Trait determinants of impulsive behavior: a comprehensive analysis of 188 rats

Ana Rosa Soares^{1,2,‡}, Madalena Esteves^{1,2,‡}, Pedro Silva Moreira^{1,2}, Ana Margarida Cunha^{1,2}, Marco Rafael Guimarães^{1,2}, Miguel Murteira Carvalho^{1,2,†}, Catarina Raposo-Lima^{1,2}, Pedro Morgado^{1,2}, Ana Franky Carvalho^{1,2,3}, Bárbara Coimbra^{1,2}, António Melo^{1,2}, Ana João Rodrigues^{1,2}, António José Salgado^{1,2}, José Miguel Pêgo^{1,2}, João José Cerqueira^{1,2}, Patrício Costa^{1,2} Nuno Sousa^{1,2}, Armando Almeida^{1,2}, Hugo Leite-Almeida^{1,2}*

- 1) Life and Health Sciences Research Institute (ICVS), School of Medicine, University of Minho, Braga, Portugal
- 2) ICVS/3B's PT Government Associate Laboratory, Braga/Guimarães, Portugal
- 3) Department of General Surgery, Hospital of Braga, Braga, Portugal

‡ equal contribution

† Present address: Kavli Institute for Systems Neuroscience and Centre for Neural Computation, Norwegian University of Science and Technology, Trondheim, Norway

^{*} Corresponding author: Hugo Leite-Almeida; School of Medicine, University of Minho; Campus de Gualtar; 4710-057 Braga, Portugal; Tel: +351-253-604931; E-mail: hugoalmeida@med.uminho.pt.

Abstract

Impulsivity is a naturally occurring behavior that, when accentuated, can be found in a variety of neuropsychiatric disorders. The expression of trait impulsivity has been shown to change with a variety of factors, such as age and sex, but the existing literature does not reflect widespread consensus regarding the influence of modulating effects. We designed the present study to investigate, in a cohort of significant size (188 rats), the impact of four specific parameters, namely sex, age, strain and phase of estrous cycle, using the variable delay-to-signal (VDS) task... This cohort included (i) control animals from previous experiments; (ii) animals specifically raised for this study; and (iii) animals previously used for breeding purposes. Aging was associated with a general decrease in action impulsivity and an increase in delay tolerance. Females generally performed more impulsive actions than males but no differences were observed regarding delay intolerance. In terms of estrous cycle, no differences in impulsive behavior were observed and regarding strain, Wistar Han animals were, in general, more impulsive than Sprague-Dawley. In addition to further confirming, in a substantial study cohort, the decrease in impulsivity with age, we have demonstrated that both the strain and sex influences modulate different aspects of impulsive behavior manifestations.

1. Introduction

Impulsive behavior manifests in normal individuals and is characterized by poorly conceived and prematurely expressed actions and choices which often lead to undesirable results ^{1,2}. Attributes include impaired action restraint and/or cancelation (impulsive action), and decision-making in the absence of adequate deliberation (impulsive choice), which is characterized by a tendency to choose smaller but immediate gratification over larger but delayed ones ²⁻⁵.

While considered an adaptive behavior impulsivity has mostly been studied in the context of neuropsychiatric diseases such as addiction, compulsive eating and aggressive behavior ^{3,6}. Evidence does suggest, however, a significant relationship between trait impulsivity and the increased propensity for the development of maladaptive behaviors ⁷⁻¹³ – see for review ¹⁴⁻¹⁶.

Two factors known to affect the manifestation of trait impulsivity are age and sex. Both in human populations ¹⁷⁻²¹ – see for review ^{22,23} – and in rodent studies ²⁴⁻³¹ – see for review ³² -, impulsive behavior has generally been shown to decline with age. Nonetheless, in some studies this trend was mediated by other factors such as strain ²⁸ or sex ³¹, while others either did not observe this association between age and impulsivity ³³ or observed an inverse relationship – i.e. young adult rats presenting greater impulsivity than adolescent counterparts ³⁴. Regarding sex, its role in action ^{25,35} and choice ^{29,31,36-39} impulsivity is much less clearly defined, often revealing contradictory results.

In an attempt to further characterize the influence of biological parameters upon action and choice impulsivity, we collected and analyzed behavioral data from a large population of 188 rats, using the variable delay-to-signal (VDS) protocol (Fig. 1). The VDS is a previously validated paradigm which provides a rapid assessment of both impulsive action and delay tolerance (associated with choice impulsivity), and, as such, is particularly adequate for the characterization of large samples ⁴⁰.

2. Results

2.1. Task acquisition

The analysis of Mauchly's test revealed that the assumption of sphericity was not met ($\chi^2_{(44)}$ =3515.1, p<0.001). As a result, degrees of freedom (df) were corrected according to the Huynh-Feldt correction (ε=0.236). There was a significant decrease in the percentage of omissions across the 10 training sessions $(F_{(2.2,309.8)}=109.40, p<0.001, \eta_p^2=0.423)$. Significant between-subjects effects were observed for age $(F_{(3.149)}=31.58, p<0.001, \eta_p^2=0.389)$, sex $(F_{(1.149)}=26.13, p<0.001, q_p^2=0.389)$ $\eta_p^2 = 0.149$) and age*sex interactions ($F_{(3.149)} = 8.73$, p<0.001, $\eta_p^2 = 0.150$). In addition, statistically significant age*session ($F_{(6.6,329.0)}$ =17.33, p<0.001, η_p^2 =0.259), sex*session ($F_{(2.2,329.0)}$ =12.12, p<0.001, η_p^2 =0.075) and age*sex*session $(F_{(6.6.329.0)}=5.53, p<0.001, \eta_0^2=0.100)$ interactions were observed (Table 1 and Supplementary Table 1). The baseline number of omissions was found to increase with age. In addition, younger animals (1-2, 2-6 m.o.) effectively reached full performance (0 omissions) in 2-4 sessions while older animals (6-12 and 12-18 m.o.) took about 6 sessions to reach the same performance (Fig. 2A). Male animals displayed an increased mean number of omissions, an attribute which was particularly evident at the baseline (Fig. 2B). When age*sex*session effects were analyzed (Fig. 2C-F), it was observed that in older animals (6-12 and 12-18 m.o.), there was a significant difference between the sexes, with males presenting higher number of omissions in the first 4-5 sessions. Graphs, stratified by age and sex, depicting the full course of training can be found in Supplementary Fig. 1.

2.2. Impulsive action

Similar to the percentage of omissions across training sessions, results from the mixed-design ANOVA violated the assumption of sphericity ($\chi^2_{(44)}$ =3312.8, p<0.001) and consequently df were adjusted in accordance with a Huynh-Feldt correction (ϵ =0.245). Significant increases (within-subjects effects) were observed for PR across training sessions ($F_{(5.3,814.7)}$ =70.72, p<0.001, η_p^2 =0.313). Between-

subjects effects of age $(F_{(3.155)}=39.56, p<0.001, \eta_p^2=0.434)$ and sex $(F_{(1.155)}=33.73, p<0.001, \eta_p^2=0.434)$ p<0.001, η_p^2 =.179) were observed, but no significant interaction term between these variables (F $_{(3,155)}$ =1.59, p=0.193, η_p^2 =.434) was evident. Post-hoc analysis revealed that the mean PRs across sessions were significantly higher for the younger animals (1-2 and 2-6 m.o.) in comparison to older animals (6-12 and 12-18 m.o.). Significantly more PRs were seen with females (M=35.49%, SE=1.24) than males (M=25.26%, SE=1.24). Analysis of within-between effects were significant for (i) age*session ($F_{(15.8.814.7)}$ =4.01, p<.001, η_p^2 =.072), (ii) sex*session $(F_{(5.2.814.7)}=4.92, p<0.001, \eta_p^2=0.031)$, and (iii) age*sex*session interactions $(F_{(15.8.814.7)}=2.13, p=0.006, \eta_p^2=0.040)$ (Table 1 and Supplementary Table 1). The assessment of the longitudinal trajectories for different groups indicated that (i) older animals displayed a constant, progressive increase in the number of PRs as session number increased, whereas younger animals displayed a disproportionate increase in the number of PRs during the first three sessions (Fig. 3A); (ii) both sexes displayed a similar number of PRs at baseline, but the rate of increase was greater in females until the third session (Fig. 3B); (iii) analysis of the age*sex*session interaction (Fig. 3C-F) revealed differences only in the 12-18 m.o. group, wherein males demonstrated a greater number of PRs ($F_{(4.2.96.5)}$ =5.36, P<0.001; Fig. 3F) compared with females. This effect was not found when the analysis was restricted to sessions at which behavior was stabilized (i.e. sessions 6-10; $F_{(3.28, 85.33)}$ =1.23 p=0.305), suggesting that the disparity may be attributed to learning (Fig. 2F), rather than to sex-related impulsivity differences.

In the VDS test, first second PR rate in the 3si block constitutes an additional readout of action impulsivity (Fig. 4 and Table 1). Although, as in training, PR rate tended to decrease in an age-dependent manner (Fig. 4A), results from the univariate analysis of variance failed to achieve statistical significance ($F_{(3,178)}$ = 2.09, p=0.103, η_p^2 =0.034). There were, however, statistically significant effects of sex on this parameter ($F_{(1,178)}$ =10.55, p=0.001, η_p^2 =0.056; Fig. 4B). No age*sex interaction was found in this measure ($F_{(3,178)}$ =0.54, p=0.659, η_p^2 =0.009).

2.3. Delay intolerance

The PR rate decreased with age in all intervals (Table 2 and Supplementary Table 2) – 3si $F_{(3,176)}$ =8.30, p<0.001 η_p^2 =0.124; 6s $F_{(3,176)}$ =13.14, p<0.001, η_p^2 =0.183; 12s $F_{(3,176)}$ =9.60, p<0.001, η_p^2 =0.141; 3sf $F_{(3,176)}$ =18.92, p<0.001, η_p^2 =0.244 (Fig. 5A) – and the 3sf ratio to baseline 3si ($F_{(3,179)}$ =7.61, p<0.001; Fig. 5C). Animals in the 12-18 m.o. group had the lowest PR rate in all intervals (Fig. 5A), and also demonstrated the greatest decrease in PR rate in the 3sf interval in comparison to baseline (log(3sf/3si); Fig. 5C). In contrast, the highest PR rate was generally seen in 2-6 m.o. animals(Fig. 5A), who also exhibited an increase in PR rate at 3sf (Fig. 5C).

Differences in performance related to sex (Table 2) were observed in the initial block, 3si ($F_{(1,176)}$ =10.85, p=0.001, η_p^2 =0.058; Fig. 5B) reflecting the same pattern observed in the training sessions, with females demonstrating higher PR rates. No other statistically significant sex differences or age vs sex interactions were observed in the intervals (Fig. 5B) or in the 3sf/3si comparison (Fig. 5D). See also Supplementary Fig. 2 for cumulative PRs throughout all intervals grouped by both age and sex.

2.4. Response latency and latency to feed

Latency to perform a (correct) response (Table 3) was affected by age in all intervals except 6s (3si $F_{(3,180)}$ =15.131 p<0.001; 6s $F_{(3,180)}$ =2.589 p=0.054; 12s $F_{(3,180)}$ =16.754 p<0.001; 3sf $F_{(3,180)}$ =4.441 p=0.005), mostly reflecting higher latencies in older (6-12 and 12-18 m.o.) in comparison with younger (1-2 and 2-6 m.o.) animals (Fig. 6A left). Sex effects were only found in the 3si interval ($F_{(1,180)}$ =9.566; p=0.002), where males showed higher latencies than females (Fig. 6A right). Age*sex interaction effects were found in all intervals, except 12s (3si $F_{(3,180)}$ =4.893 p=0.003; 6s $F_{(3,180)}$ =3.339 p=0.021; 12s $F_{(3,180)}$ =1.517 p=0.212; 3sf $F_{(3,180)}$ =3.857 p=0.010).

Latency to feed (Table 4) was affected by age ($F_{(3,179)}$ =7.25, p<0.001, η_p^2 =0.108; Fig. 6B left), with older animals (12-18 m.o.) taking on average 244.6 and 143.9 ms more to recover the reward than 2-6 m.o. and 1-2 m.o. animals, respectively.

Similarly, there was a significant association between sex and latency to feed $(F_{(1,179)}=4.00, p=0.047, \eta_p^2=0.022; Fig. 6B right)$, whereas no age*sex interaction significant effects $(F_{(3,179)}=0.569, p=0.636, \eta_p^2=0.009)$ were observed.

2.5. Sub-sampling analyses

In order to explore the effect of strain and phase of estrous cycle upon impulsive behavior, additional analyses were performed. These were restricted to specific ages in order to obtain a homogenous group where variables were equally represented.

2.5.1. Strain

Analysis of the influence of strain, as well as strain*sex interaction on impulsive behavior was restricted to 12-18 m.o. males and females (SD: N=18 (8M, 10F); WH: N=17 (12M, 5F); Tables 4 and 5). Results from the mixed-design ANOVA (Huynh-Feldt corrected df: ϵ =0.360), revealed no significant effect associated with either strain (F(1,20)<0.01, p=0.984, η_p^2 =0.148) or strain*session (F(3.2,64.9)=0.197, p=0.910, η_p^2 <0.001 upon the percentage of omissions (Fig. 7A). Regarding PR during training (Huynh-Feldt corrected df: ϵ =0.820), no significant effect of strain was demonstrated (F(1,21)=3.63, p=0.071, η_p^2 =0.147), and no statistically significant strain*session interaction effects were observed (F(7.4,154.9)=5.60, p<0.001, η_p^2 <0.001; Fig. 7B). The 1st second PR rate (F(1,33)=6.22, p=0.018, η_p^2 =0.172; Fig. 7C) did reveal that WH rats have higher rate of premature responses than SD animals. No significant effects of sex, or sex*strain interaction were seen in this group (see Table 5 section for detailed statistics).

Examination of delay tolerance parameters revealed that WH animals present higher PR rate than SD counterparts in all intervals except 3si (Fig. 7D, Table 6) – 3si $F_{(1,33)}=3.52$, P=0.071, $\eta_p^2=0.105$; 6s $F_{(1,33)}=10.00$, p=0.004, $\eta_p^2=0.250$; 12s $F_{(1,33)}=8.45$, p=0.007, $\eta_p^2=0.220$; and 3sf $F_{(1,33)}=4.55$, p=0.042, $\eta_p^2=0.231$. The log(3sf/3si) analysis failed to demonstrate any differences between strains, or influence by either sex or sex*strain interaction.

2.5.2. Estrous cycle

Females in the 2-6 m.o. age group were studied to assess the potential influence of phase of the estrous cycle (diestrus/proestrus) and impulsive behavior. Estrous cycle assessment was performed immediately after the VDS test (Diestrus: n=9; Proestrus: n=6) and only this phase was analyzed (Supplementary Fig. 3 and Supplementary Table 3). No differences were observed in the propensity for impulsive action as evaluated using the 1st second of the 3si interval ($F_{(1,14)}$ =0.75, p=0.402, η_p^2 =0.055; Supplementary Fig. 3A). Estrous cycle phase had no influence upon behavior in any of the VDS blocks (Supplementary Fig. 3B) – 3si $F_{(1,12)}$ =0.93, p=.355, η_p^2 =.072; 6s $F_{(1,13)}$ =1.92, p=0.189 η_p^2 =.129; 12s $F_{(1,12)}$ =3.50, p=.086, η_p^2 =.226; and 3sf $F_{(1,12)}$ =1.14, p=0.307, η_p^2 =0.087 - nor did it effect the variation to baseline PR rate ($F_{(1,14)}$ <0.01, F=0.997; Supplementary Fig. 3C). It is however noteworthy that, in all above-mentioned analyses, females in the proestrous phase showed a trend towards higher levels of impulsivity.

3. Discussion

In the present study we characterized action impulsivity and delay-tolerance in a group of 188 rats using the VDS test paradigm. The entire protocol lasts for 8 days, but the VDS evaluation requires only a single session making it ideal for the study of short-lived phenomena or transient life stages such as adolescence or phase of estrous cycle. An important attribute of the task is that it simultaneously captures both action impulsivity and delay tolerance. Task acquisition, reflected in the progressive decline of omission trials until complete fading, was achieved within 2-6 sessions. All animals were able to learn the task and from training sessions 7 to 10 virtually no omissions were recorded. Older animals typically required more sessions, an observation previously reported in other operant behavior protocols 41-44

Premature responses during the training phase reflect impulsive action. The behavioral construct is similar to that of the 5-choice serial reaction time task (5-csrtt) ^{45,46}. This type of impulsive behavior is also captured in the early moments of the 3si delays ⁴⁰. In both instances, impulsive action decreased with age. While a previous study in a 2-csrtt has shown that 1 m.o. rats were more impulsive than >3 m.o. animals ²⁵, the opposite was observed in a single instrumental nose poke task using animals of similar ages ³⁴. Interestingly, this last task bears some resemblances with the VDS test and results are probably more akin to delay intolerance rather than behavioral inhibition (see below). In a study confined to an older population, Muir and colleagues observed that 10-11 m.o. rats were more impulsive than 23-24 m.o. rats, though this observation was restricted to a specific condition (longer delays) ²⁴.

In the present study, sex was associated with distinct action impulsivity behavior, with females performing more premature responses than males. Similar findings have been reported in 3 m.o. rats using the 2-csrtt task, although the behavioral differences were manifested specifically in delays of longer duration ²⁵. The

available literature indeed suggests that sex-related differences are specific of delay conditions – see for instance ^{25,35,47}.

The VDS test also includes a delay (in)tolerance component which correlates with delay discounting (DD) behavior and manifests as an increment of impulsive response rate upon exposure to large delays to signal/reward ⁴⁰ – see also ⁴⁸⁻⁵⁰. In our population, 2-6 m.o. animals demonstrated increased PR rate in the 3sf block compared to baseline rate in 3si, while the remaining groups maintained (1-2 and 6-12 m.o.) or even slightly decreased (12-18 m.o.) their response rate. Our observations are consistent with DD protocols, which have demonstrated that 1 m.o. rats were less impulsive than 2 m.o. animals ^{34,51} and that 25 m.o. rats were less impulsive than 6 m.o. ²⁷. Lukkes and colleagues also have observed that early adolescent female (but not male) rats were less impulsive in a DD task than were young adult/adult females 31. It is difficult to construct a meaningful framework for the DD studies reported in the literature. On one hand, no age-related differences were demonstrated in one spatial adjusting-delay task (5, 9 and >27 m.o.) ³³, while in another study, 1 m.o. rats made more impulsive choices than 2 m.o. ²⁹. As a further confounder, 1 m.o. animals in a spatial (T-maze) DD task were shown to be more impulsive than 3 m.o. but only under very specific conditions (10 and 15 second delay) 30. One major difference between DD and VDS paradigms is that, in the former, delay and reward-size effects cannot be isolated. When the amount of reward is controlled, and the indifference point calculated over an option between an adjusted delay and a variable (random) delay, adolescent animals are found to be less impulsive than young adult animals ⁵¹, as also observed in the VDS – see also ³⁴. In this context, the relevant conclusion appears to be that reward-driven behavior in adolescents is more directed by an exogenous stimulus, while it is more goal-directed in adults ⁵² – see also ⁵³. Indeed, adolescent and adult rats differ in their reward-evoked activities of the dorsal striatum and orbitofrontal cortex ^{54,55} – see also for review ⁵⁶. Incongruence in the published results may also be partially attributed to differences in procedure, i.e. adjusting vs increasing delay ⁵⁷, ascending vs descending delay ⁵⁸ and magnitude of reinforcement ⁵⁹.

Female and male rats had similar rates of premature response in all blocks of the VDS paradigm, with exception of 3si. During that period females were seen to have a higher PR rate than males, potentially reflecting differences in action impulsivity previously observed in the training phase. Consistent with our overall findings, a number of earlier DD studies also found no significant sex-associated differences ^{39,60}. Other studies have reported small differences between male and female subjects under very particular experimental conditions ³⁷, while yet others have demonstrated an effect (females>males ³⁶), or even an opposite relationship (males>females ³⁸)³⁶. In females, another variable which might contribute to the behavior under evaluation is phase of the estrous cycle. Our results show a trend towards increased tolerance to delay during diestrus compared to proestrus. Several studies in humans also have demonstrated diminished impulsivity during the mid phase of the menstrual cycle ^{61,62}. One interpretation of these differences is that they are related to the modulating effect by sex steroids upon dopaminergic tone ⁶³⁻⁶⁵, which is known to affect impulsive behavior ^{4,5,66}.

An additional potential source of inter-study variability is the animals' strain. It has for example been shown that male Lewis rats discount faster than Fisher 344 ⁶⁷⁻⁷², although differences can be attenuated or eliminated with repeated assessment, perhaps as an effect of learning and/or aging ⁷³. Two other studies using additional rat strains ^{74,75}, support inter-strain differences. In our study, we restricted the analysis to the 12-18 m.o. group in order to evaluate groups comparable to one another. WH animals were found to have increased PR in comparison with SD rats, both during training and test, demonstrating strain differences in both action and delay intolerance components of impulsivity.

Finally, considering the heterogeneity of our population, differences in motivation and/or motor performance have necessarily to be accounted. Indeed, sex and age were associated with statistically significant differences in latency to feed, although these were in general small – maximum 244.6 ms (2-6 vs 12-18 m.o). Interestingly, younger (1-2 and 2-6 m.o.) and older (2-12 and 12-18 m.o.) groups differed substantially regarding response latencies (max. 2717.4 ms; 2-6 vs 12-18 m.o.).

This is to some extent paradoxical in the sense that in the former animals need to move in a more elaborate manner (i.e. between the nosepoking and feeding orifices), while the latter requires minimal movement (the animal can perform sequential premature responses and eventually one final correct response). Such suggests that response latencies essentially reflect animals' premature responding, i.e. animals responding at higher rates also have shorter latencies. In fact, females present both statistically significant higher PR and lower response latencies specifically at the 3si stage.

In conclusion, in this study of a sample of substantial size, we confirmed decreased action impulsivity and delay tolerance with age. Of interest was our finding that, in contrast to the prevailing view mainly derived from human studies ^{17-21,76} – see also for review ^{22,23}, delay intolerance appears to be maximal at early adulthood, not in adolescence. In our analysis of action impulsivity, we found that females demonstrated a significantly greater number of premature responses than males. No similar difference was evident for delay tolerance. Our consideration of the influence of strain suggested that, in general, WH animals acted more impulsively than SD.

4. Methods

4.1. Subjects and experimental conditions

A total of 188 rats were used in this study (see below for details). Animals were kept in a room with controlled temperature (22°C ± 1°C), humidity (50-60%) and light cycle (12 hours; lights on at 8 a.m.) and were housed in groups of 2–3 in standard plastic cages with food and water available *ad libitum*. 2–3 days prior to the initiation of the VDS protocol food availability was restricted to 1h per day. Animals' weight was controlled throughout the protocol to prevent drops below 15% of baseline values. All procedures involving animals adhered to the guidelines of the European Communities Council Directive 2010/63/EU and were approved by

the institutional ethics commission - Subcomissão de Ética para as Ciências da Vida e da Saúde (SECVS).

4.2. Data collection

A database of all VDS records obtained in our institute was compiled. The data were obtained from 3 different sources: i. animals that performed the VDS task as controls for other experiments (mostly males), excluding any animal which had undergone any form of drug treatment or whose experimental records were incomplete; ii. animals that were specifically raised for this study (mostly females); and iii. aged animals which had primarily been used for breeding purposes prior to inclusion in this experiment. The final database contained a total of 188 entries. Groups were assembled by age: 1-2, 2-6, 6-12 and 12-18 months-old (m.o.), sex and strain: Sprague-Dawley (SD) and Wistar Han (WH) (Fig. 1A). Additionally, a group of young adult females were further classified according to the phase of their estrous cycle (see below).

4.3. Variable delay-to-signal (VDS) task

The VDS protocol was administered as previously described ⁴⁰. Briefly reiterated, animals were tested 5-hole operant boxes (OB; 25x25 cm; TSE Systems, Germany) within a ventilated, sound attenuating environment. One of the OB walls contains five square apertures (#1-#5; 2.5 cm), elevated 2 cm from the grid floor. The opposing wall contains a similar aperture (#6) connected to a pellet dispenser. Each aperture contains a 3W lamp bulb and an infrared beam which detects the activity of the animals.

4.3.1. Habituation

Animals were habituated to the OB in 2 daily sessions (a.m./p.m.; 4 hours apart) for 2 consecutive days (Fig. 1B). In the first 2 sessions (habituation day 1) animals were left to explore the OB for 15 minute. For this phase, all lights were off and 10-15 sugar pellets (45 mg, Bioserv Inc., New Jersey, USA) were available at aperture #6 while apertures #1 to #5 were blocked with metallic caps. In sessions 3 and 4

(habituation day 2), animals were left to explore the OB for 30 minutes. Apertures #3 and #6 both were accessible and contained 3-5 and 10-15 pellets, respectively. During these sessions house light and aperture lights #3 and #6 were on.

4.3.2. Training

Training consisted of 2 daily sessions (a.m./p.m.; 4 hours apart) of 100 trials (or 30 minutes) each, for 5 consecutive days (Fig. 1B). These sessions were initiated by delivery of 1 pellet, after which the animals were trained to wait for 3 seconds (delay period), after which aperture #3's light was turned on (response period) up to a maximum of 60 seconds. Nose pokes in the response period (correct responses) were rewarded with the delivery of a sugared pellet at aperture #6. Responses in the delay period (premature responses, PR) and omissions (absence of response) were punished with a timeout period (3s) in complete darkness and no reward was delivered (Fig. 1C top). The house light was always on with exception of the timeout periods. In the training phase action impulsivity is measured in a manner akin to that of the 5-choice serial reaction time task ^{45,46}, i.e. by assessing the percentage of PRs.

4.3.3. Test

The VDS test consists of a single session of 120 trials (Fig.1B). The test is similar to the training phase, except that nose pokes are allowed (i.e., not punished) (Fig. 1C bottom) and the delay periods are variable. The test starts with an initial block of 25 trials at 3 second delay (3si), followed by 70 trials of randomly distributed 6 or 12 second delays (6s and 12s) and concludes with a final block of 25 trials at 3 second delays (3sf). Two aspects of impulsive behavior can be evaluated by the VDS test. Action impulsivity is captured in the 1st second of the delay and prematurity rate (PR rate) during the delays measures delay tolerance. The change in PR rate in the 3sf epoch after exposure to the longer intervals (6s and 12s) correlates with delay-discounting ⁴⁰.

PR rate is defined as the amount of PR per minute of total delay, PR/min

$$PR/min = \frac{PR_i}{N_i \times T_i} \times 60$$

where PR_i is the number of premature responses, N_i is the number of trials and T_i is the delay time for i = 3si, 6s, 12s or 3sf.

4.4. Estrous cycle assessment

To determine the stage of the estrous cycle, the vaginal cytology method was used. The vaginal smear was performed after the VDS test. Cells from the smear were transferred to a dry glass slide and were air dried and stained with the Papanicolaou staining technique. Classification by stage (proestrus, estrus, metestrus and diestrus) was based on the presence or absence of nucleated epithelial cells, cornified epithelial cells and leukocytes, according to ⁷⁷; see also ⁷⁸.

4.5. Statistical analysis

Statistical analyses were done using IBM SPSS Statistics 22 (IBM software, Inc., New York, USA). The analysis of training was performed by mixed-design ANOVA with session as the within-subjects', and sex and age as between-subjects' effects. The sphericity assumption was statistically assessed with Mauchly's test. Comparisons between groups having one level were done applying one-way ANOVA and between groups with more than one level were done using two-way ANOVA. Bonferroni post-hoc correction was performed for multiple comparisons. Findings were considered significant at p<0.05. All results are presented as mean ± SEM, unless otherwise clearly stated. Strain and estrous cycle comparisons were performed using sub-samples of the total database, as described in the results section.

4.6. Data availability

The datasets generated during and/or analyzed during the current study are available from the corresponding author upon request.

Acknowledgements

The authors would like to acknowledge Dr. Edward Ganz for carefully reading and commenting the manuscript. This work was supported by FEDER funds, through the Competitiveness Factors Operational Programme (COMPETE) and the Northern Portugal Regional Operational Programme (NORTE 2020), under the Portugal 2020 Partnership Agreement as well as national funds, through the Foundation for Science and Technology (FCT) [projects POCI-01-0145-FEDER-007038, NORTE-01-0145-FEDER-000013, NORTE-01-0145-FEDER-000023 and PTDC/NEU-SCC/5301/2014]. Researchers were supported by FCT [grant numbers SFRH/BD/52291/2013 to ME and PD/BD/114117/2015 to MRG via Inter-University Doctoral Programme in Ageing and Chronic Disease, PhDOC; PDE/BDE/113601/2015 to PSM via PhD Program in Health Sciences (Applied) and Phd-iHES; SFRH/BD/109111/2015 to AMC; SFRH/BD/51061/2010 to MMC; SFRH/SINTD/60126/2009 to AM; SFRH/BD/98675/2013 to BC; IF/00883/2013 to AJR; IF/00111/2013 to AJS; SFRH/BPD/80118/2011 to HLA].

Author contributions

AS, ME, AMC, MRG, MMC, CRL, PM, AF, BC, AM, AJR, AS, JMP, JJC and HLA performed experiments that contributed to the database; AS and HLA designed the research; AS, ME, AMC, MRG, MMC, CRL, PM, AF, BC, AM and HLM performed the experiments; AS, ME, PSM, PC and HLA analysed the data; AS, ME, PSM, PC, NS, AA and HLA discussed the results; AS, ME and HLA wrote the initial manuscript; all authors approved the final manuscript.

Competing interests

None of the authors report a conflict of interest.

References

- Daruna, J. H. & Barnes, P. A. in *The impulsive client: Theory, research, and treatment* (eds W. G. McCown, J. L. Johnson, & M. B. Shure) 23-37 (American Psychological Association, 1993).
- Evenden, J. The pharmacology of impulsive behaviour in rats V: the effects of drugs on responding under a discrimination task using unreliable visual stimuli. Psychopharmacology (Berl) 143, 111-122, doi:https://doi.org/10.1007/s002130050926 (1999).
- Bari, A. & Robbins, T. W. Inhibition and impulsivity: Behavioral and neural basis of response control. *Prog Neurobiol* **108**, 44-79, doi:https://doi.org/10.1016/j.pneurobio.2013.06.005 (2013).
- Dalley, J. W., Everitt, B. J. & Robbins, T. W. Impulsivity, compulsivity, and top-down cognitive control. *Neuron* **69**, 680-694, doi:https://doi.org/10.1016/j.neuron.2011.01.020 (2011).
- Dalley, J. W. & Roiser, J. P. Dopamine, serotonin and impulsivity.

 Neuroscience 215, 42-58, doi:https://doi.org/10.1016/j.neuroscience.2012.03.065 (2012).

- Fawcett, T. W., McNamara, J. M. & Houston, A. I. When is it adaptive to be patient? A general framework for evaluating delayed rewards. *Behav Processes* **89**, 128-136, doi:https://doi.org/10.1016/j.beproc.2011.08.015 (2012).
- Saunders, B. *et al.* Impulsive errors on a Go-NoGo reaction time task: disinhibitory traits in relation to a family history of alcoholism. *Alcohol Clin Exp Res* **32**, 888-894, doi: https://doi.org/10.1111/j.1530-0277.2008.00648.x (2008).
- 8 Kayir, H., Semenova, S. & Markou, A. Baseline impulsive choice predicts the effects of nicotine and nicotine withdrawal on impulsivity in rats. *Prog Neuropsychopharmacol Biol Psychiatry* **48**, 6-13, doi:https://doi.org/10.1016/j.pnpbp.2013.09.007 (2014).
- 9 Paine, T. A., Dringenberg, H. C. & Olmstead, M. C. Effects of chronic cocaine on impulsivity: relation to cortical serotonin mechanisms. *Behav Brain Res* **147**, 135-147, doi:https://doi.org/10.1016/S0166-4328(03)00156-6 (2003).
- Perry, J. L., Larson, E. B., German, J. P., Madden, G. J. & Carroll, M. E. Impulsivity (delay discounting) as a predictor of acquisition of IV cocaine self-administration in female rats. *Psychopharmacology (Berl)* **178**, 193-201, doi:https://doi.org/10.1007/s00213-004-1994-4 (2005).

- Anker, J. J., Perry, J. L., Gliddon, L. A. & Carroll, M. E. Impulsivity predicts the escalation of cocaine self-administration in rats. *Pharmacol Biochem Behav* **93**, 343-348, doi:https://doi.org/10.1016/j.pbb.2009.05.013 (2009).
- Diergaarde, L. *et al.* Impulsive choice and impulsive action predict vulnerability to distinct stages of nicotine seeking in rats. *Biol Psychiatry* **63**, 301-308, doi:https://doi.org/10.1016/j.biopsych.2007.07.011 (2008).
- Mendez, I. A. *et al.* Self-administered cocaine causes long-lasting increases in impulsive choice in a delay discounting task. *Behav Neurosci* **124**, 470-477, doi:https://doi.org/10.1037/a0020458 (2010).
- de Wit, H. Impulsivity as a determinant and consequence of drug use: a review of underlying processes. *Addiction biology* **14**, 22-31, doi:https://doi.org/10.1111/j.1369-1600.2008.00129.x (2009).
- Dawe, S. & Loxton, N. J. The role of impulsivity in the development of substance use and eating disorders. *Neurosci Biobehav Rev* **28**, 343-351, doi:https://doi.org/10.1016/j.neubiorev.2004.03.007 (2004).
- Pattij, T. & De Vries, T. J. The role of impulsivity in relapse vulnerability. *Curr Opin Neurobiol* **23**, 700-705, doi:https://doi.org/10.1016/j.conb.2013.01.023 (2013).

- 17 Green, L., Fry, A. F. & Myerson, J. Discounting of delayed rewards: a life-span comparison. *Psychol Sci* **5**, 33-36, doi:https://doi.org/10.1111/j.1467-9280.1994.tb00610.x (1994).
- Green, L., Myerson, J., Lichtman, D., Rosen, S. & Fry, A. Temporal discounting in choice between delayed rewards: the role of age and income. *Psychology and aging* **11**, 79-84, doi:http://dx.doi.org/10.1037/0882-7974.11.1.79 (1996).
- 19 Green, L., Myerson, J. & Ostaszewski, P. Discounting of delayed rewards across the life span: age differences in individual discounting functions. *Behav Processes* **46**, 89-96, doi:https://doi.org/10.1016/S0376-6357(99)00021-2 (1999).
- Williams, B. R., Ponesse, J. S., Schachar, R. J., Logan, G. D. & Tannock, R. Development of inhibitory control across the life span. *Dev Psychol* **35**, 205-213, doi:http://dx.doi.org/10.1037/0012-1649.35.1.205 (1999).
- Eppinger, B., Nystrom, L. E. & Cohen, J. D. Reduced sensitivity to immediate reward during decision-making in older than younger adults. *PLoS One* **7**, e36953, doi:https://doi.org/10.1371/journal.pone.0036953 (2012).
- Mather, M. The Affective Neuroscience of Aging. *Annu Rev Psychol* **67**, 213-238, doi:https://doi.org/10.1146/annurev-psych-122414-033540 (2016).

- Drobetz, R., Maercker, A. & Forstmeier, S. Delay of gratification in old age: assessment, age-related effects, and clinical implications. *Aging Clin Exp Res* **24**, 6-14, doi:https://doi.org/10.3275/8178 (2012).
- Muir, J. L., Fischer, W. & Bjorklund, A. Decline in visual attention and spatial memory in aged rats. *Neurobiol Aging* **20**, 605-615, doi:https://doi.org/10.1016/S0197-4580(99)00098-6 (1999).
- Burton, C. L. & Fletcher, P. J. Age and sex differences in impulsive action in rats: the role of dopamine and glutamate. *Behav Brain Res* **230**, 21-33, doi:https://doi.org/10.1016/j.bbr.2012.01.046 (2012).
- Adriani, W. & Laviola, G. Elevated levels of impulsivity and reduced place conditioning with d-amphetamine: two behavioral features of adolescence in mice. *Behav Neurosci* **117**, 695-703, doi:http://dx.doi.org/10.1037/0735-7044.117.4.695 (2003).
- Simon, N. W. *et al.* Good things come to those who wait: attenuated discounting of delayed rewards in aged Fischer 344 rats. *Neurobiol Aging* **31**, 853-862, doi:https://doi.org/10.1016/j.neurobiolaging.2008.06.004 (2010).
- Pinkston, J. W. & Lamb, R. J. Delay discounting in C57BL/6J and DBA/2J mice: adolescent-limited and life-persistent patterns of impulsivity. *Behav Neurosci* **125**, 194-201, doi:https://doi.org/10.1037/a0022919 (2011).

- Doremus-Fitzwater, T. L., Barreto, M. & Spear, L. P. Age-related differences in impulsivity among adolescent and adult Sprague-Dawley rats. *Behav Neurosci* **126**, 735-741, doi:https://doi.org/10.1037/a0029697 (2012).
- 30 Sonntag, K. C. *et al.* Viral over-expression of D1 dopamine receptors in the prefrontal cortex increase high-risk behaviors in adults: comparison with adolescents. *Psychopharmacology (Berl)* **231**, 1615-1626, doi:https://doi.org/10.1007/s00213-013-3399-8 (2014).
- Lukkes, J. L., Thompson, B. S., Freund, N. & Andersen, S. L. The developmental inter-relationships between activity, novelty preferences, and delay discounting in male and female rats. *Dev Psychobiol* **58**, 231-242, doi:https://doi.org/10.1002/dev.21368 (2016).
- Hunt, P. S., Burk, J. A. & Barnet, R. C. Adolescent transitions in reflexive and non-reflexive behavior: Review of fear conditioning and impulse control in rodent models. *Neurosci Biobehav Rev* **70**, 33-45, doi:https://doi.org/10.1016/j.neubiorev.2016.06.026 (2016).
- Breton, Y. A., Seeland, K. D. & Redish, A. D. Aging impairs deliberation and behavioral flexibility in inter-temporal choice. *Frontiers in aging neuroscience* **7**, 41, doi:https://doi.org/10.3389/fnagi.2015.00041 (2015).

- 34 Sturman, D. A., Mandell, D. R. & Moghaddam, B. Adolescents exhibit behavioral differences from adults during instrumental learning and extinction. *Behav Neurosci* **124**, 16-25, doi:https://doi.org/10.1037/a0018463 (2010).
- Bayless, D. W., Darling, J. S., Stout, W. J. & Daniel, J. M. Sex differences in attentional processes in adult rats as measured by performance on the 5-choice serial reaction time task. *Behav Brain Res* **235**, 48-54, doi:https://doi.org/10.1016/j.bbr.2012.07.028 (2012).
- Van Haaren, F., Van Hest, A. & Van De Poll, N. E. Self-control in male and female rats. *J Exp Anal Behav* **49**, 201-211, doi:https://doi.org/10.1901/jeab.1988.49-201 (1988).
- Koot, S., van den Bos, R., Adriani, W. & Laviola, G. Gender differences in delay-discounting under mild food restriction. *Behav Brain Res* **200**, 134-143, doi:https://doi.org/10.1016/j.bbr.2009.01.006 (2009).
- Bayless, D. W., Darling, J. S. & Daniel, J. M. Mechanisms by which neonatal testosterone exposure mediates sex differences in impulsivity in prepubertal rats. *Horm Behav* **64**, 764-769, doi:https://doi.org/10.1016/j.yhbeh.2013.10.003 (2013).
- 39 Eubig, P. A., Noe, T. E., Floresco, S. B., Sable, J. J. & Schantz, S. L. Sex differences in response to amphetamine in adult Long-Evans rats performing a

delay-discounting task. *Pharmacol Biochem Behav* **118**, 1-9, doi:https://doi.org/10.1016/j.pbb.2013.12.021 (2014).

- Leite-Almeida, H. *et al.* Variable delay-to-signal: a fast paradigm for assessment of aspects of impulsivity in rats. *Frontiers in behavioral neuroscience* **7**, 154, doi:https://doi.org/10.3389/fnbeh.2013.00154 (2013).
- Ohta, H., Matsumoto, K. & Watanabe, H. Impairment of acquisition but not retention of a simple operant discrimination performance in aged Fischer 344 rats. *Physiol Behav* **54**, 443-448, doi:https://doi.org/10.1016/0031-9384(93)90233-6 (1993).
- Port, R. L., Murphy, H. A. & Magee, R. A. Age-related impairment in instrumental conditioning is restricted to initial acquisition. *Exp Aging Res* **22**, 73-81, doi:https://doi.org/10.1080/03610739608253998 (1996).
- Roesch, M. R., Bryden, D. W., Cerri, D. H., Haney, Z. R. & Schoenbaum, G. Willingness to wait and altered encoding of time-discounted reward in the orbitofrontal cortex with normal aging. *J Neurosci* **32**, 5525-5533, doi:https://doi.org/10.1523/JNEUROSCI.0586-12.2012 (2012).
- Roux, S., Hubert, I., Lenegre, A., Milinkevitch, D. & Porsolt, R. D. Effects of piracetam on indices of cognitive function in a delayed alternation task in young

and aged rats. *Pharmacol Biochem Behav* **49**, 683-688, doi:https://doi.org/10.1016/0091-3057(94)90088-4 (1994).

- Carli, M., Robbins, T. W., Evenden, J. L. & Everitt, B. J. Effects of lesions to ascending noradrenergic neurones on performance of a 5-choice serial reaction task in rats; implications for theories of dorsal noradrenergic bundle function based on selective attention and arousal. *Behav Brain Res* **9**, 361-380, doi:https://doi.org/10.1016/0166-4328(83)90138-9 (1983).
- Bari, A., Dalley, J. W. & Robbins, T. W. The application of the 5-choice serial reaction time task for the assessment of visual attentional processes and impulse control in rats. *Nat Protoc* **3**, 759-767, doi: https://doi.org/10.1038/nprot.2008.41 (2008).
- Jentsch, J. D. & Taylor, J. R. Sex-related differences in spatial divided attention and motor impulsivity in rats. *Behav Neurosci* **117**, 76-83, doi:http://dx.doi.org/10.1037/0735-7044.117.1.76 (2003).
- Cunha, A. M. *et al.* Pawedness Trait Test (PaTRaT)—A New Paradigm to Evaluate Paw Preference and Dexterity in Rats. *Frontiers in behavioral neuroscience* **11**, doi: https://doi.org/10.3389/fnbeh.2017.00192 (2017).

- Melo, A., Leite-Almeida, H., Ferreira, C., Sousa, N. & Pego, J. M. Exposure to Ketamine Anesthesia Affects Rat Impulsive Behavior. *Frontiers in behavioral neuroscience* **10**, 226, doi:https://doi.org/10.3389/fnbeh.2016.00226 (2016).
- Carvalho, M. M. *et al.* Effect of Levodopa on Reward and Impulsivity in a Rat Model of Parkinson's Disease. *Frontiers in behavioral neuroscience* **11**, 145, doi:https://doi.org/10.3389/fnbeh.2017.00145 (2017).
- McClure, J., Podos, J. & Richardson, H. N. Isolating the delay component of impulsive choice in adolescent rats. *Frontiers in integrative neuroscience* **8**, 3, doi:https://doi.org/10.3389/fnint.2014.00003 (2014).
- Hammerslag, L. R. & Gulley, J. M. Age and sex differences in reward behavior in adolescent and adult rats. *Dev Psychobiol* **56**, 611-621, doi:https://doi.org/10.1002/dev.21127 (2014).
- Ernst, M., Daniele, T. & Frantz, K. New perspectives on adolescent motivated behavior: attention and conditioning. *Developmental cognitive neuroscience* **1**, 377-389, doi:https://doi.org/10.1016/j.dcn.2011.07.013 (2011).
- Sturman, D. A. & Moghaddam, B. Reduced neuronal inhibition and coordination of adolescent prefrontal cortex during motivated behavior. *J Neurosci* **31**, 1471-1478, doi:https://doi.org/10.1523/JNEUROSCI.4210-10.2011 (2011).

- Sturman, D. A. & Moghaddam, B. Striatum processes reward differently in adolescents versus adults. *Proc Natl Acad Sci U S A* **109**, 1719-1724, doi:https://doi.org/10.1073/pnas.1114137109 (2012).
- Simon, N. W. & Moghaddam, B. Neural processing of reward in adolescent rodents. *Developmental cognitive neuroscience* **11**, 145-154, doi:https://doi.org/10.1016/j.dcn.2014.11.001 (2015).
- Craig, A. R., Maxfield, A. D., Stein, J. S., Renda, C. R. & Madden, G. J. Do the adjusting-delay and increasing-delay tasks measure the same construct: delay discounting? *Behav Pharmacol* **25**, 306-315, doi:https://doi.org/10.1097/FBP.000000000000055 (2014).
- Tanno, T., Maguire, D. R., Henson, C. & France, C. P. Effects of amphetamine and methylphenidate on delay discounting in rats: interactions with order of delay presentation. *Psychopharmacology (Berl)* **231**, 85-95, doi:https://doi.org/10.1007/s00213-013-3209-3 (2014).
- Orduna, V., Valencia-Torres, L., Cruz, G. & Bouzas, A. Sensitivity to delay is affected by magnitude of reinforcement in rats. *Behav Processes* **98**, 18-24, doi:https://doi.org/10.1016/j.beproc.2013.04.011 (2013).
- Smethells, J. R., Swalve, N. L., Eberly, L. E. & Carroll, M. E. Sex differences in the reduction of impulsive choice (delay discounting) for cocaine in rats with

atomoxetine and progesterone. *Psychopharmacology (Berl)* **233**, 2999-3008, doi:https://doi.org/10.1007/s00213-016-4345-3 (2016).

- Smith, C. T., Sierra, Y., Oppler, S. H. & Boettiger, C. A. Ovarian cycle effects on immediate reward selection bias in humans: a role for estradiol. *J Neurosci* **34**, 5468-5476, doi:https://doi.org/10.1523/JNEUROSCI.0014-14.2014 (2014).
- Diekhof, E. K. Be quick about it. Endogenous estradiol level, menstrual cycle phase and trait impulsiveness predict impulsive choice in the context of reward acquisition. *Horm Behav* **74**, 186-193, doi:https://doi.org/10.1016/j.yhbeh.2015.06.001 (2015).
- Sun, J., Walker, A. J., Dean, B., van den Buuse, M. & Gogos, A. Progesterone: The neglected hormone in schizophrenia? A focus on progesterone-dopamine interactions. Psychoneuroendocrinology **74**, 126-140, doi:https://doi.org/10.1016/j.psyneuen.2016.08.019 (2016).
- Zheng, P. Neuroactive steroid regulation of neurotransmitter release in the CNS: action, mechanism and possible significance. Prog Neurobiol 89, 134-152, doi:https://doi.org/10.1016/j.pneurobio.2009.07.001 (2009).

- Almey, A., Milner, T. A. & Brake, W. G. Estrogen receptors in the central nervous system and their implication for dopamine-dependent cognition in females. Horm Behav 74, 125-138, doi:https://doi.org/10.1016/j.yhbeh.2015.06.010 (2015).
- Winstanley, C. A. The utility of rat models of impulsivity in developing pharmacotherapies for impulse control disorders. *Br J Pharmacol* **164**, 1301-1321, doi:https://doi.org/10.1111/j.1476-5381.2011.01323.x (2011).
- Anderson, K. G. & Woolverton, W. L. Effects of clomipramine on self-control choice in Lewis and Fischer 344 rats. *Pharmacol Biochem Behav* **80**, 387-393, doi:https://doi.org/10.1016/j.pbb.2004.11.015 (2005).
- Madden, G. J., Smith, N. G., Brewer, A. T., Pinkston, J. W. & Johnson, P. S. Steady-state assessment of impulsive choice in Lewis and Fischer 344 rats: between-condition delay manipulations. *J Exp Anal Behav* **90**, 333-344, doi:https://doi.org/10.1901/jeab.2008.90-333 (2008).
- Anderson, K. G. & Diller, J. W. Effects of acute and repeated nicotine administration on delay discounting in Lewis and Fischer 344 rats. *Behav Pharmacol* **21**, 754-764, doi:https://doi.org/10.1097/FBP.0b013e328340a050 (2010).
- 70 Garcia-Lecumberri, C. *et al.* Strain differences in the dose-response relationship for morphine self-administration and impulsive choice between Lewis

and Fischer 344 rats. *J Psychopharmacol* **25**, 783-791, doi:https://doi.org/10.1177/0269881110367444 (2011).

- Huskinson, S. L., Krebs, C. A. & Anderson, K. G. Strain differences in delay discounting between Lewis and Fischer 344 rats at baseline and following acute and chronic administration of d-amphetamine. *Pharmacol Biochem Behav* **101**, 403-416, doi:https://doi.org/10.1016/j.pbb.2012.02.005 (2012).
- Stein, J. S., Pinkston, J. W., Brewer, A. T., Francisco, M. T. & Madden, G. J. Delay discounting in Lewis and Fischer 344 rats: steady-state and rapid-determination adjusting-amount procedures. *J Exp Anal Behav* **97**, 305-321, doi:https://doi.org/10.1901/jeab.2012.97-305 (2012).
- Aparicio, C. F., Elcoro, M. & Alonso-Alvarez, B. A long-term study of the impulsive choices of Lewis and Fischer 344 rats. *Learn Behav* **43**, 251-271, doi:https://doi.org/10.3758/s13420-015-0177-y (2015).
- Wilhelm, C. J. & Mitchell, S. H. Strain differences in delay discounting using inbred rats. *Genes Brain Behav* **8**, 426-434, doi:https://doi.org/10.1111/j.1601-183X.2009.00484.x (2009).
- Richards, J. B. *et al.* Strong genetic influences on measures of behavioral-regulation among inbred rat strains. *Genes Brain Behav* **12**, 490-502, doi:https://doi.org/10.1111/gbb.12050 (2013).

- Gleich, T. *et al.* Frontal glutamate and reward processing in adolescence and adulthood. *Brain Struct Funct* **220**, 3087-3099, doi:https://doi.org/10.1007/s00429-014-0844-3 (2015).
- 77 Caligioni, C. S. Assessing reproductive status/stages in mice. *Curr Protoc Neurosci* **Appendix 4**, Appendix 4I, doi:https://doi.org/10.1002/0471142301.nsa04is48 (2009).
- Cora, M. C., Kooistra, L. & Travlos, G. Vaginal Cytology of the Laboratory Rat and Mouse: Review and Criteria for the Staging of the Estrous Cycle Using Stained Vaginal Smears. *Toxicol Pathol* **43**, 776-793, doi:https://doi.org/10.1177/0192623315570339 (2015).

Figure legends

- Fig. 1. Experimental organization. (A) Composition of the groups assessed in this work (N=188) by age, sex and strain. (B) The VDS protocol is comprised of 3 phases: habituation, training and test. Habituation and training are carried out in 4 and 10 sessions respectively, 2 sessions a day. The test session is performed in a single session on the last day. (C) Each training session (top) is comprised of 100 trials performed in a maximum of 30 minutes. During this phase, the animal learns to nosepoke following a light signal and correct responses are rewarded with a sugar pellet. If the animal does not respond to the light (omission) or responds before the light turns on (i.e., within the 3s delay; premature response) no reward is delivered and the animal is punished with a 3s timeout in complete darkness. The test session (bottom) is comprised of 120 trials performed in a maximum of 60 minutes. It is similar to training except that premature responses are allowed (i.e., not punished) and the delays vary according to the following schedule: 25 initial trials with 3s delays (3si), 70 trials with randomized 6s or 12 s delays (6s and 12s respectively) and 25 final trials with 3s delays (3sf). F – female; M – male; WH – Wistar Han; SD – Sprague-Dawley
- **Fig. 2. VDS task acquisition.** Omissions as a function of session number by age and sex. **(A)** Older animals and **(B)** males require on average more sessions to reach a steady level of performance close-to-zero omissions. The effect of sex was particularly evident in **(E)** 6-12 and **(F)** 12-18 m.o. animals. Data is presented as mean \pm SEM and statistically significant comparisons between groups are marked with *; * P<0.05; ** P<0.01; *** P<0.001; m.o. months old; %OM percentage of omissions.
- **Fig. 3. Action impulsivity (training phase).** Percentage of premature responses as a function of session number by age and sex in the training phase. **(A)** Older animals and **(B)** males demonstrated less action impulsivity during training. Differences in behavior on the basis of sex for each age group was only evident in **(F)** 12-18 m.o. animals. Data is presented as mean ± SEM and statistically

significant comparisons between groups are marked with *; * - *P*<0.05; ** - *P*<0.01; *** - *P*<0.001; m.o. - months old; %PR - percentage of premature responses.

Fig. 4. Action impulsivity (VDS). PR rate in the 1st second of the 3si block of the test session by age and sex. **(A)** No statistically significant age-dependent statistically significant differences were found. **(B)** males were found to demonstrate reduced action impulsivity in comparison with females. Sex differences for individual age groups are not plotted because no age vs sex interaction was found. Data are presented as mean \pm SEM and statistically significant comparisons between groups are marked with *; ** - P<0.01; m.o. - months old; PR rate - rate of premature responses per minute.

Fig. 5. Delay tolerance. PR rate for each block of the test by age and sex. **(A)** The rate of premature responses decreased with age in all blocks, while **(B)** a sex difference was present only in the 3si interval. When responses in the final block (3sf) are in compared to those in the initial block (3si), the same trend regarding both age **(C)** and sex **(D)** was found, i.e. there was a general decrease with age, but no effect of sex. Sex differences for individual age groups are not plotted because no age vs sex interaction was found in the 3sf block, or in the 3sf/3si. Data are presented as mean ± SEM and statistically significant comparisons between groups are marked with *; *** - P<0.05; ** - P<0.01; *** - P<0.001; m.o. - months old; PR rate - rate of premature responses per minute.

Fig. 6. Response latency and latency to feed during the VDS test. (A)

Response latency by age and sex. Response latency was influenced by age in all intervals except 6s, with older animals taking longer to perform a correct response (left). Males showed higher response latencies than females in the 3si, but no differences were found in the remaining intervals (right). **(B)** Overall latency to feed by age and sex. Latency to feed was influenced by age, with older animals (12-18 m.o.) showing increased time to retrieve the reward than younger (1-2 and 2-6 m.o. animals; left). No sex effect was found (right). Data are presented as mean \pm SEM and statistically significant comparisons between groups are marked with *; * - P<0.05; ** - P<0.01; *** - P<0.001; m.o. - months old.

Fig. 7. Influence of strain upon choice impulsivity and delay intolerance. The analysis of the effect of the strain in impulsive behavior was confined to 12-18 m.o. males and females. **(A)** Both strains learned the task equally well, progressively reducing %OM to 0. WH animals demonstrated higher action impulsivity in both **(B)** training and **(C)** PR rate in the 1st second of 3si. **(D)** During the test, WH animals consistently showed higher PR rates in 6s, 12s and 3sf blocks. Data is presented as mean ± SEM and statistically significant comparisons between groups are marked with *; * - P<0.05; ** - P<0.01; *** - P<0.001; %OM - percentage of omitted responses; %PR - percentage of premature responses; PR rate - rate of premature responses per minute; WH - Wistar Han; SD - Sprague-Dawley.

Tables

		Test			
	%OM effects		%PR effects		PR rate 1 st sec effects
	Group Group*Session Group Group*		Group*Session	Group	
Age	F(_{3,149})=31.584;	F(_{6.2,309.8})=17.33	F _(3,155) =39.556;	F _(14.6,752.33) =4.012;	F _(3,178) =2.094;
	p<0.001	1; p<0.001	p<0.001	p<0.001	p=0.103
Sex	F(_{1,149})=26.132;	F(_{2.1,309.8})=12.11	F(_{1,155})=33.726;	F(_{4.9,752.3})=4.918;	F(_{1,178})=10.55
	p<0.001	8; p<0.001	p<0.001	p<0.001	4; p=0.001
Sex*	F(_{1,149})=8.733;	F(_{6.6,309.8})=5.526;	F _(3,155) =1.594;	F(_{14.6,752.3})=2.133;	F(_{3,178})=0.535;
Age	p<0.001	p<0.001	p=0.193	p=0.008	p=0.659

Table 1. General effects of sex and age on learning and action impulsivity.

Effects on task learning were assessed through the percentage of omissions during training (%OM) while effects on action impulsivity were evaluated in the percentage of premature responses during training (%PR) and prematurity rate during the first second of the test. Main effects of group (between factor) and session/group (within/between factors), in which groups are divided by age and sex, are shown. P<0.05 was considered the threshold for statistical significance and age is measured in months.

	Test					
	PR rate effects					
	3si 6s 12s 3sf log(3sf/3si)					
Age	F(_{3,176})=8.298;	F(_{3,176})=13.138;	F(_{3,176})=9.602;	F(_{3,176})=18.919;	F(_{3,179})=7.607;	
	p<0.001	p<0.001	p<0.001	p<0.001	p<0.001	
Sex	F(_{1,176})=10.845	F(_{1,176})=0.987;	F(_{1,176})=0.051;	F(_{1,176})=0.400;	F(_{1,179})=0.499;	
	; p=0.001	p=0.322	p=0.821	p=0.528	p=0.504	
Sex*Age	F(_{3,176})=1.478;	F(_{3,176})=2.518;	F(_{3,176})=2.481;	F(_{3,176})=0.813;	F(_{3,179})=0.574;	
	P=0.222	p=0.060	p=0.063	p=0.488	p=0.632	

Table 2. General effects of sex and age on delay intolerance. Effects were assessed based on the prematurity rate (PR rate) during the test phases (3si, 6s, 12s and 3sf) and in the 3sf normalized to baseline (log(3sf/3si)). Main effects of group (age and sex) are shown. P<0.05 was considered the threshold for statistical significance and age is measured in months.

	Test						
	Response latency effects						
	3si 6s 12s 3sf						
Sex	F(_{1,180})= 9.566;	F(_{1,180})= 1.261;	F(_{1,180})= 0.003;	F(_{1,180})= 0.798;			
	p=0.002	p=0.263	p=0.954	p=0.373			
Age	F(₃ , ₁₈₀)= 15.131;	F(_{3,180})= 2.589;	F(_{3,180)} = 16.754;	F(_{3,180})=4.441;			
	p<0.001	p=0.054	p<0.001	p=0.005			
Sex*Age	F(_{3,180})= 4.893;	F(_{3,180})= 3.339;	F(_{3,180})= 1.517;	F(_{3,180})= 3.857;			
	p=0.003	p=0.021	p=0.212	p=0.010			

Table 3. General effects of sex and age on response latency. Effects were assessed based on the response latency during the test phases (3si, 6s, 12s and 3sf). Main effects of group (age and sex) are shown. P<0.05 was considered the threshold for statistical significance and age is measured in months.

	Test	
	Group	
Age	F(_{3,179})=7.249; p<0.001	
Sex	F(_{1,179})=3.997; p=0.047	
Sex*Age	F(_{1,149})=0.569; p=0.636	

Table 4. Effects of sex and age on latency to feed. Effects were assessed based on the latency to feed during the VDS test. Main effects of group (age and sex) are shown. P<0.05 was considered the threshold for statistical significance and age is measured in months.

		Test			
	%OM effects		%PR effects		PR rate 1 st sec effects
	Group Group*Session		Group	Group*Session	Group
Strain	F(_{1,20})<0.001;	F(_{3.2,64.9})=0.197;	F(_{1,21})=3.625;	F(_{7.4,154.92})=5.596;	F(_{1,33})=6.224;
	p=0.984	p=0.910	p=0.071	p<0.001	p=0.018
Sex	F _(1,20) =5.066;	F(_{3.2,64.9})=4.295;	F(_{1,21})=11.556;	F(_{7.4,154.92})=4.760;	F(_{1,33})=3.011;
	p=0.036	p=0.007	p=0.003	p<0.001	p=0.093
Strain*	F _(1,20) =2.488;	F(_{3.2,64.9})=1.518;	F(_{1,21})=2.111;	F(_{7.4,154.92})=1.089;	F _{(1,33})=0.072;
Sex	p=0.130	p=0.216	p=0.161	p=0.371	p=0.790

Table 5. General effects of strain and sex on learning and action impulsivity.

Effects on task learning were assessed through the percentage of omissions during training (%OM) while effects on action impulsivity were evaluated using the percentage of premature responses during training (%PR) and prematurity rate on the first second of the test. Main effects of group (between factor) and session/group (within/between factors), in which groups are divided by strain and sex, are shown. P<0.05 was considered the threshold for statistical significance.

	Test					
	PR rate effects					
	3si 6s 12s 3sf log(3sf/3si)					
Strain	F(_{1,33})=3.515;	F(_{1,33})=10.002;	F(_{1,33})=8.446;	F(_{1,33})=4.552;	F(_{1,33})=0.484;	
	p=0.071	p=0.004	p=0.007	p=0.042	p=0.492	
Sex	F(_{1,33})=1.095;	F(_{1,33})=0.126;	F(_{1,33})=0.788;	F(_{1,33})=0.353;	F(_{1,33})=1.321;	
	p=0.304	p=0.725	p=0.382	p=0.557	p=0.259	
Strain*	F(_{1,33})=0.418;	F(_{1,33})=4.992;	F(_{1,33})=7.881;	F(_{1,33})=0.474;	F(_{1,33})=0.081;	
Sex	p=0.523	p=0.033	p=0.009	p=0.496	p=0.778	

Table 6. General effects of strain and sex on delay intolerance. Effects were assessed based on the prematurity rate (PR rate) during the test phases (3si, 6s, 12s and 3sf) and in the 3sf normalized to baseline (log(3sf/3si)). Main effects of group (strain and sex) are shown. P<0.05 was considered the threshold for statistical significance.















