The Effects of Valproic Acid on the Skeletal Development and the Ameliorative Effects of Combined Use of Folic Acid and Vitamin E in Pregnant Albino Rats

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Abstract: Objectives: To evaluate skeletal developmental defects secondary to chronic valproic acid (VPA) use during pregnancy and the extent of ameliorative effect of combined use of vitamin E and folic acid (FA) during VPA therapy. Materials and Methods: Thirty virgin female albino rats were allowed to get pregnant and were divided into three equal groups: Control group received no medications; VPA group received oral VPA 400 mg/kg body weight (BW) starting on gestational day (GD) 1 till GD20 and Prophylaxis group received oral VPA (400 mg/kg BW) and vitamin E 250 mg/kg BW and folic acid 100 µg/kg BW using gastric tube starting on GD1 till GD20. At GD20, uterine horns were examined for resorption sites, alive or dead fetuses. Extracted living fetuses were examined for BW, crown-rump length (CRL), and antero-posterior (AP) and biparietal diameters and for congenital malformations. Results: VPA significantly reduced all body measurements of living dams compared to control dams. Prophylaxis therapy significantly increased BW, CRL and AP skull diameter compared to VPA dams. The GD20 fetus of VPA group showed delayed ossification of skull bones with wide anterior fontanel (AF) and widely separated parietal bones, very small ossification center (OC) for hyoid bone. Caudal vertebrae were unossified or showed very small OC with no evident OC for calcaneous, metatarsal bones or distal phalanges. Fetuses of prophylaxis group showed slightly wide AF than that of control animals and parietal bones are separated to lesser extent than VPA group. Mandible is developed, with an OC for hyoid bone and sternoïde are seen. Six caudal vertebrae and few OC in phalanges are seen, but no OC in calcaneous. Conclusion: VPA chronic administration during pregnancy showed deleterious effects on fetal body measurements and skeletal system development. Concomitant administration of vitamin E and FA significantly ameliorated these changes.

Keywords: Valproic Acid, Folic Acid, Vitamin E, Pregnancy, Skeletal Defects

1. Introduction

Epilepsy is the third most common chronic neurological disorder. Epilepsy has wide-ranging physiologic consequences that arise from seizures and from the use of antiepileptic drugs. The Centers for Disease Control and Prevention estimated that epilepsy currently affects 2.2 million Americans and 65 million people worldwide, while the National Institutes of Health estimates that bipolar disorder affects approximately 2.6% of the U.S. population, with the average onset of the disorder occurring at age 25, the middle of child-bearing years for women (1,2).

Valproic acid (VPA) is a widely-used first-generation antiepileptic drug, prescribed predominantly in epilepsy and psychiatric disorders to control epileptic seizures and regulate the mania associated with bipolar disorder. VPA has good efficacy and pharmaco-economic profiles (3).

Adverse drug reactions are a major public health concern and a leading cause of morbidity and mortality in the world. In the case of antiepileptic drugs (AED), adverse drug reactions constitute a barrier to successful treatment since they decrease treatment adherence and impact patients' quality of life of patients. In late 2009, the Food and Drug Administration released a warning indicating that there is an increased risk for major birth defects, including craniofacial, neural, cardiac and cognitive defects, associated with VPA therapies applied during pregnancy (4).

Valporic acid has its inherent toxicity including
hepatotoxicity and increased risk of fetal malformations if received during pregnancy. There have been 268 individual case safety reports with VPA and fatal outcome, reported a total of 156 fatalities with hepatotoxicity, 31 fatalities with pancreatitis. Hepatotoxicity was disproportionally and most commonly reported in children aged ≤6 years (104/156 reports) but affected children of all ages and polytherapy was significantly more frequently reported for VPA with fatal outcome (58%) compared with non-fatal outcome (34%) (5).

Existing literature shows that pharmacokinetic and pharmacodynamics drug-drug interactions possibly occur with the use of a combination of VPA and the atypical antipsychotics olanzapine and quetiapine (6). Between 30% and 50% of patients with brain tumors first present with a seizure, and up to 30% more will develop seizures later. Therefore, optimal management of these patients requires the use of antiseizure medications; however, VPA, despite its efficacy for seizure control, it may also increase the hematologic toxicity of antineoplastic drugs, presumably by inhibiting their metabolism, and may independently impair hemostasis, which is of some concern for patients who require surgical intervention (7).

During fetal development, in the 4th week, two cartilaginous centers appear on either side of the notochord and extend around it to form the body of the cartilaginous vertebra. A second pair of cartilaginous foci appears in the lateral parts of the vertebral bow, grow backward on either side of the neural tube to form the cartilaginous vertebral arch, and a separate cartilaginous center appears for each costal process. By the 8th week, the cartilaginous arch has fused with the body, and in the 4th month, the two halves of the arch are joined on the dorsal aspect of the neural tube. The spinous process is developed from the junction of the two halves of the vertebral arch. The transverse process grows out from the vertebral arch behind the costal process (8, 9, 10).

The ribs are formed from the ventral or costal processes of the primitive vertebral bows. The ventral ends of the ribs become united by a longitudinal bar termed the sternal plate, which is termed the membranous cranium and from its inner layer, the bones of the skull and the membranes of the brain are developed, while from its outer layer the muscles, blood vessels, true skin, and subcutaneous tissues of the scalp are developed (11).

The notochord extends as far forward as the anterior end of the mid-brain, and becomes partly surrounded by mesoderm and its posterior part corresponds with the basilar part of the occipital bone. Then, the mesoderm extends over the brain-vesicles, and thus the entire brain is enclosed by a mesodermal investment, which is termed the membranous cranium and from its inner layer, the bones of the skull and the membranes of the brain are developed, while from its outer layer the muscles, blood vessels, true skin, and subcutaneous tissues of the scalp are developed (12, 13).

The experimental animal study was designed to evaluate the skeletal developmental defects secondary to chronic use of VPA during pregnancy and the extent of the ameliorative effect of combined use of vitamin E and folic acid during VPA therapy.

### 2. Materials and Methods

#### 2.1. Animals

The study comprised 30 virgin healthy female albino rats, weighing 200-250 gm. Albino rats were chosen for low incidence of spontaneously occurring congenital defects, short-gestational span and high offspring number. Rats were purchased from the laboratories of Ministry of Agriculture, and kept under standard conditions, temperature 20°C, humidity 60% and 12-hs day/night cycle, and maintained on standard diet and free water supply till the start of study regimens.

The study also included 10 male rats used to commit intercourse with female rats to get pregnant. Intercourse was assured by getting vaginal smear and detecting sperms and this was considered as gestation day-1 (GD1).

#### 2.2. Study Protocol

The pregnant animals were divided into three groups (each in a separate cage) according to medication regimen used:

i. Control group included 10 rats kept on normal diet without any medications.

ii. VPA group included 10 rats received valproic acid (Depakene) 400 mg/kg BW orally using gastric tube starting on GD1 till GD20.

iii. Prophylaxis group included 10 rats received oral valproic acid (Depakene) 400 mg/kg BW in addition to vitamin E 250 mg/kg BW and folic acid 100 µg/kg BW orally using gastric tube starting on GD1 till GD20.

#### 2.3. Sample Obtaining and Measurements

At the 20th day of gestation, female rats were ether anesthetized and a midline longitudinal incision was made in the anterior abdominal wall to expose the abdominal cavity. Uterine horns were examined for resorption sites that appeared as small-to-large blood clots which indicate post-implantation death. Uterine horns were incised and alive or dead fetuses were extracted and their number was counted. Extracted living fetuses were examined for body measurements including body weight (BW), crown-rump length (CRL), and antero-posterior (AP) and biparietal diameters using micrometer.

#### 2.4. Technique of Skeletal Staining

Fetuses were put in ethyl alcohol 95% and totally evacuated of internal organs through midline abdominal incision and were skinned by immersion in water path heated up to 70°C for 30 sec and then were placed again in ethyl alcohol 95% for 7 days. Then, animal remnant was cleared in KOH 1% for 1-3 days until bones become clearly visible through surrounding tissues. Animals were stained by immersion for one day in freshly prepared KOH 0.5% to which 10 drops of conc. Alizarin red stain were added, then specimens were decolorized using a solution of mixture of glycerin, KOH 0.5% in distilled water of 20:1:79 volume:volume. Specimens were immersed in glycerin solution of...
ascending concentration 30, 50 and 70% and were preserved in pure glycerin to which a crystal of thymol was added to prevent fungal contamination.

2.5. Evaluation of Ossification

Skeletal ossification was assessed according Burdan et al. (14) as following:

i. Un-ossified: means no ossification as judged by absence of alizarin red stain.

ii. Incomplete ossification: means partial ossification as judged by reduced alizarin red stain.

iii. Miss-happen means abnormally shaped ossification.

2.6. Statistical Analysis

Obtained data were presented as mean±SD, ranges, numbers and ratios. Results were analyzed using Wilcoxon; ranked test for unrelated data (Z-test) and Chi-square test (X² test). Statistical analysis was conducted using the SPSS (Version 15, 2006, SPSS, Inc., Chicago, IL, USA) for Windows statistical package. P value <0.05 was considered statistically significant.

3. Results

Examination of the gravid uteri defined 138 living dams, 11 dead dams and 6 resorption sites. No dead dams or resorption sites were detected in control group, while the frequency of dead dams and resorption sites was significantly (p <0.05) higher in VPA group compared both to control and prophylaxis groups with non-significantly (p>0.05) higher frequency in prophylaxis group compared to control group (Table 1).

|                        | Control group | VPA group | Prophylaxis group | Total          |
|------------------------|---------------|-----------|-------------------|----------------|
| Living dams            | 52 (100%)     | 36 (72%)  | 50 (94.3%)        | 138 (89%)      |
| Dead dams              | 0             | 9 (18%)   | 2 (3.8%)          | 11 (7.1%)      |
| Resorption sites       | 0             | 5 (10%)   | 1 (1.9%)          | 6 (3.9%)       |
| Total                  | 52 (100%)     | 50 (100%) | 53 (100%)         | 155 (100%)     |
| Statistical significance| P1= 0.001     | P2=0.05   | P3=0.034          |                |

Data are presented as numbers; percentages are in parenthesis; P1: significance of difference between control and VPA groups; P2: significance of difference between control and prophylaxis groups; P3: significance of difference between prophylaxis and VPA groups

Exposure to VPA significantly (p<0.05) reduced all body measurements of living dams including body weight (BW), crown-rump length (CRL) and skull dimensions compared to control dams. Prophylaxis therapy ameliorated these development retardation effects of VPA manifested as significant (p<0.05) increase of BW, CRL and AP skull diameter with non-significant (p>0.05) increase of biparietal skull diameter compared to VPA dams. Moreover, dams of prophylaxis group showed non-significant (p>0.05) reduction of CRL and biparietal skull diameter compared to control dams, despite of the significant (p<0.05) reduction of BW, and skull AP diameter (Table 2, Fig. 1-4).
3.1. Assessment of Bone Staining

3.1.1. Control Group

The 20-day fetus of control rat showed normal ossification of skull bones with small anterior fontanel (Fig. 4a), well-developed ossified mandible, hyoid bone, 6 sternebrae and vertebrae (Fig. 4b). Appendicular skeleton of the fore and hind limbs appeared ossified with apparent small ossification centers for calcaneous, metatarsal bones and distal phalanges (Fig. 4c)

3.1.2. VPA Group

The 20-day fetus of experimental rat of VPA group showed delayed skeletal development shown as delayed ossification of skull bones with wide anterior fontanel and widely separated parietal bones (Fig. 5a), very small ossification center for the hyoid bone (Fig. 5b). Caudal vertebrae were either unossified or showed very small ossification centers. Moreover, no evident ossification centers for calcaneous or metatarsal bones and distal phalanges were found (Fig. 5c)

3.1.3. Prophylaxis Group

The 20-day fetus of experimental rat of prophylaxis group showed improved skeletal development shown as slightly wide anterior fontanel than that of control animals and both parietal bones are separated but to lesser extent than animals of VPA group (Fig. 6a). Mandible is developed, with an ossification center for hyoid bone and 5 sternebrae are seen (Fig. 6b), while the 5th sternebra in addition to a small ossification center in calcaneous is seen (Fig. 6c). Six caudal vertebrae and few centers in phalanges are seen, but no ossification center in calcaneous (Fig. 6d).

Table 2. Body measurements of living dams of studied groups.

|                      | Control group | VPA group | Prophylaxis group | Statistical significance |
|----------------------|---------------|-----------|-------------------|-------------------------|
| Birth weight (gm)    | 4.1±0.47      | 2.98±0.62 | 3.59±0.39         | P1=0.008                |
|                      |               |           |                   | P2=0.036                |
|                      |               |           |                   | P3=0.010                |
| Crown-rump length (cm) | 4.8±0.5      | 3.5±0.53  | 4.4±0.51          | P1=0.003                |
|                      |               |           |                   | P2=0.139                |
|                      |               |           |                   | P3=0.005                |
|                      |               |           |                   | P1=0.009                |
|                      |               |           |                   | P2=0.011                |
|                      |               |           |                   | P3=0.012                |
| AP diameter (cm)     | 1.98±0.32     | 1.46±0.25 | 1.73±0.2          | P1=0.015                |
|                      |               |           |                   | P2=0.05                 |
|                      |               |           |                   | P3>0.05                 |
| Biparietal diameter (cm) | 1.52±0.16   | 1.3±0.22  | 1.45±0.17         | P1=0.008                |
|                      |               |           |                   | P2=0.036                |
|                      |               |           |                   | P3=0.010                |

Data are presented as mean±SD; P1: significance of difference between control and VPA groups; P2: significance of difference between control and prophylaxis groups; P3: significance of difference between prophylaxis and VPA groups
Control group

VPA group

Prophylaxis group

**Fig. 4a.** A photograph of 20-day fetus of control group showing ossified skull bones; parietal (P), inter-parietal (I), supraoccipital (So) and exoccipital (Ex) bones and anterior fontanelle. Alizarin red x8.

**Fig. 5a.** A photograph of 20-day fetus of rat of AED group showing large anterior fontanelle (AF) with separation of the two parietal bones (P). Alizarin red x4.

**Fig. 6a.** A photograph of 20-day fetus of rat of prophylaxis group showing anterior fontanelle which is slightly wider than that of control animals. Both parietal bones are separated but to lesser extent than animals of AED group. Alizarin red x4.

**Fig. 4b.** A photograph of 20-day fetus of control group showing well-developed ossified mandible (M), hyoid bone (H), clavicle (Cl) and 6 sternebrae. Alizarin red x4.

**Fig. 5b.** A photograph of 20-day fetus of rat of AED group showing very small ossification center for hyoid bone (H), 5 very small vertebrae with only 4 small ossification centers for caudal vertebrae (Cv). No ossification center for calcaneous was seen (empty ring). Alizarin red x4.

**Fig. 6b.** A photograph of 20-day fetus of rat of prophylaxis group showing developed mandible, an ossification center for hyoid bone (H) and 5 sternebrae are seen. Alizarin red x8.
Fig. 4c. A photograph of 20-day fetus of control group showing ossified ribs (R), vertebrae (8 caudal vertebrae are seen). Ossified appnedicular system was shown as ossified femur (F), tibia (T), fibula (Fi). Ossification centers for calcaneous (C), metatarsal bones (Mt) and phalanges (Ph) Alizarin red x4.

Fig. 5c. A photograph of 20-day fetus of rat of AED group showing two very small ossification centers for caudal vertebrae (Cv). No ossification centers for calcaneous or phalanges were seen (empty rings). Alizarin red x4.

Fig. 6c. A photograph of 20-day fetus of rat of prophylaxis group showing absence of 5th sternebra and only 5 are shown. A small ossification center in calcaneous is seen. Alizarin red x3.

Fig. 6d. A photograph of 20-day fetus of rat of prophylaxis group showing 6 caudal vertebrae, few centers in phalanges, but no ossification center in calcaneous. Alizarin red x4.
4. Discussion

The obtained results illustrated the effect of valproic acid (VPA) if used during pregnancy manifested firstly as significant reduction of birth weight (BW) and crown-rump length (CRL) with diminution of skull dimensions of studied fetuses of VPA group compared to control group. However, combined prophylactic therapy using vitamin E and folic acid (FA) could ameliorate these effects of VPA allowing better development manifested as more weight gain and increased CRL and skull dimensions.

Secondly, VPA affected ossification and development of axial and appendicular bones manifested as reduced hyoid bone ossification center, hypoplastic mandible, reduced or absent centers for sternebrae, delayed caudal vertebral ossification and absence of ossification centers for calcaneous and metatarsals. Prophylactic therapy improved these changes with well-development of mandible, increased number of centers for sternebrae and appearance of ossification centers of calcaneous and metatarsals.

These findings go in hand with that previously reported in literature concerning the use of anti-epileptic drugs (AED) during pregnancy and with trials to evaluate prophylactic use of FA &/or antioxidants. In a similar study; Baran et al. (15) found administration of single VPA dose (300 mg/kg) resulted in weight and length loss between control and VPA groups, however, length and weight differences between groups received vitamin E or folate were not significant. Ornoy (16) documented that exposure to VPA in pregnancy is associated with approximately three-fold increase in the rate of major anomalies, mainly spina bifida and only rarely anencephaly, cardiac, craniofacial, skeletal and limb defects and a possible set of dysmorphic features, the "valproate syndrome" with decreased intrauterine growth.

Soysal et al. (17) using 25mg/kg/day phenytoin administered intraperitoneally to pregnant rats detected that fetuses were 14% shorter and the weights 13% lower compared to control animals with decreased number of the fetuses obtained in one gestation by 9%, and deteriorated ossification of skull bones, costal separation anomaly, the laterally-separated costal components were not attached to the costal arch with shape malformations in the last two ribs and wide angularity, particularly in the last six ribs.

Fadel et al. (18) detected significant increases in skeletal abnormalities, particularly in ribs in topiramate-exposed fetuses at therapeutic doses compared to control group with a direct correlation between fetal BW and the number of complete ossified vertebral centers. Koo & Zavras (19) indicated that various forms of VPA and newer generation AED agents show signals for congenital jaw or oral malformation.

Recently, in 2015; Tanoshima et al. (20) documented that at present, risks of VPA-associated congenital malformations are 2 to 7 fold higher than other common antiepileptic drugs, and so VPA should not be used as a first-line therapy in women of childbearing age unless it is the only option for the patient.

In line with the use of antioxidant and FA therapy for reduction of the frequency and severity of AED-induced teratogenic effects; Al Deeb et al. (21) found concomitant vitamin E administration significantly attenuated VPA induced decrease in fetal weight, crown rump length and malformations. Padmanabhan & Shafiullah (22) documented that plasma levels of FA and vitamin B12 must be kept substantially elevated and maintained high throughout organogenesis period to protect embryos against VPA-induced neural tube defects. Dawson et al. (23) hypothesized that FA and pantothenic acid protect embryos from VPA-induced neural tube defects possibly through prevention of VPA-induced alterations in proteins involved in neurulation. Also, Hsieh et al. (24) documented that simultaneous multiple prescriptions of FA, ascorbic acid, and N-acetyl cysteine are suggested to rescue the VPA-induced teratogenicity. Tunali et al. (25) and Gezginici-Oktayoglu et al. (26) showed that Vitamin E exerted antioxidant properties and may prevent lens (25) and kidney (26) damage caused by VPA.

Multiple studies tried to explore the mechanisms underlying the teratogenic risk of AED exposure and the prophylactic effects of vitamin E &/or FA; Ornoy et al. (16) reported that specific inhibition of histone deacetylase and changes in gene expression by VPA may explain its teratogenicity with other possible explanations as increased fetal oxidative stress induced by VPA, with the brain being more susceptible to oxidative stress in comparison to other fetal organs, or FA inhibitory action of VPA. Tung & Winn (27) experimentally found VPA exposure resulted in an increase in reactive oxygen species (ROS) staining in the head region and increased ROS levels were correlated with increased staining for apoptotic markers, and attributed developmental defects in head and neck region to these increased ROS levels.

Mallela & Hrubec (28) reported increased apoptosis as an underlying pathogenic mechanism involved in VPA teratogenicity, suggested that VPA may alter other cellular processes such as cell proliferation and differentiation in addition to apoptosis and concluded that VPA alters several biological processes in the developing embryos to induce fetal malformations. Umur et al. (29) suggested that VPA may induce apoptotic mechanisms but not through the p53 pathway and FA effectively prevents the teratogenic influence of VPA on chick embryo at neurulation stages by stopping cascade of apoptosis before caspase 3 expression.

Tiboni et al. (30) reported that after VPA (400 mg/kg) treatment 35.2% of fetuses exhibited skeletal teratogenesis, but this rate was significantly increased to 53.7% after nitric oxide synthase inhibitor N(G)-nitro-L-arginine methyl ester co-administration. Thereafter, in 2015; Tiboni & Ponzano (31) detected axial skeletal defects in 75.8% of VPA-exposed fetuses, but pre-treatment with sildenafil citrate which prolongs nitric oxide signaling significantly decreased the rate of skeletally affected fetuses to 61.6%.

Recently, in 2015; Rubinchik-Stern et al. (32), incubated human trophoblastic cells with multiple AED and found that...
compared with controls, AED-treated cells displayed half the levels of folate placental uptake carriers mRNA and protein and up to 2.7-fold increases in efflux transporters mRNA and protein expression, together with enhanced efflux transporters' activity and altered to a modest extent the transcription of nuclear receptors known to regulate transporter expression.

The obtained experimental results supported the clinical recommendations to judicious use of AED during pregnancy; the most recent are provided by the Task Force of the Commission on European Affairs of the International League Against Epilepsy and the European Academy of Neurology[33] that where possible, VPA should be avoided in women of childbearing potential, VPA should not be prescribed as a first-line treatment for focal epilepsy unless pregnancy is highly unlikely or it is the most effective treatment and the risks and benefits of VPA and other treatment alternatives should be discussed with the patients.

Moreover, Tanoshima et al.[30] performed meta-analyses of cohort studies to determine the time profiles of signal emergence and risk estimates of VPA-associated congenital malformations and identified 59 studies documented that risks of VPA-associated congenital malformations are 2-7 fold higher than other common antiepileptic drugs so it should not be used as a first-line therapy in women of childbearing age unless it is the only option for the patient.

It could be concluded that VPA chronic administration during pregnancy showed deleterious effects on fetal body measurements and skeletal system development. However, concomitant administration of vitamin E and FA showed significant ameliorative effect on these changes so, if it is strongly indicated, VPA can be used during pregnancy in combination with both vitamin E and FA.

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