Sociosexual behaviour and paternity in procarbazine-exposed rats with or without regional testicular circulatory isolation

S. A. Farr1,2,3, F. E. Johnson1,3* and G. T. Taylor2

1Department of Surgery, St Louis University Medical School, St Louis, MO 63110–0250, USA;
2Behavioral Neuroscience Laboratory, University of Missouri, St Louis, MO 63121, USA; and
3Department of Veterans Affairs, St Louis Medical Center, St Louis, MO 63106, USA

Male Sprague–Dawley rats were used in two experiments in which a procarbazine bolus (400 mg kg–1 body mass) was administered with or without testicular circulatory isolation in the form of brief clamping of the spermatic cord and gubernaculum during drug administration. Separate tests of aggressiveness, sexual motivation, copulatory performance and paternity over the subsequent 6 weeks were used to assess functional changes resulting from testicular circulatory isolation. Experiment 1 compared intermale aggression and sexual motivation of animals in groups receiving procarbazine plus testicular circulatory isolation lasting 0, 15 or 45 min with that of animals in control groups with no clamp and no drug. Experiment 2 used a 2 × 2 factorial design to evaluate sexual performance and resulting paternity in animals 2 months after testicular circulatory isolation and drug exposure compared with that in control animals. Procarbazine treatment induced minimal disruption of normal interest in a receptive female, copulatory measures (intromissions or ejaculations) and structural integrity of seminal vesicles, bulbospongiosus muscles and ventral prostate glands. Animals exposed to the drug without testicular circulatory isolation were significantly less aggressive than animals in other groups. The most profound influence of procarbazine was on paternity. Males exposed to procarbazine with or without testicular circulatory isolation impregnated notably fewer females than did control males that were not exposed to the drug. There was no evidence of recovery of normal fertility up to 10 weeks after exposure to the drug. In conclusion, the deleterious influence of procarbazine on androgen-sensitive processes appears to be specific to intermale aggression and to fertility. The testicular circulatory isolation technique, for 45 min in particular, softened the impact of the drug on social behaviour, although procarbazine suppressed fecundity even with testicular circulatory isolation.

Introduction

A primary concern in cancer treatment is long-term dysfunction in organs other than the target of chemotherapy. Procarbazine is commonly used in multi-agent regimens, particularly against lymphomas. The acute toxicity of procarbazine on bone marrow and the gastrointestinal tract usually recovers with time. Unfortunately, the acute gonadal toxicity is usually irreversible (Schilsky et al., 1980; Byrne et al., 1987; Rivkees and Crawford, 1988). Therefore, it is of considerable clinical interest to develop procedures that reduce the initial testicular injury (Ward et al., 1990; Kangasniemi et al., 1995a). Surgical procedures involving short-term regional vascular occlusion offer a potential solution. Techniques using tourniquets and occlusive intra-aortic balloons to provide regional organ protection by partial vascular occlusion have been reported (Conrad and Crosby, 1960; Gregory et al., 1982; van Vliet et al., 1988). The procedure is to isolate a vulnerable organ system from the systemic circulation for the duration of the half-life of the drug or longer.

We have reported (Farr et al., 1993) an animal model of testicular circulatory isolation (TCI), applied for the duration of the half-life of a typical chemotherapeutic drug. An arterial clamp is applied to the spermatic cord and gubernaculum immediately before injection of procarbazine, and then removed up to 45 min later. Our prior findings (Stern et al., 1990; Farr et al., 1993) suggested that TCI may provide at least partial testicular protection during chemotherapy, as measured by morphology and function of testes in rats exposed to procarbazine with or without TCI. Although the number of testicular spermatozoa may not return completely to control values, the concentration of spermatozoa in procarbazine-exposed rats with TCI may be enough to impregnate a female. Indeed, there are data pointing to successful paternity in some males that were given an unlimited time with the female (Liebscher et al., 1990), although no attempt was made to assess how ‘normal’ the sexual responses of the treated males were.
The present study used the TCI animal model to answer two questions of considerable clinical interest. (1) What are the patterns of initial and eventual disruption of androgen-sensitive sociosexual behaviour due to procarbazine administration? (2) Can TCI protect against acute and chronic behavioural dysfunction and promote full recovery following procarbazine treatment of the varied reproductive systems contributing to normal fertility?

Young adult male rats were used in two experiments using well-established paradigms for quantifying rodent social behaviour (Barfield, 1993; Taylor et al., 1993; Sachs and Meisel, 1994). Experiment 1 was used to track changes in sexual motivation and aggressive behaviour for 8 weeks after procarbazine was administered with or without TCI. Experiment 2 assessed copulatory performance and the capacity of males to impregnate females 2 months after procarbazine treatment with or without TCI. At necropsy, prostate glands, seminal vesicles and bulbospongious muscles were examined as morphological markers of normal androgenic activity.

Materials and Methods

Animals and housing

The animals were sexually naive Sprague–Dawley male rats (n = 80, 9 weeks old, Harlan Sprague–Dawley, Indianapolis, IN) at the beginning of experimentation. Animals were housed individually in hanging plastic cages measuring 48 cm × 24 cm × 20 cm. In addition, females (n = 50) and another set of males (n = 10) of the same age, strain and source as the experimental males were used as ‘stimulus animals’ for the behaviour tests. Standard rat chow and tap water were available ad libitum throughout the experiments. Temperature (21°C ± 1°C) and relative humidity (50%) were controlled automatically. Lighting was on a cycle of 12 h light and 12 h darkness each day.

Materials

Apparatus. For behavioural observations, a metal and Plexiglas apparatus measuring 120 cm × 30 cm × 30 cm, with wire mesh flooring, was constructed based on a terrarium described in detail by Taylor et al. (1993). The only difference was the designation of a section 16 cm wide in the middle of the apparatus as a neutral space to provide a clear distinction between left and right sides of the apparatus. As in the previous study (Farr et al., 1993), a Doppler flow detector (Model 812, Parks Medical Electronics, Aloha, OR) was used during surgery to confirm successful interruption of blood flow with the non-traumatic vascular clamps.

Drugs. Procarbazine hydrochloride was donated by Hoffman-LaRoche (Nutley, NJ) and freshly prepared for each administration by dissolution in 0.9% (w/v) NaCl. Oestrogen and progesterone were purchased from Sigma Chemical Co. (St Louis, MO) and suspended in olive oil for injection. The hormones were used to induce behavioural oestrus in ovariectomized females.

Surgery

Surgical procedures for TCI, as well as methods for administering anaesthesia (pentobarbital sodium) and drug (procarbazine or placebo), have been described in detail by Farr et al. (1993). Essentially, intravenous access was via internal jugular vein cutdown. TCI was accomplished by applying non-traumatic vascular clamps to the surgically exposed spermatic cord and gubernaculum. Clamps were removed after either 15 min or 45 min. Cessation and return of blood flow was confirmed by the Doppler flowmeter. Controls and drug-exposed males without TCI received sham surgery, during which the testis was exposed and returned to the scrotum using the same technique as for the TCI groups, but without clamping.

Females for Expt 1 were ovariectomized using a single abdominal incision (Taylor et al., 1989a). Uterine horns were exposed and ovaries were excised. All incisions were closed with 3-0 polyglycolic acid suture.

Experimental design

Males (n = 40) for Expt 1 were assigned at random to one of four groups (n = 10 per group). Experimental animals (groups 2–4) received a single bolus of procarbazine (N-(1-methyllethyl)-4-[(2-methylhydrazino)methyl]benzamide monohydrochloride) via a jugular vein injection (400 mg kg⁻¹) and, simultaneously, received the TCI procedure for either 15 min (group 2) or 45 min (group 3). The other experimental group (group 4) received drug without TCI. Group 1 controls received neither treatment. Because the influence of TCI alone has been the subject of previous studies from this laboratory (Stern et al., 1990), a control group receiving TCI but no drug was not included in the first experiment. Animals were left undisturbed for two weeks after surgery, and behavioural tests began during week 3 and continued into week 10.

A 2 × 2 factorial design (n = 10 per group) was used in Expt 2 with the main factors consisting of drug exposure (procarbazine or placebo) and manipulation (TCI or no TCI). Group 1 received sham surgery without either procarbazine or TCI, group 2 received TCI for 45 min without procarbazine, group 3 received procarbazine plus TCI for 45 min, and group 4 received sham surgery with procarbazine. Tests of fertility began after 2 months; during this time the animals were left undisturbed to allow the required period for new spermatogenesis in male rats (Clermont, 1972). Behavioural observations were conducted during each of two fertility tests separated by 2 weeks.

Behavioural methods and necropsies performed at the end of the two studies were procedures that we have used frequently over the past decade. A brief description for each method of data collection follows.

Procedural details

Pre-experimental experiences. Subject males were exposed to a sexually receptive female before the beginning of the experiment. An ovariectomized female was induced to oestrus with s.c. injections of 100 μg oestradiol benzoate and, 48 h later, 400 μg progesterone 3–4 h before being placed
overnight in the home cage of the male (Taylor et al., 1989a). Neither formal observations of successful copulation nor other evidence, such as the presence of a copulatory plug in the animal’s bedding (Sachs and Meisel, 1994), were obtained during this pre-experimental period. However, informal observation suggested that most males copulated during the 12 h that each male spent with a female in oestrus.

**Experiment 1: sexual motivation and intermale aggression.** In the third week after surgery, behavioural testing started and continued for the next 6 weeks. Stimulus animals for the two social behaviour paradigms were ovariectomized females ($n = 10$) and untreated adult males (also $n = 10$). Every male was observed once each week. Aggression test sessions with an unfamiliar stimulus male were conducted on 3 weeks and tests of sexual motivation with an ovariectomized female were conducted on the other 3 weeks (Taylor et al., 1994). Observations were conducted in an experimental room away from colony rooms. After each session, the apparatus was cleaned thoroughly with soap and water.

Scheduling of sessions was counterbalanced within and between groups. Stimulus animals were, by necessity, paired with several subject males during the 3 weeks of behavioural testing. However, care was taken that the same subject and stimulus animal were not paired more than once. Moreover, neither a subject male nor a stimulus animal was tested twice on the same day. Behavioural observations continued until the end of the study, when all subject males were killed.

**Experiment 2: copulatory behaviour and paternity.** Vaginal smears were used to monitor stages of the oestrous cycle of the unoperated, cyclic females ($n = 40$) used for paternity testing beginning 2 months after treatment. With colony lights off at 10:00 h, testing was conducted during the initial 4 h of darkness and in the home cages of the males. A female detected to be in late pro-oestrus to early oestrus was introduced into a male’s home cage for a mating session lasting 60 min. Measures of sexual performance were recorded for the pair. After the session was completed, the female was returned to her individual home cage to await parturition.

The procedure described by Taylor and Weiss (1987) was used to allow each male two opportunities to impregnate a female. If the female from the first mating session clearly was not pregnant, the male was given a second mating test with a different receptive female 2 weeks after the initial session; that is, only those males failing to impregnate a female during the initial 60 min session were given a second opportunity to impregnate another female.

**Behavioural measures**

Sexual motivation was evaluated apart from sexual performance. Our test of sexual motivation used a well-established paradigm (Taylor et al., 1994) for evaluating the probability of actively searching for a receptive female. Ovariectomized females were housed in small wire cages at opposite ends of an apparatus. One of the females was induced to receptivity by hormone injections; the other female was not injected. The male could move freely about the apparatus without direct interactions with the females that confound assessment of the interest of the male in making social contact. Motivation was quantified by the relative time the male spent near the inaccessible oestrous female as opposed to the non-oestrous female during test sessions lasting 20 min.

Aggressive behaviour was evaluated (Taylor et al., 1996) in a terrarium during 10 min sessions once a week between a subject male and an untreated, unfamiliar male. The two animals were introduced into the terrarium simultaneously to avoid resident–intruder complexities. Frequencies of categories of aggression in male rats described by Grant (1963) were recorded and totalled for an aggression score for each pair (Taylor et al., 1984). Each incidence of pushing, sideways kicking, aggressive grooming, aggressive posturing and attack with biting observed by the single experimenter was tallied.

Sexual performance was assessed (Taylor et al., 1989b) in Expt 2 during pairings of a male with a gonadally intact female detected as being receptive by a vaginal smear. An oestrous female was placed into the home cage of the male for the mating test. Latencies to first intromission and frequencies of both intromissions and ejaculations were recorded during a 60 min session. Frequencies of female solicitations of the male and her rejections of his sexual approaches were also measured (Taylor et al., 1989a).

**Fertility**

Paternity was established in the second experiment by delivery of live pups from female partners of the males. Tightly constrained mating sessions of 60 min were used as a limiting condition for the expression of fertility. A male was given two opportunities to impregnate different receptive females. Males were given a paternity score according to whether a female was impregnated during the initial sexual contact (score of 2), the second session (score of 1) or neither (score of 0). Numbers of pups per litter were recorded, and the entire litter was weighed.

**Physiological measures**

Sex accessory organs representing targets of androgens were examined at necropsy. We used multiple markers of the reproductive system to estimate disruption of normal function (Taylor et al., 1989b, 1996). Seminal vesicles, bulboepididymal muscles and ventral prostate glands were excised and examined for indications of pathology. Wet masses were recorded for each structure. Sperm head counts and testicular integrity were not measured in this study since these measurements were made from similar groups of animals during previous work (Stern et al., 1990; Farr et al., 1993).

**Statistical analyses**

Data from Expt 1 obtained during the weekly tests of sexual motivation and aggression were examined with analyses of variance (ANOVA). Repeated measures factorial ANOVAs, with time as the repeated factor, were performed on data from Expt 1 as a measure of recovery following experimental treatment. Tukey’s honestly significant difference tests (HSD)
were used for post-hoc, pair-wise comparisons of mean group differences. Statistical significance was set at $P < 0.05$ in each case.

Frequencies of types of behaviour and masses of tissues from Expt 2 were analysed by $2 \times 2$ factorial analyses of variance (ANOVA; $P < 0.05$), the main factors being drug (exposed to procarbazine or not) and TCI (either TCI for 45 min or not). The behavioural scores analysed were either the totals for the session for those males requiring a single session to impregnate a female or a mean average for the animals requiring two sessions of sexual contact. Finally, paternity scores reflecting whether the male impregnated a female on the first, second or neither attempt were analysed.

**Results**

Findings from the aggression tests with stimulus males and tests for interest in inaccessible oestrous females were summarized and are reported as group means ± SEM (Table 1). Females delivering pups from each group were converted to percentage values; that is, the percentages reported in Table 2 are the number of males per group that successfully impregnated a female in either of two mating sessions.

**Experiment 1: sexual motivation and internmale aggression**

There were no significant differences among main factors or the interactions for the motivation measure. The conclusion is that groups did not differ in the time spent in proximity to an oestrous as opposed to a non-oestrous female, the measure of sexual interest used here.

The repeated measure analyses of aggressive behaviour yielded non-significant values for the 'week of testing' main effect and for the group $\times$ weeks interaction. Those findings indicate that aggressive responding did not change during the weeks of testing for any of the groups. However, the value obtained for the 'group' main factor for aggressive behaviour was statistically reliable $[F(3,30) = 5.17, P < 0.05]$. Post-hoc analyses to detect which groups differed revealed that the procarbazine-exposed males without TCI were the least aggressive animals in the entire study, whereas the control group without drug or clamp and the group with drug and TCI for 45 min were most aggressive.

**Experiment 2: sexual performance and paternity**

In this paradigm, the males were allowed to interact actively with receptive females in mating tests given during weeks 8–10 after experimental treatments. There were no significant differences for the drug or the TCI main effects or for their interaction on either measures of sexual performance (copulatory intromissions and ejaculations). On the other hand, there were clear decreases in paternity due to drug treatment. Experimental and control groups were compared on a paternity score based on whether a female delivered live pups after the first, second or neither mating test. It is notable that 40% of the group without the drug or TCI failed to impregnate a female. This probably reflects the strict conditions imposed by our paradigm to limit time for mating.

Nonetheless, results of the analyses of females with live births revealed significant differences only with respect to the main factor of drug $[F(1,30) = 6.97, P < 0.05]$: that is, groups exposed to procarbazine were significantly less successful in impregnating females than the two groups not exposed to procarbazine. There were no significant differences between the two drug-exposed groups. In addition, the two groups without drug did not differ. Indeed, the group without drug but with TCI and the group without either drug or TCI fathered similar numbers of pups per litter (means = 11.0 and 12.6, respectively) and the body masses of the pups were similar (means per pup = 6.8 g and 6.2 g, respectively).
Finally, all animals were killed at the end of behavioural testing. Seminal vesicles, bulbospongious muscles and ventral prostates were excised and wet masses obtained as markers of reproductive physiology. There were no differences between drug-exposed and control groups in any of these structures.

**Discussion**

Testicular circulatory isolation is a surgical technique designed to avert local damage to testicular function during exposure to a toxin (Farr et al., 1993). An animal model of TCI was used and two experiments were conducted to answer separate but related questions about acute and chronic consequences on reproductive competence following exposure to a common chemotherapeutic agent.

Chemical treatments (Ward et al., 1990; Parchuri et al., 1993; Kangasniemi et al., 1995b) or surgical occlusion (Stern et al., 1990; van Vliet et al., 1993) have been designed to spare fertility of males exposed to procarbazine and other toxic substances used in the treatment of cancer. Those studies used indirect measures, typically sperm head counts and testicular integrity, rather than the direct measurement of paternity in the form of live births. Nevertheless, the impression left by these studies is that there are protective techniques that allow full recovery to premorbid degrees of sexuality and fertility (Stern et al., 1990; Ward et al., 1990).

However, the present study used carefully controlled contacts with females and indirect and direct measures of fertility. Young adult male rats were exposed to procarbazine with or without concurrent clamping of the spermatic cord and gubernaculum for 15 min or 45 min. Reproductive consequences were assessed using various paradigms for assessing sociosexual behaviour, androgen-sensitive tissue morphology and fertility of male rats (Taylor and Weiss, 1987; Barfield, 1993; Taylor et al., 1993). Treated males were compared with untreated controls to provide a more precise empirical test of the hypothesis that the TCI procedure would leave males with 'normal' reproductive capacity. In Expt 1, weekly behavioural tests were used to assess changes in sexual motivation and aggressiveness 3–10 weeks after surgery. In Expt 2, the study focused on copulatory performance and fertility 2 months after treatments.

The findings of Expt 1 indicated that sexual motivation, or 'libido', was not disrupted in any of the procarbazine-treated groups relative to the controls. However, there was evidence of drug-induced suppression of aggressive behaviour in males administered procarbazine without TCI. There was no evidence of recovery of this function over time. Adverse effects on a male's aggressiveness could be observed within a few weeks of surgery, and that adverse effect remained stable for up to 10 weeks after treatment. Most notably, TCI for 45 min preserved the social behaviour studied of male rats receiving the procarbazine regimen. However, males receiving the drug with only 15 min of TCI, were more similar to the unprotected, drug-exposed group than to the other groups.

The findings of Expt 2 point directly to decreased male fertility with procarbazine. Fertility was reduced to a similar extent in all drug-exposed males. The presence or absence of TCI did not improve fertility but this reduced fertility was not a result of obvious impairments to copulation. Males in procarbazine-exposed groups, particularly the group exposed to the drug plus TCI for 45 min, showed comparable sexual performance to animals from the control group with no drug treatment. In both experiments, there was no evidence for differences between groups in the integrity of seminal vesicles, bulbospongious muscles or ventral prostates. This suggests that any deleterious effects on fertility did not result from endocrine disruption to reproductive physiology. Similarly, clamping of the blood supply to the testis for up to 45 min had no lasting impact on any measure of sexual behaviour, physiology or fecundity. Results revealed no differences

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**Table 2.** Scores (means ± SEM) of sexual performance and paternity of male rats (n = 10 per group) of Experiment 2 treated with 400 mg procarbazine kg⁻¹ or placebo with or without testicular circulatory isolation (TCI)

<table>
<thead>
<tr>
<th>Group</th>
<th>Intromission frequency</th>
<th>Ejaculation frequency</th>
<th>Seminal vesicles (mg)</th>
<th>Ventral prostate gland (mg)</th>
<th>Bulbospongious muscles (mg)</th>
<th>Percentage number of males fathering live births</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 (No drug + TCI for 45 min)</td>
<td>41 ± 5</td>
<td>1.9 ± 0.3</td>
<td>34 ± 2.0</td>
<td>21 ± 2</td>
<td>27 ± 1.0</td>
<td>50%*</td>
</tr>
<tr>
<td>2 (Drug, no TCI)</td>
<td>29 ± 6</td>
<td>2.1 ± 0.4</td>
<td>34 ± 1.5</td>
<td>19 ± 2</td>
<td>29 ± 0.9</td>
<td>0%b</td>
</tr>
<tr>
<td>3 (Drug + TCI for 45 min)</td>
<td>29 ± 7</td>
<td>1.7 ± 0.3</td>
<td>35 ± 0.9</td>
<td>21 ± 2</td>
<td>28 ± 1.1</td>
<td>10%b</td>
</tr>
<tr>
<td>4 (No drug, no clamp)</td>
<td>34 ± 9</td>
<td>2.4 ± 0.2</td>
<td>37 ± 1.1</td>
<td>20 ± 1</td>
<td>27 ± 1.0</td>
<td>60%b</td>
</tr>
</tbody>
</table>

Sexual performance was measured during pairings of a male with an intact, receptive female; the frequencies of intromissions and ejaculations were recorded (Taylor et al., 1989a).

Paternity was established by the delivery of live pups from female partners of the males tested. The percentage values in this column are the number of males per group that successfully impregnated a female in either of two mating sessions.

Androgen-sensitive structures (seminal vesicles, ventral prostate glands and bulbospongious muscles) were weighed 10 weeks after treatment (mg tissue per 100 g body mass).

Superscript letters indicate statistically reliable group differences (Tukey's HSD test, P < 0.05).
between groups not exposed to procarbazine but given TCI and control groups given no drug and no clamp.

The overall conclusion is that procarbazine causes no detectable change in the interest of a male rat for a receptive female nor in his copulatory effectiveness. However, the drug suppresses intermale aggressive behaviour and, perhaps more important, the capacity to impregnate a female. There was no recovery over time in animals experiencing impairment soon after the drug exposure, suggesting that losses in normal social behaviour and fertility are probably permanent. Previous work (Farr et al., 1993) points to failure of testicular spermatogenesis to recover fully as a likely source of the infertility in males exposed to procarbazine. The mechanism for the reduced aggressiveness in procarbazine-exposed animals is not obvious. It is unlikely to be based on androgens; androgen-sensitive sexual behaviour and secondary sex structures recovered normally in both drug and TCI groups. Certainly, the differences in aggressive behaviour suggest modification of the central nervous system. That aggressiveness and fertility of the group not exposed to procarbazine but given TCI for 45 min was equal to that of controls suggests full restoration of both peripheral and central functions.

Conclusions on the value of the TCI procedure are mixed. Although the procedure failed to prevent the infertility common with procarbazine, TCI reduced the impact of the drug on behaviour. Compared with drug-exposed males without TCI, procarbazine-exposed males given 45 min of TCI showed less disruption of responses to another male. Of clinical relevance is the confirmation from our animal findings of prior reports (Devlen et al., 1987) for men who have undergone chemotherapeutic treatments. The suggestion is that procarbazine will reduce their capacity to father children, even with attempts to block toxicity by using procedures such as TCI.

Nevertheless, the findings give reason for optimism. The data suggest that procarbazine-treated men might be expected to show rapid recovery of their capacity to initiate and complete mutually satisfying copulation. In addition, a notable finding was the safety of the TCI procedure, which did not produce dysfunction in behaviour, physiology or fertility.

The research was supported, in part, by grants from the Veterans Administration (to F. E. Johnson) and a UM-St Louis Research Award (to G. T. Taylor).

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