Aripiprazole and Riluzole treatment alters behavior and neurometabolites in young ADHD rats: a longitudinal $^1$H-NMR spectroscopy study at 11.7T

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Attention deficit hyperactivity disorder (ADHD), Tourette syndrome (TS) and obsessive compulsive disorder (OCD) are neurodevelopmental psychiatric disorders known to be accompanied by alterations of neurometabolites in the frontocortical area affecting mainly glutamatergic circuits. These disorders are often comorbid in TS and converge during school age and adolescence in a phenotype with hyperkinetic symptoms, motor and/or phonic tics.1–5 A general cortical disinhibition has been hypothesized as a possible explanation of these shared pathophysiological features, however, a well-defined etiology of the co-occurrence is still missing.6

There is no cure for TS, however, there are treatments to help manage the tics including pharmacological interventions. In this respect there has been a long-lasting debate about a potential worsening of tic symptoms during stimulant medication.7 Although current guidelines recommend the treatment of ADHD and co-occurring tics with stimulants, German guidelines for tic management in Tourette patients8–10 recommend the use of aripiprazole due to its therapeutic influence on ADHD symptoms and mild side effect profile.11,12 Aripiprazole is a ‘dopaminergic stabilizer’ in use for the treatment of schizophrenia patients but can also be used as first choice to treat ADHD in patients with TS. Riluzole is an approved drug for the treatment of amyotrophic lateral sclerosis (ALS) leading to a prolongation of patient survival and providing a long-term safety profile. It has already been used in open and randomized clinical trials for OCD symptoms13–17 and very recently also for tic management.18

Co-occurring ADHD, OCD and TS symptoms are oftentimes worsening during adolescence leading to changes in drug regimens and to polypharmacy. The changes of symptoms might be caused by neurochemical alterations affecting only certain brain regions or systems. To analyze these developmental changes longitudinal studies are needed. With respect to neurometabolic adaptions, modern techniques like Proton magnetic resonance spectroscopy ($^1$H-MRS)19 are unique in their ability to gain repeated access to multiple brain regions at defined developmental stages providing a longitudinal set of data on an extensive neurometabolite battery. $^1$H-MRS is a non-invasive in vivo method capable of measuring neurometabolite concentrations in a selected voxel (volumetric pixel) placed within a specific region of interest (ROI). By this, even subtle concentration changes can be detected that might have a potential pathophysiological significance for brain development.20,21

Our study provides for, we believe, the first time a detailed overview of the absolute concentration changes during the adolescent period of 12 neurometabolites within the juvenile male rat striatum and prefrontal cortex. These regions resemble the main inhibitory and excitatory areas of the cortico-striatal-thalamic-cortical circuit thought to be affected during neurodevelopment in TS, ADHD and OCD patients.1 Moreover, an age- and strain-related detailed neurochemical profile of the spontaneously hypertensive rat (SHR), the most-commonly used animal model for ADHD, and its normotensive control, the Wistar Kyoto rat (WKY)22 is provided. Interestingly, we could identify a neurometabolomic
fingerprint for each of the strains that can be used as reference for further studies. Finally, we investigated the effect of aripiprazole and riluzole, two newly proposed compounds to treat the hyperkinetic symptomatology in young patients affected by TS. To that end we generated neurochemical profiles of hyperactive and control juvenile animals and could correlate those with a thorough analysis of behavior. In summary, this study provides (I) a detailed developmental neurochemical profile of the strains investigated under control conditions, (II) under subchronic pharmacological treatment and (III) a correlation analysis between the absolute quantification of neurometabolites and the behavioral profiles.

MATERIALS AND METHODS

Animals

Behavioral assessments and MRS measurements were carried out within a period of 13 months on six cohorts of juvenile male Wistar Kyoto (WKY) and SHR (Charles River Laboratories, Sulzfeld, Germany). Up to 4 pups per cage were housed together with the mother animals until post-natal day (PND) 21. For the entire duration of the experiment the rats had free access to standard laboratory rodent chow and water, and were housed in a room with 12 h light-dark cycle. Temperature and humidity were maintained at 22 ± 1 °C and 45–55%, respectively.

The study design is to analyze treated and untreated animals in a longitudinal manner (PND 35, 42, 46 and 50) in terms of behavior and neurochemical alteration during the period of adolescence. All experiments were approved by the Committee for Animal Experimentation of the University of Ulm and the regional administrative authority in Tübingen (TVA No. 1200).

Drug treatment

In this gender- and age-matched vehicle-controlled designed study, each cohort randomly underwent a subchronic pharmacological intervention with vehicle (saline, 1% Tween80, Sigma Aldrich, Taufkirchen, Germany, cat. No. P17541), aripiprazole (Sigma Aldrich, cat. No. SML0935; 1.5 mg kg$^{-1}$ i.p.) or riluzole (Sigma Aldrich, cat. No. R3772; 6 mg kg$^{-1}$ i.p.). The subchronic daily drug treatment started on PND 35, when rats were weighing 100 g ± 30 g, and ended on PND 50. Investigations were carried out within both WKY (9 vehicle, 7 aripiprazole and 8 riluzole) and SHR (10 vehicle, 9 aripiprazole and 9 riluzole) strain. The sample size was determined on the WKY (9 vehicle, 7 aripiprazole and 8 riluzole) and SHR (10 vehicle, 9 aripiprazole and 9 riluzole) strain. The sample size was determined on the WKY (9 vehicle, 7 aripiprazole and 8 riluzole) and SHR (10 vehicle, 9 aripiprazole and 9 riluzole) strain. The sample size was determined on the WKY (9 vehicle, 7 aripiprazole and 8 riluzole) and SHR (10 vehicle, 9 aripiprazole and 9 riluzole) strain. The sample size was determined on the WKY (9 vehicle, 7 aripiprazole and 8 riluzole) and SHR (10 vehicle, 9 aripiprazole and 9 riluzole) strain. The sample size was determined on the WKY (9 vehicle, 7 aripiprazole and 8 riluzole) and SHR (10 vehicle, 9 aripiprazole and 9 riluzole) strain. The sample size was determined on the WKY (9 vehicle, 7 aripiprazole and 8 riluzole) and SHR (10 vehicle, 9 aripiprazole and 9 riluzole) strain. The sample size was determined on the WKY (9 vehicle, 7 aripiprazole and 8 riluzole) and SHR (10 vehicle, 9 aripiprazole and 9 riluzole) strain.

Behavioral assessment

Spontaneous locomotor activity in standard cage type III analysis. All behavioral experiments were conducted on PND 35, 42, 46 and 50 immediately before the MRS acquisition session. On each test day animals were weighed before testing and placed in the center of the individual test cage (Plexiglas standard cage type III, 425 × 266 × 155 mm) to record locomotor activity. The recording was performed for 10 min. at 50 fps rate using two cameras (Panasonic HC-V160 High Definition Video Camera) placed above (60 cm distance) and on the side (50 cm distance) of the cage, respectively. Each video was analyzed using the Ethovision 10.XT software (Noldus, Wageningen, The Netherlands) for animal behavior quantification. Horizontal locomotor activity was tracked during 5 of the 10 videotaped minutes from minute 2 to 7 and it was analyzed in terms of mean distance moved (cm), velocity (cm s$^{-1}$), time spent moving/not moving (s).

Repetitive behavior analysis. The same videos were used for stereotypic behavior quantification blindly conducted by a trained observer. Within the same 5 min. of locomotion tracking, repetitive behaviors were counted. Grooming (behaviors composed of paw licking, paw rubbing over the head, licking and rubbing the side of the body, the ano-genital region and the tail), rearing (vertical locomotion up or down a vertical surface using the forepaws only) and climbing behavior (vertical locomotion up on a vertical surface using both forepaws or pull and hind paws or push to grasp footholds and haul themselves up). Each behavior was analyzed in terms of frequency (counts) and total time spent (s) performing the specific behavior.

Proton ($^1$H) magnetic resonance spectroscopy session

MRS was carried out on a 11.7T animal scanner (Bruker, Ettlingen, Germany) equipped with a volume resonator for excitation and a phased-array surface coil for signal reception. Two regions of interest of 18.5 and 27.5 mm$^3$ volume respectively positioned in the left striatum and the left medial prefrontal cortex were investigated. MRS was performed with a short-echo-time STEAM (Stimulated Echo Acquisition Mode) sequence at TR/TE/TM: 5000/3.5/10 ms, 5 kHz spectral width, 2048 data points and 256 averages. Water-suppressed spectra were collected over a total acquisition time of 21 min. Molar metabolite concentration was quantified by LC Model (Version 6.3-1C, S. Provencher, Oakville). The unsuppressed water signal was used as an internal reference.

All spectra were visually inspected for the presence of spurious resonances or artefacts prior to the statistical analysis. Cramer-Rao lower bounds (CRLB) of LC Model analysis were used to assess accuracy and reliability of the CRLB higher than 50% was considered as exclusion criterion for metabolite evaluation. Other rejection criteria, such as poor SNR (< 6), existence of strong baseline distortions and line widths (full-width at half-maximum peak height, FWHH) were applied. MRS investigation immediately followed behavioral analysis. Spectra were acquired longitudinally from the same subject on PND 35, 42, 46 and 50 under anesthesia (0.8–1.8% isoflurane: air mixture supplied by face mask). Consistent with published recommendations for prolonged anesthesia exposure, breath rate was monitored throughout the scanning and isoflurane was adjusted to maintain respiration within specified target range (35–45 c.p.m.). Body temperature was maintained using a heated hydro system. The total duration of MRS including animal preparation was 90–110 min.

Statistical analysis

Data are reported as mean ± s.e.m. unless otherwise stated. Behavioral data analysis was done by Mann–Whitney U-test since the normal assumption was questionable. For each metabolite detected using 1H MRS technique, two-tailed paired t-test was used to compare changes in neurometabolite absolute concentrations between the two rat strains treated with vehicle. Respective data checks supported the normal assumption here. Finally, time- and treatment-related changes in neurometabolite absolute concentrations were assessed separately for each metabolite by two-way analysis of variances with treatment as grouping factor and time (PND 35, 42, 46 and 50) as within-group factor. The presence of variance heterogeneity was excluded by the Levene-test. Data were again tested for normal distribution and Bonferroni post hoc tests were applied to account for multiple testing arising from pairwise comparisons. Specifically, each time point of the drug treated groups were compared to the respective vehicle-treated group within the same strain. All differences were considered significant when $P < 0.05$. The analyses were conducted in GraphPad Prism 6.

RESULTS

Strain differences of WKY and SHR rats under vehicle and drug treatment conditions during adolescence.

Body weight during treatment and brain weight at the end of the treatment period. The body weight of the rats steadily increased during the whole experimental course according to the age of both strains and there were no significant differences in body weight between SHR and WKY at any developmental time point. Rat brains were collected at the end of the experimental period and weighed and there were no statistically related differences (Supplementary Figure S1a). Moreover, at any developmental stage both drug treatments did not affect the physiological body weight gain, the daily drug treatment did also not affect the total brain weight after 15 days of subchronic administration (Supplementary Figure S1b).

Spontaneous locomotor activity. During the entire duration of the experiment the spontaneous locomotor activity was analyzed. The SHR strain showed the typical hyperactive phenotype compared...
to WKY rats ($P < 0.0001$, Supplementary Figure S2a–d). Treatment with both drugs did not affect the strain-specific spontaneous locomotor activity (mean distance travelled as well as mean velocity). After an initial and significant increase of activity in the SHR strain treated with riluzole the effect was lost to the end of the treatment period (Figures 1a and b).

**Repetitive behavior analysis.** Spontaneous repetitive behavior like grooming, climbing and rearing behavior (mean frequency and duration) were analyzed. As shown in the Supplementary Material (Supplementary Figure S3a and b) the SHR strain showed an age-related, significantly increased frequency of rearing (PND 35, 46, 50, $P < 0.001$; PND 42, $P < 0.01$) and climbing (PND 35, 42, 50, $P < 0.05$; PND 46, $P < 0.01$) behavior compared to WKY rats in which such a climbing behaviour cannot be recognized. Grooming frequency was shown to be significantly different between strains only on PND 42 ($P < 0.01$) and 46 ($P < 0.05$). SHRs also show a significant increased time spent rearing (PND 35, $P < 0.05$; PND 42, 46, 50 $P < 0.001$), climbing (PND 42, 50 $P < 0.05$, PND 46, $P < 0.01$), but not grooming compared to WKY (Supplementary Figure S4).

As shown in Figures 2a and b, drug treatment during adolescence with both drugs significantly reduced grooming (PND 35, $P < 0.01$; PND 42, $P < 0.05$) and climbing (PND 35, $P < 0.001$; PND 42 and 46, $P < 0.05$) frequency in the SHRs but not in the WKY strain. Rearing frequency was not affected. The same treatment does not affect the time spent grooming, climbing or rearing (Supplementary Figure S4).

**In vivo ($^1$H) MRS absolute quantification of neurometabolites during adolescence**
Typical locations and size of the volumes of interest (VOIs) centered in the striatum and prefrontal cortex that were selected for spectroscopic studies and a representative spectrum for each of the analyzed VOI is shown in Figures 3a and b. Metabolites used in the final analysis were GABA, glutamate, glutamine, aspartate, inositol, taurine, PE, alanine, and GSH as absolute concentration, while combined values were reported for NAA+NAAG (total NAA), GPC+PCh (total choline), and Cr+PCr (total creatine) due to the inability to consistently resolve individual components. Data from prefrontal cortex and striatum are shown respectively in Supplementary Figure 6a and b.

Region-specific neurochemical profile of WKY and SHR rats under vehicle, aripiprazole and riluzole treatment during adolescence. Neurometabolite levels quantified in both striatum and prefrontal cortex within the WKY and SHR groups treated with vehicle showed time course dependent trajectories. They display a brain region-specific profile that for some components was also strain-specific. In the striatal region, PND 50 reveals the largest strain-related differences: SHR show significant higher striatal glutamine and aspartate concentrations while in the prefrontal region higher glutamine levels are accompanied by increased glutamate, total NAA, taurine and alanine concentrations. Total creatine shows higher concentrations starting from late adolescence, while the low aspartate concentration present at the early- and mid-adolescence reach WKY levels toward the latest time points.
Data are shown in the Supplementary Material (Supplementary Figure S5).

Under daily, subchronic drug treatment conditions during rats’ adolescence, we did not detect any significant neurochemical alteration within the brain region-specific profile of WKY rats, both with aripiprazole and riluzole administered intraperitoneally. On the contrary, in SHR rats both drug treatments exerted a decrease of the concentrations of neurometabolites and in general the effect was stronger in the prefrontal cortex compared to the striatum. In particular, aripiprazole treatment of the juvenile SHR strain induced a strong reduction of prefrontal glutamine, glutamate and GABA that persisted for all time points analyzed as well as a strong and stable reduction of total NAA, GSH, inositoles and creatine concentration. PE, taurine and total choline also showed initially significantly reduced prefrontal levels that then tend to stabilize over time, while alanine and aspartate levels seem not to be affected by treatment with aripiprazole.

On the other hand, riluzole treatment exerts a strong, stable and specific reduction of the prefrontal levels of glutamate over time, while glutamine levels start to decrease at PND 46 and GABA levels are not severely affected. Total creatine and inositole levels are also heavily reduced during the entire period of riluzole treatment. A reduction of glutamine, total NAA, PE, GSH and aspartate could be detected towards PND 46. Alanine levels, again, seem not to be affected by the treatment.

Finally, within the striatum, both drugs showed an overall weaker effect and significantly lower neurometabolite concentrations could only be randomly detected. Worth noticing is that

Figure 2. Grooming, climbing and rearing frequency of juvenile WKY- (a) and SHR- rats (b) treated with vehicle (black filled line), aripiprazole (red dotted line) or riluzole (green dotted line) during adolescence. Stars indicate levels of significance (*p < 0.05, **p < 0.01, ***p < 0.001) treated animals compared to the vehicle-treated group tested with analysis of variance 2 ways followed by Bonferroni post hoc test.
following aripiprazole treatment striatal glutamate alone shows a significant and stable reduction that was not seen at PND 50 anymore. A specific and stable reduction of any of the striatal neurometabolites analyzed could not be detected as indicator of the riluzole treatment within the striatum. Graphic representation of the data related to the absolute quantification of the neurometabolites is shown in Supplementary Figure S6b.

Brain region-specific correlation overview between SHR characteristic phenotype and neurometabolite absolute concentration.

After assessing the hyperactivity in the SHR experimental group, we wanted to test whether altered concentrations of the metabolites detected correlate to specific motor activity characteristics such as total distance moved, velocity and spontaneous climbing frequency (Supplementary Figure S3). We used these three parameters that were shown to be significantly different between the two strains (Supplementary Figure S3 and 4).

In Figures 4a and b, we provide an overview of the correlations between hyperactivity parameters and mean concentrations of each metabolite detected within the specific VOI over time in both strains under vehicle treatment conditions. In brief, in all of the three behavioral parameters a negative correlation of striatal aspartate levels was found to be significant ($P < 0.005$). Striatal total creatine levels also show a significant negative correlation with the mean total distance moved and velocity, but not with the climbing behavior. This behavior, however, was correlated positively with prefrontocortical total creatine levels ($P < 0.05$). Prefrontal cortex levels of glutamine positively correlate with hyperlocomotion, while its levels in the striatum show a significant positive correlation with spontaneous climbing frequency ($P < 0.05$). We also found a negative correlation of prefrontal GSH with the mean distance moved ($P < 0.005$) and mean velocity ($P < 0.05$). Finally, the mean climbing frequency, in our study a peculiarity of juvenile SHR rats’ repetitive behavior repertoire shows a unique positive correlation with total creatine in the prefrontal cortex ($P < 0.05$).

DISCUSSION

The comorbidity of TS with ADHD and/or OSD is—at least in part—caused by genetic predispositions. Within their own complex well-described clinical phenomenology, they share especially ‘hyperkinetic’ features that results in tics, motor disinhibition/hyperactivity and compulsive/ritualistic movements suggesting possible malfunctions within overlapping neuronal circuits. A review of transcranial magnetic stimulation (TMS) studies revealed a general cortical disinhibition but a pathophysiological mechanism that explains their co-occurrence is not yet defined. As hypothesized by Mol Debes all entities may be considered as ‘disorders of disinhibition’ to be explained by fronto-striatal dysfunctions that go along with a co-shared genetically susceptibility as well as neurotransmitter release alterations. Due to the unresolved pathophysiology and overlapping clinical phenomenology, the pharmacological interventions are still limited to symptomatic treatments. In light of recent clinical trials (for review see Ghanizadeh et al.), we were motivated to study the in vivo effect of aripiprazole and riluzole on behavior especially on hyperkinetic movements by subchronic daily application during adolescence in an animal model. We employed magnetic resonance spectroscopy ($^{1}$H-MRS) to analyze the concentration of brain region-specific neurometabolites of individual rats over time to also foster clinical translatability of experimental paradigms using animal models and patient cohorts. $^{1}$H-MRS ability of non-invasively quantifying the distribution of molecules of interest in the brain has been validated by ex vivo tissue samples in both, animals and humans. However, depending on the specific metabolite, in some cases concentrations are difficult to compare due to the rapid degradation processes starting immediately after cell death.

We used the SHR and its normotensive control (WKY) since the SHR strain meets many of the criteria defining it as an adequate animal model of ADHD as it presents the behavioral characteristics of inattention, impulsivity and hyperactivity. Our results show that neither aripiprazole nor riluzole increase significantly the body weight of SHR and WKY rats during adolescence, maintaining the strain-specific, slightly higher weight for the SHR compared to controls. Increased body weight is a common side
effect often reported in studies investigating antipsychotic chronic treatments in patients, however, animals were only treated for about 2 weeks so that longer treatment periods are needed to finally proof that the compounds are not affecting body and/or brain weight.

The spontaneous locomotor activity measurements under control conditions readily confirmed the known strain-related locomotor patterns showing SHRs travelling a threefold increased distance at a fivefold increased velocity when compared to the WKY strain. Treatment with both drugs did not affect the juvenile spontaneous pattern of locomotion showed by control (WKY) animals. Interestingly, also the SHR strain did not show any changes in locomotion, however grooming and climbing was significantly reduced while rearing was not. These results indicate that both compounds exert a selective action on the repetitive behavioral traits but not on the spontaneous hyperlocomotion that are described as typical features of this ADHD model. The repetitive climbing behavior, that was observed to be specific for the SHR strain only, was also significantly affected.

Our attempt to quantify and correlate our $^1$H-MRS brain-specific metabolite correlations to distance moved, sorted by brain region. WKY and SHR strains are indicated by circles and triangles respectively and colored in grey scale according to age. All mean values shown refer to the vehicle-treated groups.

| Metabolite | PFC | STR | PFC | STR | PFC | STR |
|------------|-----|-----|-----|-----|-----|-----|
| Ala        | ns  | ns  | ns  | ns  | ns  | ns  |
| Asp        | **  | **  | ns  | **  | **  | **  |
| tot. Cho   | ns  | n.s.| ns  | ns  | ns  | ns  |
| tot. Cr    | ns  | *   | ns  | *   | ns  | *   |
| GABA       | ns  | ns  | ns  | ns  | ns  | ns  |
| Glu        | ns  | *   | *   | ns  | *   | ns  |
| Gln        | *   | ns  | ns  | ns  | ns  | *   |
| GSH        | **  | ns  | *   | ns  | ns  | ns  |
| Ins        | ns  | ns  | ns  | ns  | ns  | ns  |
| tot. NAA   | ns  | ns  | ns  | ns  | ns  | ns  |
| Tau        | ns  | ns  | ns  | ns  | ns  | *   |
| PE         | ns  | ns  | ns  | ns  | ns  | ns  |

**Figure 4.** Correlation overview. (a) List of the $^1$H-MRS detected 12 metabolites and their correlation values to mean distance moved, mean velocity and mean climbing frequency, sorted by brain region (PFC: prefrontal cortex; STR: striatum). Red and blue colors indicate positive and negative correlations respectively. Symbols legend: *$P < 0.05$; **$P < 0.005$; NS, $P > 0.05$. (b) Representative graphs of the most significant metabolite correlations to distance moved, sorted by brain region. (c) Representative graph metabolites that do not significantly correlate to mean distance moved, sorted by brain region. WKY and SHR strains are indicated by circles and triangles respectively and colored in grey scale according to age. All mean values shown refer to the vehicle-treated groups.
presence/absence of anesthesia, age or diet regime of the participants, acquisition period, readout of results (as ratio or absolute concentration).

On the neurochemical level our study revealed strain-related and brain region-specific differences. Within the striatum of vehicle-treated SHR rats, glutamine and aspartate levels were significantly higher at the end of the adolescence period compared to control animals; also glutamate concentrations are slightly higher. This finding is well in line with ADHD medication-free children that were reported to have elevated striatal glutamate levels. The prefrontal cortex of SHRs we found glutamate, glutamine, total NAA, taurine and alanine to be significantly higher when compared to the control strain. In children with ADHD, an MRS study also revealed increased glutamate/glutamine in both frontal cortical areas and increased N-acetyl aspartate but also choline in the right frontal area of the ADHD-H subjects. N-acetylaspartate/creatinine (NAA/Creatine) ratio in the right frontal region, and myoinositol/creatinine (Myo inositol/Creatine) ratio in the right and left frontal regions also appeared to be highly associated with the regulation of sensorimotor, language, and memory and learning functioning in children with ADHD. Carrey et al. also showed higher striatal glutamate, glutamate/glutamine (Glx) but also creatine (Cr) concentrations in children with ADHD compared to controls. In general, our results are very much in line with the possible increase of Glx (combination of glutamate, glutamate and GABA peaks) within the striatum and cortex in pediatric patients with ADHD and other neurodevelopmental disorders suggesting changes in fronto-striatal glutamatergic circuits across the lifespan. Finally, 1H-MRS investigations analyzing bilateral dorsolateral prefrontal cortex and white matter in children affected by ADHD showed a correlation between several metabolite concentrations and symptoms. This, together with the higher concentration of excitatory neurotransmitters found within the frontal brain area could represent one of the reasons explaining the hyperactive nature of the SHR animal model investigated.

The neurochemical scenario that has been measured in treated animals showed a strain-specific effect on the SHR rats that was in general stronger in the prefrontal cortex compared to the striatum. Aripiprazole treatment induced a strong reduction of prefrontocortical glutamine, glutamate and GABA as well as a stable reduction of total NAA, GSH, inositoles and creatine. Values for PE, taurine and total choline were also lower and got more stable over time. In contrast to these results, a spectroscopic study in rats performed by McLoughlin et al. demonstrated that antipsychotic drugs (including aripiprazole) consistently increased N-acetyl aspartate (NAA) levels in at least one brain area, suggesting a common therapeutic response. Children with ADHD, however, treated with stimulants and healthy children were found to have higher choline ratios in the left prefrontal region and lower N-acetyl-aspartate ratios in the left striatum region as well as lower glutamate–glutamine ratios in the left cerebellum when compared with drug naive children with ADHD. In these three regions, there was no difference between treated children with ADHD and typically developed children. In line with our result, decreased NAA in the dorsolateral prefrontal cortex was found by Husarova et al. after atomoxetine medication supporting the hypothesis that atomoxetine could decrease hyperactivity of dorsolateral prefrontal cortex neurons. Our results should help to understand the mechanism by which aripiprazole exerts its function as a dopaminergic stabilizer being also able to influence the glutamine-glutamate-GABA cycle. This effect has also been shown in a recently published paper showing that aripiprazole treatment normalized patient values of striatal glutamate to control levels.

In good correlation with the data shown by the in vivo MRS study on adults female Sprague Dawley rats, in our study riluzole heavily reduced glutamate in the PFC and to some extent also glutamine, but not GABA. Together with its function as glutamatergic modulator, riluzole was found to readily reduce also total creatine and inositoles level indicating its role in cell plasticity. Given the higher levels of choline, total NAA and creatine were found within the right prefrontal white matter of OCD patients, correlating with the severity of symptoms, our results support the hypothesis that riluzole is an effective drug for the treatment of OCD symptoms.

The main result in vehicle-treated condition is the significant negative correlation of striatal aspartate with distance moved, velocity and climbing behavior, as well as a positive correlation between prefrontal glutamine levels and distance moved and velocity. In general, correlations seems to be heavily driven by strain differences, but results do not change when we observe the strains separately even with the given small sample size. In particular, GSH prefrontal levels negatively correlated with distance moved and velocity, reminding us of the results of Brennan’s study in which lower posterior cingulate cortex GSH/Cr ratio were found in OCD participants compared with non-OCD participants.

In conclusion we were able to provide an in vivo MRS protocol showing specific temporal and spatial changes of metabolites during brain maturation and under different drug treatments in a rat model for ADHD indicating a ‘hyperactive’ state of cortico-striatal-thalamo-cortical circuitries. The non-invasive use of preclinical 1H-MRS technique might serve as reference for future studies and encourage the 1H-MRS-based neurochemical analysis of human brain regions to further define neuropathology and control compound testing in neuropsychiatric diseases.

CONFLICT OF INTEREST

During the last 5 years Professor JMF received research funding from the EU, DFG (German Research Foundation), BMFT (Federal Ministry of Health), BMBF (Federal Ministry of Education and Research), BMFSFJ (Federal Ministry of Family, Senior Citizens, Women and Youth), German armed forces, several state ministries of social affairs, State Foundation Baden-Württemberg, Volkswagen Foundation, European Academy, Pontifical Gregorian University, RAZ, CJD, Cantias, Diocese of Rotterdam-Stuttgart. Moreover, he received travel grants, honoraria and sponsoring for conferences and medical educational purposes from DFG, AACAP, NIMH/NH, EU, Pro Helvetia, Shire, several universities, professional associations, political foundations, and German federal and state ministries. Clinical trials for Lundbeck, BMBF, Servier. Professor Fegert holds no stocks of pharmaceutical companies. The remaining authors declare no conflict of interest.

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