Research Article

Characterization of four types of tail abnormalities in rats treated prenatally with valproic acid

Caracterización de cuatro tipos de anormalidades en la cola de ratas tratadas prenatalmente con ácido valpróico

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Received: June 10, 2014 Accepted: July 07, 2014
Find this paper at: http://www.uv.mx/enneurobiologia/vols/2014/9/9.html

Abstract
Valproic acid (VPA) is an anticonvulsant drug used mainly for the treatment of epilepsy, bipolar disorder and schizophrenia. The VPA has been shown to be a potent teratogen that causes birth defects and malformations. Likewise, there are reports that suggest a relationship between the use of VPA during pregnancy and an increased incidence of children with neurological disorders such as autism. In humans, prenatal exposure to VPA produces malformations including dimorphic facial features suggestive of a lesion in the neural tube, as in the case of spina bifida, heart disease, limb defects and craniofacial anomalies as well as genital abnormalities. Herein we describe four tail abnormalities found in rats treated prenatally with valproic acid, which has been used as an animal model for the study of autistic features. These malformations may be associated with neural damage, but further studies are needed in order to correlate each tail abnormality with the kind of neural alteration.

Key words: Teratogen, Valproic acid, Animal model, Malformations, Neural tube defects.

Resumen
El ácido valpróico (AVP) es un fármaco anticonvulsivo usado principalmente para el tratamiento de la epilepsia, trastorno bipolar y esquizofrenia. Se ha demostrado que el AVP es un teratógeno potente que causa defectos de nacimiento y malformaciones. Así mismo, hay informes que sugieren una relación entre el uso de AVP durante el embarazo y un aumento de la incidencia de niños con trastornos neurológicos tales como el espectro autista. En los seres humanos, la exposición prenatal a la AVP produce malformaciones que sugieren un defecto durante el cierre del tubo neural, como es el caso de la espina bífida, enfermedades del corazón, defectos de las extremidades, anomalías craneofaciales, así como anomalías genitales. En el presente trabajo se describen cuatro anormalidades de la cola en ratas tratadas prenatalmente con ácido valpróico. Este tratamiento se ha utilizado como modelo animal para el estudio de rasgos autistas. Estas malformaciones pudieran estar asociadas con algún daño en el sistema nervioso, sin embargo se requieren más estudios para correlacionar cada anormalidad de la cola con el tipo de alteración neural.

Palabras clave: Teratógeno, Ácido valpróico, Modelo animal, Malformaciones, defectos del tubo neural.

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1. Introduction

Valproic acid (VPA) was first synthesized in 1882 by B. S. Burton, and is commonly used as both an antiepileptic drug and in some cases to treat personality disorders such as schizophrenia and bipolar disorder, or migraine. Its name is derived from the original purification of *Valeriana officinalis*, which is commonly used to treat insomnia and anxiety.

Shortly after VPA was out in the market, it was demonstrated to cause various malformations in newborns of mothers that were administered VPA during pregnancy. A recent study revealed the association of the treatment with anticonvulsants with at least 14 types of malformations. Interestingly, treatment with VPA caused significantly more malformations as compared to carbamazepine, the main anticonvulsant used in humans for the treatment of seizures. These types of malformations have been grouped in a fairly specific embryopathy called fetal valproate syndrome. The percentages of malformations in humans are: 12.7% spina bifida, 2.5% atrial septal defect, 5.2% cleft palate, 4.8% hipospadix, 2.2% polidactyly, 6.8% craniophalangeal synostosis.

Children with prenatal exposure to VPA showed malformations similar to those exposed to thalidomide *in utero*, but with a reduced severity of the symptoms. These included abnormal features indicative of an injury at the time of neural tube closure, such as heart disease, craniofacial abnormalities, genital abnormalities and limb defects, and spina bifida which is the most frequent disorder related to incomplete closing of the neural tube. Some teratogenic effects of VPA have been also been reported in animals. For example, mice exposed *in utero* to VPA can develop cardiovascular malformations, limb defects, skeletal malformations, and cerebellar abnormalities such as significantly fewer Purkinje cells in the cerebellar vermis.

In addition to its teratogenic effects, the use of VPA in pregnant women has been associated with a higher incidence of autism in the offspring. A recent study reported that 60% of children who were prenatally exposed to VPA had more than two autistic characteristics but only 11.7% of them met the diagnostic criteria of the autistic spectrum. To our knowledge and despite the information presented above, physical malformations in the rat exposed to VPA *in utero* remain undescribed. Therefore, in the present study we assessed the teratogenic effects of VPA upon the physical condition of the rat body.

2. Materials and methods

**Animals**

A total of 6 adult females and 6 males, of healthy Wistar rats were used under laboratory conditions. Inverted cycle 12/12h light/dark cycle, (lights off at 08:00 h), food (Rodent Diet, Rismart nu3-225) and fresh water were provided *ad libitum*.

We determined the late proestrus of every female rat through vaginal cytology to establish the moment in which females were receptive and able to copulate with a male. The couple was left undisturbed for a whole day and then females were isolated in single home cages with a thin layer of aspen chip. The day of isolation was considered as day 1 of gestation. Pregnant rats received a single intraperitoneal injection of 600mg/kg of sodium valproate (NaVPA) on gestational day 12. The expected day of delivery animals were monitored and pups were identified by sex and grown until postnatal day 21. Tail malformations were reported until day 21 when it was easy to classify them.

All procedures involving animals were reviewed by and conducted in compliance with the guidelines of Centro de Investigaciones Cerebrales, Universidad Veracruzana, Xalapa, Mexico, based on the standard of Official Mexican Norm NOM-062-ZOO-1999.

3. Results

A total of 72 pups whose mothers were treated with VPA were obtained from the 6
litters, in which 25 were observed with a tail abnormality. Hence, 34% of animals showed abnormalities. We also did not find any other body malformation besides tail (Figure 1).

Tail description

Four types of tail malformations were observed and each tail type was established according to the distance from the crooked point in the tail to the rat’s body, so the tales were divided in 3 sections: 1) proximal, 2) medial, 3) distal.

Sex differences

We found 16 male and 9 female pups with tail abnormalities out of the 25 total tail bended pups. The proportion was 64% and 36% respectively (Figure 2).

Tail classification

The tail abnormalities were divided in 4 categories according with the features of the length curve and bending shape tail. The categories were: Short tail (A), simple bent in the middle (B), short and bent tail (C), double flexure tail (D) (Figure 3).

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Figure 3. Categories of bent tails. A Short tail, B simple bent in the middle, C short and bent tail, D double flexure tail.

Figure 4. Types of tail abnormalities by sex.
4. Discussion

The results confirm that prenatal exposure to valproic acid at critical stages produce teratogenic effects. Specifically, abnormal tail were classified into four groups, the tail as A, B, C, D. The anomalies found in this report illustrate the potent teratogenic effect of valproic acid in these prenatally treated animals. Considering that the method used in this study was first reported as an animal model of autism, a disorder with a frequency ratio of 4:1 between human males and females, in this sense it is interesting is that in general the found abnormalities were more frequent in males than in females, so may be due to a protective mechanism against VPA present in females. On other hand, studies about motor behavior level should consider these possible anomalies during test batteries to validate as an animal model of autism, since that abnormalities in organs such as the tail may cause deterioration in functions such as thermoregulation, as the tail of the rat is a body heat loss; this is due to its length and width, in addition, it lacks skin and nutrients and has not vascularized connections between arteries and veins. Another consideration is related to the equilibrium, rats used to distribute weight while climbing or walking through an edge surface. Although the mechanisms of teratogenic action of VPA are not yet known and the explanation thereof is beyond the present study, our results originate outlook study possible interactions of prenatal treatment with VPA and genes as RORA which has associated with the higher proportion of males and females with autism due to the higher proportion of abnormalities in males than in females (Figure 2). Also allows exploring the hormonal connection because of VPA side effects on steroid hormone levels could be distributed not uniformly across the placenta of embryos, and the differences in the tails shape could be due intrauterine position of the fetus. This may have a greater effect in males, as observed in this work. In fact, androgen prenatal exposure affects anogenital distance which is a feature of sexual dimorphism as well fingers length. As similar process occur with drugs such as cocaine therefore, future studies could determine whether intrauterine position leads to different levels of prenatal exposure to VPA, like other drugs.

5. Conclusion

Valproic acid affects normal embryonic development. When an embryo is exposed to VPA at the time of neural tube closure, has a high risk of malformations as twisted and / or bent tail. These malformations are observed in four different types and are more frequent in males. This phenotype could be an indicator of alterations in the central nervous system. These defects must be considered in studies of prenatal treatment of VPA as an animal model of autism, its possible relationship to features of the spectrum and its influence upon assessments for batteries and motor behavioral tests. Further studies are needed to determine the relationship of the anomalies in appendages and structures of the central nervous system.

6. Acknowledgements

Centro de Investigaciones Cerebrales Universidad Veracruzana Xalapa México CONACyT scholarship 235940 (PSL) 240116 (BBA) Promep Grant 103.5/07/2753 (RT).

7. References


