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A STUDY OF ANTIPSYCHOTIC INDUCED TARDIVE DYSKINESIA IN INDIAN POPULATION

Gupta Ravi, Bhatia Manjeet S.

# ABSTRACT

**Objective**: This study was aimed at, to assess the point prevalence and risk factors in Indian population.

**Design:** Cross sectional comparative study.

**Place and duration of study:** The sample population was chosen from patients attending Psychiatry OPD of GTB Hospital during two months of the study (October 2004 to November, 2004).

**Subjects and Methods:** 62 subjects with history of exposure to antipsychotic drugs were included in this study. Their demographic data, illness related factors, and drug history were assessed. Tardive Dyskinesia (TD) was diagnosed according to DSM-IV-TR criteria. Significance of categorical variables was assessed by Chi-Square test and independent t-test was applied for comparison of numerical variables. Odds ratio was calculated using logistic regression.

**Results:** Point prevalence of TD in this sample was 16%. Advancing age in female gender predicted development of TD. Other known risk factors did not show any statistical significant effect.

**Conclusion:** Elderly females are at increased risk for development of TD. Point prevalence rate of 16% in mixed sample is fairly high and underscores the need for further studies in this area.

**Key words:** Tardive Dyskinesia, Neuroleptics, Risk factors.

# INTRODUCTION

Tardive dyskinesias (TD) are involuntary move- ments that usually after long-term neuroleptic therapy. These movements commonly involve orofacial muscu- lature, and appear as puckering, lip-smacking, repeated tongue protrusion, pouting, chewing, facial grimacing etc1.

Prevalence of TD has been described ranging from 3% to 70% depending upon the sample chosen1 and the length of exposure to the drugs or higher cumulative doses2. Besides neuroleptic exposure there are some other factors that increase the chances of TD if they co- occur in the subjects exposed to antipsychotics. These are older age, female sex, ethnicity (lower chances in Asians than Americans), family history of mood disor-

**Gupta Ravi,** MD Senior Resident

**Bhatia Manjeet S.**, MD, MNAMS

Professor and Head of Psychiatry Department, University College of Medical Sciences and GTB Hospi- tal, Dilshad Garden, Shahdara, Delhi-110095

**Correspondence:**

**Dr. M.S. Bhatia,**

D-1, Naraina Vihar, Delhi-110028

E-mail: [manbhatia1@rediffmail.com](mailto:manbhatia1@rediffmail.com) 0091-98101-61790

ders, use of neuroleptic in mood disorders patients, brain damage, substance use, and diabetes1.

Moreover, presence of TD may impair social rela- tionships, dental problems, weight loss, ambulatory dif- ficulties, speech difficulty1, therefore it is mandatory to take the informed consent from the patient before ex- posing him to neuroleptic treatment.

Despite being such a sensitive issue, Indian litera- ture on this subject is scarce and only few reports are available and western data that we use at present to ad- dress this issue have ethnic/ genetic bias. There- fore, present study was designed to assess the preva- lence of TD in Indian subjects exposed to neuroleptics and to find out the role of various risk factors in their occurrence.

# SUBJECTS AND METHODS

The sample population was chosen from patients attending Psychiatry OPD of GTB Hospital during two months of the study (October 2004 to November, 2004). Subjects were screened for exposure to neuroleptic drugs and those who have been using any of the antipsychotic drugs regularly for the past two years were included in this study after obtaining consent.

Information was gathered by history provided by the patient and a reliable informant along with physical

examination of the patient and past medical records. Pa- tients who were not having the drug prescriptions for last two years were excluded from the study so as the pa- tients with spontaneous dyskinesias and withdrawal dyskinesias.

Final sample consisted of 62 patients and their age, sex, age of onset of illness, primary diagnosis, co- morbid psychiatric and medical illnesses, drug history with doses and family history of psychiatric disorders were noted.

To compare the effect of mean daily drug doses, mean doses of past two years were calculated by sum- ming up each dose with the number of days of exposure for each drug and then dividing it by 730. e.g., if a pa- tient has been exposed to risperidone 2 mg for 180 days, then to 1 mg for 180 days, then to 3 mg for past year, the mean dose was (2 X 180 + 1 X 180 + 3 X 365 / 730). Thus it provided the average daily dose of the drug taken be person. Two years period was taken because in the initiation and continuation phase of treatment drug doses were relatively higher than subsequent dosing. However, exposure to any drug for less than a month was ignored for the sake of statistical analysis. Presence of TD was diagnosed according to DSM-IV-TR criteria in all the sub- jects included in this study.

*Statistical analysis:*

For analysis subjects were divided into two groups- those having tardive dyskinesia and those without tardive dyskinesia. The age of onset of primary psychiatric illness, present age and differences in mean doses of drugs were compared by the independent t-test. Other categorical variables were assessed using Fisher’s exact test . Finally, logistic regression was run taking tardive dyskinesia as dichotomous variable and co-morbidity, positive family history, exposure to risperidone, olanzapine, trihexyphenidyl and antidepressants as independent vari- ables to predict the effect of these variables on occur- rence of TD. SPSS version 13.0 for Windows was used to carry out analysis.

# RESULTS

Prevalence of TD in our sample was 16.12%; they were more common in females *(P< 0.001)*; however there was no statistical significant difference *(P> 0.05)* between both the groups on the basis of age of onset of primary illness, age of onset of TD, primary diagnosis, co-mor- bid psychiatric or medical disorders and family history of psychiatric illness (Table 1).

Table 1

Distribution of Demographic and Illness Related Risk Factors

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **S.No.** | **Variable** | **Non TD (N=52)** | **TD (N=10)** |  |
| 1. | Sex |  |  |  |
|  | Male | 38 (73%) | 1 (10%) |  |
|  | Female | 14 (27%) | 9 (90%) | P < .001 |
| 2. | Age | 36.9 + 13.93 | 45 +10.68 | p= 0.08\* |
| 3. | Age of onset of illness | 29.6 + 13.7 | 33.8 + 9.89 | p=0.36\* |
| 4. | Diagnosis |  |  |  |
|  | Schizophrenia | 21 (41%) | 2 (20%) |  |
|  | BPD | 11 (21%) | 3 (30%) |  |
|  | Depression with psy Feat | 11 (21%) | 2 (20%) |  |
|  | Other psychotic disorders | 7 (13%) | 2 (20%) |  |
|  | Somatoform & Anxiety disorder | 2 (4%) | 1 (10%) | *P* = 0.55 |
| 5. | Co-morbidity |  |  |  |
|  | HT | 2 (4%) | 0 |  |
|  | DM | 0 | 2 (20%) |  |
|  | Depression | 7 (13%) | 0 |  |
|  | Substance Use | 5 (10%) | 0 |  |
|  | Anxiety Disorders | 3 (6%) | 0 |  |
|  | Epilepsy | 1 (2%) | 0 | *P*= 0.01 |
| **6.** | Family History |  |  |  |
|  | Depression | 4 (8%) | 1 (10%) |  |
|  | MDP | 3 (6%) | 1 (10%) |  |
|  | Schizophrenia | 2 (4%) | 1 (10%) |  |
|  | Epilepsy | 1 (2%) | 0 | *P*= 1.00 |

* Independent sample t test, Rests are Fisher’s Exact test

Table 2

Average Daily Doses of antipsychotics and antiparkinson drugs in both groups

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **S. No.** | **Drug** | **Non TD (N=52)** | **TD (N=10)** | **p\*** |
| 1. | Risperidone | 3.3 + 2.2 (44%) | 3.5 +2.0 (60%) | 0.8 |
| 2. | Olanzepine | 6.7 + 5.3 (35%) | 7.5 + 3.5 (20%) | 0.8 |
| 3. | THP | 4.0 +1.6 (29%) | 3.8 + 1.3 (60%) | 0.8 |

* Independent sample t test; Figures aside indicate number of people exposed to respective drugs

Table 3

Odds Ratio Values of Risk Factors on Occurrence of TD

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| **S. No.** | **Variable** | **Significance** | **Odds Ratio** | **95% CI** | |
| Lower | Upper |
| 1. | Co-morbidity | 0.714 | 0.70 | 0.10 | 4.71 |
| 2. | Family History | 0.206 | 1.66 | 0.75 | 3.63 |
| 3. | Risperidone | 0.216 | 3.15 | 0.51 | 19.41 |
| 4. | Olanzapine | 0.118 | 2.28 | 0.81 | 6.42 |
| 5. | Trihexyphenidyl | 0.102 | 3.74 | 0.76 | 18.22 |
| 6. | Antidepressant | 0.583 | 0.59 | 0.09 | 3.77 |

We did not find any patient using conventional neuroleptics because we stressed on the medical records of past two years. However, most of the patient had long term illness (average approximately 10 years), exposure to conventional neuroleptics could not be ruled out and result must be interpreted with caution. We also analyzed whether different daily doses of Risperidone, Olanzapine, and Trihexyphenidyl affect the development of TD, but did not find any significant difference *(P> 0.05)* between groups (Table 2). Other drugs used by samples were as follows- Aripiprazole to 14% and 20%, clozapine to 2% and 0; Valproate to 27% and 2%; Lihium to 8% and

30%, Fluoxetine to 20% and 10%; Escitalopram to 15% and 20%; and other antidepressants to 15% and 0 sub- jects in Non-TD and TD group respectively.

Classification of the study group was 87.1% dur- ing the logistic regression. Risk of development of tar- dive dyskinesia was negatively associated with co-mor- bidity and exposure to antidepressant while positively associated with family history of psychiatric disorders, and exposure to risperidone, olanzapine and trihexyphenidtl (Table 3). However, none of these vari- ables reached statistical significance.

# DISCUSSION

Previous studies on Indian population have re- ported a prevalence of 10%3 and 29%4 for drug induced tardive dyskinesia. Prevalence found in present study is lower than described by Bhatia et al4, probably because they have included schizophrenic patients which usu- ally require higher doses and are often exposed to con-

ventional antispyhotics. In this study, exposure to neuro- leptic was the only criteria for inclusion without being affected by diagnosis and duration of treatment. There- fore average doses of antipsychotics were relatively lower and all the subjects were on atypical drugs for at least past two years thus contributing to lower incidence5,6 . Moreover, our findings go along with the previous reports of ethnic variation7 where lower rates of TD have been described in Asian subjects1. However at this point, we are unable to comment which of these factors played major role for such low incidence, and it requires further study.

Female subjects in this study suffered higher rates of dyskinesias. It has been described as an important risk factor in literature but few studies did not find any evidence in its favor8,9 and one even reported higher in- cidence in males4. In a review article, Sachdev1 suggests that it is not only the gender but ‘age- gender interac- tion’ that works behind increased prevalence of TD in females. Post-hoc analysis of our data found that females were older (44.13+ 14.69) than males (34.15 + 11.97) *(P=0.008)*. Thus we also opine that age-gender interac- tion may be more important for development of TD, rather than any of the factors alone.

Age has been described as the most important risk factors across studies1,9-12. Higher prevalence with in- creasing age can be attributed to age related brain de- generation13, neurological pathologies14, co-morbid medi- cal disorders14, longer years of treatment8,12 and thus higher cumulative neuroleptic doses1 and spontaneous dyskinesias14. However, we did not find any effect of age on the prevalence of dyskinesia in the whole sample. Similar results were reported in other studies4,8. It is pos- sible that there is a cut-off point of age beyond which the

risk of TD increases as described by Woerner et al2 who demonstrated that risk of TD increases after 50 years even on the lower doses of antipsychotics and despite the facts that younger subjects had longer exposure to neuroleptics. The mean age of subjects in our sample was lower than this cut off point. Secondly, as we have mentioned, according to us, age and gender interaction is the root cause for development of TD, rather than any of the factors alone.

Illness related factors that increase the risk of TD are early age of onset of primary illness15, presence of mood disorders in the patient1,4,11, history of substance use12 and family history of mood disorders1. However, we could not confirm any of the findings (in-fact co-mor- bidity and exposure to antidepressants were negatively associated with risk of TD) except for the family history of psychiatric illness which enhanced the risk (but statis- tically insignificant). Similar results have been found in other studies which could not correlate it with primary diagnosis8-10, and age of first exposure to antip- sychotic10.

Not only the psychiatric disorder but presence of medical disorder- particularly diabetes mellitus may be associated with increased risk16. However, our findings do not support it and confirm findings of Miller et al12.

Lastly, drug related factors e.g., exposure to con- ventional neuroleptics8, higher cumulative doses or longer duration of treatment8,10,12 and use of antiparkinsons drug15 have been reported to increase the risk of TD. In this study we found that the doses of antipsychotics and trihexyphenidyl did not affect TD, Though those with TD had more chances to be exposed to trihexyphenidyl and atypical neuro- leptics (but statistically insignificant). However, due to past history of exposure to conventional antipsy- chotic for pretty long period, validity of this finding can be challenged.

This study had few methodological limitations- ex- clusion criteria cut down the sample to small size in present study. Though in this sample we did not have any subject with the history of conventional drug use in past two years, such possibility can not be ruled out be- fore that. Moreover, the mean daily doses of drugs were also low due to sample with mixed diagnosis. Our re- sults have probably been affected by the lower mean doses of the drugs, use of atypical antipsychotics in past two years that are known to reverse TD5-6 and use of con- comitant trihexyphenidyl that can have a protective ef- fect. As hypothesized at inception, we did not find any evidence that antidepressants increase rate of TD and go along with previous finding10.

In conclusion, this study confirmed that female gender with advancing age is the only risk factors for TD and it is unaffected by presence of other dis- orders, doses of atypical neuroleptics and exposure to other psychotropic drugs. Due to small size of

sample, results must be generalized with caution and further research is required with methodological im- provements.

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# STATEMENT OF INTEREST

There are no conflicting interests to declare as study was not funded by any agency.

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