Perspective: Gestational Tryptophan Fluctuation Altering Neuroembryogenesis and Psychosocial Development

Xiaohong Huang 1,2,*, Zhendong Feng 1 and Heng-wei Cheng 2,3

1 Department of Pathophysiology, Institute of Neuroregeneration & Neurorehabilitation, Qingdao University, Qingdao 266071, China; zhendongfeng@hotmail.com
2 Department of Animal Sciences, Purdue University, West Lafayette, IN 47907, USA; hwcheng@purdue.edu
3 Livestock Behavior Research Unit, USDA-ARS, West Lafayette, IN 47907, USA
* Correspondence: ysrred@gmail.com

Abstract: Tryptophan, as the sole precursor of serotonin, mainly derived from diets, is essential for neurodevelopment and immunomodulation. Gestational tryptophan fluctuation may account for the maternal-fetal transmission in determining neuroembryogenesis with long-lasting effects on psychosocial development. Personality disorders and social exclusion are related to psychosocial problems, leading to impaired social functioning. However, it is not clear how the fluctuation in mother-child transmission regulates the neuroendocrine development and gut microbiota composition in progeny due to that tryptophan metabolism in pregnant women is affected by multiple factors, such as diets (tryptophan-enriched or -depleted diet), emotional mental states (anxiety, depression), health status (hypertension, diabetes), and social support as well as stresses and management skills. Recently, we have developed a non-mammal model to rationalize those discrepancies without maternal effects. This perspective article outlines the possibility and verified the hypothesis in bully-victim research with this novel model: (1). Summarizes the effects of the maternal tryptophan administration on the neuroendocrine and microbial development in their offspring; (2). Highlights the inconsistency and limitations in studying the relationship between gestational tryptophan exposure and psychosocial development in humans and viviparous animals; and (3). Evidences that embryonic exposure to tryptophan and its metabolite modify bullying interactions in the chicken model. With the current pioneer researches on the biomechanisms underlying the bully-victim interaction, the perspective article provides novel insights for developing appropriate intervention strategies to prevent psychosocial disorders among individuals, especially those who experienced prenatal stress, by controlling dietary tryptophan and medication therapy during pregnancy.

Keywords: tryptophan; psychosocial development; gestation; neuroendocrine; microbiota-gut-brain axis; hypothalamic-pituitary-adrenal axis

1. Psychosocial Development in Adolescence

The Centers for Disease Control and Prevention (CDC, USA) reported that suicide is the second leading cause of death among children and youth between the ages of 10 and 24 during 2000–2017 [1]. The undisclosed suicide attempt is the major risk factor for completed suicides, which is correlated with the psychosocial factors in adolescents, especially among bullying victims [2]. Psychosocial problems are associated with personality disorders, social exclusion, and consequentially impaired social functioning [3,4]. The majority of the psychological disorders are thought to be caused by a person’s nature (genetic factors influencing personality characteristics), nurture (all environments affecting behavioral development), and their interactions (integrated genetic inheritance and environmental factors to physical, physiological, and behavioral development) [5]. Amongst the pregnancy-specific environments, maternal prenatal depression is associated with poor psychosocial functioning [6], and smoking during pregnancy increases the risk...
of attention-deficit/hyperactivity disorder (ADHD) in adolescent boys [7]. Moreover, sociological studies note that babies born to stressed mothers are easier prey for bullies [8], i.e., gestational experiences may reprogram fetal brain development and increase vulnerability to bullying. Thus, the adverse maternal-fetal transmission, e.g., maternal distress [9,10], malnutrition [11], or drug abuse [12], affecting tryptophan (Trp) metabolism, may be involved in the psychosocial disorders in offspring.

2. Tryptophan in Pregnancy

Tryptophan, an essential amino acid, is mainly derived from the diets in the small intestines and is particularly important for gestation. As the sole precursor of serotonin (5-hydroxytryptophan, 5-HT), the maternal circulating Trp is involved in the fetal brain development directly and indirectly via the 5-HT pathway: delivered to the fetal brain and synthesized into 5-HT by tryptophan hydroxylase (TPH) 2, and synthesized into 5-HT at the placenta by TPH1, then delivered to the fetal brain (Figure 1) [13]. Serotonergic (5-HTergic) dysfunction, as one of the cellular mechanisms, is underlying the psychopathological changes including suicidality [14]. In addition, the placental Trp degrades along the cytokine-induced activated Trp catabolite (TRYCAT) pathway, producing kynurenines (KYNs) for suppression of T cell responses [15]. The altered Trp metabolism is one of the core biological signatures of prenatal maternal stress, e.g., lower plasma concentrations of Trp and KYN have been found in severe gestational anxiety and depression [10]. Antenatal stress adversely affects neuroembryogenesis via reprogramming the placental Trp-KYN metabolism, which contains various neuroactive metabolites such as kynurenic acid (KYNA), 3-hydroxykynurenine (3-HK), and quinolinic acid (QUIN), laying the foundation for the neuropathology and related psychiatric disorders [9]. Moreover, urinary Trp and purine metabolites are consistently upregulated in the gestational diabetes mellitus patients [16]. Excessive increase in Trp availability compromises pregnancy by undermining T cell suppression, e.g., in pre-eclampsia [15] which is the most significant individual obstetric risk factor for schizophrenia [17]. Schizophrenia is characterized by a disturbed TRYCAT pathway in the fetal brain [18] and increased activity of systemic cortisol metabolism [19].

![Figure 1. The mechanisms underlying the gestational Trp fluctuation on prenatal, perinatal, and postnatal development. The maternal circulating Trp is affected by multiple factors, such as drug administrations and diets. The placental Trp can be synthesized to 5-HT, degraded to KYN, or transferred to the embryo. The fetus is exposed to the Trp and its metabolite, 5-HT, and KYN, which regulates the neurogenesis and neuroendocrine development as well as the gut microbiota composition and diversity in the progeny. The mother-child microbe transmissions during the 40-week pregnancy...](image-url)
pregnancy, labor, and feeding constitute the initial microbiota community in the progeny and consequentially, regulate decision making and behavioral exhibition via the MGB axis. The development and activity of the MGB axis are regulated by the maternal-fetal transfers of nutrients and microbes together with multiple environmental factors, including the disease and medicinal history, culture, eating habits, education, etc. 5-HT: serotonin; KYN: kynurenine; MGB axis: microbiota-gut-brain axis; Trp: tryptophan.

The excess cortisol by maternal stress crosses the placenta, affects the neuroembryogenesis, reprograms the development and activity of the neuroendocrine, and consequently induces long-lasting changes in cognitive and behavioral disorders throughout life [20]. Moreover, prenatal stress reduces the abundances of the gut Trp-metabolizing microbes in the dam and offspring, which alters the fetal exposure to Trp and its metabolites, and mediates the aberrant fetal neurogenesis, consequently reprograms microbial development in the progeny [21]. Thus, diets and or drug administrations during pregnancy may cause placental Trp fluctuation, by which it consequently alters the placental 5-HT synthesis, maternal-fetal Trp and 5-HT supply, fetal 5-HT concentrations, and gut microbiota composition as well as the endocrine and neuroendocrine developments (Figure 1). Once there, the outcomes alter the psychosocial development in offspring.

3. Maternal Trp Fluctuation Alters Neuroendocrine and Gut Microbiome in Offspring

To propose the perspective of gestational Trp fluctuation affecting psychosocial development in offspring, the existing evidence has been reviewed and presented in Table 1. Maternal Trp fluctuation by the Trp-enriched or -depleted diet during pregnancy alters the fetal neuroendocrine and microbiome, which may reprogram the development of the hypothalamic-pituitary-adrenal (HPA) and hypothalamic-pituitary-gonadal (HPG) axes and mediate the activity of the microbiota-gut-brain (MGB) axis in offspring, consequently reprogramming the psychosocial development.

Table 1. Non-comprehensive review of the effects of maternal Trp administration on the physiological and behavioral exhibitions in offspring.

| Species | Treated Time | Control | Treatment | Exhibitions in Offspring | Refs |
|---------|--------------|---------|-----------|--------------------------|------|
| Wistar rats | 14 days prior to mating–P4mo * | Standard chow powder (3.5 g Trp/kg) | 10 g Trp mixed with the diet (13.5 g Trp/kg) | Decreased BW of the male offspring at P4mo; Decreased 5-HT concentration, TPH2 activity, and 5-HT uptake in the frontal cortex and brain stem. | [22] |
| SHRs or DOCA-salt hypertensive rats | 7 continued days prior to mating | Stock chow diet | 30 mg Trp/kg/day mixed with the diet | Increased BW and blood pressure during P5wk–P15wk; Increased brain weight at P20wk; Increased total 5-HT metabolite content (5-HT plus 5-HIAA) in the medulla at P20wk. | [23] |
| Humans | 210 min | Continued fasting | 1 g Trp orally | Increased incidence of fetal breathing movements; Unchanged breathing rates and breath interval variability. | [24] |
| SD rats | E17 | Saline vehicle | 200 mg Trp/kg oral gavage | Increased Trp, 5-HT, and 5-HIAA concentrations in the fetal brain at E17 and E18. | [25] |
| SD rats | E15–E21 | Saline vehicle | 200 mg Trp/kg oral gavage | Increased serum PRL at P40d and P70d; Increased serum LH at P70d; Increased forebrain 5-HT and 5-HIAA at P70d. | [26] |
| Wistar rats | E19 and E21 | 0.1 N-HCl vehicle | 250 mg Trp/kg i.p. | Increased intracerebral concentrations of Trp at E19; Decreased valine, methionine, leucine, tyrosine, phenylalanine, and histidine at E19; Increased phosphoserine, threonine, serine, glutamic acid, and Trp at E21; Decreased methionine, leucine, and histidine at E21; Increased protein synthesis activity indicated by [3H] Leucine incorporation at E19 and E21. | [27] |
| SD rats | E14.5-late puberty | Control chow (0.22% Trp) | Trp free diet (0.00% Trp) | Dwarfism pups; Decreased serum GH concentration in male and female offspring; Severe hypoprolactinemia; Normal right-timed onset of puberty in both male and female rats. | [28,29] |
Table 1. Cont.

| Species     | Treated Time | Control | Treatment | Exhibitions in Offspring | Refs |
|-------------|--------------|---------|-----------|--------------------------|------|
| Wistar rats | E5–E21       | Regular chow diet | Trp-free diet (0.2% Trp) | Unchanged Brain weights in newborn pups; Decreased BW in newborn pups; Reduced numbers of 5-HTergic neurons at the dorsal raphe, especially at the medial and caudal sections of dorsal raphe, which contains the majority of 5-HTergic neurons; Unchanged brain 5-HT concentration. | [30] |
| SD rats     | E1–E21       | Control | 200 mg Trp/kg oral gavage | Increased kidney weight-to-BW ratio at P12wk; Increase blood pressure in male offspring at P4wk, P6wk, P8wk, P10wk, and P12wk; Decreased plasma level of L-citrulline, a precursor of L-arginine and SDMA, an indirect inhibitor of NO synthase; Increased gene expressions in the AHR pathway. | [31] |
| CKD SD rats | E1–early postnatal life * | Control | 200 mg Trp/kg oral gavage | Decreased systolic blood pressure, mean arterial pressure, and creatinine at P12wk; Decreased plasma level of L-citrulline and SDMA; Altered the abundance of the Trp-metabolizing microbes, i.e., increased the abundance of the genus *Intestinimonas* and decreased the abundance of *Turibacter*. | [31] |
| SD rats     | E1–late puberty | Control rat chow (0.22% Trp) | Trp free diet (0.00% Trp) | Pronounced dwarfism pups; Decreased serum GH concentration in males and females; Marked hypoprolactinemia and severe hypoprolactinemia in males; Hypoprolactinemia in females; Right-timed pubertal maturation in both sexes. | [28,29] |
| SD rats     | E1–P12wk *   | Control rat chow (0.22 g Trp/100 g of pellets) | high-Trp diet (1 g Trp/100 g of pellets) | Increased blood 5-HT, i.e., hyperserotonemia during P1wk–P2wk; Decreased blood GH; Decreased TPH1 activity in gastrointestinal tracts tissue; Decreased IGF-I expression in hepatic and muscle tissue. | [32] |
| SD rats     | E1–weaning   | 500 mg Trp/100 g diet | 75 mg Trp/ 100 g diet | Decreased average BW at weaning; Unaffected opacities at P22d. | [33] |
| SD rats     | E1–P25d *   | TD.99366 control diet (1.8 g Trp/kg) | TD.08125 Trp-deficient diet (1 g Trp/kg) | Decreased birth healthy pig rate and birth weight of piglet per pen with similar total birth weight per pen; Decreased serum phosphoserine, taurine, cysteine, proline in newborns and increased liver n-6:n-3 PUFA ratio; Altered gene expressions, including the genes related to cytotoxic effector regulation, NADH oxidation, ROS metabolism, and tissue development. | [34] |
| Pigs        | Third trimester of gestation–delivery | 2 x Trp diet (0.26% Trp fed in the morning and afternoon) | High-low Trp diet (0.3% Trp fed in the morning and 0.13% Trp fed in the afternoon) | Decreased birth healthy pig rate and birth weight of piglet per pen with similar total birth weight per pen; Decreased serum phosphoserine, taurine, cysteine, proline in newborns and increased liver n-6:n-3 PUFA ratio; Altered gene expressions, including the genes related to cytotoxic effector regulation, NADH oxidation, ROS metabolism, and tissue development. | [35] |
| Outbred CD-1 mice | Lactation (P0d–P8d) | Standard laboratory diet (0.14% Trp) | Trp-deficient diet (0.00% Trp) | Unchanged time spent in open sectors in the 0-maze test in adolescent daughters (P189d–P193d); Unchanged time spent in floating in the forced-swim test in adolescent daughters; Unchanged achieved breakpoint in the progressive ratio operant procedure in adolescent daughters; Decreased plasma CORT concentrations and similar BDNF concentrations following restraint stress in adolescent daughters. | [36] |

* The pups are fed the same diet as mothers throughout postnatal life. 5-HIAA: 5-hydroxyindoleacetic acid; 5-HT: serotonin; AHR: aryl hydrocarbon receptor; BDNF: brain-derived neurotrophic factor; BW: body weight; d: day; CKD: chronic kidney disease; CORT: corticosterone; DOCA: deoxycorticosterone acetate; E: embryonic day; GH: growth hormone; IGF-1: insulin-like growth factor-1; i.p.: intraperitoneal; LH: luteinizing hormone; mo: month; NADH: nicotinamide adenine dinucleotide; P: postnatal; PRL: prolactin; PUFA: polyunsaturated fat; SD: Sprague-Dawley; SDMA: symmetric dimethylarginine; SHRs: spontaneously hypertensive rats; TPH: tryptophan hydroxylase; Trp: tryptophan; ROS: reactive oxygen species; wk: week.
3.1. Maternal Trp Fluctuation Alters Psychosocial Development via Reprogramming Neuroendocrine in Offspring

Serotonin and its receptors appear early during prenatal development [37]. The fetal 5-HT, transferred from the placenta or synthesized from Trp, functions as a morphogen and tropic factor regulating brain development during embryogenesis, then as a neurotransmitter mediating the neuroendocrine functions and as a potent immune cell modulator regulating the immune system to influence multiple psychosocial events [37–39]. The 5-HT turnover rate in the fetal central nervous system (CNS) is at the highest stage in an animal’s entire lifespan [40]. During the prenatal and early postnatal development, 5-HT influences neural maturation and synaptogenesis [37], and disrupting 5-HT signaling leads to neurological and psychiatric disorders [41–43]. The alterations in the 5-HTergic system during embryogenesis have been well exemplified in pregnant women who take selective serotonin reuptake inhibitors (SSRIs), with the concerns of affecting fetal development and associated postnatal behavioral exhibitions, such as depression, anxiety, and aggressiveness [44,45].

Maternal Trp-deficient diets in pregnant mice decrease the plasma corticosterone (CORT) concentrations following restraint stress in the female adolescent offspring [36]. Thus, the maternal Trp fluctuation may alter the development and function of the HPA axis. The HPA axis is crucial in regulating the stress coping strategy confronting conflict during peer interactions [46]. The 5-HTergic neurons in the raphe nuclei regulate the HPA axis by modulating the impulses activity in the hypothalamus; ontogenetically, the 5-HTergic system regulates the synthesis of arginine vasopressin (AVP, or its non-mammalian homologue AVT, arginine vasotocin) via innervating the periventricular nucleus (PHN) and ventromedial hypothalamus (VMH) [47,48] to functionally mediate aggressiveness [49]. Equally important, maternal Trp administration in mice alters the development and function of the HPG axis during the process of pubertal maturation in offspring via regulating the productions of prolactin (PRL) and luteinizing hormone (LH) in the pituitary gland [26,28,29]. Trp-free diets from the gestation to puberty cause hypoandrogenism and hypoprolactinemia in the progeny [28,29], while maternal administration of Trp-enriched diets increases serum LH at postnatal (P) 70 d [26]. Serotonin also modulates PRL production by inducing the release of vasoactive intestinal peptide (VIP) and oxytocin (OT) from the hypothalamus [50]. A high serum PRL level is associated with psychological stress responses in humans [51]. In addition, acute psychosocial stress disrupts the LH surge in female mice [52] and central precocious puberty exerts significant effects on psychosocial development [53].

3.2. Maternal Trp Fluctuation Alters Psychosocial Development via Reprogramming the Gut Microbiota in Offspring

Multi-hits early-life stress alters gut microbiome, brain gene expression, and social communication, laying the foundation of the physical and mental health in adulthood [54,55]. The gut microbiota works as a virtual endocrine organ [56]. Investigating gestational Trp’s role in preventing hypertension programmed by maternal chronic kidney disease (CKD), Hsu et al. [31] found that the gestational Trp oral gavage in the CKD rats restored the abundance of microbes in offspring at P12wk, such as Lactobacillus and Turicibacter; and altered the Trp-metabolizing microbes, i.e., increased and decreased the abundance of the genus Intestinimonas and Turicibacter, respectively. The gut microbes may mediate the activity of the HPA axis via reprogramming the MGB axis. In addition, the decreased abundance of Lactobacillus has been characterized in the infant rhesus monkeys who experienced maternal separation and show increased stress reactivity [57]. Lactobacillus casei strain Shirota, a widely used probiotic strain, releases the stress-associated symptoms by suppressing the hypothalamic corticotropin-releasing hormone (CRH) and plasma CORT in the rats under water-avoidance stress [58]. Moreover, a low abundance of Turicibacter has been characterized in the patients with autism spectrum disorder (ASD) [59], associated with the 5-HT metabolic disorder [60]. Daily intake of Lactobacillus helveticus CCFM1076
restores the balance of the 5-HTergic system in both the gastrointestinal tract and brain, thereby ameliorating ASD-like behaviors [61].

3.3. The Accompanied Physiological Alterations Are Associated with the Neuropsychological Impairment in Offspring

The physiological alterations during the offspring growth following the maternal Trp administration, including body weight (BW), blood pressure, and breath movement [22–24,28–31,33–35], may be underlying the Trp’s and 5-HT’s roles in controlling neuroembryogenesis and related 5-HTergic development in the pontomedullary brainstem [62,63]. The National Collaborative Perinatal Project (NCPP, New England) revealed a relationship between obstetrical complications and neuropsychological deficits in children aged 7 years [64]. A low birth weight is associated with neuropsychological impairments, following an index of inferred hypoxic insults [64] caused by mild to severe pre-eclampsia, maternal hyper- or hypo-tension, or maternal diabetes [15], which has been linked to the maternal Trp fluctuation [15,16,23,31]. Moreover, a survey conducted in Scania among 2000 healthy young women aged 18–34 years revealed that the women underweight have poorer psychological health compared with ones with normal BW, while the overweight women are more likely to have poor health-related behaviors and lack of internal locus of control [65]. Thus, the dissatisfaction with body image followed by improper Trp administrations may have counterbalanced mental health, causing negative influences on the psychosocial development in offspring.

4. Inconsistency in the Current Findings and Gaps in Knowledge

Ultimately understanding the effects of prenatal malnutrition or medicine during pregnancy on a child’s psychosocial development is a complex story. Issues are likely to cause inconsistency in the current findings, including the pregnancy stage of Trp taken, its intensity, and pregnant women’s age, eating habits, lifestyle, and healthy status. Huether et al. [61] reported that feeding a 10 g Trp-enriched standard chow powder diet daily from two weeks prior to mating through postnatal life, P4mo, resulted in a lower BW, brain 5-HT concentration, TPH2 activity, and 5-HT uptake in Wistar rats [22]. However, the administration of 30 mg/kg/day of Trp mixed with a stock chow diet for one week prior to mating increases BW at P15wk and the total 5-HT metabolite content in the medulla at P20wk in spontaneously hypertensive rats (SHRs) or deoxycorticosterone acetate (DOCA)-salt hypertensive rats [23]. Moreover, the considerations about maternal seasonality and season of conception are advocated when studying the causality between maternal Trp fluctuation and fetal neurodevelopment. Because the differences in the plasma Trp concentrations have been detected across maternal seasonality and season of conception [66]. The longitudinal studies of the Trp impacts also are needed, because the early life adversity can be buffered and masked by the favorable experience at a later age, like family support, culture, and education [67]. Taken together, a number of challenges in human subject and experimental non-human mammalian researches, including the inevitable background noise during the embryogenesis and multiple stimulations during the postnatal development, exist in unveiling the association between the maternal Trp metabolism and the psychosocial development in their progeny (Figure 1).

The National Center for Educational Statistics (NCES, Washington, DC, USA) in 2015 revealed that 21% of the students aged 12–18 years are being bullied on school premises [68]. The studies about the underlying mechanisms of bullying mainly focus on genetics and environments, while the contribution of the maternal-fetal transmission is not well clear. The 40-week pregnancy, labor (natural labor vs. cesarean section), feeding (breast-feeding vs. formula-feeding), and the fluctuation in the individual pregnancy-specific environments may affect the neurogenesis, neuroendocrine development, and gut microbiota composition and diversity in progeny, which consequently alters the development and activity of the MGB, HPA, and HPG axes, affecting the sociometric status in a peer relationship. For instance, in aberrant fetal neurodevelopment, the reduced abundances of the Trp-
metabolizing microbes have been associated with the stress-disrupted placental Trp [21].
However, the mother-to-child transmission of the dietary Trp and 5-HT synthesized in
the placenta as well as microbes during pregnancy, labor delivery, and breastfeeding
is irresistible in viviparous animals [69], which causes inevitable background noise in
studying the influences of gestational Trp fluctuation on the physiological and behavioral
developments in offspring. To date, no specific animal model has been established for
investigating the underlying mechanisms of the effects of mother-to-child transmission
on bullying.

5. Evidences in a Chicken Model

5.1. Chicken Model

With the significant, fundamental similarities between the human and chicken
genomes [70,71], avian species have been central in investigating the subsequent cog-
nitive, social, and neural development influenced by the experiences occurring during the
ontogenic stages due to the directly controllable embryogenesis in the oviparous animals.
Chicken have been used as an animal model in various clinical and psychopharmaco-
logical studies [72,73] as well as for assessing the effects of genetics, environments, and
genetic-environmental interactions on psychopathological disorders [74,75]. Noteworthy,
chickens are currently an elective model system in investigating the ontogenic origins of
behaviors [70,74,75], like ASD [76,77]. Social behavioral deficits have been reported in the
chicken model, including impaired early social predispositions [76,78], low aggregation
and belongingness, and weak vocalization, when exposed to sodium valproate (VPA)
during E14–E21 [77]. A laying hen produces more than 300 eggs a year with a potential
for producing numerous embryos with similar genetic backgrounds and robust microbial
communities at once [79] but independence of maternal influences on the neuroembryo-
genesis. In doing so, the chicken model has been employed in investigating the effects of
embryonic exposure of Trp and its metabolite, 5-HT, on aggression and bullying in our lab
(Figure 2) [74,80,81]. Within flocks, chickens have a social hierarchy referred to as a pecking
order: peckers within a flock always start feather pecking at others, as a reason to avoid
adding young birds, small birds, or strange birds to the flock. Hence, peckers within a flock
could be a comparison to bullies within a social group.

5.2. Embryonic Trp Exposure Yields Bullying Victim

The photograph represents the gestational tryptophan fluctuation in the chicken model
for bullying research. Embryonic Trp or 5-HT exposure regulates neuronal development, neuro-
transmission, intestinal histomorphology, and microbiota composition, consequently altering the
postnatal behavioral and physiological exhibitions, which determines an individual’s psychosocial
development during the lifespan. 5-HT: serotonin; BW: body weight; Trp: tryptophan.
5.2. Embryonic Trp Exposure Yields Bullying Victim

In ovo Trp administration (500 µg/egg) yields bullying victims in White Leghorn birds, a commercial strain [75], reducing the BW and aggressiveness in the male offspring before and during adolescence [80], resulted from the reprogrammed MGB axis. The gut histomorphology has been altered by the Trp exposure, i.e., the crypt depth is increased while the villus/crypt ratio is decreased [80]. The abundant genus, Olsenella, in the Trp group [80] has been identified as the core microbe in the crypt of the human colon [81]. The correlations between the gut morphological alteration and high abundance of Olsenella are consistent with the findings in patients with depression [82] and ASD [83]. Moreover, the abundance of Olsenella is increased in the lean people in the Japanese population [84], which is positively correlated with the Yale Food Addiction Scale (YFAS) used for identifying signs of addictive-like eating behavior, hedonic food intake [85]. Thus, the alteration in the gut bacterial community may be involved in determining the bodily force and regulating the neuropsychological activity facing bullying.

Peer victimization exhibits more stress with altered HPA axis activity during the trier social stress test (TSST) [86]. The HPA axis is developmentally cross-linked with the gut microbiota [87], which hints at the gut microbes’ role in neurogenesis and related behavioral development in offspring. The chicken HPA axis is functionally similar in stress response to that of mammals [88–91]. The increased catecholamine concentrations in the chicken hypothalamus, including dopamine, epinephrine, and norepinephrine, post embryonic Trp exposure may be associated with the alterations in the gut microbiome and MGB axis function, laying the foundation of their mental status confronting conflictual peer interactions [80]. Taking as an example skatole, one of the gut Trp metabolites produced by Olsenella [92], shows a positive association with the anatomical connectivity of the amygdala [85], by which it is involved in reward learning and addiction [93]. The amygdala is constantly scanning our environments for initiating the HPA to regulate emotional and mental reactions in responding to internal and external stimulations [94,95].

5.3. Embryonic Serotonin Exposure Reduces Aggressiveness in Bullies

Identical twins with ASD differ significantly in the severity of social traits [96], which is probably due to the unequal blood supply from the placenta during embryogenesis. The placenta is the primary source of 5-HT for fetal development. The promoter of 5-HT transporter (5-HTT) contains two common alleles, which are associated with the dysregulation of 5-HTergic neurotransmission: the short (S) allele has a lower transcriptional efficiency compared to the long (L) allele [97]. Bullied children with the SS genotype are at greater risk for developing emotional problems at age 12 than those children with the SL or LL genotype [97]. In humans and various animals, the 5-HTergic system regulates aggression via innervating the midbrain and hypothalamic dopaminergic neurons [98–100]. The midbrain dopamine storage capacity is negatively correlated with aggression [101]; individuals who have a low dopamine transmission capacity show more impulsive aggression in response to provocation [101]. The midbrain and hypothalamic 5-HTergic innervations have been evidenced in the chicken brain as well as the dopaminergic neuronal distribution and neurotransmission during mid-late embryogenesis [48]. Moreover, in ovo 5-HT injection (10 µg/egg) reduces aggressive behaviors at a cost of increased fearfulness during adolescence and before sex maturation in the White Leghorn birds [102]. The Dekalb XL birds, a highly aggressive strain [75,103], the inborn bullies, are picked to explore the effect of prenatal 5-HT exposure (10 and 20 µg/egg) on the behavioral development and underlying mechanisms [104]. Both dosages of 5-HT exposure reduce aggression at P7wk, and the 5-HT exposure effects can be achieved via the different paths by modifying the embryonic 5-HTergic and dopaminergic systems and altering fetal 5-HTergic influence on the thalamic circuit and HPA axis. Briefly, the 10 µg 5-HT exposure reduces the 5-HT turnover rate, increases 5-HT1a receptor expression, and facilitates the ventral tegmental area (VTA) neuronal development, suggesting an increased 5-HT availability but not 5-HT pool potentially facilitates the midbrain neuronal development. The 20 µg 5-HT exposure enhances
the 5-HTergic regulation to the hypothalamus and VMH, indicated by the increased both the 5-HTergic and dopaminergic neurotransmission [104]. Hyper-excitability of the neurons in the ventrolateral part of the VMH exaggerates aggression under certain pathological conditions, like ASD [105]. These findings shed light on the mechanisms underlying hypo- and hyper-serotonemia as the potential risk factors of ASD [106].

6. Conclusions

Tryptophan is required for protein synthesis for fetal development and participates in programming the neural circuits and development of the neuroendocrine system; on the other hand, excess Trp is responded to maternal inflammation to initiate signaling cascades, such as KYN metabolic pathway, to stimulate embryonic brain development. Through the mother-child transmission, maternal circuiting Trp, as one of the major gestational Trp resources, is critical for the development of MGB, HPA, and HPG axes in offspring with long-lasting impacts on physical, physiologic, and behavioral outcomes. The abnormal Trp metabolism during embryogenesis has been recognized as one of the reasons in patients with emotional problems and mental diseases. Maintaining an appropriate Trp level by controlling dietary Trp and rigorous medication therapy during pregnancy has been advocated as a biotherapeutic targeting strategy for preventing psychosocial disorders, emotional problems, and mental diseases in adolescence, like bullying.

Author Contributions: Conceptualization, X.H. and H.-w.C.; literature review, X.H. and Z.F.; and writing, X.H. and H.-w.C. All authors have read and agreed to the published version of the manuscript.

Funding: This study was supported by the grant award (No: 2017-67015-26567) of the NIFA-AFRI, USDA.

Institutional Review Board Statement: Not applicable.

Informed Consent Statement: Not applicable.

Acknowledgments: We thank the technicians of the USDA-ARS Livestock Behavior Research (West Lafayette, IN) for technical assistance. Mention of trade names or commercial products in this article is solely for the purpose of providing specific information and does not imply recommendation or endorsement of the USDA. The USDA is an equal opportunity provider and employer.

Conflicts of Interest: The Authors declare that there is no conflict of interest.

References

1. Curtin, S.C.; Heron, M. Death Rates Due to Suicide and Homicide Among Persons Aged 10-24: United States, 2000-2017. NCHS Data Brief 2019, 352, 1–8.
2. Levi-Belz, Y.; Gavish-Marom, T.; Barzilay, S.; Apter, A.; Carli, V.; Hoven, C.; Sarchiapone, M.; Wasserman, D. Psychosocial Factors Correlated with Undisclosed Suicide Attempts to Significant Others: Findings from the Adolescence SEYLE Study. Suicide Life-Threat. Behav. 2019, 49, 759–773. [CrossRef] [PubMed]
3. Oltmanns, T.F.; Melley, A.H.; Turkheimer, E. Impaired social functioning and symptoms of personality disorders assessed by peer and self-report in a nonclinical population. J. Pers. Disord. 2002, 16, 437-452. [CrossRef] [PubMed]
4. El Bouhaddani, S.; Van Domburgh, L.; Schaefer, B.; Doreleijers, T.A.H.; Veling, W. Peer status in relation to psychotic experiences and psychosocial problems in adolescents: A longitudinal school-based study. Eur. Child Adolesc. Psychiatry 2017, 27, 701–710. [CrossRef] [PubMed]
5. Ball, H.; Arsenault, L.; Taylor, A.; Maughan, B.; Caspi, A.; Moffitt, T. Genetic and environmental influences on victims, bullies and bully-victims in childhood. J. Child Psychol. Psychiatry 2008, 49, 104–112. [CrossRef] [PubMed]
6. Korhonen, M.; Luoma, I.; Salmelin, R.; Tamminen, T. A longitudinal study of maternal prenatal, postnatal and concurrent depressive symptoms and adolescent well-being. J. Affect. Disord. 2012, 136, 680–692. [CrossRef]
7. Schmitz, M.; Denardin, D.; Silva, T.L.; Pianca, T.; Hutz, M.H.; Faraone, S.; Rohde, L.A. Smoking During Pregnancy and Attention-Deficit/Hyperactivity Disorder, Predominantly Inattentive Type: A Case-Control Study. J. Am. Acad. Child Adolesc. Psychiatry 2006, 45, 1338–1345. [CrossRef]
8. Lereya, S.T.; Wolke, D. Prenatal family adversity and maternal mental health and vulnerability to peer victimisation at school. J. Child Psychol. Psychiatry 2013, 54, 644–652. [CrossRef]
9. Notarangelo, F.M.; Schwarz, R. Restraint Stress during Pregnancy Rapidly Raises Kynurenic Acid Levels in Mouse Placenta and Fetal Brain. Dev. Neurosci. 2016, 38, 458–468. [CrossRef]
10. Keane, J.M.; Khashan, A.S.; McCarthy, F.P.; Kenny, L.C.; Collins, J.M.; O’Donovan, S.; Brown, J.; Cryan, J.F.; Dinan, T.G.; Clarke, G.; et al. Identifying a biological signature of prenatal maternal stress. *JCI Insight* 2021, 6, 143007. [CrossRef]

11. Tsuji, A.; Nakata, C.; Sano, M.; Fukushima, T.; Shibata, K. L-Tryptophan Metabolism in Pregnant Mice Fed a High L-Tryptophan Diet and the Effect on Maternal, Placental, and Fetal Growth. *Int. J. Tryptophan Res.* 2013, 6, 21–33. [CrossRef] [PubMed]

12. Edvinsson, Å.; Hellgren, C.; Kallak, T.K.; Åkerud, H.; Skalkidou, A.; Stener-Victorin, E.; Fornes, R.; Spigset, O.; Lager, S.; Olivier, J.; et al. The effect of antenatal depression and antidepressant treatment on placental tissue: A protein-validated gene expression study. *BMC Pregnancy Childbirth* 2019, 19, 479. [CrossRef] [PubMed]

13. Bonnin, A.; Goeden, N.; Chen, K.; Wilson, M.L.; King, J.; Shih, J.C.; Blaky, R.D.; Deneris, E.S.; Levitt, P. A transient placental source of serotonin for the fetal forebrain. *Nature 2011*, 472, 347–350. [CrossRef] [PubMed]

14. Russo, S.; Kema, I.; Bosker, F.; Haavik, J.; Korf, J. Tryptophan as an evolutionarily conserved signal to brain serotonin: Molecular evidence and psychiatric implications. *World J. Biol. Psychiatry* 2007, 10, 1–11. [CrossRef]

15. Badawy, A.A.-B. The tryptophan utilization concept in pregnancy. *Obstet. Gynecol. Sci.* 2012, 55, 155–159. [CrossRef]

16. Law, K.P.; Han, T.-L.; Mao, X.; Zhang, H. Tryptophan and purine metabolites are consistently upregulated in the urinary metabolome of patients diagnosed with gestational diabetes mellitus throughout pregnancy: A longitudinal metabolomics study of Chinese pregnant women part 2. *Clin. Chim. Acta* 2017, 468, 126–139. [CrossRef]

17. Jones, P.; Cannon, M. The New Epidemiology of Schizophrenia. *Psychiatr. Clin. N. Am.* 1998, 21, 1–25. [CrossRef]

18. Anderson, G.; Maes, M. Schizophrenia: Linking prenatal infection to cytokines, the tryptophan catabolite (TRYCAT) pathway, NMDA receptor hypofunction, neurodevelopment and neuroprogression. *Prog. Neuro-Psychopharmacol. Biol. Psychiatry* 2013, 42, 5–19. [CrossRef]

19. Steen, N.E.; Tesli, M.; Kähler, A.K.; Methlie, P.; Barrett, E.A.; Larsson, S.; Mork, E.; Levås, K.; Rossberg, J.I.; et al. SRD5A2 is associated with increased cortisol metabolism in schizophrenia spectrum disorders. *Prog. Neuro-Psychopharmacol. Biol. Psychiatry* 2010, 34, 1500–1506. [CrossRef]

20. Glover, V.; O’Connor, T.; O’Donnell, K. Prenatal stress and the programming of the HPA axis. *Neurosci. Biobehav. Rev.* 2010, 35, 17–22. [CrossRef]

21. Galley, J.D.; Chen, H.J.; Antonson, A.M.; Gur, T.L. Prenatal stress-induced disruptions in microbial and host tryptophan metabolism and transport. *Behav. Brain Res.* 2021, 414, 113471. [CrossRef] [PubMed]

22. Huether, G.; Thömke, F.; Adler, L. Administration of tryptophan-enriched diets to pregnant rats retards the development of the ventricular system of their offspring. *Dev. Brain Res.* 1992, 68, 175–181. [CrossRef]

23. Ito, H.; Shiokawa, H.; Torii, M.; Suzuki, T. Effects of Tryptophane on Shrsp Offspring Growth. *J. Appl. Physiol.* 1981, 50, 318–328. [CrossRef]

24. Devoe, L.D.; Castillo, R.A.; Searle, A.; Searle, J.R. Maternal dietary substrates and human fetal biophysical activity. *Am. J. Obstet. Gynecol.* 1986, 155, 135–139. [CrossRef]

25. Arevalo, R.; Alfonso, D.; Castro, R.; Rodriguez, M. Fetal brain serotonin synthesis and catabolism is under control by mother intake of tryptophan. *Life Sci.* 1991, 49, 53–66. [CrossRef]

26. Martin, L.; Diaz, M.R.; Santana-Herrera, C.; Milena, A.; Santana, C. Tryptophan ingestion by gestant mothers alters prolactin and luteinizing hormone release in the adult male offspring. *Brain Res.* 1997, 774, 265–268. [CrossRef]

27. Fando, J.L.; Dominguez, F.; Herrera, E. Tryptophan Overload in the Pregnant Rat: Effect on Brain Amino Acid Levels and In Vitro Protein Synthesis. *J. Neurochem.* 1981, 37, 824–829. [CrossRef]

28. Imbesi, R.; Castrogiovanni, P. Embryonic and postnatal development in experimental tryptophan deprived rats. A preliminary study. *Histochim.* 2008, 39, 487–498. [CrossRef]

29. Imbesi, R.; Mazzone, V.; Castrogiovanni, P. Is Tryptophan ‘more’ Essential than Other Essential Aminoacids in Development? A Morphologic Study. *Anat. Histol. Embryol.* 2008, 38, 361–369. [CrossRef]

30. Flores-Cruz, G.M.; Escobar, A. Reduction of serotonergic neurons in the dorsal raphe due to chronic prenatal administration of a tryptophan-free diet. *Int. J. Dev. Neurosci.* 2012, 30, 63–67. [CrossRef]

31. Hsu, C.-N.; Lin, H.-C.; Yu, H.-R.; Huang, L.-T.; Tiao, M.-M.; Tain, Y.-L. Maternal Tryptophan Supplementation Protects Adult Rat Offspring against Hypertension Programmed by Maternal Chronic Kidney Disease: Implication of Tryptophan-Metabolizing Microbiome and Aryl Hydrocarbon Receptor. *Int. J. Mol. Sci.* 2020, 21, 4552. [CrossRef] [PubMed]

32. Castrogiovanni, P.; Musumeci, G.; Trovato, F.M.; Avola, R.; Magro, G.; Imbesi, R. Effects of high-tryptophan diet on pre- and postnatal development in rats: A morphological study. *Z. für Ernährungswissenschaft* 2013, 52, 297–308. [CrossRef] [PubMed]

33. Bunce, G.E.; Hess, J.L. Lenticular Opacities in Young Rats as a Consequence of Maternal Diets Low in Tryptophan and/or Vitamin E. *J. Nutr.* 1976, 106, 222–229. [CrossRef] [PubMed]

34. Penatti, E.M.; Barina, A.E.; Raju, S.; Li, A.; Kinney, H.C.; Commons, K.G.; Nattie, E.E. Maternal dietary tryptophan deficiency alters cardiorespiratory control in rat pups. *J. Appl. Physiol.* 2011, 110, 318–328. [CrossRef]

35. Xu, K.; Bai, M.; Bin, P.; Duan, Y.; Wu, X.; Liu, H.; Yin, Y. Negative effects on newborn piglets caused by excess dietary tryptophan in the morning in sows. *J. Sci. Food Agric.* 2019, 99, 3005–3016. [CrossRef]

36. Zoratto, F.; Berry, A.; Anzidei, F.; Fiore, M.; Alleva, E.; Laviola, G.; Macri, S. Effects of maternal l-tryptophan depletion and corticosterone administration on neurobehavorial adjustments in mouse dams and their adolescent and adult daughters. *Prog. Neuro-Psychopharmacology Biol. Psychiatry* 2011, 35, 1479–1492. [CrossRef]
37. Baumgarten, H.G.; Göthert, M. Serotonergic Neurons and 5-HT Receptors in the CNS; Springer Science & Business Media: Berlin/Heidelberg, Germany, 2012; Volume 129.
38. Yu, Q.; Teixeira, C.; Mahadevia, D.; Huang, Y.-Y.; Balsam, D.; Mann, J.J.; A Gingrich, J.; Ansorge, M.S. Dopamine and serotonin signaling during two sensitive developmental periods differentially impact adult aggressive and affective behaviors in mice. Mol. Psychiatry 2014, 19, 688–698. [CrossRef]
39. Wan, M.; Ding, L.; Wang, D.; Han, J.; Gao, P. Serotonin: A Potent Immune Cell Modulator in Autoimmune Diseases. Front. Immunol. 2020, 11, 186. [CrossRef]
40. Hamon, M.; Bourgojn, S.; Artaud, F.; El Mestikawy, S. The respective roles of tryptophan uptake and tryptophan hydroxylase in the regulation of serotonin synthesis in the central nervous system. J. Physiol. 1981, 77, 269–279.
41. St-Pierre, J.; Laurent, L.; King, S.; Vaillancourt, C. Effects of prenatal maternal stress on serotonin and fetal development. Placenta 2016, 48, S56–S71. [CrossRef]
42. Suri, D.; Teixeira, C.; Cagliostro, M.K.C.; Mahadevia, D.; Ansorge, M.S. Monoamine-Sensitive Developmental Periods Impacting Adult Emotional and Cognitive Behaviors. Neuropsychopharmacology 2015, 40, 88–112. [CrossRef] [PubMed]
43. Field, T. Prenatal depression effects on early development: A review. Infant Behav. Dev. 2011, 34, 1–14. [CrossRef] [PubMed]
44. Smit-Rigter, L.A.; Noorlander, C.W.; von Oerthel, L.; Chameau, P.; Smidt, M.P.; van Hooft, J.A. Prenatal fluoxetine exposure induces life-long serotonin 5-HT3 receptor-dependent cortical abnormalities and anxiety-like behaviour. Neuropharmacology 2012, 62, 865–870. [CrossRef] [PubMed]
45. Nulman, I.; Koren, G.; Rovet, J.; Barrera, M.; Streiner, D.L.; Feldman, B.M. Neurodevelopment of Children Prenatally Exposed to Selective Reuptake Inhibitor Antidepressants. J. Clin. Psychiatry 2015, 76, e842–e847. [CrossRef]
46. von Klitzing, K.; Perren, S.; Klein, A.M.; Stadelmann, S.; Helbling, J.C.; Levy, E.; Chevalier, G.; Hatzinger, M. The interaction of social risk factors and HPA axis dysregulation in predicting emotional symptoms of five- and six-year-old children. J. Psychiatr. Res. 2012, 46, 290–297. [CrossRef]
47. D’Hondt, E.; Eelen, M.; Berghman, L.; Vandensande, F. Colocalization of arginine-vasotocin and chicken luteinizing hormone-releasing hormone-1 (cLHRH-I) in the preoptic-hypothalamic region of the chicken. Brain Res. 2000, 856, 55–67. [CrossRef]
48. Huang, X.; Kuang, S.; Applegate, T.J.; Lin, T.-L.; Cheng, H.-W. The development of the serotonergic and dopaminergic systems during chicken mid-late embryogenesis. Mol. Cell. Endocrinol. 2019, 493, 110472. [CrossRef]
49. Morrison, T.R.; Melloni, R.H. The Role of Serotonin, Vasopressin, and Serotonin/Vasopressin Interactions in Aggressive Behavior. In Behavioral Neurobiology of Alcohol Addiction; Springer Science and Business Media LLC: Berlin/Heidelberg, Germany, 2014; Volume 17, pp. 189–228.
50. Ashbury, J.E.; Lennartsson, A.-K.; Jonsdottir, I.H. Prolactin in response to acute psychosocial stress in healthy men and women. Psychoneuroendocrinology 2011, 36, 1530–1539. [CrossRef]
51. Lennartsson, A.-K.; Jonsdottir, I.H. Prolactin in response to acute psychosocial stress in healthy men and women. Psychoneuroendocrinology 2011, 36, 1530–1539. [CrossRef]
52. Wagenmaker, E.R.; Moenter, S.M. Exposure to Acute Psychosocial Stress Disrupts the Luteinizing Hormone Surge Independent of Estrous Cycle Alterations in Female Mice. Endocrinology 2017, 158, 2593–2602. [CrossRef]
53. Yang, D.; Zhang, W.; Zhu, Y.; Liu, L.; Gao, X.; et al. Initiation of the Hypothalamic–Neuroendocrine–Behavioral Neural Axis in Young Girls Undergoing Early Precocious Puberty Exerts Remodeling Effects on the Prefrontal Cortex. Front. Psychiatry 2019, 10, 332. [CrossRef] [PubMed]
54. Usui, N.; Matsuzaki, H.; Shimada, S. Characterization of Early Life Stress-Induced Gut Microbiota. Brain Sci. 2021, 11, 913. [CrossRef] [PubMed]
55. Rincel, M.; Aubert, P.; Chevalier, J.; Grohard, P.-A.; Basso, L.; de Oliveira, C.M.; Helbling, J.C.; Levy, E.; Chevalier, G.; Leboyer, M.; et al. Multi-hit early life adversity affects gut microbiota, brain and behavior in a sex-dependent manner. Brain Behav. Immun. 2019, 80, 179–192. [CrossRef] [PubMed]
56. Cheng, H.-w.; Jiang, S.; Hu, J. Gut-Brain Axis: Probiotic, Bacillus subtilis, Prevents Aggression via the Modification of the Central Serotonergic System. In Oral Health by Using Probiotic Products; Mahmoudi, R., Moosazad, S., Aghaei, K., Eds.; IntechOpen: London, UK, 2019.
57. Bailey, M.T.; Coe, C.L. Maternal separation disrupts the integrity of the intestinal microflora in infant rhesus monkeys. Dev. Psychobiol. 1999, 35, 146–155. [CrossRef]
58. Takada, M.; Nishida, K.; Kataoka-Kato, A.; Gondo, Y.; Ishikawa, H.; Suda, K.; Kawai, M.; Hoshi, R.; Watanabe, O.; Igarashi, T.; et al. Probiotic Lactobacillus casei strain Shirota relieves stress-associated symptoms by modulating the gut–brain interaction in human and animal models. Neurogastroenterol. Motil. 2016, 28, 1027–1036. [CrossRef]
59. Liu, F.; Li, J.; Wu, F.; Zheng, H.; Peng, Q.; Zhou, H. Altered composition and function of intestinal microbiota in autism spectrum disorder: A systematic review. Transl. Psychiatry 2019, 9, 1–13. [CrossRef] [PubMed]
60. Fung, T.C.; Vuong, H.E.; Luna, C.D.G.; Pronovost, G.N.; Aleksandrova, A.; Riley, N.G.; Vavilina, A.; McGinn, J.; Rendon, T.; Forrest, L.R.; et al. Intestinal serotonin and fluoxetine exposure modulate bacterial colonization in the gut. Nat. Microbiol. 2019, 4, 2064–2073. [CrossRef] [PubMed]
61. Kong, Q.; Wang, B.; Tian, P.; Li, X.; Zhao, J.; Zhang, H.; Chen, W.; Wang, G. Daily intake of Lactobacillus alleviates autistic-like behaviors by ameliorating the 5-hydroxytryptamine metabolic disorder in VPA-treated rats during weaning and sexual maturation. Food Funct. 2021, 12, 2591–2604. [CrossRef] [PubMed]
62. Badawy, A.A.-B. Tryptophan metabolism, disposition and utilization in pregnancy. *Biosci. Rep.* 2015, 35, e00261. [CrossRef]

63. Hilaire, G.; Voituron, N.; Menuet, C.; Ichiyama, R.; Subramanian, H.H.; Dutschmann, M. The role of serotonin in respiratory function and dysfunction. *Respir. Physiol. Neurobiol.* 2010, 174, 76–88. [CrossRef]

64. Seidman, L.J.; Buka, S.L.; Goldstein, J.M.; Horton, N.J.; Rieder, R.O.; Tsuang, M.T. The relationship of prenatal and perinatal complications to cognitive functioning at age 7 in the New England Cohorts of the National Collaborative Perinatal Project. *Schizophr. Bull.* 2000, 26, 309–321. [CrossRef] [PubMed]

65. Ali, S.M.; Lindström, M. Socioeconomic, psychosocial, behavioural, and psychological determinants of BMI among young women: differing patterns for underweight and overweight/obesity. *Eur. J. Public Health* 2005, 16, 324–330. [CrossRef] [PubMed]

66. Levitan, R.D.; Sqaipi, M.; Atkinson, L.; Murphy, K.; Levitt, A.; Bocking, A.; Post, M.; Knight, J.A.; Matthews, S.G. Seasonality of plasma tryptophan and kynurenine in pregnant mothers with a history of seasonal affective disorder: Vulnerability or adaptation? *World J. Psychiatry* 2020, 21, 529–538. [CrossRef] [PubMed]

67. Barker, D. The fetal origins of adult disease. *Fetal Matern. Med. Rev.* 1994, 6, 71–80. [CrossRef]

68. Balayan, K.; Verma, M. Understanding Bullying: Roles, Types, Gender Difference and Effect on Mental Health. *Int. J. Indian Psychol.* 2021, 9, 2217–2223. [CrossRef]

69. Hornef, M.; Penders, J. Does a prenatal bacterial microbiota exist? *Mucosal Immunol.* 2017, 10, 598–601. [CrossRef]

70. International Chicken Genome Sequencing Consortium. Sequence and comparative analysis of the chicken genome provide unique perspectives on vertebrate evolution. *Nature* 2004, 432, 695–716. [CrossRef]

71. Beacon, M.T.H.; Davie, J.R. The chicken model organism for epigenomic research. *Genome* 2021, 64, 476–489. [CrossRef]

72. Ahmed, A.A.; Ma, W.; Ni, Y.; Zhou, Q.; Zhao, R. Embryonic exposure to corticosterone modifies aggressive behavior through alterations of the hypothalamic pituitary adrenal axis and the serotonergic system in the chicken. *Horm. Behav.* 2014, 65, 97–105. [CrossRef]

73. Hymel, K.A.; Salmeto, A.L.; Loria, M.J.; White, S.W.; Sufka, K.J. Strain vulnerability and resiliency in the chick anxiety–depression model. *Physiol. Behav.* 2013, 120, 124–129. [CrossRef]

74. Cheng, H.; Singleton, P.; Muir, W. Social stress differentially regulates neuroendocrine responses in laying hens: I. Genetic basis of adrenal responses under three different social conditions. *Psychoneuroendocrinology* 2004, 29, 961–971. [CrossRef] [PubMed]

75. Cheng, H.; Muir, W. Chronic social stress differentially regulates neuroendocrine responses in laying hens: II. Genetic basis of neuroendocrine responses under three different social conditions. *Psychoneuroendocrinology* 2004, 29, 961–971. [CrossRef] [PubMed]

76. Lorenzi, E.; Pross, A.; Rosa-Salva, O.; Versace, E.; Sgadó, P.; Vallortigara, G. Embryonic Exposure to Valproic Acid Affects Social Predispositions for Dynamic Cues of Animate Motion in Newly-Hatched Chicks. *Front. Physiol.* 2019, 10, 501. [CrossRef] [PubMed]

77. Nishigori, H.; Kagami, K.; Takahashi, A.; Tezuka, Y.; Sanbe, A.; Nishigori, H. Impaired social behavior in chicks exposed to sodium valproate during the last week of embryogenesis. *Psychopharmacology* 2013, 227, 393–402. [CrossRef]

78. Sgadó, P.; Rosa-Salva, O.; Versace, E.; Vallortigara, G. Embryonic Exposure to Valproic Acid Impairs Social Predispositions of Newly-Hatched Chicks. *Sci. Rep.* 2018, 8, 5919. [CrossRef]

79. Lee, S.; La, T.-M.; Lee, H.-J.; Choi, I.-S.; Song, C.-S.; Park, S.-Y.; Lee, J.-B.; Lee, S.-W. Characterization of microbial communities in the chicken oviduct and the origin of chicken embryo gut microbiota. *Sci. Rep.* 2019, 9, 6838. [CrossRef] [PubMed]

80. Huang, X.; Hu, J.; Peng, H.; Cheng, H.-W. Embryonic Exposure to Tryptophan Yields Bullying Victimization via Reprogramming the Microbiota-Gut-Brain Axis in a Chicken Model. *Nutrients* 2022, 14, 661. [CrossRef]

81. Saffarian, A.; Mulet, C.; Regnault, B.; Amiot, A.; Tran-Van-Nhieu, J.; Ravel, J.; Sobhani, I.; Sansonetti, P.J.; Pédro, T. Crypt- and Mucosa-Associated Core Microbiotas in Humans and Their Alteration in Colon Cancer Patients. *mBio* 2019, 10, 01315–01319. [CrossRef]

82. Barandouzi, Z.A.; Starkweather, A.R.; Henderson, W.; Gymafi, A.; Cong, X.S. Altered Composition of Gut Microbiota in Depression: A Systematic Review. *Front. Psychiatry* 2020, 11, 541. [CrossRef]

83. Lasheras, I.; Seral, P.; Latorre, E.; Barroso, E.; Gracia-García, P.; Santabárbara, J. Microbiota and gut-brain axis dysfunction in autism spectrum disorder: Evidence for functional gastrointestinal disorders. *Asian J. Psychiatry* 2020, 47, 101874. [CrossRef]

84. Gophna, U.; Konikoff, T.; Nielsen, H.B. Oscillospira and related bacteria—From metagenomic species to metabolic features. *Environ. Microb.* 2017, 19, 81–841. [CrossRef] [PubMed]

85. Osadchiy, V.; Labus, J.S.; Gupta, A.; Jacobs, J.; Ashe-McNalley, C.; Hsiao, E.Y.; Mayer, E.A. Correlation of tryptophan metabolites with connectivity of extended central reward network in healthy subjects. *PLoS ONE* 2018, 13, e0201772. [CrossRef] [PubMed]

86. Knack, J.M.; Jensen-Campbell, L.A.; Baum, A. Worse than sticks and stones? Bullying is associated with altered HPA axis function and dysfunction. *Brain Cogn.* 2011, 77, 183–190. [CrossRef] [PubMed]

87. Berthoud, H.-R.; Albarga, V.L.; Neuhuber, W.L. Gut-brain communication and obesity: Understanding functions of the vagus nerve. *J. Clin. Investig.* 2021, 131, 139. [CrossRef]

88. Gastón, M.; Cid, M.; Salvatiera, N. Bicuculline, a GABA_A-receptor antagonist, blocked HPA axis activation induced by ghrelin under an acute stress. *Behav. Brain Res.* 2017, 320, 464–472. [CrossRef] [PubMed]

89. Gualtieri, F.; Armstrong, E.A.; Longmoor, G.K.; D’Eath, R.B.; Sandilands, V.; Boswell, T.; Smulders, T.V. Unpredictable Chronic Mild Stress Suppresses the Incorporation of Neurons at the Caudal Pole of the Chicken Hippocampal Formation. *Sci. Rep.* 2019, 9, 7129. [CrossRef]

90. Smulders, T. The Avian Hippocampal Formation and the Stress Response. *Brain Behav. Evol.* 2017, 90, 81–91. [CrossRef]
91. Smulders, T.V. Telencephalic regulation of the HPA axis in birds. *Neurobiol. Stress* 2021, 15, 100351. [CrossRef]

92. Li, X.; Jensen, R.L.; Højberg, O.; Canibe, N.; Jensen, B.B. Olsenella scatoligenes sp. nov., a 3-methylindole- (skatole) and 4-methylphenol- (p-cresol) producing bacterium isolated from pig faeces. *Int. J. Syst. Evol. Microbiol.* 2015, 65, 1227–1233. [CrossRef]

93. Wassum, K.M.; Izquierdo, A. The basolateral amygdala in reward learning and addiction. *Neurosci. Biobehav. Rev.* 2015, 57, 271–283. [CrossRef]

94. Gouveia, F.V.; Hamani, C.; Fonoff, E.T.; Brentani, H.; Alho, E.J.L.; de Morais, R.M.C.B.; de Souza, A.L.; Rigonatti, S.P.; Martinez, R.C.R. Amygdala and Hypothalamus: Historical Overview With Focus on Aggression. *Neurosurgery* 2019, 85, 11–30. [CrossRef] [PubMed]

95. Francis, D.D.; Caldji, C.; Champagne, F.; Plotsky, P.M.; Meaney, M.J. The role of corticotropin-releasing factor–norepinephrine systems in mediating the effects of early experience on the development of behavioral and endocrine responses to stress. *Biol. Psychiatry* 1999, 46, 1153–1166. [CrossRef]

96. Castelbaum, L.; Sylvester, C.M.; Zhang, Y.; Yu, Q.; Constantino, J.N. On the Nature of Monozygotic Twin Concordance and Discordance for Autistic Trait Severity: A Quantitative Analysis. *Behav. Genet.* 2020, 50, 263–272. [CrossRef] [PubMed]

97. Sugden, K.; Arseneault, L.; Harrington, H.; Moffitt, T.E.; Williams, B.; Caspi, A. Serotonin Transporter Gene Moderates the Development of Emotional Problems Among Children Following Bullying Victimization. *J. Am. Acad. Child Adolesc. Psychiatry* 2010, 49, 830–840. [CrossRef] [PubMed]

98. Moukhles, H.; Bosler, O.; Bolam, J.P.; Vallée, A.; Umbriaco, D.; Geffard, M.; Doucet, G. Quantitative and morphometric data indicate precise cellular interactions between serotonin terminals and postsynaptic targets in rat substantia nigra. *Neuroscience* 1997, 76, 1159–1171. [CrossRef]

99. Navailles, S.; De Deurwaerdère, P. Presynaptic control of serotonin on striatal dopamine function. *Psychopharmacology* 2010, 213, 213–242. [CrossRef]

100. De Deurwaerdère, P.; Di Giovanni, G. Serotonergic modulation of the activity of mesencephalic dopaminergic systems: Therapeutic implications. *Prog. Neurobiol.* 2017, 151, 175–236. [CrossRef]

101. Schlüter, T.; Winz, O.; Henkel, K.; Prinz, S.; Rademacher, L.; Schmaljohann, J.; Dautzenberg, K.; Cumming, P.; Kumakura, Y.; Rex, S.; et al. The Impact of Dopamine on Aggression: An [18F]-FDOPA PET Study in Healthy Males. *J. Neurosci.* 2013, 33, 16889–16896. [CrossRef]

102. Dennis, R.L.; Fahey, A.G.; Cheng, H.W. Alterations to Embryonic Serotonin Change Aggression and Fearfulness. *Aggress. Behav.* 2013, 39, 91–98. [CrossRef]

103. Dennis, R.; Cheng, H. The dopaminergic system and aggression in laying hens. *Poult. Sci.* 2011, 90, 2440–2448. [CrossRef]

104. Huang, X.; Kuang, S.; Applegate, T.J.; Lin, T.-L.; Cheng, H.-W. Prenatal Serotonin Fluctuation Affects Serotonergic Development and Related Neural Circuits in Chicken Embryos. *Neuroscience* 2021, 473, 66–80. [CrossRef] [PubMed]

105. Jiang-Xie, L.-F.; Liao, H.-M.; Chen, C.-H.; Chen, Y.-T.; Ho, S.-Y.; Lu, D.-H.; Lee, L.-J.; Liou, H.-H.; Fu, W.-M.; Gau, S.S.-F. Autism-associated gene Dlgap2 mutant mice demonstrate exacerbated aggressive behaviors and orbitofrontal cortex deficits. *Mol. Autism* 2014, 5, 32. [CrossRef] [PubMed]

106. Anderson, K.N.; Lind, J.N.; Simeone, R.M.; Bobo, W.V.; Mitchell, A.A.; Riehle-Colarusso, T.; Polen, K.N.; Reelhuis, J. Maternal Use of Specific Antidepressant Medications During Early Pregnancy and the Risk of Selected Birth Defects. *JAMA Psychiatry* 2020, 77, 1246. [CrossRef] [PubMed]