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The Behavior of APO-SUS Rats in Animal Models with Construct Validity for Schizophrenia

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Schizophrenic patients are known to suffer from a number of information processing disturbances, including deficits in both prepulse inhibition of startle and latent inhibition. Since these behavioral phenomena can also be observed in animals, they represent an ideal starting point for developing animal models having construct validity for specific deficits observed in schizophrenia. The principal question is how to induce a condition in animals most similar to the schizophrenic deficit. In the present study, we have selected rats on the basis of their response to an open field or to the dopaminergic agonist apomorphine, and evaluated their prepulse inhibition and latent inhibition. We used three different selection procedures (open field selection for novelty response, gnawing cage selection for apomorphine response, and pharmacogenetic selection for apomorphine response).

The results show that, irrespective of the selection procedure used, rats with a high response to novelty or apomorphine susceptible (collectively called APO-SUS rats) show diminished prepulse inhibition of the acoustic startle response as compared to rats with a low response to novelty or apomorphine unsusceptible (collectively called APO-UNSUS rats). This difference was apparent only at low prepulse intensities. Moreover, these APO-SUS rats show diminished latent inhibition in a conditioned taste aversion paradigm as compared to APO-UNSUS rats. Given the fact that the pharmacogenetically bred APO-SUS rats show several central nervous, endocrinological, and immunological similarities to schizophrenic patients, they are hypothesized to represent an interesting nonpharmacological animal model for schizophrenia-prone patients.

[Key words: schizophrenia, APO-SUS rats, animal model, prepulse inhibition, latent inhibition, acoustic startle response]

Research into the neurobiological deficits involved in schizophrenia has long been hampered by a lack of adequate animal models for this severe psychiatric disorder. Thus, most animal models have not established more than predictive validity (Ellenbroek, 1993), which limits their ability to enhance our understanding of schizophrenia. The development of animal models has, however, gained momentum in the last number of years, due to the results of neuropsychological and psychophysiological research showing information processing deficits in schizophrenic patients (Braff, 1993). Thus, schizophrenic patients show, among others, deficits in prepulse inhibition of the acoustic startle response and in latent inhibition. Prepulse inhibition refers to the diminished response to an acoustic startle stimulus when it preceded by a less intense acoustic stimulus. Latent inhibition refers to the detrimental effect of prior stimulus preexposure to the subsequent condition of that specific stimulus. In comparison to control subjects, schizophrenic patients are known to have decreased prepulse inhibition (Braff and Geyer, 1990) and diminished latent inhibition (Baruch et al., 1988). The advantage of these paradigms is that they can be measured with almost identical methods in humans and rats, making them suitable for studying the neuronal substrates of information processing deficits observed in schizophrenic patients. Indeed, these paradigms represent important examples of animal models with construct validity for specific deficits observed in schizophrenia (Ellenbroek and Cools, 1990; Geyer and Markou, 1995), for instance, models in which the psychopathological construct of the disease is modeled (Willner, 1984).

Nevertheless, an important question in relation to schizophrenia is how to induce a "schizophrenia-like" condition. So far, research has focussed predominantly on the dopamine hypothesis. Thus, the indirect dopamine agonist amphetamine is known to induce deficits in prepulse inhibition (Mansbach et al., 1988) and latent inhibition (Weiner et al., 1984). In this respect, one could argue that the usefulness of these models is not much higher than that of animal models with predictive validity (Ellenbroek, 1993). An alternative strategy may be to search for differences among rat lines or strains (Glowa and Hansen, 1994). Thus, the Maudsley Low reactive rats (Commissaris et al., 1990) and the Flinders resistant rats show startle habituation deficits (Markou et al., 1993). As yet, however, no strain of rats has been found to show deficits in prepulse inhibition or latent inhibition. A number of years ago we started to breed Wistar rats selected on their stereotyped gnawing response to apomorphine (Cools et al., 1990). These rats (the apomorphine susceptible APO-SUS and unsusceptible APO-UNSUS lines) not only show a differential response to apomorphine, but also differ in a large number of behavioral, endocrinological, and immunological parameters (Cools et al., 1990, 1993, 1994). Interestingly, the APO-SUS rats show a heightened response to external stimuli, which is reminiscent of schizophrenic patients (McGhie, 1970).
This observation led us to suggest that the APO-SUS rats might also show deficits in information processing reminiscent of schizophrenia. In order to investigate this hypothesis, we used animals selected from our normal outbred Wistar colony, as well as animals from our pharmaco-genetically bred lines and tested them in two experimental paradigms: prepulse inhibition of the acoustic startle response and latent inhibition with the conditioned taste aversion paradigm.

**Materials and Methods**

The experiments were performed in accordance with the Helsinki Declaration and institutional guidelines.

**Animals and selection procedure**

All Wistar rats were obtained and bred in the Central Animal Laboratory of the Univ of Nijmegen. They weighed between 200 and 250 g at the time of the experiment and had water and food freely available except during the experiment. They were housed in temperature-controlled rooms with a standard 12 L:12 D cycle: lights on from 0700 to 1900 hr. Three different types of selection procedures were used (Cools et al., 1990):

1. **Open-field selection.** For this selection procedure, drug and experiment naive male Wistar rats were obtained from our normal outbred strain. Three days before the selection procedure they were individually housed in MacrolonR cages (40 x 25 cm). At the day of the experiment each rat was placed in the experimental room 30 min before testing to adapt to the environment. After this period the rat was placed on a large open field (160 x 160 cm) without walls. The behavior of the rat was recorded for 30 min with the help of a video-tracking system developed in our laboratory. This system allows the continuous recording of locomotor behavior and gives information on (among others) habituation time (defined as the time until the rat stops for at least 90 sec), total locomotor activity as well as locomotor patterns. Previously, we have shown how this type allows for the discrimination between high responders (HR) and low responders (LR) to novelty (Cools et al., 1990). LR rats are defined as animals that locomote more than 6000 cm and habituated in more than 840 sec. LR rats are defined as animals that locomote less than 4800 cm and habituate in less than 840 sec. In the past we have shown that the behavior in this open field closely correlates with apomorphine susceptibility (Cools et al., 1990). More specifically, LR show a high susceptibility to apomorphine, whereas LR show a low susceptibility to apomorphine.

2. **Gnawing cage selection.** For this selection procedure, male Wistar rats were obtained from our normal outbred population. The animals received subcutaneous injections of 1.5 mg/kg apomorphine HCl (Brodacels ACF) in the neck and immediately placed in the gnawing box for 45 min. This box is virtually identical to that of Ljungberg and Ungefeldt (1978). It consists out of a perspex hole board (60 x 60 cm) with a central cubicle (25 x 25 cm). The box contains 32 holes, each of which is surrounded by 10 concentric ridges. A microphone is placed underneath the central cubicle to allow registration of sounds.

Through this microphone and a large number of infrared beams, a number of behavioral activities can be automatically recorded, like locomotor activity, frequency, and duration of hole dipping, etc. For our purpose, only the stereotyped gnawing response is important. The gnawing on the ridges surrounding the holes produces a characteristic sound that is detected by the microphone, fed into the computer, and scored as one gnaw. Based on this gnawing response, we have selected two types of rats: GnAW rats, i.e., rats that gnaw more than 500 times per 45 min and NONGNAW rats, i.e., rats with gnaw less than 10 times per 45 min. Half of the animals were subjected to the gnawing test 1 week before being placed in the prepulse inhibition or the latent inhibition paradigm and the other half 1 week after the prepulse inhibition. Since there were no statistically significant differences in either prepulse inhibition scores or in gnawing scores, the results of both groups were added together.

3. **Pharmaco-genetic selection.** As discussed in the introductory paragraphs, several years ago we started a breeding program to pharmaco-genetically select apomorphine susceptible and unsusceptible rats (Cools et al., 1990). After an initial selection in the gnawing box (described above), the nine males and females with the highest score and the nine males and females with the lowest score were paired (the FO generation). Their offspring was again tested in the gnawing box and for each new generation, the four best male and female litters were selected. The best litters are defined as those litters showing the highest (APO-SUS) and the lowest (APO-UNSUS) mean gnawing response per gender. Out of these four best male and female litters nine new pairs of rats were selected to produce the next generation, with the condition that brother/sister pairing is not allowed. The litters are weaned 28 d after birth. After weaning, the fathers are returned to the mothers to allow for two more litters to be made. These litters are being used for other experiments (such as the ones described in this article). The present experiments were performed with naive male APO-SUS and APO-UNSUS rats belonging to the 17th and 18th generation. APO-SUS rats are defined as animals born from an APO-SUS mother and father. APO-UNSUS rats are likewise defined as animals born from an APO-UNSUS father and mother. The animals were individually housed 3 d before the experiments in macronol cages (40 x 25 cm) with food and water ad libitum.

**Prepulse inhibition of the acoustic startle response**

The prepulse inhibition experiments were performed in a acoustic startle chamber of San Diego Instruments. Basically, the cage consists of a Plexiglas tube (8.2 cm in diameter, 25 cm in length) resting on a plastic frame. A piezoelectric accelerometer mounted under the tube detected and transduced the motion of the tube. Stimulus delivery was done using the SR-LAB software, via a speaker mounted 10 cm above the cylinder. The computer software also digitized, rectified, and recorded the response of the accelerometer, with 100 m sec readings collected beginning at stimulus onset. Startle amplitude was defined as the average of the 100 readings. The whole system was mounted within a sound attenuating chamber. Throughout the startle session a background level of 70 dB was maintained.

The startle session started with a 5 min habituation session in the startle system. After this habituation period, five startle pulses (120 dB[A], 40 m sec broad band burst, the A refers to the A weighing scale for sound measurement) were delivered to test basal startle responsiveness. Next, five blocks of seven trials were delivered to measure prepulse inhibition. Each of these blocks consisted of two startle trials, one no-stimulus condition and one of four different prepulse-startle pairing administered pseudorandomly. In these pairings the prepulse was either 2, 4, 8, or 16 dB[A] above background. These prepulses were always 20 m sec broad band bursts and were always followed by a startle pulse (120 dB[A]) 100 m sec later. The session was terminated by five additional startle pulses to check a measure for startle habituation. The interval between two trials was between 10 and 20 sec.

The degree of prepulse inhibition (in percentage) was calculated according to the formula:

\[
\text{Prepulse inhibition} = \frac{100 - \frac{\text{startle amplitude on prepulse trial}}{\text{startle amplitude on startle trial}} \times 100}{\text{startle amplitude on startle trial}}
\]

**Latent inhibition**

Latent inhibition was measured using the conditioned taste aversion paradigm. Rats were housed individually. Food was freely available throughout the experiment. Water bottles were removed from their cages 24 hr before the start of the experiment. On the first day of the experiment rats were subdivided into two groups: a preexposed group and a nonpreexposed group. The whole experiment was performed in the home cage of the rats. The preexposed group received a bottle with 50 ml of a 5% sucrose solution, whereas the nonpreexposed group received a bottle with 50 ml of plain tap water for 30 min. After these 30 min, the bottles were weighed to determine the amount of solution drunk. This procedure was repeated at day 2 and 3. At day 4, all rats were given a bottle of 50 ml of a 5% sucrose solution, again for 30 min. Immediately after the 30 min each rat was treated with 50 mg/kg of LiCl (5 ml/kg i.p.). On the final test day (day 5) the rats received one bottle with 50 ml of tap water and one bottle with 50 ml of 5% sucrose. The degree of conditioned taste aversion (in %) is determined by the formula:

\[
\text{CIA} = \frac{\text{ml of sucrose consumed}}{\text{ml of sucrose consumed} + \text{ml of tap water consumed}} \times 100.
\]

In the series of experiments with the APO-SUS and APO-UNSUS rats, an additional experiment was performed with 1 d preexposure. In this case, all rats (preexposed and nonpreexposed) received 50 ml of tap water at days 1 and 7 for 30 min. Only at day 4 the preexposed
**PREPULSE INHIBITION**

**Open Field Selection**

![Graph](image)

Figure 1. The effects of different prepulse intensities on the acoustic startle response of animals selected on the open field. The percentage of inhibition (± SEM) is calculated as described in the Materials and Methods section. *p < 0.05 one-way ANOVA HR versus LR rats.

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Statistics

In the prepulse inhibition experiment a mixed ANOVA was used with the different prepulse intensities as within-subject factors and the groups of rats as between-subject factors. In case of statistical differences between the two groups of rats, one-way ANOVA was used to analyze the difference between the groups for each prepulse intensity. In the latent inhibition experiment, a two-way ANOVA was performed with factors preexposure and rat group. All p-values mentioned are two-tailed.

Results

**Prepulse inhibition**

Three prepulse inhibition experiments were performed with rats of each of the three selection procedures. For sake of clarity, we will discuss each of these separately.

**HR and LR rats**

The high- and low responders to novelty selection was done on 20 male Wistar rats. Six rats fulfilled the criteria for LR (mean distance 3711 ± 343 cm, mean habituation time 392 ± 87 sec) and six fulfilled the criteria for HR (mean distance 8456 ± 496 cm, mean habituation time 1273 ± 380 sec).

The prepulse inhibition response of the HR and LR rats is displayed in Figure 1. There were no differences in basal startle reactivity, as measured by the first five startle responses: HR, 789 ± 103; LR, 685 ± 78. However, there were differences in prepulse inhibition. The mixed ANOVA showed an overall significant effect as well as a significance interaction between group and prepulse condition [F(4,54) = 4.6, *p < 0.05]. One-way ANOVA showed that HR had significantly less prepulse inhibition at the lowest prepulse intensity (2 dB[A] above background) [F(1,58) = 4.71, *p < 0.05]. There were no significant differences between groups at other prepulse intensities.

**NONGNAW and GNAW rats**

As described in the Materials and Methods section, two groups of 20 rats were tested. Group 1 was tested in the gnawing box 1 week before the prepulse inhibition session; the other group first received a prepulse inhibition session followed, 1 week later, by a gnawing box experiment. Since there were no statistically significant differences between the two groups on either test, the data were pooled. Out of these 40 rats tested, 10 fulfilled the criteria for non-gnawing rats (mean gnawing score of 2.4 ± 2.7) and 13 fulfilled the criteria for gnawing rats (mean gnawing score 1022 ± 417).

The prepulse inhibition response of the NONGNAW and GNAW rats is displayed in Figure 2. Again, there were no differences in basal startle reactivity, as measured by the first five startle responses: NONGNAW rats, 471 ± 30; GNAW Rats, 544 ± 30. However, there were differences in prepulse inhibition. The mixed ANOVA showed an overall significant effect as well as a significant interaction between group and prepulse condition [F(4,109) = 2.83, *p < 0.05]. As in the case of the open-field selection, the difference observed between the groups selected in the gnawing box could be ascribed to a significant difference at the lowest prepulse intensities [one-way ANOVA, *F*(1,113) = 2.94, *p < 0.05]. There were, again, no significant differences between groups at other prepulse intensities.

**APO-UNSUS and APO-SUS rats**

For this experiment, 10 male APO-UNSUS and 9 male APO-SUS rats were used. In contrast to the open-field and the gnawing box selection, these pharmacogenetically selected animals differed in basal startle reactivity: APO-SUS mean 1444 ± 161; APO-UNSUS mean 883 ± 107. This difference was highly significant [F(1,93) = 10.16, *p < 0.002].

Moreover, these animals differed in the degree of prepulse inhibition. As can be seen from Figure 3, there was a significant prepulse condition with group interaction [F(4,89) = 3.12, *p < 0.05]. One-way ANOVA showed that this was due to differences between the groups at both the 2 dB[A] prepulse trials [F(1,93)
PREPULSE INHIBITION
Gnawing Box Selection

Figure 2. The effects of different prepulse intensities on the acoustic startle response of animals selected in the gnawing box. The percentage of inhibition (± SEM) is calculated as described in the Materials and Methods section. *p < 0.05 one-way ANOVA GNAW versus NONGNAW rats.

Latent inhibition
NONGNAW and GNAW rats. For this experiment, 50 rats were subdivided into two groups of 25. One group was first tested in the gnawing box with apomorphine, followed 1 week later by the latent inhibition paradigm. The reverse order was used for the second group. Since no significant differences were seen on either the latent inhibition or the gnawing box scores, the data were pooled.

The 50 rats were subdivided into one preexposure group and one nonpreexposure group in the latent inhibition paradigm. The results of this experiment are displayed in Figure 4. There were no differences in overall intake of water and sucrose of the test day (nonpreexposed group 11.8 ± 1.8 ml; preexposed 13.2 ± 1.0 ml). However, there were clear differences between the two groups with respect to sucrose preference. The rats receiving 3 d of water prior to the conditioning trial (the nonpreexposed group) showed a strong conditioned taste aversion, as is seen by the very low sucrose preference (20.1 ± 2.4%). On the other hand, the animals that received 3 d of preexposure of 5% sucrose solution show a much smaller degree of conditioned taste aversion as is shown by the much higher sucrose consumption. In other words, prior preexposure reduced conditioning, for instance, latent inhibition had occurred.

When the percentage of sucrose consumption on the test day was plotted as a function of the apomorphine induced gnawing score an interesting picture occurred (Fig. 4b). Although there was no relationship for the nonpreexposed group (data not...

PREPULSE INHIBITION
Pharmacogenetic Selection

Figure 3. The effects of different prepulse intensities on the acoustic startle response of animals selected from the pharmacogenetic breeding program. The percentage of inhibition (± SEM) is calculated as described in the Materials and Methods section. *p < 0.05 one-way ANOVA APO-SUS versus APO-UNSUS rats.
shown), there was a significant negative correlation between apomorphine induced gnawing and percentage of sucrose consumed (r(25) = -0.62; p < 0.01). In other words, the animals with the highest apomorphine gnawing score had the least latent inhibition. This can also be seen in Figure 4c. Subdividing the animals according to the criteria described in the Materials and Methods section leads to 10 NONGNAW rats (mean gnawing score 3.0 ± 0.8) and 8 GNAW rats (mean gnawing score 728 ± 65). These animals show a significant difference in degree of latent inhibition (i.e., drank less sucrose) both at 1 d and at 3 d preexposure. Moreover, whereas the APO-UNSUS showed a significant degree of latent inhibition already at 1 d preexposure, the APO-SUS rats did not.

Discussion

Animal models with construct validity for schizophrenia are generally based on information processing deficits (Ellenbroek and Cools, 1990). However, an important problem is that we do not know how these processes can be disturbed in a way resembling schizophrenia. In the present study, we have analyzed two information processing aspects in rats selected on the basis of their apomorphine response or their response to novelty. In the past, we have shown that these selection paradigms lead to very similar groups of rats (Cools et al., 1990). The present data also show that animals with a high susceptibility for apomorphine-induced gnawing or a high response to novelty have a strongly reduced prepulse inhibition and a strongly diminished latent inhibition. Indeed, the data are almost identical for all three groups of rats. Nevertheless, there are some differences. Thus, whereas the HR and GNAW rats show diminished prepulse inhibition only at 2 dB[A] prepulse intensity, the APO-SUS rats showed disruption at 2 and 4 dB[A]. Likewise, whereas the GNAW rats showed diminished latent inhibition with a 3 d preexposure time, the APO-SUS rats a diminished latent inhibition at both 1 and 3 d preexposure. Taking these data together, it appears that the difference between APO-SUS and APO-UNSUS rats is larger than between HR and LR rats or between GNAW and NONGNAW rats. This result is not surprising, given the fact that APO-SUS and APO-UNSUS rats have been bred for 17 generations. During the course of this breeding scheme, the mean differences in apomorphine-induced gnawing also continuously increased (Cools et al., 1990; Ellenbroek and Cools, unpublished data).

Apart from these differences in degree of prepulse and latent inhibition, there were also differences in baseline startle amplitude. Thus, APO-SUS rats had a much higher baseline startle response (1444 ± 161) than the other two rat types [ANOVA F(2,137) = 28.37, p < 0.001] and rat type [F(1,32) = 9.3, p < 0.005]. The interaction between preexposure and rat type was not significant [F(2,31) = 2.12, p > 0.15]. Post hoc Duncan tests showed that APO-SUS rats showed less latent inhibition (i.e., drank less sucrose) both at 1 d and at 3 d preexposure. Moreover, whereas the APO-UNSUS showed a significant degree of latent inhibition already at 1 d preexposure, the APO-SUS rats did not.
Thus, local infusion of dopamine (Swerdlow et al., 1990) or the dopamine D_{1} agonist quinpirole (Wan et al., 1994) into the nucleus accumbens disrupts prepulse inhibition. Moreover, early social isolation (at weaning), which is accompanied by an increase in mesolimbic dopamine, reduces prepulse inhibition (Geyer et al., 1993). From these data one would expect an increase in dopaminergic (especially D_{1}) transmission in APO-SUS rats, given their diminished sensorimotor gating. However, although we have not yet measured the dopamine content or transmission in these animals, behavioral data suggest a low functional dopamine D_{1} activity in the nucleus accumbens of APO-SUS rats (Cools et al., 1994), which would also explain their increased sensitivity to apomorphine. A possible explanation for this apparent conflict is provided by a recent study of Schwarzkopf and his colleagues (1992). They showed that rats, neonatally treated with 6-OHDA, showed disrupted prepulse inhibition, despite a virtually complete loss (> 90%) of dopamine in the dorsal striatum. These authors suggested that these animals are characterized by a low “tonic” level of dopamine, but an enhanced responsiveness of the dopaminergic system. Interestingly, APO-SUS rats are also characterised by an increased responsiveness of both the hypothalamus–pituitary–adrenal axis and the dopaminergic system (Cools et al., 1994; Rots et al., 1995). Moreover, neonatally 6-OHDA lesioned rats (Schwarzkopf et al., 1992) and APO-SUS rats (Cools et al., 1994) have increased D_{1} dopamine receptor binding in the dorsal striatum. In other words, the reduced prepulse inhibition of these animals may have been due to an enhanced “phasic” release of dopamine in response to startle testing.

Latent inhibition

Latent inhibition refers to the detrimental effect of prior stimulus preexposure upon the subsequent conditioning of that stimulus. Schizophrenic patients, in an acute stage of their illness, have a reduced latent inhibition (Baruch et al., 1989). Gray et al. (1992) have proposed that the reduced latent inhibition of schizophrenic patients is due to their decreased ability to use previously stored information. As with prepulse inhibition, we found that GNAW rats and APO-SUS rats have reduced latent inhibition. Given the fact that HR and APO-SUS rats behaved virtually identical to each other (cf Cools et al., 1990), we did not test HR rats. Moreover, we could show that there was an inverse relationship between degree of latent inhibition and apomorphine gnawing response (Fig. 4b).

There is ample evidence for a role of dopamine in the mediation of latent inhibition. Thus, amphetamine reduces latent inhibition when given in both the preexposure and the conditioning phase (Weiner et al., 1984; Gray et al., 1992; Killcross and Robbins, 1993). In spite of the clear evidence for a role of dopamine in latent inhibition, the neuronal mechanisms underlying this phenomenon are still very unclear. Solomon and Stanton (1982) have presented evidence that local application of dopamine into the nucleus accumbens but not the dorsal striatum disrupts latent inhibition. However, this could not be replicated by Killcross and Robbins (1993). Moreover, early social isolation, which leads to an increase in dopamine transmission within the ventral striatum, does not affect latent inhibition (Wilkinson et al., 1994). On the other hand, early maternal separation, which affects the dopaminergic transmission within the dorsal striatum and which enhances the sensitivity for apomorphine, also reduces latent inhibition (Ellenbroek and Cools, 1995).

APO-SUS rats as a model for schizophrenia-prone patients

Overall, the results of the present study suggest that APO-SUS rats resemble schizophrenic patients in at least two paradigms: latent inhibition, i.e., a cognitive deficit characteristic of acute schizophrenic, and prepulse inhibition, i.e., a psychophysiological deficit that seems to occur both in treated and nontreated schizophrenic. These data suggest that the APO-SUS rats may represent an interesting animal model for at least certain information processing aspects of schizophrenia. However, disturbances in prepulse inhibition does not only occur in schizophrenic patients, but also in schizotypal patients (Cadenhead et al., 1993) and in patients suffering from obsessive compulsive disorders (Swerdlow et al., 1993). Less is known about the specificity of latent inhibition, although disturbances have been reported in psychotic-prone subjects (DeLaCasa et al., 1993). In other words, it is not yet clear how well APO-SUS rats model...
the schizophrenic condition specifically. One difference between schizophrenic patients and APO-SUS rats is related to the prepulse intensity effect. Thus, whereas APO-SUS rats only show disruption of prepulse inhibition with low intensity prepulses (Fig. 3), schizophrenic patients have disturbances at all intensities tested (Grillon et al., 1992). It is at present unclear whether this represents a qualitative difference (i.e., different mechanisms acting at different prepulse intensities), or a quantitative difference (i.e., with increasing prepulse intensities the differences between APO-SUS and UMSUS is reduced). There is, however, some evidence for the latter explanation. Thus, the pharmacogenetically selected APO-SUS show deficits at 2 and 4 \text{dB[A]} prepulse intensity, whereas the animals selected with the gnawing cage only show deficits at 2 \text{dB[A]}. Moreover, we have recently shown that the dopaminergic agonist 7-OHDPAT disrupts prepulse inhibition in a dose-dependent manner (Ellenbroek and Cools, in preparation): 0.1 mg/kg only affects 2 \text{dB[A]}, 0.33 mg/kg affects 2 and 4 \text{dB[A]}, and 1 mg/kg affects 2, 4, and 8 \text{dB[A]}. If the difference is, indeed, of a quantitative nature, it seems logical to assume that further breeding should eventually lead to disruption of prepulse inhibition at all prepulse intensities.

The validity of APO-SUS rats as animal models for (aspects of) schizophrenic patients can also be studied in other paradigms like P50 gating, startle habituation, Kamin Blocking. This is currently under investigation. However, APO-SUS rats show more similarities with schizophrenia. Thus, APO-SUS rats (Cools et al., 1994) and schizophrenic patients (Toru et al., 1984) have elevated level of mRNA for tyrosine hydroxylase in the substantia nigra pars compacta. Likewise, enhanced metabolism in the globus pallidus has been described in schizophrenic patients (Early et al., 1987) and in APO-SUS rats (Cools et al., 1994). APO-SUS rats have also been found to have enhanced levels of dopamine D2 binding in the dorsal striatum (Cools et al., 1994). Whether increased binding of D2 antagonist also occur in the caudate-putamen of drug naive schizophrenic patients remains unclear. Thus Wong et al. (1986) did report increases, which could, however, not be replicated in at least two other studies (Farde et al., 1990; Martinvet al., 1990). It has been hypothesized for a long time that schizophrenic patients suffer from a state of hyperarousal (Venables, 1964), which seems to parallel the heightened response to novelty in APO-SUS rats (see above). There have been several reports of decreased numbers of natural killer cells in schizophrenic patients (DeLisi et al., 1983; Sasaki et al., 1994). We have recently also found decreased numbers of NK cells in APO-SUS rats (A. R. Cools, C. Heynen, A. Kavelaars, and B. A. Ellenbroek, unpublished data). A very interesting finding is the reduced sensitivity of APO-SUS rats for rheumatoid arthritis (vandellangerijt et al., 1994). This agrees very well with the known negative association between rheumatoid arthritis and schizophrenia (Vinogradov et al., 1991).

Overall conclusions

The results of the present set of experiments showed that rats with an enhanced sensitivity to apomorphine showed deficits in information processing also seen in several psychomotor diseases, especially schizophrenia. Thus, irrespective of selection procedure (open field, gnawing cage, or pharmacogenetic selection) rats with a high susceptibility to apomorphine showed a decreased prepulse inhibition of the acoustic startle response (especially with weak prepulses) and a diminished latent inhibition, although the strongest effects were seen in the pharmacogenetically selected group. However, the data also suggest that the changes in prepulse inhibition in APO-SUS rats are not as strong as those seen in schizophrenic patients. This suggests that APO-SUS rats may represent an interesting model for psychosis-prone (or schizophrenia-prone) patients, especially since these rats show several other (biochemical, endocrinological, and immunological) features also reported for schizophrenic patients.

The present findings of distinct changes in prepulse inhibition and latent inhibition in rats that have never been treated with a drug also may have implications for possible new approaches to the identification of antipsychotic drugs. Most screening tests for antipsychotic treatments rely upon drugs such as dopamine agonists to induce the behavioral deficit of interest. The ability to study the effects of putative antipsychotics in a nonpharmacological behavioural model may help to identify antipsychotic drugs with a novel mechanism of action. In contrast to the effectiveness of antipsychotics in latent inhibition (Dunn et al., 1993), relatively few studies have found antipsychotics to improve prepulse inhibition in normal rats (Hoffman et al., 1993; Swardlow and Geyer, 1993). Based on the present results, one would predict that the APO-SUS rats would be more sensitive to the facilitatory effects of antipsychotics on prepulse inhibition.

References


