Abstract

Although methamphetamine (METH) is the most commonly abused drug during pregnancy, there is surprisingly little data on METH effects on gestational tissues. This study, building on our recent research describing the effects of METH on antioxidant status and catecholamine signaling in gestational tissues, assesses serotonin levels in the umbilical cord during pregnancy and effects of demographics and neonatal variables. We hypothesized that METH ingestion and/or smoking may alter serotonin signaling in the umbilical cord. Umbilical cords (n = 85) were collected, processed to total lysates and standardized to 2 mg/mL protein. Serotonin levels were determined by ELISA then correlated to different demographic and pregnancy variables, antioxidant and catecholamine levels then the effects of METH and/or smoking determined. Levels of serotonin in umbilical cords were not significantly different between METH users nor smokers and control cords. Additionally, there were no significant correlations with delivery type, parity, gravidity neonatal baby weight, or maternal age. Serotonin levels were positively correlated with maternal diabetes (P=0.04) and negatively correlated with Gestational age (P = 0.05, r = 0.2310) and placental weight (P = 0.04, r = -0.3527). Serotonin levels may also be affected by maternal hypertension (P = 0.06) as well as preeclampsia (P = 0.08). While umbilical serotonin was not related to levels of noradrenaline and dopamine nor to the antioxidant enzymes glutathione-S-transferase, catalase, super-oxide dismutase or xanthine oxidase levels; serotonin increased as thioredoxin reductase (TRX, P = 0.0003, r = 0.40) and glutathione peroxidase (GPx, P = 0.01 , r = 0.28) activities increased. A potential association between serotonin and with Vitamin C levels was also noted (P = 0.07, r = 0.2942). When stratified for METH and smoking, patients with diabetes showed that umbilical serotonin may be further lowered by smoking but not METH (P = 0.06 non smokers vs. smokers, P = 0.08 smokers vs. Meth+smokers). Furthermore, THC caused a significantly higher gravida (P = 0.01) than non-smokers. When stratified, smokers showed a significantly higher chance in getting pregnant than those that do not smoke (P=0.0002). Those that smoke in addition to THC, however, have a significantly lower chance of getting pregnant than those that only smoke (P=0.0173). THC also suggestively causes a lower neonatal baby weight (P = 0.02). Whilst THC did not show any significant differences in CAT levels with respect to different types of smokers, those who smoke THC have significantly higher levels of Vitamin C (P = 0.0383) and a significantly lower level of GPx activity than those who do not smoke (P = 0.05) and a significantly higher level of noradrenaline (P = 0.02). Serotonin signaling in umbilical cords is affected by hypertension and diabetes as well as oxidative stress, which may itself be caused by the two clinical syndromes.

Keywords: Serotonin; Fetal; Physical abnormalities; Phenylephrine

Abbreviations: METH: Methamphetamine; THC: Tetra Hydro Cannabinol

Introduction

Any compound ingested during pregnancy can, depending on its physico-chemical characteristics, pass into the fetal circulation. After transiting the placenta in blood, chemicals emerging into the fetal circulation are delivered directly to the fetal heart by the umbilical artery. The fetal heart bypasses the pulmonary circuit and pumps blood around the fetal circulation then returns de-oxygenated blood carrying metabolic wastes to the placenta through the umbilical vein [1]. Hence, the umbilical cord plays a vital role in both delivering and removing oxygen, essential nutrients and other chemicals to and from the fetus. Therefore, the placental and umbilical vessels are, also the major route for harmful drugs and chemicals to reach the fetus. These harmful chemicals may cause direct deleterious fetal effects and may also interfere with placental or umbilical function thereby disrupting with the delivery and removal of blood and other essential substances [1].

Most physical birth defects occur during the embryonic period (up to 8 weeks from conception) [2] when the embryo is differentiating and organ structure developing [3]. Also during the embryonic period, the umbilical cord and placenta are not
Does Methamphetamine Abuse Affect Serotonin Signaling in the Maternal-Placental-Fetal System?

Tamayo RAC, Wright TE, Collier AC (2015) Does Methamphetamine Abuse Affect Serotonin Signaling in the Maternal-Placental-Fetal System? Pharm Pharmacol Int J 2(6): 00042. DOI: 10.15406/ppij.2015.02.00042

Citation: Tamayo RAC, Wright TE, Collier AC (2015) Does Methamphetamine Abuse Affect Serotonin Signaling in the Maternal-Placental-Fetal System?

Pharm Pharmacol Int J 2(6): 00042. DOI: 10.15406/ppij.2015.02.00042

fully functional and cannot detoxify substances transiting from the maternal circulation [3]. After the embryonic period, physical abnormalities of teeth, palate and ears can occur up to 16 weeks [2]. Furthermore, functional and physical abnormalities of the eyes, central nervous system and genitalia can be caused at any time from conception to term [2].

Most women do not know that they are pregnant until the second missed menstrual cycle, hence; often the embryonic period has passed without any conscious knowledge or change in behavior in the mother [4]. Consequently many women accidentally continue recreational or medical drug use and keep ingesting foods and other substances that they would otherwise avoid. In other cases, some mothers either do not (by choice) or cannot (through addiction or medical need) stop their ingestion of potentially harmful substances [4]. Methamphetamine (METH) and to a lesser extent Marijuana addictions belong to the latter case where, more often than not, the mother is physically and/or psychologically unable to stop ingesting these drugs. METH use in pregnancy is associated with a number of substandard reproductive and pregnancy outcomes including pre-eclampsia, pre-term birth, intra-uterine growth restriction, small-for-gestational-age babies and neonatal addicts. Other studies have reported that marijuana use in pregnancy is also associated with low birth weight and small for gestational age infant, as well as preterm delivery; other relationships involve affecting the neural and morphological development of the fetus [5].

Both METH and the active ingredient in marijuana (d9-tetrahydrocannabinol, THC) have interactions with serotonin receptors and serotoninergic signaling. As early as 1974 interactions between amphetamine effects and serotoninergic signaling were being reported [5]. Amphetamines are potent inhibitors of 5HT binding in the brain [6] but little is known about METH and 5HT interactions in the maternal-placental-fetal unit. The one study in this area demonstrated that amphetamines can inhibit placental monoamine transporters [7] but there is a dearth of knowledge regarding placental, umbilical or fetal interactions between METH and 5HT. Similarly, it has been shown that cannabinoids can directly inhibit the 5HT3 and 5HT1a receptors [8,9] and that patients with a clinical deficiency in endocannabinoids and to a lesser extent cross talk between THC and 5HT signaling, may explain therapeutic benefits of marijuana use in some syndromes [10]. However, investigations of the effects of THC on umbilical cord serotonin levels and pregnancy outcomes are likewise, absent from the literature.

It has long been known that serotonin and other compounds that bind serotonin receptors (such as histamine) an alter tone in the umbilical artery and vein [11]. The human umbilical artery and vein have both been reported to express 5HT1D beta, 5-HT2A, 5-HT2B, 5-HT4 and 5-HT7 [12,13]. It has subsequently been demonstrated that only 5HT1B/D and 5HT2A cause functional contraction of the human umbilical artery and vein [14-16]. Serotonin is of interest for altering umbilical contractility since studies have shown that noradrenaline, the predominant sympathetic neuro transmitter, as well as phenylephrine and clonidine can only cause contraction in the umbilical vein of 9%, 14% and 11% respectively of the maximum caused by 5-HT [16]. In the umbilical artery, the maximal contractility caused by noradrenaline and phenylephrine was higher, but still only 18% and 29% of that for 5-HT and clonidine had no effect [16]. Additionally, histamine and 5HT produced strong, dose-dependent vasoconstrictions but epinephrine and norepinephrine cause only slight vasoconstriction [17]. Meanwhile early studies also demonstrated that the human fetal heart can respond to serotonin [18] Subsequently, 5HT2B and 5HT4 receptors have been identified in fetal heart structures [19,20]. Evidence suggests that the responses of the umbilical endothelia to 5HT are oxygen-dependent and may only occur under conditions of low oxygen tension [21]. This raises interesting questions for pregnancy conditions where oxidative stress may be present such as during pre-eclampsia and/or when medial or illicit drugs with highly oxidative properties (such as COX-2 inhibitors or METH) are ingested during pregnancy.

In addition to providing a conduit for METH to directly reach the fetus, changes in vascular tone in the umbilicus may adversely affect fetal development by restricting the delivery of essential compounds and/or removal of wastes. Secondly, since the umbilical artery leads directly to the heart, if METH causes high levels of catecholamines to be produced in the umbilical artery, may adversely affect the fetal cardiovascular system. Whilst the net effect of this is unlikely to alter fetal circulatory dynamics, since changes in heart muscle dynamics tend to be balanced by changes in vascular tone [22]. The implications of higher levels of catecholamines produced in the umbilical artery and delivered directly to the fetal heart include increased work by the heart and the potential for dysrhythmias, which may cause fetal distress and/or death. Finally, since catecholamines affect gland cells and change neuro-endocrine signaling [22]. METH may affect normal endocrine signaling in the developing fetus, either through direct delivery of METH to fetal endocrine tissues or through increased catecholamine release in the umbilical cord which are then delivered to fetal tissues. In this paper we have assessed the effects of METH ingestion, Marijuana and tobacco smoking, independently and together on serotonin in umbilical endothelia. We hypothesized that ingestion of METH, THC and/or tobacco smoke may cause increased production of serotonin or alter serotonergic signaling in the umbilical vessels.

Materials and Methods

Tissue collection and processing

One hundred umbilical cord pieces (10 cm) were collected with informed consent from women undergoing Caesarian section at term. 85 were available for this study. Each piece of cord was coded with a study number, then stored at -80 °C until use. This study was approved by the University of Hawaii Institutional Review Board for Human Subjects and the Kapiolani Hospital Ethical Review Board.

For processing to lysates, cord pieces were thawed, wet weight recorded then cords homogenized mechanically with an IKA Labortechnik Laboratory homogenizer (XXX, Germany) at 6,500 RPM for 30 s each in a 1:4 (w:v) Tris-HCl buffer containing 5 mM MgCl2 and 2 mM PMSF (pH 7.4). Total lysates were normalized to 2 mg/mL protein using the Bicinchoninic acid method, then aliquoted and frozen at -80 °C until use [23]. ELISA for catecholamine neurotransmitters and serotonin. Levels of

Citation: Tamayo RAC, Wright TE, Collier AC (2015) Does Methamphetamine Abuse Affect Serotonin Signaling in the Maternal-Placental-Fetal System? Pharm Pharmacol Int J 2(6): 00042. DOI: 10.15406/ppij.2015.02.00042
serotonin were determined with a commercial Serotonin ELISA from Rocky Mountain Diagnostics (Colorado Springs, CO) as per the manufacturer's instructions.

Data analysis and statistics

Quantification of serotonin was performed blinded using only the study number for identification of samples. After ELISAs were completed and concentrations assigned, the study key was accessed and results were sorted into four groups: cords from non-smoking, non-METH patients (N), cords from METH using patients (M), cords from tobacco smoking patients (S), cords from THC positive patients (T) and cords from patients who both smoked tobacco and used METH (T/M), cords from those who both smoked tobacco and were THC positive (T/S) and cords from mothers who used all three (M/T/S). Data on METH and tobacco status were based on self reported METH or smoking status and were also confirmed with ELISA detection of METH parent drug and metabolites and/or cotinine levels in umbilical cord lysates [24]. Statistical analyses were performed using Prism 5.0 with significance set at \( \alpha = 0.05 \). (GraphPad Prism, San DiegoCA).

Since these data are discrete, two-tailed student t-tests were used to assess differences between groups. Stratification and comparisons between groups were analyzed using multi-variate analyses [25].

Results

When levels of serotonin in umbilical cords were compared with demographic variables, there were no significant differences between METH users nor smokers and control cords (figure not shown). Additionally, there were no significant correlations with delivery type, parity, gravidity, neonatal baby weight, or maternal age (figure not shown). Serotonin levels were, however, positively correlated with maternal diabetes (\( P = 0.04 \)) and negatively correlated with Gestational age (\( P = 0.05, r = 0.2310 \); Figure 1A & 2A, respectively). Moreover, Serotonin levels were negatively correlated to placental weight (\( P = 0.04, r = 0.3527 \); Figure 2B).

Serotonin levels may also be affected by maternal hypertension (\( P = 0.06 \)) as well as Preeclampsia (\( P = 0.08 \); Figure 1B & 1C, respectively). Serotonin levels are suggestively not related to levels of noradrenalin and dopamine nor to antioxidant enzymes glutathione-S-transferase, catalase, super-oxide dismutase, or xanthine oxidase levels (figures not shown); serotonin increased as thioredoxin reductase (TRX, \( P = 0.0003, r = 0.40 \)) and glutathione peroxidase (GPx, \( P = 0.01, r = 0.28 \)) activities increased (Figure 2D & 2E, respectively). There is also a potential association between serotonin and with Vitamin C levels and serotonin in levels (\( P = 0.07, r = 0.2942 \); Figure 2C). When stratified for METH and smoking, patients with diabetes showed that umbilical serotonin may be further lowered by smoking but not METH (\( P = 0.06 \) non smokers vs. smokers, \( P = 0.08 \) smokers vs. Meth+smokers; figures not shown).

Furthermore, THC caused a significantly higher gravida (\( P = 0.01 \)) than non-smokers (Figure 3). When stratified, smokers showed a significantly higher chance in getting pregnant than those who do not smoke (\( P = 0.0002 \); Figure 4). Those that smoke in addition to THC, however, have a significantly lower chance of getting pregnant than those that only smoke (\( P = 0.0173 \), Figure 4). THC also suggestively causes a lower neonatal baby weight (\( P = 0.02 \), Figure 5). Whilst THC did not show any significant differences in CAT levels with respect to different types of smokers, those who smoke THC have significantly higher levels of Vitamin C (\( P = 0.0383 \), Figure 6), a significantly lower level of GPx activity than those who do not smoke (\( P = 0.05 \), Figure 6) and a significantly higher level of noradrenalin (\( P = 0.02 \), Figure 7). Serotonin signaling in umbilical cords is affected by hypertension and diabetes as well as oxidative stress, which may itself be caused by the two clinical syndromes.

Citation: Tamayo RAC, Wright TE, Collier AC (2015) Does Methamphetamine Abuse Affect Serotonin Signaling in the Maternal-Placental-Fetal System? Pharm Pharmacol Int J 2(6): 00042. DOI: 10.15406/ppij.2015.02.00042
Does Methamphetamine Abuse Affect Serotonin Signaling in the Maternal-Placental-Fetal System?

**Figure 2:** Serotonin levels in the Antioxidant Defense Network and Pregnancy variables. 2A, 2D, 2E: Gestation (P = 0.05), TRx (P = 0.0003) and GPx (P = 0.0132) levels increase with serotonin levels. 2B: Serotonin levels significantly decrease as placental weight increases (P = 0.0441). 2C: Vitamin C levels may increase with serotonin levels (P = 0.0772). Significance was assigned using correlation, linear regression P values and 95% Confidence lines.

**Figure 3:** Gravida levels with respect to TCH. Those who smoke THC have a significantly higher chance of getting pregnant than those who do not smoke THC (P = 0.0149). Significance was assigned using T-tests (α = 0.05). * = P < 0.05, ** = P < 0.01, *** P < 0.001. Bars are means ± SEM.

**Figure 4:** Gravida and different types of smokers. Those that smoke a significantly higher chance in getting pregnant than those that do not smoke (P=0.0002). Those that smoke in addition to THC have a significantly lower chance of getting pregnant than those that only smoke (P=0.0173). Significance was assigned using T-tests (α = 0.05). * = P < 0.05, ** = P < 0.01, *** P < 0.001. Bars are means ± SEM.

Citation: Tamayo RAC, Wright TE, Collier AC (2015) Does Methamphetamine Abuse Affect Serotonin Signaling in the Maternal-Placental-Fetal System?. Pharm Pharmacol Int J 2(6): 00042. DOI: 10.15406/ppij2015.02.00042
Does Methamphetamine Abuse Affect Serotonin Signaling in the Maternal-Placental-Fetal System?

Figure 5: THC and neonatal baby weight. Neonatal baby weights are significantly lower in those that smoke THC ($P=0.0246$) than those that do not smoke at all. Significance was assigned using T-tests ($\alpha = 0.05$). * = $P < 0.05$, ** = $P < 0.01$, *** $P < 0.001$. Bars are means ± SEM.

Figure 6: Antioxidant Defense Network enzymes versus different types of smokers. 3A: There are no significant differences in CAT levels with respect to different types of smokers. 3B: Those who smoke THC have significantly higher levels of Vitamin C ($P = 0.0383$). 3C: Those who smoke have a significantly lower level of GPx activity than those who do not smoke ($P = 0.05$). Significance was assigned using multiple t-tests ($\alpha = 0.05$). * = $P < 0.05$, ** = $P < 0.01$, *** $P < 0.001$. Bars are means ± SEM.

Figure 7: GST, SOD and NA levels versus different types of smokers. 4A: There are no significant differences in GST levels with respect to different types of smokers. 4B: Those who smoke have significantly higher levels of SOD ($P = 0.0023$). 4C: Those who smoke THC and other illicit drugs have significantly higher levels of noradrenaline ($P = 0.0217$). Significance was assigned using T-tests ($\alpha = 0.05$). * = $P < 0.05$, ** = $P < 0.01$, *** $P < 0.001$. Bars are means ± SEM.

References
1. Benirschke K (2012) Pathology of The Human Placenta (2nd edn), Springer-Verlag, New York, USA.
2. Koren G, Cohen MS (2006) Special aspects of perinatal and pediatric pharmacology. In: Katzung BG (Ed.), Basic and Clinical Pharmacology (9th edn), East Norwalk: Appleton and Lange, USA, pp. 913-923.
3. Sastry BV (1999) Techniques to study Human Placental Transport. Advanced Drug Delivery Reviews 38(1): 17-39.

Citation: Tamayo RAC, Wright TE, Collier AC (2015) Does Methamphetamine Abuse Affect Serotonin Signaling in the Maternal-Placental-Fetal System?. Pharm Pharmacol Int J 2(6): 00042. DOI: 10.15406/ppij.2015.02.00042
Does Methamphetamine Abuse Affect Serotonin Signaling in the Maternal-Placental-Fetal System?

4. Barbara L Thompson, Pat Levitt, Gregg D Stanwood (2009) Prenatal exposure to drugs: effects on brain development and implications for policy and education. Nature Reviews Neuroscience 10(4): 303-312.

5. Caldwell J, Sever PS (1974) The biochemical pharmacology of abused drugs. I. Amphetamines, cocaine, and LSD. Clinical Pharmacology and Therapeutics 16(4): 625-638.

6. Cascio CS, Kellar KJ (1983) Characterization of [3H]tryptamine binding sites in brain. European Journal of Pharmacology 55(1-2): 31-39.

7. Ramamoorthy JD, Ramamoorthy S, Leibach FH, Ganapathy V (1995) Human placental monoamine transporters as targets for amphetamines. American Journal of Obstetrics and Gynecology 173(6): 1782-1787.

8. Russo EB, Burnett A, Hall B, Parker KK (2005) Agonistic properties of cannabidiol at 5-HT1a receptors. Neurochemical Research 30(8): 1037-1043.

9. Barann M, Moklerings G, Bruss M, Bonisch H, Urban BW (2002) Direct inhibition by cannabinoids of human 5-HT3A receptors: probable involvement of an allosteric modulatory site. British Journal of Clinical Pharmacology 137(5): 589-596.

10. Russo EB (2004) Clinical endocannabinoid deficiency (CECD): Can this concept explain therapeutic benefits of cannabis in migraine, fibromyalgia, irritable bowel syndrome and other treatment-resistant conditions? Neuro Endocrinology Letters: 25(1-2): 31-39.

11. Altura BM, Malaviya D, Reich CF, Louis R Orkin (1972) Effects of Vasoactive agents on isolated human umbilical arteries and veins. American Journal of Physiology 222: 345-355.

12. MacLennan SJ, Whittle MJ, McGrath JC (1989) 5-HT1-like receptors requiring functional cyclo-oxygenase and 5-HT2 receptors independent of cyclo-oxygenase mediate contraction of the human umbilical artery. British Journal of Pharmacology 97(3): 921-933.

13. Ullmer C, Schmuck K, Kalman HD, Lubbert H (1995) Expression of serotonin receptor mRNAs in blood vessels. FEBS Letters 370(3): 215-221.

14. Karlsson C, Bodelesson G, Bodelesson M, Stjernquist M (1999) Characterization of 5-hydroxytryptamine receptors mediating circular smooth muscle contraction in the human umbilical artery. Gynecologic and Obstetric Investigation 47: 102-107.

15. Santos Silva AJ, Cairnso E, Marques B Verde I (2009) Regulation of Human Umbilical Artery Contractility by Different Serotonin and Histamine Receptors Reproductive Sciences in press 16(12): 1175-1185.

16. Tuncer M, Dogan N, Oktay S & Kayaalp SO (1985) Receptor mechanisms for 5-hydroxytryptamine in isolated human umbilical artery and vein. Archives of International Pharmacodynamics and Therapeutics 276(1): 17-27.

17. Yoshikawa F & Chiba S (1991) Pharmacological analysis of vasoconstrictor responses of isolated and perfused human umbilical arteries. Heart Vessels 6(4): 197-202.

18. McMurphy DM, Boreus LO (1971) Studies on the pharmacology of the perfused human fetal ductus arteriosus. American Journal of Obstetrics and Gynecology 109(6): 937-942.

19. Buyon JP, Clancy R, Di Donato F, Miranda-Carus ME, Askanae AD, et al. (2007) Cardiac 5-HT(4) serotoninergic receptors, S2Kd SSA/ Ro and autoimmune-associated congenital heart block. Journal of Autoimmunity 19(1): 79-86.

20. Nebigil CG, Etienne N, Schaeferling B, Hickel P, Launay JM, et al. (2001) Developmentally regulated serotonin 5-HT2B receptors. International Journal of Developmental Neuroscience 19(4): 365-372.

21. Xie H, Triggle CR (1994) The endothelium contributes to the contractile responses of the human umbilical artery to 5-hydroxytryptamine and endothelin-1 under low but not high P02 conditions. Canadian Journal of Physiology and Pharmacology 72(10): 1171-1179.

22. Katzung BG (2006) Introduction to autonomic Pharmacology. In: Katzung BG (Ed.), Basic and Clinical Pharmacology (9th edn), Philadelphia: McGraw Hill, USA, p. 75-92.

23. Smith PK, Krohn RI, Hermanson GT, Mallia AK, Gartner FH, et al. (1985) Measurement of protein using bicinchoninic acid. Analytical Biochemistry 150(1): 76-85.

24. Wright TE, Milam KA Rougee L, Collier AC (2010) Measuring drugs of abuse (including cotinine) in umbilical cord samples. Journal of Perinatology.

25. Hatch E, Bracken M (1986) Effect of Marijuana use in pregnancy on fetal growth. American Journal of Epidemiology 124(6): 986-993.

Citation: Tamayo RAC, Wright TE, Collier AC (2015) Does Methamphetamine Abuse Affect Serotonin Signaling in the Maternal-Placental-Fetal System? Pharm Pharmacol Int J 2(6): 00042. DOI: 10.15406/ppij.2015.02.00042