



The vicious cycle towards violence: focus on the negative feedback mechanisms of brain serotonin neurotransmission

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Violence can be defined as a form of escalated aggressive behavior that is expressed out of context and out of inhibitory control, and apparently has lost its adaptive function in social communication. Little is known about the social and environmental factors as well as the underlying neurobiological mechanisms involved in the shift of normal adaptive aggression into violence. In an effort to model the harmful acts of aggression and violence in humans, we recently (re)developed an animal model that is focused on engendering uncontrolled forms of maladaptive aggressive behavior in laboratory-bred feral rats and mice. We show that certain (8–12%) constitutionally aggressive individuals gradually develop, over the course of repetitive exposures to victorious social conflicts, escalated (short-latency, high-frequency and ferocious attacks), persistent (lack of attack inhibition by defeat/submission signals and perseverance of the aggressive attack-biting bout), indiscriminating (attacking female and anesthetized male intruders) and injurious (enhanced vulnerable-body region attacks and inflicted wounding) forms of offensive aggression. Based on the neurobiological results obtained using this model, a revised view is presented on the key role of central serotonergic (auto)regulatory mechanisms in this transition of normal aggression into violence.

Keywords: aggression, violence, rodents, serotonin, 5-HT_{1A} receptor

INTRODUCTION

The human capacity for unchecked aggressiveness and violence inflicts an awful and costly burden on society. Unfortunately, the current intervention strategies and treatment options for curbing these problematic behavioral expressions are largely inadequate. Hence, a more fundamental knowledge about the social and neurobiological determinants of aggression is desperately needed. In particular, the interaction between environmental factors and the neurochemical substrates that specifically underlies the shift towards escalated and maladaptive forms of aggressive behavior (e.g., violence) need to be unraveled. Experimental laboratory models of aggression in rodents and other animal species are indispensable to obtain this goal. However, most animal studies of aggression have traditionally been concerned with the ultimate and proximate mechanisms of normal adaptive aggressive behavior, while clinically the focus is predominantly on violent individuals and excessive or inappropriate forms of human aggression. Besides several ethical, funding and translational constraints, the lack of biologically relevant and valid animal models of these pathological forms of aggressive behavior is one important reason for the gap in our knowledge about the neurobiological roots and mechanisms of violence in humans. Therefore, new experimental models in preclinical research are now being developed that focus more on provoking escalated and uncontrolled forms of aggressive behavior in order to approximate the behavior of direct concern in public health and criminal justice (Miczek et al., 2004, 2007; Haller and Kruk, 2006; Nelson and Trainor 2007).

ANIMAL MODELS

One major methodological obstacle for developing valid animal models of violence is the fact that most laboratory animal studies of aggression are conducted in rodent strains that are very placid and docile. In virtually all laboratory mouse and rat species, the aggressive behavioral traits (including the underlying genetic components) have been dramatically compromised due to selection and inbreeding in the course of the century-long domestication process. Consequently, to obtain appreciable levels of offensive aggression in these timid and predominantly serene laboratory strains, several procedural manipulations have been employed to enhance their general tendency to display offensive aggressiveness. Obviously, some of these procedures have been adopted with the intent to mimic the conditions under which violent behavior in humans occur. Generally, these procedures can be categorized into manipulations at the (A) behavioral/environmental, (B) neurophysiological (C) neuropharmacological and (D) genetic level. An overview of these various procedural manipulations/paradigms that heighten aggression, and its similarity to the symptomatology, etiology, neurobiology and pharmacology of human violent behavior according to the general validity criteria of animal models (face, construct and predictive), are listed in **Table 1** and the reader is referred to another paper in this special issue (Natarajan et al., in review) for a brief description of these various animal models.

Clearly, several of these models have a reasonable construct-validity with the violence-provoking conditions and mechanisms in humans. However, while the listed experimentally-heightened forms of aggressive behavior may to some extent resemble more intense forms when compared to the already low species-typical rates of

Table 1 | Procedural manipulations to induce increased levels or pathological forms of aggressive behavior in experimental animals (rats, mice and hamsters).

Procedural manipulations	Validity			References
	Face	Construct	Predictive	
BEHAVIORAL/ENVIRONMENTAL				
Social isolation/deprivation-enhanced aggression				
• In early life stages (maternal separation)	Poor	Fair	NA	Veenema (2009)
• During adolescence/young adult age	Poor	Fair	Fair	Toth et al. (2008), Guidotti et al. (2001)
Stress provocation-elicited aggression				
• Electric-footshock exposure	Poor	Poor	Fair	Blanchard et al. (1978)
• Frustrative non-reward	Fair	Good	Fair	Miczek et al. (2002)
• Social instigation/provocation or priming	Fair	Good	Fair	Miczek et al. (2002)
• Chronic unpredictable mild stress	Poor	Fair	NA	Mineur et al. (2003)
Anticipation/experience-heightened aggression				
• Instrumental conditioning	Fair	Fair	Fair	Miczek et al. (2002)
• Repetitive winning	Fair	Good	NA	Hsu et al. (2005)
NEUROPHYSIOLOGICAL				
Electrical/chemical brain stimulation – induced				
• Hypothalamus	Fair	Poor	Fair	Kruk (1991)
• PAG	Fair	Poor	NA	Siegel (2004)
• Amygdala	Fair	Poor	NA	Keele (2005)
• Vasopressin	Fair	Poor	NA	Ferris (1992)
Electrolytic/chemical brain lesion-induced				
• Prefrontal cortex	Poor	Fair	NA	de Bruin et al. (1983)
• Raphe or general 5-HT depletion	Fair	Fair	NA	Olivier et al. (1990), Olivier (2004)
(NEURO)PHARMACOLOGICAL				
Glucocorticoid hypofunction-enhanced	Fair	Fair	Fair	Haller and Kruk (2006)
Adolescent chronic drug exposure–enhanced				
• Anabolic steroids	Fair	Good	NA	Melloni et al. (1997)
• Cocaine	Fair	Good	Fair	Harrison et al. (2000)
• Alcohol	Fair	Good	NA	Ferris et al. (1998)
• Glucocorticoids	Fair	Poor	NA	Wommack et al. (2005)
Adult acute drug treatment-heightened				
• Alcohol	Fair	Good	Fair	Miczek et al. (2004)
• Amphetamine-(derivatives)	Fair	Fair	NA	Grimes et al. (2006)
GENETIC				
Artificial selection for high-aggressiveness				
• SAL mice	Good	Good	Fair	Caramaschi et al. (2008)
• TA mice	Fair	Good	NA	Natarajan et al. (2009a,b)
• NC900 mice	Poor	Good	NA	Natarajan et al. (2009a,b)
Targeted gene deletion/overexpression-enhanced				
• > 45 gene candidates	Fair	Good	NA	Maxson and Canastar (2003), Maxson (2009)

For every paradigm its putative similarity with human violence is roughly indicated (good, fair or poor, NA = not assessed yet), as to its symptomatology (face validity), etiology and neurobiology (construct validity), and pharmacology (predictive validity). For every paradigm, a reference to one key (review) article describing the employed methodology is given.

aggression in these laboratory animals, they mostly still fall into the normative range when compared to the patterns and levels of their wild ancestors. Indeed, much higher levels and broader-ranges of spontaneous and normal adaptive resident-intruder offensive aggression are encountered in feral or semi-natural populations of rats and mice as compared to their laboratory-bred conspecifics (de Boer et al., 2003). Therefore, an increase in the intensity of aggression as reflected by shorter attack latencies and/or enhanced

frequency and duration of attacks is only one quantitative aspect of an individual's aggressive behavioral tendency and does not automatically reflect abnormal forms of aggression that are characteristic of violence. More productive and clinically relevant animal models of pathological expression of aggression should demonstrate intense and injurious aggression that *exceeds normal species-typical levels and patterns* (Miczek et al., 2004; Haller and Kruk, 2006; Nelson and Trainor, 2007). In other words, forms of aggressive behavior

that is not subject to inhibitory control anymore and has lost their adaptive function in social communication. This out of control, out of content and out of context aggressive behavior is expressed in:

1. The disappearance of the normal investigatory and threatening sequence of acts and postures, and immediately engage in the ultimate consummatory phase of aggression i.e., the physical attack-bite bout (reflected in decreased attack-bite latencies and high attack/threat ratios).
2. Persistence in the aggressive attack-biting mode even though the intruder displays submissive supine and crouching/defeat postures that normally terminates the aggressive attack bout in the resident (quantifiable by disorganized temporal and sequential structure of social and aggressive behavior or perseverance of the aggressive attack-biting bout).
3. Orienting the attack-bites toward vulnerable-body regions (head and ventral surface) of the opponents resulting in severe wounding and eventually death of the intruder if not stopped by the experimenter (indicated by increased vulnerable attack ratios and inflicted wounds).
4. Losing the ability to discriminate context and/or the type of opponent (resulting in attacking dominant males, females or even anesthetized/dead conspecifics). Inclusion of these core behavioral abnormality criteria for deviant aggressiveness in rodent aggression models will certainly add to the validity of a “violence” model and expansion of our much needed knowledge about the neurobiology of pathological and violent forms of human aggression.

FROM AGGRESSION TO VIOLENCE

Recent experiments in our lab using feral (wild-derived) rats and mice have shown that the above described violent characteristics can be engendered in certain medium to high-aggressive wild-type rats (approximately 8–12% of the animals) upon repeatedly permitting them to dominate intruder conspecifics during resident-intruder contests. Enhanced levels of offensive aggression and an increased probability of winning an aggressive encounter following previous victories (the so-called “trained fighter” or “winner” effect) was originally described by Ginsberg and Allee (1942), and since then has been demonstrated frequently in a wide variety of animal species (see Hsu et al., 2005, for review). Furthermore, this effect is reported to be most pronounced in the more aggressive individuals (Oyegbile and Marler, 2006). Clearly, some of our constitutionally aggressive male wild-type rats gradually develop, over the course of 10–25 repetitive exposures of victorious social conflicts escalated (short-latency, high-frequency and –intensity attacks), persistent (lack of attack inhibition by defeat/submission signals), indiscriminating (attacking female and anesthetized male intruders) and injurious (enhanced vulnerable-body region attacks) forms of offensive aggression (Figure 1).

Similarly, male wild-derived house mice that were artificially selected for short-attack latencies (i.e., high-aggressive SAL mice) show virtually all of the above-mentioned signs of violent aggressive behavior already after 3–5 repeated winning experiences (Haller et al., 2006; Caramaschi et al., 2008; Natarajan et al., 2009a,b). Compared to non-aggressive LAL or low/normal-aggressive aLAL males, SAL mice display very short-latency and high-frequency of biting attacks with approximately half of those aimed at vulnerable-body regions (neck,

head and belly). Furthermore, 80–90% of the fighting-experienced SAL males started to attack anaesthetized male intruders as well as (oestrus) females (Figure 2).

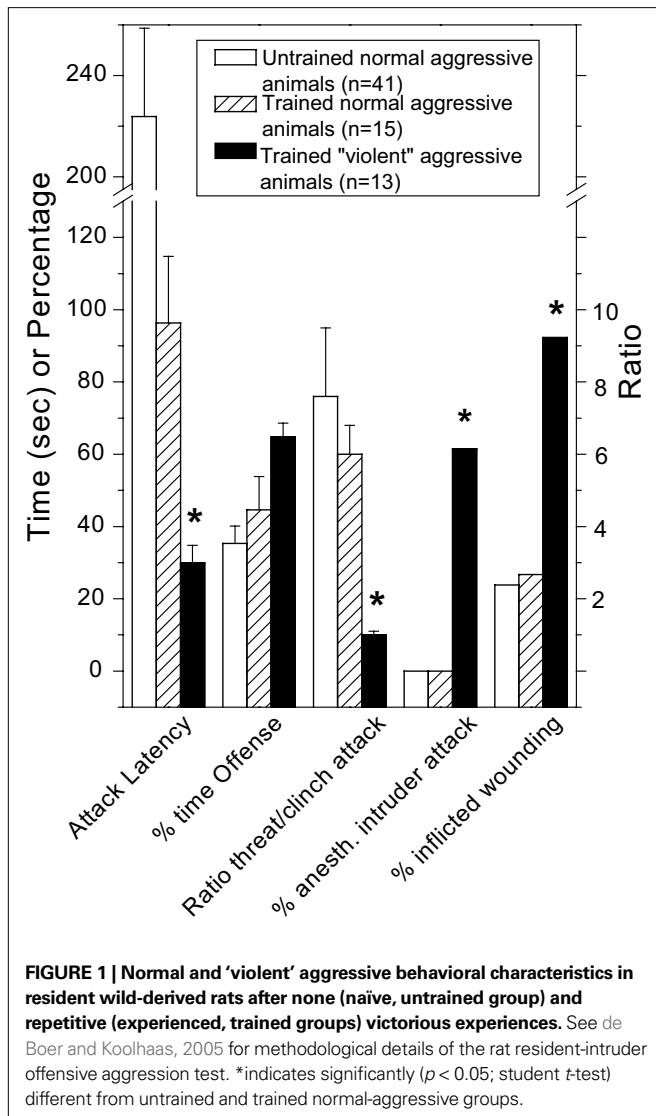
Thus, upon positive reinforcing or “pleasurable” victorious social experiences, these constitutionally high-aggressive rats and mice are very prone to show a breakdown of the aggressive behavioral inhibition mechanisms, and transform their initial functional adaptive aggressive behavior into a more violent and pathological form.

SEROTONIN

Obviously, this animal model affords us the opportunity to identify the neural plastic changes in the neurochemical “aggression” control systems that are hypothesized to underlie such a transformation. To date, we have focused in particular on the (auto)-regulatory components of brain serotonin (5-HT) neurotransmission. More than any other neurochemical system, this evolutionary ancient and very well conserved neurotransmitter system is generally considered the primary molecular orchestrator of aggression in a wide variety of animal species (Kravitz and Huber 2003; Miczek et al., 2004; Summers et al., 2005). However, the nature of this linkage is complex and it has proven difficult to unravel the precise role of this amine in the predisposition for and execution of aggressive behavior in both its normal and pathological forms. The big dogma that 5-HT generally inhibits aggression is obsolete now and the much-favored 5-HT deficiency hypothesis of aggression has to be revised. Ongoing research from our and other research groups on the functional status of the 5-HT system before, during and after the execution of normal adaptive and abnormal pathological forms of aggression has led to the following view: display of normal adaptive aggressive behavior aimed at territorial control, social dominance and coherence is positively related to 5-HT neuronal (re)activity, whereas an inverse relationship develops between tonic, trait-like 5-HT activity and pathological forms of aggression (e.g., violence). For example, in our wild-type rats, a clear positive correlation was found between the level of adaptive aggressiveness and basal CSF concentrations of 5-HT and/or its metabolite 5-HIAA (van der Vegt et al., 2003a). Moreover, local tissue as well as microdialysate levels of 5-HT and 5-HIAA in the frontal cortex did not differ significantly between high- and low-aggressive animals, although there was a slight tendency for lower levels in high-aggressive animals (de Boer et al., 2003; Figure 3). This negative correlation between aggression and frontal cortical 5-HT levels became highly significant upon inclusion of samples from the abnormal and excessively aggressive trained fighter animals (Figure 3).

Apparently, tonic brain 5-HT activity decreases as a *consequence* of acquiring repeated victorious experiences and adopting unchecked forms of aggression. A similar observation was made recently in our feral, artificially-selected high-aggressive mice: Only *after* repeated resident-intruder fighting experience, serotonin levels in the prefrontal cortex were found to be significantly lower in violent SAL mice (Figure 4; Caramaschi et al., 2008).

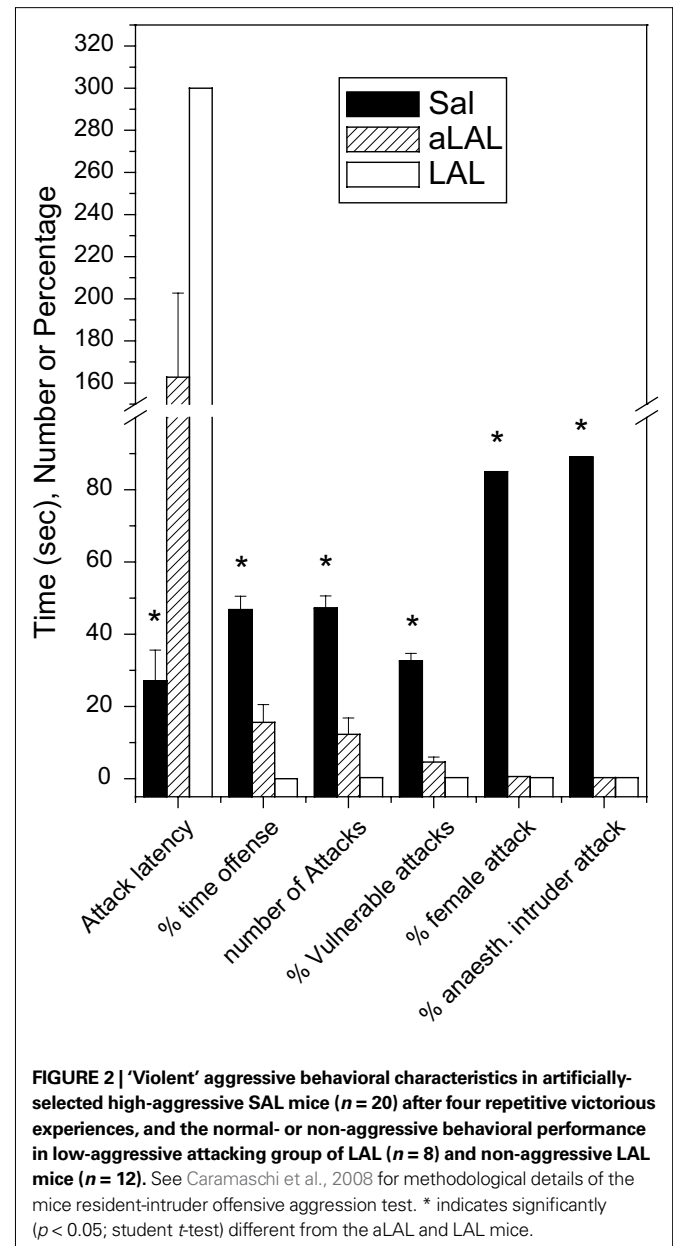
Moreover, these data are consistent with microdialysis studies showing lower prefrontal 5-HT values in aggressive individuals only *after* the fighting had been initiated and when it was terminated (Van Erp and Miczek, 2000) or merely anticipating aggression in individuals who are extensively trained/primed for aggression (Ferrari et al., 2003). Furthermore, these findings accommodate and bridge the aggressive behavior stimulatory role of 5-HT described



for several invertebrate species (for review see Kravitz and Huber, 2003) and the putative inhibitory role of 5-HT in rodent and primate species. Indeed, a critical evaluation of the csf 5-HIAA data in aggressive humans and monkeys revealed that the serotonergic deficiency appears to be most readily detected in well-circumscribed groups of individuals who had engaged in impulsive and violent forms of aggressive behavior rather than in individuals with more instrumental (functional) forms of aggression (Coccaro, 1989; Raleigh et al., 1991; Mehlman et al., 1994; Tuinier et al., 1995).

5-HT AUTORECEPTORS

Profound functional changes in the key autoregulatory sites that importantly control firing and 5-HT release of the serotonergic neurons, i.e., presynaptic 5-HT_{1A} and 5-HT_{1B} autoreceptors and serotonergic reuptake transporters (5-HTT), are hypothesized to causally underlie this transition of normal adaptive aggressive behavior into abnormal excessive forms that cause harm and injury. Previously, we and others clearly showed that acute treatment with 5-HT_{1A} and 5-HT_{1B} receptor agonists potently and selectively



suppressed aggressive behaviour (see Miczek et al., 2004 and Olivier and van Oorschot, 2005 for reviews). Moreover, we showed that these anti-aggressive effects are largely expressed via their action on the inhibitory somatodendritic and terminal autoreceptors, presumably attenuating intruder-activated 5-HT neurotransmission (see de Boer and Koolhaas, 2005 for review). This strongly indicates that the actual display of aggressive behavior, or at least the intent thereof, is associated with (short-lasting) increased 5-HT neurotransmission, a paradoxical view but supported by increased molecular neuronal activation markers (immediate-early gene expression) in 5-HT-positive raphe neurons after performance of aggressive behavior (van der Vegt et al., 2003b; Veening et al., 2005). This view agrees well with the general hypothesis that activation of the 5-HT system is most strongly related to general motor activity (Jacobs and Fornal, 1999).

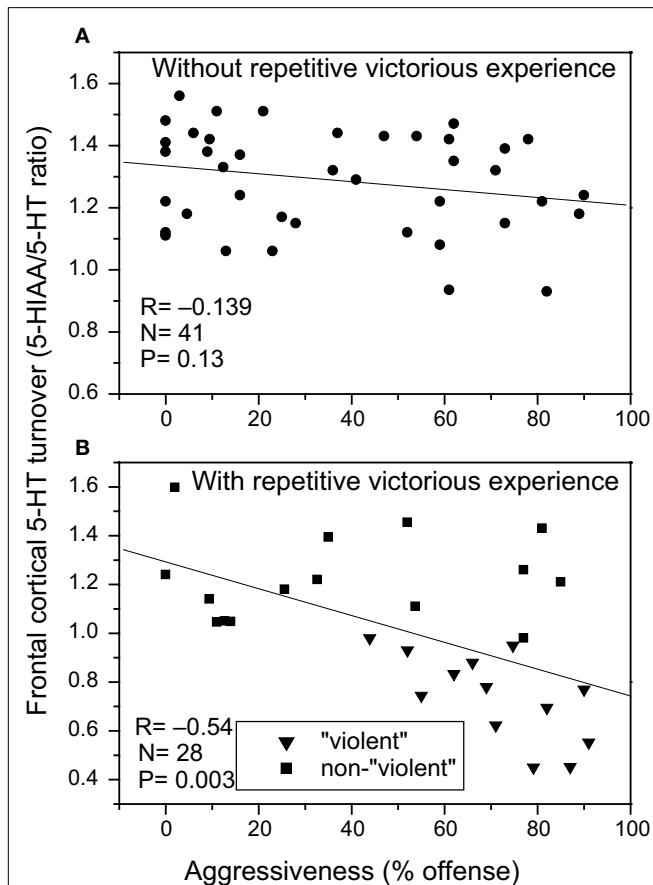


FIGURE 3 | Relationship between the individual level of aggressiveness (measured as % time spent on offensive aggressive behaviors during a 10 min resident-intruder conflict test) and frontal cortical serotonin turnover (measured as the ratio between tissue concentration 5-HIAA and 5-HT) in (A) untrained residents and (B) trained fighter rats that either show abnormal and/or violent aggressive characteristics (filled triangles) or normal forms of aggressiveness (filled squares). Note that 5-HT turnover deficiency occurs in only those individuals that developed excessive and abnormal forms of aggressiveness upon repeatedly winning fights. Data partly taken from de Boer and Koolhaas (2005), see Caramaschi et al. (2007) for technical details of prefrontal tissue 5-HT/5-HIAA measurements.

More importantly, we found that the broad individual differences in offensive aggression in our wild-type animals are associated with pronounced functional, but not structural, differences in 5-HT_{1A} and 5-HT_{1B} properties: compared to low-aggressive animals, high-aggressive animals are characterized by enhanced somatodendritic 5-HT_{1A} and terminal 5-HT_{1B} autoreceptor activity (de Boer et al., 2001; van der Vegt et al., 2001; Caramaschi et al., 2007).

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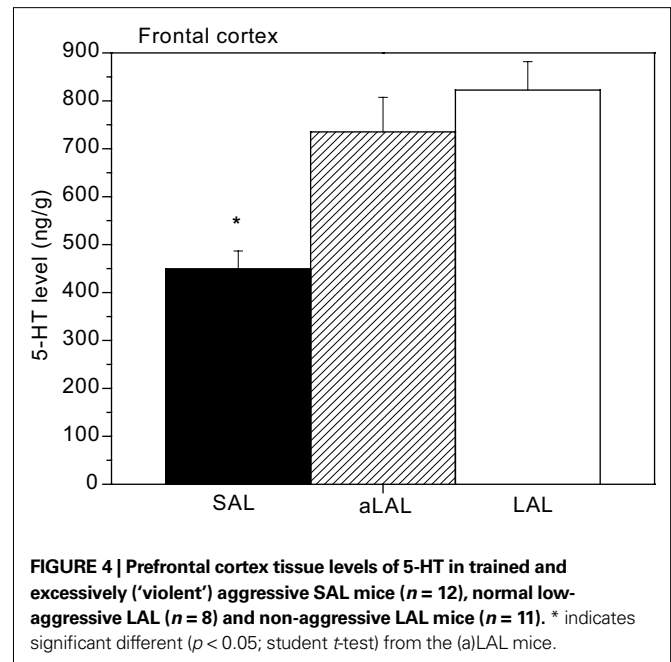


FIGURE 4 | Prefrontal cortex tissue levels of 5-HT in trained and excessively ('violent') aggressive SAL mice ($n = 12$), normal low-aggressive LAL ($n = 8$) and non-aggressive LAL mice ($n = 11$). * indicates significant different ($p < 0.05$; student t -test) from the (a)LAL mice.

Furthermore to signify the causality of this correlation, 5-HT_{1A} autoreceptor sensitivity increased or decreased upon enhancing (by repeated victory experiences) or attenuating (by repeated defeat experiences) aggressiveness, respectively.

Currently, we scrutinize the various cellular and molecular mechanisms (i.e., receptor internalization, G_i/G_o-effector protein coupling, GIRK-channel activation/expression, TREK-1 potassium channel activation/expression) that importantly regulate this inhibitory autoreceptor function to ascertain the main cause of the detected 5-HT_{1A} autoreceptor supersensitivity.

CONCLUDING REMARKS

Our results support the need to delineate aggression as a highly functional form of social communication from violence. Display of normal adaptive aggressive behavior aimed at territorial control, social dominance and coherence is positively related to 5-HT neuronal (re)activity, whereas an inverse relationship develops between tonic, trait-like 5-HT activity and pathological forms of aggression. Based on our findings we hypothesize that the enhanced inhibitory autoreceptor function is a normal compensatory adaptive response to the more reactive state of the brain 5-HT system in high-aggressive animals. However, an excessive activation or perhaps overshooting of this normal autoreceptor brake function may be a causative link in the cascade of neurochemical events leading to the 5-HT hypofunction characterizing violence as the pathological form of aggression.

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