RESULTS: Out of 3025 patients 35 were prescribed Clozapine as monotherapy and 5 patients had clozapine plus psychopharmacological augmentation. Ages ranged from 21-86. Out of the 39 patients, there were 13 male and 26 female. The predominant diagnosis was mood disorder or MDD with psychotic features followed by schizophrenia. The augmentation antipsychotics used were aripiprazole and risperidone. In the literature, the most frequent augmentation strategy for TRS is adding another antipsychotic with more D2 receptor blockade. Other strategies involve identifying and treating the symptoms not controlled by clozapine.

CONCLUSIONS: Currently augmentation of Clozapine in TRS is highly individualized due to lack of supporting evidence to state the contrary. When working with treatment resistant patients who are not responding to clozapine alone, it is imperative to thoroughly review and consider all treatment options and augmentation strategies. More studies should be done in controlled settings to better evaluate possibilities as well as more evaluations to be done on other ways of augmentation of clozapine. Literature has stated between 20-60% of patients are defined as TRS. Clozapine is considered as one of the most effective treatment available at present time for TRS. Recent literature suggests despite its superior efficacy, as many as 70% of those suffering from TRS on clozapine continue to suffer from positive, negative or cognitive symptoms. The literature has abundant adjunctive treatment strategies such as the addition of antipsychotics, mood stabilizers, antidepressants, or even with the use of electroconvulsive therapy. We emphasize the importance of correctly identifying TRS patients who may benefit from the initiation of clozapine, what would be beneficial for them if they do not respond, how to tailor their treatment to target symptoms not being ameliorated, and recommend treatment in these complex cases be multidisciplinary.

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RE-KINECT: Real-World Dyskinesia Screening Study and Registry in Patients Taking Antipsychotic Agents: Interim Baseline Burden of Illness Results

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ABSTRACT: Background: Tardive dyskinesia (TD) is associated with prolonged exposure to dopamine receptor blockers including antipsychotics. This registry describes the prevalence and impact of involuntary movements (possible TD) in a real-world population of patients taking antipsychotics.

METHODS: RE-KINECT (NCT03062033) aims to enroll 1,000 patients from 70 US psychiatric practices. Adults with ≥3 months lifetime exposure to antipsychotic(s) and ≥1 psychiatric disorder are eligible for two-tier screening: informal observation, and then clinician observation of abnormal involuntary movements in general body regions (head/face, neck/trunk, upper/lower limbs) and confirmation of possible TD. Based on clinician assessment, patients are assigned to Cohort 1 or Cohort 2 (without or with abnormal involuntary movements, respectively). In both cohorts, the following baseline assessments are included: clinician’s assessment of clinical psychiatric severity, patient perceived health-related quality of life (EuroQOL 5-Dimensions), social burden/disability questionnaire (Sheehan Disability Scale), and 12-month retrospective chart review of medical and treatment history. Cohort 2 also participate in 12-month longitudinal evaluation. Interim baseline data are available from four sites.

RESULTS: Baseline data are currently available for 116 patients—mean age, 49.6 years; female, 60.3%; schizophrenia/schizoaffective disorder, 32.8%; at least 1 mood disorder, 84.5%, and 10.4 years mean cumulative lifetime exposure to antipsychotic(s). The most concerning health condition for both cohorts is their mental health (69.0%), followed by physical activity and nutrition (33.6%). 32.8% of subjects had clinician confirmation of possible TD.

CONCLUSION: This novel registry aims to evaluate the real-world potential impact/burden of TD. Preliminary analyses suggest that TD is common in patients with schizophrenia and mood disorders taking antipsychotics. Further analyses will explore the burden of illness in this population.

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