CYP2D6 Genotype in Relation to Liver Toxicity Due to Tetrabenazine in Iraqi Patients with Hyperkinetic Movement Disorders

Zainab A. Abboud*, Shatha H Ali* and Nawfal M Sheaheed**

*Department Clinical Pharmacy, College of Pharmacy, University of Baghdad, Baghdad, Iraq.
**Department of Clinical and Laboratory Science, College of Pharmacy, University of Baghdad, Baghdad, Iraq.
***Ministry of Health and Environment, Baghdad Teaching Hospital, Medical City, Baghdad, Iraq.

Abstract

The common types of movement disorders are; dystonia which is a syndrome of repetitive muscle contractions. While, Huntington disease is autosomal dominant progressive neurodegenerative disorder, which is characterized by involuntary movements ("chorea"). Tetrabenazine therapy has been shown to effectively control these movements compared to placebo. The most commonly reported side effects of tetrabenazine increase liver enzymes and these side effects are dose dependent.

This is a prospective case controlled study was carried on 50 patients whom divided into 2 groups: group 1 involved 25 chorea patients, and group 2 included patients with dystonia, whom treated with (tetrabenazine) for three months. In addition to control group involved 25 healthy subjects to estimate the prevalence of genetic polymorphism of CYP 450 2D6 enzyme in related to tetrabenazine efficacy and toxicity. Blood samples were collected at the beginning to perform a genotyping assay for CYP 450 2D6 enzyme by PCR and to assess liver function and after three months of treatment to assess liver and measuring the plasma concentration of tetrabenazine, alpha and beta dihydrotetrabenazine by HPLC.

The results show a significant CYP 450 2D6 enzyme polymorphism. And elevations of liver enzymes in the patient indicate hepatotoxicity of tetrabenazine and its metabolites.

Keyword: Dystonia, Chorea, Tetrabenazine, Alpha and Beta dihydrotetrabenazine, CYP 450 2D6 Enzyme Polymorphism, Liver Function.


cumpyورثائي لـ CYP2D6 الذين يتعاطون علاج التترابينازين

وظائف الكبد

الخلاصة

من بين الأنواع الشائعة لاضطرابات الكبد خلل التوتر العضلي، وهو متلازمة اضطرابات الحركة العصبية التي تؤدي إلى تقلصات العضلات. قد ثبت أن علاج التترابينازين يتحكم بشكل فعال في هذه الحالات مقارنة بعلاج الهرمن. إن أكثر الأثر الجيني شيعًا للتترابينازين في ارتفاع إزيمات الكبد وهذه الأثر الجيني تعتمد على زيادة الجرعة. أجريت الدراسة على 50 مريضاً تم تطبيقهم إلى مجموعة واحدة، المجموعة الأولى شملت 25 مريضاً يعانون من حمل التوتر العضلي، والمجموعة الثانية شملت 25 مريضة مصابين بعلاج التترابينازين لمدة ثلاثة أشهر. بالإضافة إلى مجموعة السيطرة. صحي تم جمع عينات في بداية الدراسة لقياس وظيفة الكبد. بالإضافة إلى تشخيص الانتشار الجيني لenzyme CYP 2D6 في المرضى. تم قياس تركيز التترابينازين، ألفا وبيتا التترابينازين في البلازما مع وظائف الكبد. هذه الدراسة توضح أن هناك تقدم مع الأشكال الجينية كبيرة لenzyme CYP 2D6 هذه الأشكال، مع ارتفاع كبير في أنزيمات الكبد في المرضى مقارنة مع مجموعة السيطرة. الكلمات الافتتاحية: خلل التوتر العضلي، تترابينازين، يعانون، ذو ماي هابي، التترابينازين، آلفا وبيتا، CYP 2D6، وظائف الكبد.

1Corresponding author E-mail: zainabaa87@yahoo.com
Received: 28/11/2019
Accepted: 15/2/2020

Iraqi Journal of Pharmaceutical Science
Introduction
The term "movement disorders" refers to a group of nervous system (neurological) conditions that cause abnormal increase in the movements, which may be voluntary or involuntary. Common types of movement disorders include:

Dystonia is a syndrome of neurological movement disorder in which constant or repetitive muscle contractions result in twisting and repetitive movements or abnormal fixed postures. Dystonia is often intensified or aggravated by physical activity, and symptoms may progress into adjacent muscles. Huntington disease is primarily an adult-onset hereditary autosomal dominant progressive neurodegenerative disorder, which is characterized by involuntary movements ("chorea"), psychiatric symptoms, and cognitive dysfunction that can lead to dementia.

Presently, the backbone of treatment for movement disorders is symptomatic and supportive care — no drugs are available to stop or prevent the progression of hyperkinetic movement disorders. Tetrabenazine therapy (TBZ), a benzoquinoline derivative, is an oral monoamine-depleting agent with selectivity for dopamine. It specially prevents the presynaptic monoamine storage by inhibiting the vesicular monoamine transporter type-2, a presynaptic transporter found mainly in the central nervous system.

The most commonly reported side effects of tetrabenazine increase liver enzymes (ALP, ALT, and AST). Side effects of tetrabenazine are dose dependent.

This study aimed to predict phenotypes of CYP2D6: poor metabolizers, intermediate metabolizers, and extensive metabolizers or ultra-rapid metabolizers among Iraqi subjects. And to investigate whether CYP2D6 gene polymorphism effect the tetrabenazine, and it is two metabolites , α-H-tetabenazine and β-H-tetabenazine and their relation to liver toxicity.

Patient and Methods
Patient selection and study design
Fifty movement disorder patients participated in the current prospective case controlled study where divided into 2 groups: Group 1 involved 25 patients suffer from chorea (12 males and 13 females) with mean age 41.36±2.39 years, and Group 2 includes patients with dystonia (8 males and 17 females) with mean age 38.04±2.62 years, treated with (tetrabenazine) for three months. The starting dose of tetrabenazine was 12.5 mg once daily in the morning; titrate up at weekly intervals by 12.5 mg/day; doses of 37.5 to 50 mg was divided into 3 doses. Ethical approval was obtained from the department of clinical pharmacy at the College of the Pharmacy.

The study is carried out in Baghdad medical city during the period from September 2018 to June 2019. Patients undergoing clinical examination by measuring Unified Huntington's Disease Rating Scale (UHDRS) and Unified Dystonia Rating Scale (UDRS) in the movement disorder unit of the hospital as well as in private clinic.

Ten millimeter Blood sample were collected at the beginning of the study to measure, liver function tests (serum ALP, ALT, and AST) and genetic polymorphism of CYP 450 2D6 enzyme for both group and after three months of the study to measure the plasma concentration of tetrabenazine , alpha and beta tetrabenazine and liver function tests with the clinical examination for side effects.
**Genotyping of CYP2D6 gene**

After DNA isolation from venous blood by Wizard Genomic DNA purification Kit (Promega, USA). Genotyping of CYP2D6 gene performed by polymerase chain reaction (PCR) conventional (allele specific method) according to method described by Taimour Langae, Issam Hamadeh, Arlene B. Chapman, et al (13). The following primers was used for PCR amplification.

1. CYP2D6 *2 (2850 C>T rs16947)
   - Forward: 5’ GGCCCCTGCACTGTTTCC 3’
   - Reverse: 5’ AAGGGGAACCCTGAGAGC 3’

2. CYP2D6 *4 (1846 G>A rs3892097)
   - Forward: 5’ TGCCGCCTTCGCCAACCACT 3’
   - Reverse: 5’ GCAGAGACTCCTCGGTCTCTC 3’

3. CYP2D6 *10 (100 C>T rs1065852)
   - Forward: 5’ TGTCCAGAGGAGCCCATTT 3’
   - Reverse: 5’ GTCGAAGCAGTATGGTGTGTTCT 3’

**Measure plasma level of tetrabenazine by HPLC**

About (20 µl) of patient plasma was injected into the liquid chromatographic system consisting of a C18 µ Bondapak column and fluorescence detector. The mobile phase was acetonitrile-1% acetate buffer, pH 4.5 (50:50) at a flow-rate of 1 ml/min. The fluorescence of the eluent was quantified using an excitation wave length of 265 nm and an emission filter (KV418)(14),(15).

**Biochemical assay for liver function**

Serum ALP level was measured by Cobas c 311 chemistry analyzer made by Roche Diagnostics in cooperation with Hitachi High-Technologies Corporation (16). While serum AST, ALT, and TSB level evaluated using a ready-made kit for this purpose, according to the method of Kirsch JF, et al (17), Kim WR, et al (18), and Kao TW, et al (19) respectively.

**Statistical analysis**

Data will be analyzed by using SAS (Statistical Analysis System) (version 24.0) program (SPSS Inc., Chicago, Illinois, USA) and Minitab version 17 software. In all comparisons, a p-value <0.05 was considered statistically significant.

**Results**

**Distribution and demographic data of the patients**

Fifty patients completed the course of study successfully, there were non-significant differences between all parameters at baseline as shown in (Table 1).

---

**Table 1. Demographic data and baseline characteristics of the patients.**

| Data                     | Group 1       | Group 2       | P-value |
|--------------------------|---------------|---------------|---------|
| Age (yrs.)               | 41.36±2.39    | 38.04±2.62    | 0.468   |
| No. of subjects          | 25            | 25            | -----   |
| Gender                   | 13 females    | 17 females    | 0.374   |
|                          | 12 males      | 8 males       |         |
| disease Duration(yrs.)   | 6.148±0.599   | 5.400±0.580   | 0.374   |
| BMI                      | 21.120±0.590  | 22.760±0.751  | 0.259   |
| Serum Total Bilirubin(mg/dl)| 0.61±0.052   | 0.64±0.060    | 0.909   |
| Serum Alkaline Phosphatase U/L | 60.03±3.59 | 58.26±4.44 | 0.570   |
| Serum Aspartate Transaminase U/L | 28.96±1.49 | 30.48±1.70 | 0.435   |
| Serum Alanine Transaminase U/L | 42.72±2.05 | 41.89±2.17 | 0.666   |

Data are expressed as Mean±SEM.

**Distribution of patients with (CYP 450 2D6) gene polymorphism**

Seventy four percent (37) of patients have CYP 450 2D6*2 (normal allele) while 26% (13) of patients have allele polymorphisms (CYP 450 2D6*10) . No patients were observed with CYP 450 2D6*4 polymorphisms in this study(Figure 1).
Figure 1. Histogram showing the distribution of patients with (CYP 450 2D6) gene Polymorphism plasma concentration of tetrabenazine, alpha and beta dihydrotetrabenazine for patients with CYP 450 2D6 polymorphism versus the patients without CYP 450 2D6 polymorphism in patients using tetrabenazine:

Tetrabenazine, alpha and beta dihydrotetrabenazine concentration were increased significantly in the patients with CYP 450 2D6*10 polymorphisms compared to patients with CYP 450 2D6*2 using tetrabenazine in chorea and dystonia (Figure 2, 3).

Figure 2. Plasma concentration of tetrabenazine, alpha and beta dihydrotetrabenazine for patients with CYP 450 2D6 polymorphism versus the patients without CYP 450 2D6 polymorphism in chorea patients using tetrabenazine.

Figure 3. Plasma concentration of tetrabenazine, alpha and beta dihydrotetrabenazine for patients with CYP 450 2D6 polymorphism versus the patients without CYP 450 2D6 polymorphism in dystonia patients using tetrabenazine.
Effect of tetrabenazine on liver enzymes

Patients was subdivided into A subgroup (patients with normal allele and B subgroup (patients with allele polymorphism). There was a non-significant difference in serum Alkaline Phosphatase (ALP), Alanine Transaminase (ALT), Aspartate Transaminase (AST), and total bilirubin (TSB) among treated subgroups at pretreatment. While after three months of treatment with tetrabenazine, the study showed that was a significant elevated in serum ALP, ALT, and AST while non-significant difference TSB in both subgroup A and B compared to subgroups at pretreatment, which is highly increased in liver enzymes in subgroup B as showed in (Table 2).

Table 2. Effect of tetrabenazine on liver enzymes in patients with chorea and dystonia after three months of treatment.

| Liver Enzyme | Chorea | P-value | Dystonia | P-value |
|--------------|--------|---------|----------|---------|
|              | Pretreatment | After treatment | Pretreatment | After treatment |
| ALP (U/L) A  | 63.93±4.09 | 80.11±4.71 | 0.014* | 59.35±5.36 | 74.02±4.67* | 0.047* |
| B            | 49.99±6.25 | 98.57±7.53 | 0.001* | 54.80±7.93 | 91.67±5.75* | 0.004* |
| ALT (U/L) A  | 44.02±2.58 | 50.92±2.18 | 0.049* | 42.99±2.58 | 54.55±3.08 | 0.007* |
| B            | 39.40±2.95 | 61.74±4.22 | 0.001* | 38.40±3.87 | 67.55±4.77 | 0.001* |
| AST (U/L) A  | 29.50±1.63 | 33.72±1.17 | 0.045* | 29.95±1.83 | 39.47±1.89 | 0.001* |
| B            | 27.57±3.46 | 39.86±2.35 | 0.015* | 32.17±4.35 | 52.50±3.45 | 0.005* |
| TSB (mg/dl) A | 0.594±0.06 | 0.557±0.033 | 0.583 | 0.658±0.08 | 0.679±0.06 | 0.826 |
| B            | 0.643±0.11 | 0.614±0.059 | 0.820 | 0.567±0.08 | 0.583±0.06 | 0.876 |

Data are expressed as Mean±SEM; *significantly difference compare to pretreatment within the same group (p<0.05). A (patients with normal CYP 450 2D6 gene) and B (patients with CYP 450 2D6 polymorphism).

The correlation of plasma concentration of alpha dihydrotetabenazine and plasma concentration of tetrabenazine with liver enzymes (ALP, AST, ALT, and TSB) in patients group:

The study showed a significant correlation between tetrabenazine and alpha dihydrotetabenazine plasma concentrations and serum AST (Figure 4) and ALT (Figure 5) and TSB (Figure 6). The ALP, and TSB serum concentrations were non-significant with serum ALP and TSB as in (Table 3) in chorea patients.

Table 3. Summarizes the relationship between tetrabenazine and alpha dihydrotetabenazine plasma concentration and serum concentration of ALP, AST, ALT, and TSB in chorea patients.

| Concentration | ALP | AST | ALT | TSB |
|---------------|-----|-----|-----|-----|
| TBZ R_value   | 0.344 | 0.399 | 0.591 | 0.173 |
| P_value       | 0.092 | 0.048* | 0.002* | 0.409 |
| Alpha DTBZ R_value | 0.303 | 0.343 | 0.599 | 0.189 |
| P_value       | 0.141 | 0.093 | 0.002* | 0.366 |

* Significant difference
While in dystonia the study showed a non-significant correlation between tetrabenazine and alpha dihydrotetrabenazine plasma concentrations and the ALP, AST, and TSB serum concentrations and only was significant between the tetrabenazine plasma concentrations and serum AST and ALT as in (Table 4) and (Figure 7, 8).

Table 4. Summarizes the relationship between tetrabenazine and alpha dihydrotetrabenazine plasma concentration and serum concentration of ALP, AST, ALT, and TSB in dystonia patients.

| Concentration | ALP  | AST  | ALT  | TSB   |
|---------------|------|------|------|-------|
| TBZ           | R_value| 0.210| 0.460| 0.430 | -0.158|
|               | P_value| 0.314| 0.021*| 0.032*| 0.450 |
| Alpha DTBZ    | R_value| 0.235| 0.288| 0.249 | -0.108|
|               | P_value| 0.259| 0.162| 0.230 | 0.609 |

Figure 4. Plasma concentration of tetrabenazine related to serum AST in chorea patients.

Figure 5. Plasma concentration of tetrabenazine related to serum ALT in chorea patients.

Figure 6. Plasma concentration of alpha dihydrotetrabenazine related to serum ALT in chorea patients.

Figure 7. Plasma concentration of tetrabenazine related to serum AST in dystonia patients.

Figure 8. Plasma concentration of tetrabenazine related to serum ALT in dystonia patients.
Discussion

Distribution of patients with (CYP 450 2D6) gene polymorphism

Figure (1) shows that the number (percentage) of patients with CYP 450 2D6*2 was 37 (74%), whereas, patients with CYP 450 2D6 *10 was 13 (26%). No patients were observed with CYP 450 2D6 *4 polymorphisms in this study. To avoid tetrabenazine side effects, genotyping tests before the initiation of therapy could identify patients with unacceptable mortality and morbidity risks. The CYP2D6 activity ranges considerably within a population and includes ultrarapid metabolizers (UMs), extensive metabolizers (EMs), intermediate metabolizers (IMs) and poor metabolizers (PMs). The allele *10 give rise to substrate-dependent decreased activity. It is clear that alleles *3, *4, *5, *6 and *7 have no enzyme activity (20). No patients were observed with CYP 450 2D6 *4 polymorphisms in this study.

plasma concentration tetrabenazine, Alpha and Beta dihydrotetrabenazine for all patients:

There has been promoting from monitoring a drug level in plasma of tetrabenazine and its metabolites for hyperkinetic movement disorders and take with precaution in the polymorphism of CYP 450 2D6 enzyme. The plasma concentration of tetrabenazine after one and half hour for therapeutic efficacy suggested as less than 2.5 ng/ml and Alpha and Beta dihydrotetrabenazine 40.5 ng/ml and 25.7 ng/ml respectively (21).

Tetrabenazine is rapidly metabolized via first-pass metabolism into two main metabolites known as alpha and beta-dihydrotetrabenazine (DTBZ). Of these two compounds, alpha-DTBZ is pharmacologically active, whereas beta-DTBZ is pharmacologically inert. Peak plasma concentrations of alpha-DTBZ and beta-DTBZ are achieved within one to 1.5 hours following an oral dose, and these compounds have a half-life of approximately 10 hours. In contrast, TBZ exhibits a half-life of about six hours. DTBZ is further metabolized by CYP2D6 to O-dealkylated DTBZ, which is subsequently excreted via the urine and feces. The metabolites are primarily renal eliminated (22).

There was a significant increase in plasma concentration of alpha and beta-dihydrotetrabenazine in patients with CYP2D6 polymorphism in comparison with those with non_mutant gene-phenotype that is due to the low activity of the enzyme (31.91± 0.56 vs. 52.69± 3.7) (19.96± 0.34 vs. 25.01± 0.97) respectively for chorea patients, while (33.59± 0.58 vs. 52.5± 4.7) (18.25± 0.87 vs. 24.85± 1.1) for dystonia patients, which results in elevation in the plasma concentration of tetrabenazine (2.467±0.086ng/ml vs. 4.657± 0.37) for chorea and (1.932± 0.10 vs. 4.325±0.34) for dystonia.

Effect of tetrabenazine on liver enzymes

The results regarding ALP, ALT, and AST as shown in (Table 2) indicate that a significant elevation in liver enzymes in both groups. Moreover, a non-significant difference in serum TB among groups post-treatment.

The present results indicate that a highly significant effect in both subgroup A and B of both group compared to pretreatment, and statistically significant difference observed after three months between subgroups this increment in groups B related to CYP 450 2D6 polymorphism which lead to increase the drug concentrations.

In this study, highly significant elevations of liver enzymes in the patients compared to pretreatment, indicating hepatotoxicity…

A positive non-significant correlation between tetrabenazine plasma concentrations and ALP in both chorea and dystonia were observed, while it was significantly correlated with ALT and AST in both chorea and dystonia. Regarding alpha dihydrotetrabenazine concentration, a (positive) non-significant correlation was found with ALP, ALT, and AST in both groups.

Consequently, there was no significant correlations were detected between plasma concentrations of tetrabenazine and alpha DTBZ levels and TSB level in both chorea and dystonia

Conclusions

According to the data of the present study, we can conclude that:
1. There was high incidence (26%) of CYP 450 2D6 gene polymorphisms in Iraqi patients with movement disorders and mainly CYP 450 2D6*10 which is intermediate metabolizers (IMs)
2. Higher plasma concentration of tetrabenazine among patient with CYP 450 2D6*10 polymorphism in comparison with patient without CYP 450 2D6 polymorphism.
3. Positive significant correlations were detected between serum levels of tetrabenazine and its metabolites with liver function parameters indicating hepatic toxicity due to tetrabenazine metabolites.
Acknowledgement

The present work was abstracted from PhD thesis submitted to the Department of Clinical Pharmacy, College of Pharmacy, University of Baghdad. The authors are greatly thankful to Baghdad Teaching Hospital, Medical City and AL-Zahraa Teaching Hospital for supporting this project.

Reference

1. Fahn S, Jankovic J. Principal & Practice of Movement Disorders, 1st ed., Churchill Livingstone, Elsevier, Philadelphia, USA; 2007:p. 1-652.
2. "Dystonias Fact Sheet - National Institute of Neurological Disorders and Stroke". Archived from the original on 23 April 2018.
3. Balint B, Bhatia KP. Dystonia: an update on phenomenology, classification, pathogenesis and treatment. Current opinion in neurology. 2014;27(4):468-476.
4. Samuel Frank. Treatment of Huntington’s Disease. Neurotherapeutics. 2014; 11(1): 153-160.
5. Frank S, Jankovic J. "Advances in the pharmacological management of Huntington's disease”. Drugs.2010; 70 (5): 561-571.
6. Jankovic J. Medical treatment of dystonia. Movement Disorders. 2013;28(7):1001-1012.
7. Pettibone DJ, Pflueger AB, Totaro JA. Tetrabenazine-induced depletion of brain monoamines: mechanism by which desmethylimipramine protects cortical norepinephrine. European journal of pharmacology. 1984;102(3-4):431-436.
8. Ingelman-Sundberg M, Sim SC, Gomez A, et al. Influence of cytochrome P450 polymorphisms on drug therapies: pharmacogenetic, pharmacocoeigenetic and clinical aspects. Pharmacol Ther. 2007;116(3):496-526.
9. Kwong EH. A novel genotyping algorithm for the CYP2D6*10 allele in Asians using real-time, rapid-cycle PCR and multiplex PCR (Doctoral dissertation, University of British Columbia).
10. Chan A, Pirrmohamed M, Comabella M. Pharmacogenomics in neurology: current state and future steps. Annals of neurology. 2011;70(5):684-697.
11. Jankovic J, Clarence-Smith K. Tetrabenazine for the treatment of chorea and other hyperkinetic movement disorders. Expert Review of Neurotherapeutics. 2011 ;11 (11) :1509 - 1523.
12. Scott, L. J. Tetrabenazine. CNS drugs. 2011; 25(12): 1073-1085.
13. Whyte, M. P., Kempa, L. G., McAlister, W. H., et al. Elevated serum lactate dehydrogenase isoenzymes and aspartate transaminase distinguish Albers-Schönberg disease (chloride channel 7 deficiency osteopetrosis) among the sclerosing bone disorders. Journal of Bone and Mineral Research.2010; 25(11):2515-2526.
14. Langaee, T., Hamadeh, I., Chapman, A. B., et al. A novel simple method for determining CYP2D6 gene copy number and identifying allele (s) with duplication/multiplication. PLoS One, 2015; 10(1), e0113808.
15. Mehvar R, Jamali F. Concentration-effect relationships of tetrabenazine and dihydrotetrabenazine in the rat. Journal of pharmaceutical sciences. 1987;76(6):461-465.
16. Ramazani A, Rezaei M, Rouhani M. An Applicable Method for the Estimation of Tetrabenazine by Simple RP-HPLC in Tablet Dosage Form. Chemical Methodologies. 2017 Oct 1;1:136-44.
17. Patton C. J, Crouch S .R. Spectrophotometric and kinetics investigation of Berthelot reaction for the determination of ammonia . Annual. Chem 1977; 49(3):464-469
18. Kirsch JF, Eichele G, Ford GC, et al. Mechanism of action of aspartate aminotransferase proposed on the basis of its spatial structure. Journal of molecular biology. 1984;174(3):497-525.
19. Kim WR, Flamm SL, Di Biscaglie AM, Bodenheimer HC. Serum activity of alanine aminotransferase (ALT) as an indicator of health and disease. Hepatology. 2008 ;47(4):1363-1370.
20. Kao TW, Chou CH, Wang CC, et al. Associations between serum total bilirubin levels and functional dependence in the elderly. Internal medicine journal. 2012;42(11):1199-1207.

21. Shu-Feng Zhou. Polymorphism of Human Cytochrome P450 2D6 and Its Clinical Significance Part II. Clinical Pharmacokinetics. 2009; 48(12): 761-804.

22. Scherman D, Jaudon P, Henry JP. Characterization of the monoamine carrier of chromaffin granule membrane by binding of [2-3H]dihydrotetrabenazine. Proceedings of the National Academy of Sciences. 1987;80(2):584-8.

23. Kenney C, Jankovic J. Tetrabenazine in the treatment of hyperkinetic movement disorders. Expert Review of Neurotherapeutics. 2006;6:7-17.