38)
		10	4.15 (1.55-8.07)	2.70 (1.19-0.58)
Diseases of the nervous system (G00	30	262	1	1 == 6
No [†] Yes [‡]	30 95	263	1 ref.	1 ref.
		237	3.55 (2.27-5.56)	2.4 (1.36-4.24)
Extrapyramidal and movement disor		471	1 maf	1
No [†]	92	471	1 ref.	1 ref.
Yes [‡]	33	29	5.93 (3.42-10.29)	3.51 (1.89-6.52)
Parkinson disease (G20-G21)	101	100	4	1 5
No [†]	101	488	1 ref.	1 ref.
Yes [‡]	24	12	9.87 (4.76-20.46)	5.68 (2.54-12.68)
Epilepsy and recurrent seizures (G40	,	100		
No [†]	114	483	1 ref.	1 ref.
Yes [‡]	11	17	2.79 (1.26-6.15)	1.27 (0.53-3.04)
Sleep disorders (G47)				
No	118	459	1 ref.	1 ref.
Yes [‡]	7	41	0.66 (0.29-1.52)	0.56 (0.22-1.40)
Other diseases of the nervous system				
No	109	401	1 ref.	1 ref.
Yes [‡]	16	99	0.59 (0.34-1.05)	0.61 (0.32-1.16)
Diseases of the circulatory system (I	00-199, excl. 160-169)			
No [†]	24	200	1 ref.	1 ref.
Yes [‡]	101	300	2.82 (1.74-4.56)	1.18 (0.66-2.09)
Cerebrovascular diseases (I60-I69)				
No [†]	75	413	1 ref.	1 ref.
Yes‡	50	87	3.18 (2.07-4.87)	0.98 (0.57-1.70)
Subarachnoid or Intracerebral hemo	rrhage (I60-I62)			
No [†]	118	492	1 ref.	1 ref.
Yes [‡]	7	8	3.65 (1.30-10.28)	1.73 (0.55-5.51)
Cerebral infarction (I63)			. , ,	, , ,
No [†]	87	439	1 ref.	1 ref.
Yes‡	38	61	3.17 (1.99-5.06)	0.89 (0.5-1.60)

AOR, adjusted odds ratio; ICD-10, International Classification of Diseases, Tenth Revision; OR, odds ratio; ref. referent.

Controls were matched to cases on index date (± 1 month) and days between the admission date and the index date. All ORs were calculated with unconditional logistic regression and adjusted for matching factors. Main categories are depicted in bold.

*Adjusted on age, sex, and rehabilitation discipline (neurology/nonneurology).

[†]Defined as no-read *ICD-10* code record of the respective disorder within the claims data.

[‡]Defined as a read *ICD-10* code record of the respective disorder at admission.

Supplementary Table 5 Odds Ratios of Selected Drug Classes Among Cases With Incident Delirium and Matched Controls, by Users or Nonusers

Nonsers' 66 256 1 ref. 1 ref. 1 ref. Users' 59 244 0.94 (0.63-1.39) 1.4 (0.88-2.25) Nonsers' 34 70 2.30 (1.44-3.68) 2.11 (1.25-3.58) Nonsers' 15 473 1 ref. 1 ref. Nonsers' 15 473 1 ref. 1 ref. Nonsers' 105 436 1 ref. 1 ref. Nonsers' 105 436 1 ref. 1 ref. Nonsers' 105 436 1 ref. 1 ref. Users' 20 64 0.30 (0.75-2.24) 0.62 (0.28-1.35) Nonsers' 115 458 0.95 (0.64-1.95) 0.62 (0.28-1.36) Parating protp (801A8) 1 1 ref. 1 ref. 1 ref. Nonsers' 54 122 0.95 (0.64-1.95) 0.62 (0.28-1.36) Nonsers' 50 1 ref. 1 ref. 1 ref. Users' 50 1 ref. 1 ref. 1 ref. 1 ref.	Drug Classes (ATC Codes)	Cases, n (%) ($n = 125$)	Controls, n (%) ($n = 500$)	OR (95% CI)	AOR (95% CI)*
Users' hor	Proton pump inhibitors (A02BC)				
bases bases <th< td=""><td>Nonusers[†]</td><td>66</td><td>256</td><td>1 ref.</td><td>1 ref.</td></th<>	Nonusers [†]	66	256	1 ref.	1 ref.
Nonsers 1 1 1 ref. 1 ref. 1 ref. Lines' 10 20 1.53 1.67		59	244	0.94 (0.63-1.39)	1.4 (0.88-2.25
Useri3470230 (144-368)211 (125-38)Nonuseri154731 ef.1 ef.Nonuseri154731 ef.1 ef.Nonuseri0261 ef.1 ef.Nonuseri0461 ef.1 ef.Nonuseri15481 ef.1 ef.Nonuseri15481 ef.1 ef.Nonuseri15481 ef.1 ef.Nonuseri17781 ef.2 (202-13)Nonuseri71781 ef.1 ef.Useri541 ef.1 ef.1 ef.Nonuseri75331 ef.1 ef.Nonuseri75331 ef.1 ef.Nonuseri75391 ef.1 ef.Nonuseri75391 ef.1 ef.Nonuseri75391 ef.1 ef.Nonuseri71191 ef.1 ef.Nonuseri73391 ef.1 ef.Nonuseri7339<					
nonumers' 10 (11) 15 473 14 (12) 13 (12) 14 (10) 12 (14) 13 (12) 14 (10) 12 (14) 13 (12) 14 (10) 12 (14) 14 (10) 12 (14) 14 (1					
Nonuserie 115 473 1 ref. 1 ref. 1 ref. lood glucos lovering drugs ext, lusuillins (ADD) Nonuserie 1 add (ADD 24) 1 Add (ADD		34	70	2.30 (1.44-3.68)	2.11 (1.25-3.58
Users' 10 27 15.1 (0.72-3.27) 14.4 (0.02-3.34) Nonusers' 105 436 1 ef 1 ef Nonusers' 105 436 1 ef 1 ef Nonusers' 10 436 1 eff 1 eff Nonusers' 10 430 0.55 (0.461-155) 0.62 (0.281-135) Nonusers' 71 737 1 eff 1 eff 1 eff Users' 54 122 236 (1.572-56) 244 (1.292-245) Nonusers' 71 737 1 eff 1 eff 1 eff Users' 54 122 236 (1.572-56) 244 (1.292-245) Nonusers' 75 331 1 eff 1 eff 1 eff Users' 90 310 1 247 (0.72-24) 1 25 (0.79-240) 1 26 (0.79-240) 1 26 (0.79-240) 1 26 (0.79-240) 1 26 (0.79-240) 1 26 (0.79-240) 1 26 (0.79-240) 1 26 (0.79-240) 1 26 (0.79-240) 1 26 (0.79-240) 1 eff 1 eff 1 eff 1 eff 1 eff 1 eff		115	472	1	1
block place lowering drugs, excl. Insulins (ADD) Insunsers' 105 476 1 ref. Liters' 20 64 Liters' 20 64 Liters' 20 64 Liters' 20 64 Liters' 20 75 64 Liters' 20 75 64 Liters' 100 ADD Liters' 71 75 78 Nonusers' 71 77 78 Liters' 75 78 Li					
Nomiser's 105 436 1 ref. 1 ref. 1 ref. User's 20 64 130 (075-24) 107 (035-20) 0.62 (028-1.93) Nemuser's 115 458 1 ref. 1 ref. 1 ref. User's 54 122 236 (157-356) 204 (129-324) Water's 71 378 1 ref. 1 ref. 1 ref. User's 54 122 236 (157-356) 204 (129-324) Nomiser's 73 333 1 ref. 1 ref. 1 ref. Nomiser's 73 339 2 18 (137-351) 1 25 (07-204) 1 25 (07-204) Nomiser's 9 319 2 13 (137-351) 1 26 (07-204)			21	1.55 (0.72-5.27)	1.44 (0.02-5.54
Utersi 20 64 1.30 (0.75-224) 1.07 (0.55-204) Nonusersi 115 458 1 ref. 1 ref. Nonusersi 115 458 1 ref. 1 ref. Paratir group (107/AB) 71 78 1 ref. 1 ref. Usersi 54 1 22 236 (157-350) 232 (0.72-204) Vaceri 55 33 1 ref. 1 ref. 1 ref. Usersi 50 1 47 1 61 (107-242) 1 25 (0.73-250) Vaceri daro X athibitors (201/A) 72 1 20 (0.73-351) 1 28 (0.74-220) 1 23 (0.73-220) Vaceri daro X athibitors (201/A) 74 1 20 (0.73-351) 1 28 (0.74-220) 1 28 (0.74-220) Vaceri daro X athibitors (201/A) 74 1 20 (0.73-351) 1 28 (0.74-20) 1 28 (0.74-20) Vaceri daro X athibitors (201/A) 74 1 28 (0.72-106) 1 28 (0.74-20) 1 28 (0.72-20) Vaceri daro X athibitors (201/A) 74 1 28 (0.72-20) 1 28 (0.72-20) 1 28 (0.72-20) Vaceri daro X athibitors (201/A) 74 <td< td=""><td></td><td>, ,</td><td>436</td><td>1 ref</td><td>1 ref</td></td<>		, ,	436	1 ref	1 ref
Itamin Kanagonists (B01AA) Iref. I					
Users' 10 42 0.52 (0.45-1.95) 0.62 (0.23-1.32) Nonuscri 71 378 1 r.f. 1 r.f. Nonuscri 71 378 1 r.f. 1 r.f. Nonuscri 75 337 1 r.f. 1 r.f. 1 r.f. Nonuscri 75 337 1 r.f.	/itamin K antagonists (B01AA)				
leparts group (B01AB) Nonusers' 71 378 72 236 (157.356) 204 (129.24 Meeter agregation inhibitors, exc. heparin (B01AC) Nonusers' 75 373 373 1ref. 1 (Leers' 75 100 174) Nonusers' 75 373 1ref. 1 (Leers' 75 100 174) Nonusers' 75 373 1ref. 1 (Leers' 75 100 174) Nonusers' 75 75 75 75 174 Nonusers' 75 75 75 75 75 174 175 75 175 175 175 175 175 175 175 175		115	458	1 ref.	1 ref.
Nonusers 1 278 1 ref. 1 ref. 1 ref. 1 ref. 1 ref. Mattel aggregation inhibitors, sext. beparin (001AC) 333 1 ref. 1 ref. 1 ref. Nonusers' 50 147 1 (1072-42) 1 25 (0.73-20) Nonusers' 26 181 1 ref. 1 ref. Nonusers' 29 19 219 (137-351) 1 28 (0.74-223) Area in thing (C01) 16 25 1 ref. 1 ref. Nonusers' 9 306 1 ref. 1 ref. Nonusers' 19 25 1 48 (007-325) 1 39 (0.83-232) Nonusers' 73 39 1 ref. 1 ref. 1 ref. Users' 73 39 1 ref. 1 ref. 1 ref. Nonusers' 73 39 1 ref. 1 ref. 1 ref. Vers' 74 1 82 (121-23) 1 52 (0.96-2.35) 1 52 (0.96-2.35) 1 52 (0.96-2.35) Jation fing regents (C07) 10 1 ref. 1 ref. <	Users [‡]	10	42	0.95 (0.46-1.95)	0.62 (0.28-1.39
Users'5412223 (157-3.6)24 (1.20-3.24)Nonusers'75331.ef.1.ef.1.ef.Nonusers'501471.61 (107-2.42)1.25 (107-3.05)Nonusers'9311.ef.1.ef.1.ef.Users'93191.91 (137-3.51)1.28 (0.74-2.05)Nonusers'1.664751.ef.1.ef.1.ef.Users'9301.81 (137-3.51)1.28 (0.74-2.05)Nonusers'1.664751.ef.1.ef.1.ef.Nonusers'92.61.48 (0.57-3.35)1.93 (0.53-2.35)Nonusers'85301.ef.1.93 (0.53-2.35)Nonusers'733591.ef.1.ef.1.ef.Nonusers'9.94.261.ef.1.ef.1.ef.Users'7.11.35 (0.92-2.49)0.74 (0.42-1.31)1.26 (0.72-1.86)Alcian change blocker (0.08)1.11.ef.1.ef.1.ef.Users'7.11.35 (0.92-2.49)0.74 (0.42-1.31)1.26 (0.72-1.86)Alcian change blocker (0.08)1.11.ef.1.ef.1.ef.Users'7.11.35 (0.92-2.49)0.74 (0.42-1.31)1.26 (0.72-1.86)Alcian change blocker (0.08)1.11.ef.1.ef.1.ef.Users'7.11.36 (0.72-1.86)1.55 (0.82-2.49)1.55 (0.82-2.49)1.75 (0.82-2.49)Nonusers'1.11.ef.1.ef.1.ef.1.ef.1.ef.Users'7.11.36	Heparin group (B01AB)				
Tablet agregation inhibitors, excl. heparin (001AC) 1 ref. 1 ref. <td>Nonusers†</td> <td>71</td> <td>378</td> <td>1 ref.</td> <td>1 ref.</td>	Nonusers†	71	378	1 ref.	1 ref.
Names 75 253 1 ref. 1 ref. <th1 ref.<="" th=""> 1 ref. 1 ref.</th1>	Users‡	54	122	2.36 (1.57-3.56)	2.04 (1.29-3.24
Users' 50 147 161 (107-242) 125 (0.79-200 Nonusers' 26 181 1 ref. 1 ref. <td>Platelet aggregation inhibitors, exc</td> <td>cl. heparin (B01AC)</td> <td></td> <td></td> <td></td>	Platelet aggregation inhibitors, exc	cl. heparin (B01AC)			
Direct factor Xa inhibitors (801AF) Description Description Nonusers' 99 319 2.19 (1.37-3.51) 1.28 (0.74-2.23 Nonusers' 116 475 1ref. 1 ref. 1 ref. Nonusers' 9 25 1.48 (0.67-3.25) 1.37 (0.58-3.25 Nonusers' 85 396 1 ref. 1 ref. 1 ref. Users' 40 1.81 (1.17-2.80) 1.39 (0.58-3.25 Nonusers' 73 359 1 ref. 1 ref. Users' 40 1.81 (1.17-2.80) 1.39 (0.58-3.25 Nonusers' 52 141 1.82 (1.21-2.73) 1.16 (0.72-1.88 Nonusers' 54 311 1 ref. 1 ref. 1 ref. Users' 26 74 1.51 (0.92-2.49) 0.74 (0.42-1.31 Users' 26 74 1.52 (0.95-2.30) 1.76 (0.51-53) Nonusers' 18 465 1 ref. 1 ref. 1 ref. Nonusers' 18 465 1 ref. 1 ref.		75	353	1 ref.	1 ref.
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be a blocking agents (C07) Nonusers' 73 59 1 ref. 1 ref. Users' 52 141 182 (1.21-2.73) 1.16 (0.72-1.86 Jacum channel blockers (C08) Users' 26 74 151 (0.02-2.49) 0.74 (0.42-1.31 Users' 26 74 151 (0.02-2.49) 0.74 (0.42-1.31 Users' 54 311 1 ref. 1 ref. Users' 71 189 2.18 (1.46-3.25) 1.52 (0.96-2.39 ipdi modifying agents (C10) Nonusers' 63 221 1 ref. 1 ref. Users' 62 179 1.78 (1.19-2.65) 1 (0.63-1.58) Tygs for urinary frequency and incontinence (C04BD) Nonusers' 64 321 1 ref. 1 ref. Users' 62 179 1.78 (1.19-2.65) 1 (0.63-1.58) Tygs for urinary frequency and incontinence (C04BD) Nonusers' 118 486 1 ref. 1 ref. Users' 1 10 470 1 ref. 1 ref. Users' 1 10 470 1 ref. 1 ref. Nonusers' 110 470 1 ref. 1 ref. Nonusers' 15 30 2.14 (1.11-4.13) 0.97 (0.45-2.04 Viers' 6 27 0.88 (0.36-2.19) 0.69 (0.26-1.85 Nonusers' 1 18 463 1 ref. 1 ref. Nonusers' 1 18 7 7 37 0.74 (0.32-1.71) 0.7 (0.28-1.75 Nonusers' 1 18 463 1 ref. 1 ref. Nonusers' 1 18 7 7 0.93 2.17 (0.93-9.20 Nonusers' 1 18 7 7 37 0.74 (0.32-1.71) 0.7 (0.28-1.75 Nonusers' 1 18 463 1 ref. 1 ref. Users' 7 7 37 0.74 (0.32-1.71) 0.7 (0.28-1.75 Nonusers' 1 18 7 481 1 ref. 1 ref. Nonusers' 1 17 481 1 ref. 1 ref. Users' 8 1 17 0.93 2.2 2.2 2.3 10 0.86 (0.45-1.03) 0.86 (0.51-1.33) Nonusers' 1 17 481 1 ref. 1 ref. Users' 2 2 2.2 2.3 10 0.86 (0.51-1.03) 0.86 (
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Alcium channel blockers (C08)Nonusers'994261 ref.1 ref.Nonusers'26741.51 (0.92-2.49)0.74 (0.42-1.31gents acting on the renin-angiotensin system (C09)					
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Nonusers 105 466 1 ref. 1 ref. Users 20 34 2.65 (1.46-4.82) 1.97 (0.99-3.92 Intigout preparations (M04)			37	0.74 (0.32-1.71)	0.7 (0.28-1.75
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Users ¹ 43 217 0.68 (0.45-1.03) 0.86 (0.53-1.39 Opioid drugs (N02A) 7 1 ref. 1 ref. 1 ref. Nonusers ¹ 103 397 1 ref. 1 ref. 1 ref. Users ¹ 22 103 0.82 (0.50-1.37) 1.22 (0.68-2.17) Other analgesic and antipyretics (N02B) 7 7 0.88 (0.47-1.30) Nonusers ¹ 92 333 1 ref. 1 ref. Users ¹ 33 167 0.71 (0.46-1.11) 0.78 (0.47-1.30) Intiepileptic drugs (N03) 7 80 1 ref. 1 ref.		8 2	202	1 rof	1 ref
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Nonusers 103 397 1 ref. 1 ref. Users 22 103 0.82 (0.50-1.37) 1.22 (0.68-2.17) ther analgesic and antipyretics (N02B) 333 1 ref. 1 ref. Nonusers [†] 92 333 1 ref. 1 ref. Users [‡] 33 167 0.78 (0.47-1.30) Intiepileptic drugs (N03) Nonusers [†] 98 420 1 ref. 1 ref. Nonusers [†] 98 420 1 ref. 1 ref. 1.32 (0.75-2.34) Users [±] 27 80 1.46 (0.89-2.38) 1.32 (0.75-2.34)		45	217	0.00 (0.45-1.03)	0.80 (0.53-1.39
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Nonusers 92 333 1 ref. 1 ref. Users 33 167 0.71 (0.46-1.11) 0.78 (0.47-1.30) Interpileptic drugs (N03)			103	0.02 (0.30-1.37)	1.22 (0.08-2.17
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Nonusers [†] 98 420 1 ref. 1 ref. Users [‡] 27 80 1.46 (0.89-2.38) 1.32 (0.75-2.34)			107	0.71 (0.40-1.11)	0.70 (0.47-1.30
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	Nonusers			1 101.	1 1 1 1,
				1 46 (0 89-2 38)	

Supplementary Table 5 (continued)

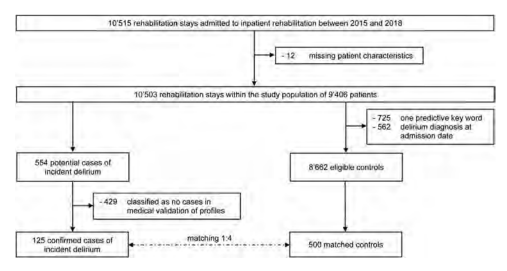
Drug Classes (ATC Codes)	Cases, n (%) ($n = 125$)	Controls, n (%) ($n = 500$)	OR (95% CI)	AOR (95% CI)*
Nonusers [†]	102	477	1 ref.	1 ref.
Users‡	23	23	4.70 (2.54-8.72)	2.86 (1.42-5.77)
Antipsychotic drugs (N05A)				
Nonusers [†]	86	471	1 ref.	1 ref.
Users‡	39	29	7.46 (4.37-12.74)	8.06 (4.26-15.22
Benzodiazepine derivatives (N05)	BA)			
Nonusers†	119	475	1 ref.	1 ref.
Users‡	6	25	0.96 (0.38-2.39)	1.18 (0.42-3.30)
Hypnotics and sedatives (N05C)				
Nonusers [†]	117	468	1 ref.	1 ref.
Users [‡]	8	32	1.00 (0.45-2.23)	0.95 (0.39-2.31)
Antidepressant drugs (N06A)				
Nonusers [†]	81	367	1 ref.	1 ref.
Users [‡]	44	133	1.52 (0.99-2.31)	1.88 (1.14-3.10)
Dementia drugs (N06D)				
Nonusers [†]	114	493	1 ref.	1 ref.
Users [‡]	11	7	7.02 (2.64-18.69)	2.59 (0.90-7.47)
Other nervous system drugs (N07	7)			
Nonusers [†]	119	483	1 ref.	1 ref.
Users [‡]	6	17	1.44 (0.55-3.73)	1.74 (0.57-5.30)
Drugs for obstructive airways dis	eases (R03)			
Nonusers [†]	114	468	1 ref.	1 ref.
Users [‡]	11	32	1.41 (0.69-2.89)	0.84 (0.37-1.90)

AOR, adjusted odds ratio; ATC, Anatomical Therapeutic Chemical; OR, odds ratio.

Controls were matched to cases on index date (±1 month) and days between the admission date and the index date. All ORs were calculated with unconditional logistic regression and adjusted for matching factors.

*Adjusted on age, sex, rehabilitation discipline (neurology/nonneurology). *Defined as no administration at any time prior the index date.

[‡]Defined as at least 1 administration at any time from the admission date until the index date.



Supplementary Figure 1. Flowchart of case and control selection. Cases were patients with at least 2 recorded delirium predictive keywords (commonly used terms to describe delirious patients) who were classified as incident delirium episodes by 2 to 3 independent physicians as defined in a previous validation study.³⁰ Eligible controls were patients in the study population who did not have any record of delirium predictive keywords in their medical notes or a diagnosis of prevalent delirium on admission. Each case was matched to 4 controls on calendar time [by assigning the index date (±1 month) of the cases to their controls] and time between admission date and index date.