Health resource utilization and cost before versus after initiation of second-generation long-acting injectable antipsychotics among adults with schizophrenia in Alberta, Canada: a retrospective, observational single-arm study

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Research Article

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Abstract

Background

Long-acting injectable (LAI) antipsychotics, along with community treatment orders (CTOs), are used to improve treatment effectiveness through adherence among individuals with schizophrenia. Understanding real-world medication adherence, and healthcare resource utilization (HRU) and costs in individuals with schizophrenia overall and by CTO status before and after second generation antipsychotic (SGA)-LAI initiation may guide strategies to optimize treatment among those with schizophrenia.

Methods

This retrospective observational single-arm study utilized administrative health data from Alberta, Canada. Adults (≥18 years) with schizophrenia who initiated a SGA-LAI (no use in the previous 2-years) between April 1, 2014 and March 31, 2016, and had ≥1 additional dispensation of a SGA-LAI were included; index date was the date of SGA-LAI initiation. Medication possession ratio (MPR) was determined, and paired t-tests were used to examine mean differences in all-cause and mental health-related HRU and costs (Canadian dollars), comprised of hospitalizations, physician visits, emergency department visits, and total visits, over the 2-year post-index and 2-year pre-index periods. Analyses were stratified by presence or absence of an active CTO during the pre-index and/or post-index periods.

Results

Among 1,211 adults with schizophrenia who initiated SGA-LAIs, 64% were males with a mean age of 38 (standard deviation [SD] 14) years. The mean overall antipsychotic MPR was 0.39 (95% confidence interval [CI] 0.36, 0.41) greater during the 2-year post-index period (0.84 [SD 0.26]) compared with the 2-year pre-index period (0.45 [SD 0.40]). All-cause and mental health-related HRU and costs were lower post-index versus pre-index (p<0.001) for hospitalizations, physician visits, emergency department visits, and total visits; mean total all-cause HRU costs were $33,788 (95% CI -$38,993, -$28,583) lower post- versus pre-index ($40,343 [SD $68,887] versus $74,131 [SD $75,941]), and total mental health-related HRU costs were $34,198 (95% CI -$39,098, -$29,297) lower post- versus pre-index ($34,205 [SD $63,428] versus $68,403 [SD $72,088]) per-patient. Forty-three percent had ≥1 active CTO during the study period; HRU and costs varied according to CTO status.

Conclusions

SGA-LAIs are associated with greater medication adherence, and lower HRU and costs however the latter vary according to CTO status.

1. Background

Schizophrenia is a serious chronic mental illness affecting approximately 0.3–1.0% of the global population [1, 2]. Although prevalence is relatively low, the burden of disease is substantial [2]. In Europe, the annual direct healthcare cost per patient has been reported to range from €533 to €13,704, with inpatient hospitalizations representing the largest proportion of costs in the majority of countries [3]. The total annual economic burden attributed to schizophrenia in Canada was estimated at $6.85 billion CDN in 2004 [1].
Pharmacotherapy with antipsychotics is considered to be the cornerstone of symptom treatment and relapse prevention among individuals with schizophrenia [4–6]. However, nonadherence to antipsychotics is common and associated with worsening functional outcomes, higher admission to the hospital and emergency department (ED), and higher rates of relapse and suicide [7–9]. Conversely, adherence to antipsychotics has been found to be associated with better long-term clinical outcomes [7]. Long-acting injectable (LAI) formulations of first- and second-generation antipsychotics (FGA and SGA, respectively) were developed with the goal of improving medication adherence [10, 11]. Clinical guidelines recommend LAIs over oral formulations of antipsychotics for individuals who display nonadherence, have frequent relapses, or pose a safety risk to others [6, 12, 13]. Although guidelines also recommend LAI antipsychotics be offered as an option in line with personal preference, it is noted that certain circumstances may not allow for this option, such as in the case of community treatment orders (CTOs) [6]. CTOs are available for use in over 75 jurisdictions worldwide; admissibility criteria and provisions of care can vary. In Alberta, a CTO is intended to assist with treatment adherence, including medications and attending health care appointments and other community support services, in those with mental illness who would otherwise decompensate, likely cause harm to themselves or others, or suffer substantial deterioration or impairment in the absence of continued treatment.

While CTOs are a common component of clinical practice in the treatment of schizophrenia, few studies investigating the effects of LAI antipsychotics on healthcare resource utilization (HRU) and associated costs have incorporated CTO information [14]. Additionally, it has been suggested that nonadherence and subsequent associated consequences may take time to develop, and that a 2-year observation period may result in a markedly higher likelihood of detecting a difference compared with a 1-year period [15]. Therefore, the aim of this observational study was to examine and compare real-world antipsychotic medication adherence, and HRU and associated costs between the 2-year period before and after SGA-LAI initiation among adults with schizophrenia; subgroup analysis was performed based on CTO status.

2. Methods

This retrospective, non-interventional, single arm study is reported according to the STROBE guidelines [16]. Ethics approval (Pro00086993) was obtained from the institutional review board at the University of Alberta, Canada. This is a study of administrative data without any intervention. No study participants were placed at risk as a result of this study, and informed consent was not required.

2.1 Data source

Administrative data from the Discharge Abstract Database (DAD), National Ambulatory Care Reporting System (NACRS), Practitioner Claims, and Pharmaceutical Information Network (PIN) were used. CTO data was obtained from Alberta Health Services. Data were linked to the Population Registry, which contains demographic information for all Albertans with Alberta Health Care Insurance Plan (AHCIP) coverage; all Alberta residents are eligible for AHCIP and over 99% participate [17]. The DAD includes demographic, administrative, diagnostic, procedural, and resource intensity weighting (RIW) information on all patients discharged from hospital; diagnostic data contains a most responsible diagnosis and up to 24 secondary diagnostic codes using the International Classification of Disease - Version 10 - Canadian Enhancement (ICD-10-CA) codes. NACRS includes patient and RIW information in facility-based ambulatory care centers, including the ED; ICD-10-CA codes are used and contains a most responsible diagnosis field and up to 9 secondary diagnostic codes. Practitioner Claims includes patient, provider, service and direct billing information on fee-for-service, alternative payment plan physician billing and shadow billing; up to 3 ICD - Version 9 - Clinical Modification (ICD-9-CM; Alberta specific) diagnostic codes can be used per visit. PIN contains information on dispensed prescription medications from community pharmacies.
2.2 Cohort selection

The population of interest were adults with schizophrenia who initiated SGA-LAIs in Alberta within the inclusion period from April 1, 2014 to March 31, 2016. Eligibility criteria included: 1) ≥18-years and initiated a SGA-LAI (≥1 dispensation within no use in the previous 2-years) during the inclusion period (index date was the first date of dispensation of either aripiprazole, paliperidone, or risperidone, which were the three approved SGA-LAIs in Alberta during this timeframe), 2) ≥1 additional SGA-LAI dispensation during the 2-year post-index period, 3) met the case definition for schizophrenia within 9 years prior to the index date [18], 4) had AHCIP coverage over the study period (between April 1, 2014 and March 31, 2018). The following case algorithm was used to identify schizophrenia: ≥1 hospitalization or ≥3 physician visits with a diagnostic code for schizophrenia and/or related condition within a 3-year period [19]; the specific diagnostic codes used (see Additional file 1) were based on previous case validation studies and input from clinical experts [18, 19].

CTO status was defined by the presence or absence of ≥1 active CTO during the 2-year pre-index and/or post-index periods. The resultant four CTO categories included individuals: 1) without any active CTOs in both the pre- and post-index periods (pre=no / post=no), 2) with ≥1 active CTO in both the pre- and post-index periods (pre=yes / post=yes), 3) with ≥1 active CTO during the pre-index, but not the post-index period (pre=yes / post=no), and 4) with no active CTO during the pre-index period, but ≥1 active CTO during the post-index period (pre=no / post=yes).

2.3 Study measures

Study measures were presented for the overall cohort and by CTO status. Baseline characteristics on the index date included age, sex, urban or rural residence (determined by the second digit of the postal code), and the SGA-LAI that was initiated. A Charlson Comorbidity Index (CCI) score was determined during the 2-year period before the index date; the score was based on the presence or absence of 17 individual comorbid conditions that were weighted according to their potential for influencing mortality [20].

Study outcomes included antipsychotic medication adherence, all-cause and mental health-related HRU, and associated healthcare costs that were measured and compared between the 2-year pre-index and post-index periods for all subjects within the overall cohort and the CTO subgroups. Within the overall cohort, the number of individuals who received ≥1 dispensation for an antipsychotic medication was determined, and medication possession ratio (MPR) was calculated based on the number of antipsychotic medication days of supply during the 2-year pre- and post-index periods, and presented overall, as well as according to formulation (oral or injectable) and generation (first or second; see Additional file 2 for a complete listing of antipsychotic medications that were included). HRU included all-cause and mental health-related (ICD-9 290-319 from any diagnostic field; ICD-10 F01-F99 from the most responsible diagnostic field) hospitalizations, physician visits, ED visits, and total visits [21, 22]. Multiple physician visits per day per person were considered as one visit per day per person. The cost for each hospitalization and ED visit was determined by multiplying the RIW value of the visit with the 2016/2017 Canadian Institute for Health Information cost of a standard hospital stay (CSHS) for Alberta [23]. RIW is a measure to estimate HRU and represents the relative value of resources that a given patient, contingent on diagnostic case-mix, would be expected to consume relative to a standard patient; CSHS provides standardized average costs incurred through the direct care of a standard hospital or ambulatory care visit. Direct billing costs were used for physician visits. Costs are presented in Canadian dollars ($CDN) and reflect healthcare costs incurred over the study period.

2.4 Statistical analyses
Descriptive statistics are reported as means and standard deviations (SD) or counts and percentages, where appropriate. Paired t-tests were used to test for statistically significant differences between the pre- and post-index periods for HRU and associated costs; costs were also compared using Wilcoxon signed-rank tests and medians and interquartile ranges (IQR) were reported. A conventional alpha of 0.05 and a two-tailed level of significance were used; accompanying 95% confidence intervals (CI) are reported. Statistical analyses were performed using Python version 3.6.5 and STATA version 17.0.

3. Results

3.1 Cohort selection

Of the 126,162 individuals who received an antipsychotic dispensation during the inclusion period, 1,211 met eligibility criteria (Figure 1). Within the cohort, 43% received at least one active CTO during the study period, among whom the majority (80%) persisted with the same CTO status throughout the pre- and post-index periods (pre=no / post=no [n=689; 57%]; pre=yes / post=yes [n=275; 23%]), and 20% had a CTO status change (pre=yes / post=no [n=133; 11%]; pre=no / post=yes [n=114; 9%]).

3.2 Characteristics

Table 1 details the baseline subject characteristics of the overall cohort and by CTO status. Overall, the mean age of individuals with schizophrenia was 38 (SD 14) years, there were twice as many males (64.3%) as females (35.7%), 89.0% lived in urban areas, and the overall mean CCI score was 0.6 (SD 1.3). Compared with the other CTO cohorts, the pre=no / post=yes cohort had a higher proportion of males (71.9%) and were younger in age (34 years [SD 14]), with almost one third of individuals between 18 and 24 years of age (32.5%). On the index date, the majority of individuals initiated paliperidone (67.6% - 71.4%), followed by aripiprazole (16.7% - 21.1%), followed by risperidone (11.3% - 13.4%).

Table 1. Subject characteristics presented for the overall cohort and by CTO status.
| Age, years | Overall Cohort (n=1211; 100%) | CTO status | pre=no/post=no (n=689; 57%) | pre=yes/post=yes (n=275; 23%) | pre=yes/post=no (n=133; 11%) | pre=no/post=yes (n=114; 9%) |
|------------|-------------------------------|-------------|-------------------------------|--------------------------------|-------------------------------|-------------------------------|
| Mean (SD)  | 38 (14)                       | 39 (14)     | 38 (14)                       | 38 (12)                        | 34 (14)                       |
| Categories, n (%) |                        |             |                               |                                |                               |
| 18-24      | 239 (19.7%)                   | 134 (19.5%) | 44 (16.0%)                    | 24 (18.1%)                     | 37 (32.5%)                    |
| 25-29      | 196 (16.2%)                   | 101 (14.7%) | 58 (21.1%)                    | 18 (13.5%)                     | 19 (16.7%)                    |
| 30-34      | 150 (12.4%)                   | 95 (13.8%)  | 27 (9.8%)                     | 14 (10.5%)                     | 14 (12.3%)                    |
| 35-39      | 132 (10.9%)                   | 64 (9.3%)   | 43 (15.6%)                    | 15 (11.3%)                     | 10 (8.8%)                     |
| 40-44      | 124 (10.2%)                   | 64 (9.3%)   | 26 (9.5%)                     | 24 (18.1%)                     | 10 (8.8%)                     |
| 45-49      | 105 (8.7%)                    | 66 (9.6%)   | 17 (6.2%)                     | 15 (11.3%)                     | <10 (<8.8%)                   |
| 50-54      | 101 (8.3%)                    | 63 (9.1%)   | 22 (8.0%)                     | 12 (9.0%)                      | <10 (<8.8%)                   |
| 55-59      | 69 (5.7%)                     | 43 (6.2%)   | 15 (5.5%)                     | <10 (<7.5%)                    | <10 (<8.8%)                   |
| 60-64      | 40 (3.3%)                     | 25 (3.6%)   | <15 (<5.5%)                   | <10 (<7.5%)                    | <10 (<8.8%)                   |
| ≥65        | 55 (4.5%)                     | 34 (4.9%)   | <15 (<5.5%)                   | <10 (<7.5%)                    | <10 (<8.8%)                   |

| Sex, n (%) |                        |             |                               |                                |                               |
| Male       | 779 (64.3%)               | 436 (63.3%) | 177 (64.4%)                   | 84 (63.2%)                     | 82 (71.9%)                    |
| Female     | 432 (35.7%)               | 253 (36.7%) | 98 (35.6%)                    | 49 (36.8%)                     | 32 (28.1%)                    |

| Residence, n (%) |                        |             |                               |                                |                               |
| Urban       | 1078 (89.0%)             | 608 (88.2%) | 252 (91.6%)                   | 118 (88.7%)                    | 100 (87.7%)                   |
| Rural       | 133 (11.0%)              | 81 (11.8%)  | 23 (8.4%)                     | 15 (11.3%)                     | 14 (12.3%)                    |

| CCI, mean (SD) | 0.6 (1.3) | 0.7 (1.4) | 0.5 (1.0) | 0.6 (1.3) | 0.6 (1.0) |
|---------------|-----------|-----------|-----------|-----------|-----------|

| Index SGA LAI, n (%) |                        |             |                               |                                |                               |
| Paliperidone     | 833 (68.8%)           | 471 (68.4%) | 186 (67.6%)                   | 95 (71.4%)                     | 81 (71.1%)                    |
| Aripiprazole     | 229 (18.8%)           | 126 (18.3%) | 58 (21.1%)                    | 26 (19.5%)                     | 19 (16.7%)                    |
| Risperidone      | 149 (12.3%)           | 92 (13.4%)  | 31 (11.3%)                    | 14 (11.3%)                     | 14 (12.3%)                    |
Abbreviations: CCI = Charlson comorbidity index; CTO = community treatment order; SD = standard deviation; SGA LAI = second generation antipsychotic long-acting injectable. Subject numbers less than 10 need to be redacted as per data privacy standards, and prevented from being back-calculated.

3.3 Antipsychotic medication use and adherence

Table 2 details the antipsychotic medication use and MPR of the overall cohort during the 2-year pre-index and post-index periods. In addition to the study-defined eligibility criteria of ≥1 SGA-LAI dispensation during the post-index period and none during the pre-index period, SGA oral medications were dispensed to 79.8% of the overall cohort during the pre-index period and 69.5% during the post-index period. Respectively, FGA oral and LAIs were dispensed to 10.7% and 15.9% of the overall cohort during the pre-index period, and 9.9% and 5.5% during the post-index period. The overall mean antipsychotic MPR was 0.39 (95% confidence interval [CI] 0.36, 0.41) greater during the 2-year post-index period (0.84 [SD 0.26]) compared with the 2-year pre-index period (0.45 [SD 0.40]). Among the oral antipsychotics, the MPR of FGA and SGA medications were not significantly different between the pre- and post-index periods. Although the MPR of FGA-LAI was low during the study period (0.07 [SD 0.20] pre-index; 0.02 [SD 0.11] post-index), it was 0.05 (95% CI -0.060, -0.038) less in the post-index period. The MPR of SGA-LAI was 0.64 (SD 0.33) during the post-index period.

Table 2: Antipsychotic medication use and mean medication possession ratio of the overall cohort during the pre-index and post-index periods.

|                        | Pre-index Period (n=1,211; 100%) | Post-index Period (n=1,211; 100%) | Mean difference [95% CI] |
|------------------------|----------------------------------|----------------------------------|--------------------------|
| Received ≥1 dispensation, n (%) |                              |                                  |                          |
| Oral                   |                                  |                                  |                          |
| First generation       | 130 (10.7%)                      | 120 (9.9%)                       |                          |
| Second generation      | 966 (79.8%)                      | 841 (69.5%)                      |                          |
| Long acting-injectable |                                  |                                  |                          |
| First generation       | 192 (15.9%)                      | 67 (5.5%)                        |                          |
| Second generation      | N/A (N/A)                        | 1,211 (100%)                     |                          |
| Medication possession ratio, mean (SD); mean difference [95%CI] |                                  |                                  |                          |
| Overall                | 0.45 (0.40)                      | 0.84 (0.26)                      | **0.39 [0.36, 0.41]**   |
| Oral                   |                                  |                                  |                          |
| First generation       | 0.03 (0.14)                      | 0.03 (0.13)                      | -0.0046 [-0.014, 0.0043]|
| Second generation      | 0.40 (0.39)                      | 0.38 (0.41)                      | -0.019 [-0.044, 0.0065] |
| Long acting-injectable |                                  |                                  |                          |
| First generation       | 0.07 (0.20)                      | 0.02 (0.11)                      | **-0.05 [-0.060, -0.038]** |
| Second generation      | N/A (N/A)                        | 0.64 (0.33)                      | N/A                      |
**Bolded** mean difference indicates statistically significant difference (p<0.001) between the 2-year post- and 2-year pre-index using paired t-tests. Abbreviations: CI = confidence interval; N/A = not applicable; SD = standard deviation.

### 3.4 Healthcare resource utilization

Figure 2 show the mean per patient number and comparative differences of all-cause and mental health-related total visits, as well as hospitalizations, physician visits, and ED visits for the overall cohort between the pre-index and post-index periods. During the study period, all-cause physician visits were most common (97.8 [SD 75.5] pre-index, 74.7 [SD 69.1] post-index), followed by ED visits (5.1 [SD 7.5] pre-index, 3.6 [SD 8.5] post-index), and hospitalizations (2.5 [SD 2.3] pre-index, 1.2 [SD 2.0] post-index) within the overall cohort. The mean number of total all-cause HRU visits was significantly lower by 23.6% (74.7 [SD 69.1] versus 97.8 [SD 75.5] visits; mean difference -23.1 [95% CI -27.7, -18.5]) in the post-index period compared with the pre-index period. The mean difference in specific types of HRU visits between the post-index and the pre-index periods was also significantly lower for hospitalizations (-1.3 [95% CI -1.4, -1.1]), physician visits (-20.3 [95% CI -24.7, -15.9]), and ED visits (-1.5 [95% CI -1.9, -1.1]). For the most part, mental health-related HRU visits within the overall cohort comprised the majority of all-cause total (87% and 81%), hospital (92% and 83%), physician visits (88% and 82%), and ED visits (53% and 47%) during pre-index and post-index periods, respectively. The total mean number of mental health-related visits within the overall cohort was significantly lower in the post-index period compared with the pre-index period (84.8 [SD 72.6] versus 60.2 [SD 63.6]; mean difference -24.6 [95%CI -29.1, -20.0]), as well as hospitalizations, physician visits, and ED visits (p<0.001).

Among the CTO cohorts, all-cause and mental health-related total, hospital, physician, and ED visits were significantly lower in the post-index period compared with the pre-index period, with the exception of the pre=no / post=yes CTO cohort that displayed no significant differences in HRU visits between the pre- and post-index periods (Additional file 3). The pre=yes / post=no CTO subgroup consistently showed the largest difference in visits across all-cause and mental health-related total HRU, as well as hospitalizations, physician visits, and ED visits (Additional file 3).

### 3.5 Health resource utilization costs

Figure 2 also shows the mean cost ($CDN) and comparative differences of all-cause and mental health-related total HRU costs, as well as hospitalization, physician visit, and ED visit costs for the overall cohort between the pre-index and post-index periods. All-cause hospitalizations incurred the highest costs ($54,560 [SD $63,681] pre-index, $27,453 [SD $56,298] post-index), followed by physician visits costs ($17,077 [SD $13,582] pre-index, $11,253 [SD $12,911] post-index), and ED costs ($2,494 [SD $3,413] pre-index, $1,638 [SD $3,615] post-index) during the study period. The mean cost difference between the post-index period and the pre-index period was significantly lower for all-cause total HRU (-$33,788 [95% -$38,993, -$28,583]), and the specific costs for hospitalizations (-$27,108 [95% CI -$31,526, -$22,690]), physician visits (-$5,824 [95% CI -$6,699, -$4,949]), and ED visits (-$857 [95% CI -$1,034, -$679]) within the overall cohort. Mental health-related HRU costs within the overall cohort comprised the majority of all-cause total (92% and 85%), hospital (94% and 87%), physician visit (89% and 84%), and ED visit (64% and 56%) costs during pre-index and post-index periods, respectively. The total cost of mental health-related visits within the overall cohort was significantly lower in the post-index period compared with the pre-index period ($34,205 [SD $63,428] CDN versus $68,403 [SD $72,088] CDN; mean difference -$34,198 [95%CI -$39,098, -$29,297]), as well as hospitalizations, physician visits, and ED visits (p<0.001). Statistical results were the same when median costs were compared using the Wilcoxon signed-rank test (Additional file 4).

Among the CTO cohorts, all-cause and mental health-related total, hospital, physician, and ED visit costs were significantly lower in the post-index period compared with the pre-index period (p<0.001), with the exception of the pre=no / post=yes CTO cohort that displayed no significant differences in HRU costs (Table 3). The pre=yes / post=no...
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Table 3. Mean healthcare costs during the pre- and post-index periods among the CTO cohorts.
| CTO status     | pre=no / post=no (n=689; 57%) | pre=yes / post=yes (n=275; 23%) | pre=yes / post=no (n=133; 11%) | pre=no / post=yes (n=114; 9%) |
|----------------|--------------------------------|----------------------------------|---------------------------------|--------------------------------|
|                | pre-index                      | post-index                       | pre-index                       | post-index                       |
| All-Cause, $CDN |                                |                                  |                                 |                                 |
| Total, mean (standard deviation) | $61281 (72902) | $34387 (65056) | $91342 (80380) | $100699 (82557) | $15987 (22746) | $79285 (72358) | $78268 (78204) |
| difference [95%CI] | -$26894 [-33472, -20316] | -$40018 [-51363, -28673] | -$84712 [-98938, -70486] | -$1017 [-19082, 17048] |
| Hospitalizations, mean (standard deviation) | $44871 (62059) | $23016 (52911) | $66326 (60263) | $36392 (67225) | $76597 (72378) | $7913 (17777) | $59026 (59563) | $55499 (63312) |
| difference [95%CI] | -$21855 [-27526, -16185] | -$29934 [-39438, -20430] | -$68684 [-81158, -56210] | -$3527 [-18210, 11156] |
| Physician visits, mean (standard deviation) | $14013 (12111) | $9759 (12641) | $22347 (15303) | $13266 (13551) | $21861 (12627) | $7300 (5654) | $17297 (13275) | $20039 (14477) |
| difference [95%CI] | -$4254 [-5286, -3222] | -$9081 [-11093, -7069] | -$14561 [-16684, -12439] | $2742 [-682, 6166] |
| ED visits, mean (standard deviation) | $2396 (3768) | $1612 (4258) | $2669 (2892) | $1666 (2400) | $2241 (1988) | $774 (1412) | $2962 (3593) | $2730 (3335) |
| difference [95%CI] | -$785 [-1038, -531] | -$1002 [-1310, -695] | -$1467 [-1811, -1123] | -$232 [-933, 469] |
| Mental health-related, $CDN |                                |                                  |                                 |                                 |
| Total, mean (standard deviation) | $55360 (68443) | $27965 (59719) | $86532 (73380) | $92753 (74477) | $13974 (22046) | $75092 (71688) | $71361 (70648) |
| difference [95%CI] | -$27395 [-33667, -21122] | -$42311 [-53047, -31575] | -$78779 [-91476, -66082] | -$3731 [-20470, 13008] |
| Hospitalizations, mean (standard deviation) | $41827 (59543) | $19300 (49347) | $64106 (59601) | $31838 (61986) | $70814 (63850) | $7188 (17376) | $57283 (59364) | $51652 (57815) |
| difference [95%CI] | -$22527 [-27964, -17090] | -$32268 [-41216, -23319] | -$63626 [-74636, -52616] | -$5631 [-19328, 8067] |
| Physician visits, mean (standard deviation) | $12128 (10663) | $78456 (11155) | $20546 (12711) | $11340 (12104) | $20243 (6316) | $6316 (12989) | $15773 (13139) | $17972 (13319) |
| (standard deviation) | (14677) | (5398) |
|----------------------|---------|--------|
| difference [95%CI]  | -$4282 [-5248, -3316] | -$9206 [-11138, -7274] | -$13927 [-15957, -11897] | $2200 [-996, 5395] |
| ED visits, mean (standard deviation) | $1405 (2080) | $819 (2314) | $1881 (1978) | $1044 (1978) | $1696 (1417) | $470 (967) | $2036 (2588) | $1736 (2066) |
| difference [95%CI]  | -$586 [-742, -430] | -$837 [-1055, -619] | -$1226 [-1472, -980] | -$300 [-799, 198] |

**Bolded** mean difference indicates statistically significant difference (p<0.001) between the 2-year post- and 2-year pre-index periods using paired t-tests. Abbreviations: CDN = Canadian; CI = confidence interval; CTO = community treatment order; ED = emergency department.

### 4. Discussion

In this retrospective, non-interventional, single arm study of 1,211 adults with schizophrenia, real-world antipsychotic medication adherence, and HRU and associated costs were compared between the 2-year period before and after SGA-LAI initiation using administrative data from April 1, 2012 to March 31, 2018 in Alberta, Canada; subgroup analysis was performed based on CTO status. Overall, results showed that among adults with schizophrenia, adherence to antipsychotic medication was greater, and total HRU and associated costs, including all-cause and mental health-related hospitalizations, physician visits, and ED visits were lower during the 2-year period following initiation of a SGA-LAI compared to the 2-year period before. Regarding CTO status, 80% of individuals persisted with the same CTO status throughout the pre- and post-index periods (pre=no / post=no; pre=yes / post=yes), and had similar decreases in HRU and costs as the overall cohort. Among those with a change in CTO status during the study period, the pre=yes / post=no CTO cohort consistently showed the largest differences in HRU and costs between the pre- (higher) and post- (lower) index periods; the pre=no / post=yes CTO cohort displayed no significant differences in HRU visits and costs between the pre- and post-index periods. Collectively, results of this study indicate that SGA-LAIs may be an effective strategy for assisting with long-term antipsychotic medication adherence, and while causality cannot be established, lower HRU and associated costs to the healthcare system occurred over a 2-year period after SGA-LAI initiation; however, these can vary depending on CTO status.

Results from the overall study cohort are consistent with recent observational studies, which have reported lower HRU and costs after initiation of LAI antipsychotic medication compared with before among individuals with schizophrenia [24, 25]. Among individuals who initiated SGA-LAI antipsychotic medication in Quebec, Canada, Stip et al. (2018) found that the number of hospitalizations and ED visits were lower in the 1-year period after initiation compared to the year prior, and that the observed 64% reduction in total HRU costs was primarily driven by the pre-post difference in hospitalization costs [24], which has been reported to represent the largest proportion of direct healthcare costs for schizophrenia [3]. However, in contrast to Stip et al. (2018) who also reported no significant pre-post differences in physician visits, we found that the number of physician visits were significantly lower in the post-versus pre-index period, which may be partially due to the longer observation period employed in the current study, as a 2-year observation period has been suggested to result in a higher likelihood of detecting a difference compared with 1-year periods [15]. Collectively, our results support and extend previous findings by showing that HRU and associated costs, made up primarily of hospitalization costs, were lower during the 2-year period after SGA-LAI initiation compared with before, which is a longer observation period than most previous reports [15].
Another unique aspect of this study was that analysis was also performed based on CTO status during the study period, of which 43% of individuals received at least one active CTO. The intention of a CTO is to assist with improving treatment adherence in patients with greater illness severity. To this end, O’Brien et al. (2009) found that in Ontario, Canada, patients who were placed on a CTO had severe and persistent mental illness [26]. Not surprisingly, varying patterns of HRU and associated costs were found among the different CTO cohorts. While 80% of individuals persisted with the same CTO status throughout the pre- and post-index periods (57% pre=no / post=no; 23% pre=yes / post=yes) and had similar decreases in HRU and costs as the overall cohort, 20% of individuals had a CTO status change during the study period (11% pre=yes / post=no; 9% pre=no / post=yes) along with associated HRU and costs that varied from the overall cohort. The pre=yes / post=no CTO cohort displayed the largest mean difference in all measured HRU and associated costs between the pre-index period, which was higher, and the post-index period, which was lower. It is possible that after SGA-LAI initiation these individuals may have attained a stable phase of illness where symptoms and functioning significantly improved [6]. The pre=no / post=yes CTO cohort displayed HRU and costs that were distinctive from the overall cohort, as well as the other CTO cohorts. This cohort most likely persisted with nonadherence for a time after SGA-LAI initiation, as demonstrated by the introduction of a CTO during the post-index period to provide additional assistance with treatment adherence. Considering that nonadherence to antipsychotics has been found to be associated with higher inpatient admissions and ED visits [7-9], and naturalistic studies examining HRU before and after initiation of a CTO in Canada have found lower numbers of in-patient admissions and days of hospitalization, and ED visits following CTO activation [26-30], these two factors may have contributed to the finding of no significant pre-post differences in HRU and associated costs within this CTO cohort; the smaller sample size of this cohort may have also reduced power to detect true differences.

This study has several important strengths, including the large size, population-based design, incorporation of CTO status, and long observation periods. However, this study is also subject to a number of limitations that should be taken into consideration when interpreting results. 1) While individuals with schizophrenia were identified using a validated case definition, administrative health data was used as opposed to medical records, and therefore there is a potential for misclassification of the study groups or measures. 2) Pre-post study designs lack a control group, hence the outcome patterns in the counter-factual patients who are similar to our patient group in all aspects except SGA-LAI initiation are unknown; this is particularly important when interpreting outcomes in patients who had a change in CTO status, which may be an indicator of other factors including disease instability. 3) PIN data only captures dispensations of prescription medications, and therefore actual medication up-take by patients is unknown. However, since LAI antipsychotics are injected by healthcare professionals, the dispensation of SGA-LAIs in this study should be highly representative of the actual uptake. 4) Use of over-the-counter medications, prescription medications provided in a hospital or secondary care setting, and other non-pharmacotherapy treatments are not captured within provincial administrative data and therefore, not reported.

5. Conclusions

Among adults with schizophrenia, antipsychotic medication adherence was higher, and HRU and associated costs primarily comprised of hospitalization costs, were lower during the 2-year period after SGA-LAI initiation as compared to before, extending findings of previous mirror-image and naturalistic studies. CTOs should be considered in research studies involving individuals with schizophrenia as they are commonly used in this population and our results show that HRU and associated costs can vary depending on CTO status.

Abbreviations
Declarations

Ethics approval and consent to participate

Ethics approval (Pro00086993) was obtained from the institutional review board at the University of Alberta. This is a study of administrative data without any intervention. No study participants were placed at risk as a result of this study, and informed consent was not required.

Consent for publication

Not applicable

Availability of data and materials

The data that support the findings of this study are not available. The data custodians, Alberta Health Services and Alberta Health do not allow users of the data to publish the data. Please contact the corresponding author for requests related to the data used in this study.

Competing interests

The author(s) declared the following potential conflicts of interest with respect to the research, authorship, and/or publication of this article: SD, HS, and MS have no competing interests to declare. SK, KW, KM, AG, HL, and LR are members of the Real World Evidence Unit, an academic entity at the University of Alberta that conducts research including investigator-initiated industry-funded studies. PC has acted as a consultant and received honoraria and research travel grants from Alkermes, Allergan, Boehringer Ingelheim, HLS Therapeutics, Janssen, Lundbeck, Otsuka, and Sunovion between 2015 and 2021. No other conflict of interest is declared. All authors of this study had complete autonomy over the content and submission of the manuscript, as well as the design and execution of the study.

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Authors’ contributions

SK, KW, KM, and LR designed the study, and KW, AG, and HS analysed the data. KM and KW wrote the main manuscript text and KM and HL prepared the figures and tables. All other authors revised it critically for important intellectual content and approved the final version to be published. All authors are accountable for the work and
integrity of the work. The corresponding author and guarantor accepts full responsibility of the work and/or conduct of the study, had access to the data, and controlled the decision to publish. SK and LR attest that all listed authors meet authorship criteria and that no others meeting the criteria have been omitted.

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Figures

Figure 1

Flow diagram of cohort selection.
Figure 2

Mean per-patient health care resource utilization and cost among the overall cohort. All comparative differences (mean difference [95% confidence interval]) between the post-index and pre-index periods were statistically significant (p<0.001) using paired t-tests. Abbreviations: CDN = Canadian; ED = emergency department; HRU = healthcare resource utilization.

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