



Strain-specific battery of tests for domains of mania: effects of valproate, lithium and imipramine

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The lack of efficient animal models for bipolar disorder (BPD), especially for the manic pole, is a major factor hindering the research of its pathophysiology and the development of improved drug treatments. The present study was designed to identify an appropriate mouse strain for modeling some behavioral domains of mania and to evaluate the effects of drugs using this strain. The study compared the behavior of four strains: Black Swiss, C57Bl/6, CBA/J and A/J mice in a battery of tests that included spontaneous activity; sweet solution preference; light/dark box; resident-intruder; forced-swim and amphetamine-induced hyperactivity. Based on the 'manic-like' behavior demonstrated by the Black Swiss strain, the study evaluated the effects of the mood stabilizers valproate and lithium and of the antidepressant imipramine in the same tests using this strain. Results indicated that lithium and valproate attenuate the 'manic-like' behavior of Black Swiss mice whereas imipramine had no effects. These findings suggest that Black Swiss mice might be a good choice for modeling several domains of mania and distinguishing the effects of drugs on these specific domains. However, the relevance of the behavioral phenotype of Black Swiss mice to the biology of BPD is unknown at this time and future studies will investigate molecular differences between Black Swiss mice and other strains and assess the interaction between strain and mood stabilizing treatment.

Keywords: behavioral domains, mood stabilizers, strain differences, reward seeking, vigor, aggression

INTRODUCTION

Bipolar disorder (BPD) is a chronic and debilitating illness which includes alternate episodes of depression and mania (Sadock and Kaplan, 2002). The estimated lifetime prevalence rate of BPD is 3.9% and the median age of onset is 25 years. It is currently recognized that treatment outcomes for many bipolar patients are relatively poor, as significant side effects, high rates of relapse and lingering residual symptoms are evident (for review, see Miklowitz and Johnson, 2006).

The paucity of appropriate animal models for BPD hinders the research of the pathophysiology and the development of new treatments for this condition (Tecott and Nestler, 2004; Gould and Einat, 2007). A number of possible approaches for the development of new models for BPD and for mania include models based on molecular or genetic manipulations [e.g., Ralph-Williams et al., 2003; Machado-Vieira et al., 2004; Einat and Manji, 2006; Roybal et al., 2007; Malkesman et al., 2009), models based on pharmacological interventions (Antelman et al., 1998; Gould et al., 2007; Riegel et al., 2009)] and models induced by environmental stressors [e.g., sleep deprivation (Gessa et al., 1995), dominant-submissive behavior (Knapp et al., 2002; Malatynska and Knapp, 2005), swim stress (Flaisher-Grinberg and Einat, 2009b)].

An additional possible approach emphasizes the identification or development of model animals (Insel, 2007). A model animal can be a specific species or a strain (either standard or with targeted mutation) which demonstrates behavioral or biological similarities with a disorder and may therefore be more homologous with it (Insel, 2007; Ashkenazy et al., 2008; Flaisher-Grinberg et al., 2008;

Malkesman et al., 2009). This approach was the conceptual basis for the development of strains with direct mutations to be used as models for mania (Ralph-Williams et al., 2003; Roybal et al., 2007). However, the identification of appropriate standard mice strains was also found productive. For example, strain variations in the effects of amphetamine and lithium suggest that only a few strains are suitable for use in the amphetamine-induced hyperactivity model of mania (Gould et al., 2007). Preliminary experiments in our laboratory suggest that based on their behavioral repertoire, mice from the Black Swiss (BS) strain may be more suitable for use in modeling a number of the domains of mania compared with other strains. Specifically, BS mice were found to demonstrate 'manic-like' increased risk-taking and reward-seeking behaviors as well as an elevated response to psychostimulants compared with C57Bl/6 mice (Hiscock et al., 2007; Flaisher-Grinberg and Einat, 2009a).

The present study was designed to continue and explore the utility of the BS strain for modeling domains of mania. In Stage 1, the behavior of BS mice was assessed in a continuous battery of tests relevant to the manic pole of BPD, including spontaneous activity, sweet solution preference, light/dark box, forced swim test (FST), resident-intruder and amphetamine-induced hyperactivity. These tests are part of a broader battery of tests that was suggested to model separate domains of mania (Einat, 2006). From this proposed battery we chose to utilize the tests that have better validation in the context of affective disorders (for elaborated details, see Flaisher-Grinberg and Einat, 2009a). In Stage 2, in order to test the pharmacological (predictive) validity of the battery, the effects of

the prototypic mood stabilizers valproate and lithium, and of the tricyclic antidepressant imipramine, on the behavior of BS mice were evaluated. Given that pharmacological validity is supported when drugs that are effective in the disorder (i.e., mood stabilizers such as lithium and valproate) are effective in the model while drugs that are not effective in the disorder (i.e., prototypic antidepressant such as imipramine) are not effective in the model (Willner, 1991) and in accordance with their effects in manic patients (Miklowitz and Johnson, 2006), it was hypothesized that valproate and lithium, but not imipramine, would induce an anti-manic effect.

MATERIALS AND METHODS

SUBJECTS

For Stage 1, Black Swiss (Taconic, NY, USA), C57Bl/6 (Harlan, IN, USA), CBA/J and A/J (Jackson laboratory, ME, USA) mice were used. The selection of these strains was based on previous results in the amphetamine-induced hyperactivity model (Monleon et al., 1995; Agmo et al., 1997; Anjaneyulu et al., 2003; Arban et al., 2005; Gould et al., 2007; Hiscock et al., 2007) and on previous experiments performed in our laboratory. For Stage 2, only BS mice were utilized. All mice were transported to our laboratory, where experimentation started no less than 1 week later to allow for appropriate acclimatization time. Male mice, 10 to 12 weeks old and 25 ± 7 g at the start of treatment period, were singly housed in a colony room with constant temperature ($22 \pm 1^\circ\text{C}$), 12/12 light dark cycle (lights on/off at 0730/1930) and ad-lib food and water. For the resident-intruder test (see Behavioral tests), strain-matched mice served as intruders. Intruders were 7 to 8 weeks old, 22 ± 5 g at the start of treatment period and were group housed 3–4 in a cage under the same conditions as described above. All experiments were conducted during the early light phase of the light/dark cycle, in dedicated experimental rooms and under standard fluorescent lights (except for the sweet solution preference test, conducted in the colony room). All experimental procedures followed NIH guidelines and were approved by the University of Minnesota Institutional Animal Care and Use Committee (Protocol No. 0610A94146). The number of animals per group in all experiments was 10–16 except for the amphetamine experiments where group size was half of the original number. Intruder mice were used twice and care was taken to counterbalance first and second exposures across test groups.

DRUGS AND DRUG ADMINISTRATION

Valproate, lithium and imipramine (SIGMA, St. Lewis, MI, USA) were dissolved in saline to a dose of 50, 100 and 200 mg/kg (valproate), 100 and 200 mg/kg (lithium) or 10, 20 and 30 mg/kg (imipramine) and administered via I.P. injection in a volume of 10 ml/kg. Administration started 1 day before the beginning of the test battery and continued throughout the days of testing. Drugs were administered twice daily, with morning injections 30 min before testing and evening injections at 8 PM. For the amphetamine-induced hyperactivity test, d-amphetamine (SIGMA, St. Lewis, MI, USA) was dissolved in saline to a dose of 1 mg/kg and administered via I.P. injection in a volume of 10 ml/kg immediately before the beginning of the test. Vehicle groups received an equivalent volume of saline. Drug doses were chosen based on their effects in animal models of affective disorders (e.g. Monleon et al., 1995; Agmo et al., 1996; Gould et al., 2007) and in studies performed in our laboratory

(Hiscock et al., 2007; Flaisher-Grinberg and Einat, 2009b; Flaisher-Grinberg et al., 2009). Schedule of administration was selected based on previous work demonstrating its efficacy in some of the models included in the current study (Hiscock et al., 2007; Flaisher-Grinberg and Einat, 2009b; Flaisher-Grinberg et al., 2009).

EQUIPMENT AND PROCEDURES

Sequence of experiments

To minimize the effects of previous tests on later ones in the battery the order of tests was arranged from the less to the more intrusive (McQuade et al., 2003). In Stage 1, one cohort of mice was subjected to two sets of tests separated by 1 week of rest: spontaneous activity test; light/dark box; forced swim test; amphetamine-induced hyperactivity (set 1) and sweet solution preference and resident-intruder tests (set 2). Specific tests were separated by 48-h rest period. In Stage 2, one cohort of mice was subjected to all six tests conducted on consecutive days in the following sequence: spontaneous activity test; sweet solution preference; light/dark box; resident-intruder test; forced swim test and amphetamine-induced hyperactivity test. The unification of the two test sets to one and the switch from 48 to 24 h separation between tests followed a number of pilot tests which indicated no difference in the behavioral response of BS mice under these two conditions (data not shown).

Behavioral tests

Spontaneous activity. Mice were tested for spontaneous activity in $50 \times 25 \times 20$ cm transparent plastic automated activity monitors (Opto3, Columbus Instruments, Columbus, OH, USA). Infrared beam crossings were recorded for 60 min in 10-min intervals and total ambulatory activity was calculated across the entire session. At the end of the session mice were returned to their home cages and the boxes were wiped clean with a 10% alcohol solution.

Sweet solution preference test. Mice were supplied with a bottle of 1% saccharin solution (SIGMA, St. Lewis, MI, USA) on top of the regular supply of water and food. Saccharin concentration was selected as it is in the high end of the concentration-intake curve (Flaisher-Grinberg et al., 2009) and based on previous work done with BS mice (Hiscock et al., 2007). The saccharin solution bottle was available to the mice throughout the entire sweet solution preference test period. Weights of saccharin solution and water bottles were taken at the beginning of the experiment and every 24 h thereafter. Sweet solution preference was calculated daily as the ratio of saccharin out of total liquid consumption. For Stage 1, saccharin preference was measured across 4 days and for Stage 2 it was measured across 2 days because it was demonstrated that differences between groups were established within the first 2 days.

Light/dark box. The light/dark box apparatus consisted of a wooden box, $60 \times 40 \times 45$ cm, divided by a partition to 1/3 covered, black-painted segment and 2/3 open, well lit, white-painted segment. Mice were placed in the light compartment with the partition slightly ajar to permit free transitions between compartments. Behavior was digitally recorded from above for a 5-min session and recordings served to manually score the time spent in the light compartment. An entry into the light compartment was scored when all four paws

of the mouse were in that compartment. At the end of the session mice were returned to their home cages and the area was wiped clean with a 10% alcohol solution.

Resident-intruder test. Resident mice were transferred in their home cages to an experimental room where cage covers were removed. After a 5-min adaptation period, a younger, previously group-housed mouse (i.e., intruder), was placed into the resident's cage and behavior was digitally recorded from above for a 10-min session. Recordings served to manually score resident's aggressive interactions (defined as attempts to bite, actual bite, boxing postures and wrestling postures) and non-aggressive interactions (defined as other types of body contact including sniffing, allogrooming and body contact). Behaviors performed when not interacting were not scored. The resident's aggression score was calculated as the ratio of aggressive interactions out of total (aggressive + non-aggressive) interactions. At the end of the session the intruder was removed and placed back in its home cage, the resident's cage was covered and both mice were returned to the colony room. To minimize harm to animals, mice were briefly separated when attacks became vicious and included significant biting. Hence, the total time of attacks could not be scored and only numbers of aggressive and non-aggressive interactions were scored from recordings. This method was previously shown to be sensitive enough to demonstrate the effects of mood stabilizers in this test (Einat, 2007b).

Forced swim test (FST). Mice were placed for a 6-min session in a vertical cylindrical plastic container (25 cm tall × 18 cm diameter), filled to a depth of 15 ± 1 cm with tap water at $22 \pm 1^\circ\text{C}$. This depth was sufficient to ensure that mice could not escape or touch the floor of the container. At the end of the session, mice were removed from the water, dried with a paper towel, and placed back in their home cage. Water in the container was changed after each session. Sessions were digitally recorded from the side and the last 4 min of each session were scored using an automated acquisition and analysis software (Biobserve, Bonn, Germany). Due to technical problems, sessions from Experiment 3, Stage 2, were manually scored by a trained experimenter, blind to the experimental conditions. Scored behaviors were defined as active (swim and struggle) versus passive (floating with only minimal movements needed to keep head above water) behaviors.

Amphetamine-induced hyperactivity. Mice were administered with amphetamine or vehicle and immediately placed in activity monitors for a 60-min session, where activity was detected by infrared beams and total ambulatory activity was calculated across the entire session. At the end of the session mice were returned to their home cages and the boxes were wiped clean with a 10% alcohol solution.

STATISTICAL ANALYSIS

For Stage 1 of the study, the spontaneous activity, light/dark box, resident-intruder and FST data were analyzed using analyses of variance (ANOVA) with Strain (BS/C57Bl/6/CBA/J/A/J) as a main factor. For the sweet solution preference test, data were analyzed using mixed ANOVA with Strain as a main factor and Day as a repeated measures factor. For the amphetamine-induced hyperactivity test, data were analyzed using a two way ANOVA with Strain and Stimulant (amphetamine/vehicle) as main factors. Significant

main or interaction effects were followed by Fisher LSD *post hoc* comparisons. Significance was assumed at $p < 0.05$. For Stage 2 of the study, similar statistics were used except that the main factor in the different ANOVA tests was Drug dose (valproate 0, 50, 100, 200/ lithium 0, 100, 200/imipramine 0, 10, 20, 30) instead of strain.

RESULTS

STAGE 1: STRAIN DIFFERENCES IN A BATTERY OF MODELS FOR MANIA

As previously reported (for details see Flaisher-Grinberg et al., 2009), BS mice demonstrated a significantly higher sweet solution preference compared with all other strains and across all 4 days of the test (ANOVA: Strain, $F(3,44) = 22.93$, $p < 0.0001$, Day, $F(3,132) = 42.63$, $p < 0.0001$, Strain × Day, $F(9,132) = 4.11$, $p < 0.001$). As shown in **Table 1**, in comparison with the other strains, BS mice exhibited the highest level of aggression in the resident-intruder test, the lowest immobility time in the FST, the highest amount of time in the light compartment of the light/dark box (except for the CBA/J strain) and the most intense response to amphetamine. In the spontaneous activity test, both BS and C57Bl/6 mice demonstrated similar activity levels which were higher compared with A/J and CBA/J mice.

STAGE 2: THE EFFECTS OF VALPROATE, LITHIUM AND IMIPRAMINE ON THE BEHAVIOR OF BS MICE TESTED IN A BATTERY OF MODELS FOR MANIA

Experiment 1: The effects of valproate in a battery of models for mania

As previously demonstrated (Flaisher-Grinberg and Einat, 2009b), 200 mg/kg valproate reduced sweet solution preference across the 2 days of the test and 100 mg/kg valproate reduced sweet solution preference on Day 1 with a similar trend on Day 2 (**Figure 1A**, Dose, $F(3,36) = 9.98$, $p < 0.0001$, Day, $F(1,36) = 4.01$, $p < 0.05$, Dose × Day interaction, $F(3,36) = 0.65$, $p = 0.58$, see Figure for results of *post hoc* comparisons). This effect was found despite the general increase in total liquid consumption following valproate administration (data not shown). As expected (Flaisher-Grinberg et al., 2009), 200 mg/kg valproate increased immobility time in the FST with a similar trend at the 100 mg/kg dose (**Figure 1B**, Dose, $F(3,34) = 3.16$, $p < 0.05$, see Figure for results of *post hoc* comparisons). In the amphetamine-induced hyperactivity test, a trend for valproate-induced attenuation of amphetamine-induced hyperactivity was demonstrated, resulting from an attenuation at the 200 mg/kg dose (**Figure 1C**, Dose, $F(3,30) = 2.66$, $p = 0.06$, Stimulant, $F(1,30) = 38.34$, $p < 0.0001$, Dose × Stimulant interaction, $F(3,30) = 2.04$, $p = 0.13$, see Figure for results of *post hoc* comparisons). Valproate had no effect in the non-amphetamine treated groups.

Valproate had no effect on spontaneous activity across the entire session ($F(3,35) = 1.63$, $p = 0.20$) and across the first 10-min session ($F(3,35) = 0.04$, $p = 0.98$). Valproate also had no effects on behavior in the light/dark box ($F(3,36) = 0.06$, $p = 0.97$) or the resident-intruder test ($F(3,36) = 1.84$, $p = 1.57$). Interestingly, in contrast with the results of Stage 1, mice from the vehicle group demonstrated a very low aggression ratio (mean = 0.25, SE = 0.06) in the resident-intruder test. $N = 10$ /group in each of these three tests.

Experiment 2: The effects of lithium in a battery of models for mania

As previously demonstrated (Flaisher-Grinberg et al., 2009), 200 mg/kg lithium reduced sweet solution preference across the 2 days of the test (**Figure 2A**, Dose, $F(2,44) = 8.14$, $p = 0.001$, Day,

Table 1 | Strain differences in tests for domains of mania.

Test Measurement collected	Strain	N	Mean score ± SE	ANOVA analysis	
Resident- intruder test (Ratio of aggressive/total interactions)	BS	11	0.72 ± 0.05	Strain: $F(3,43) = 15.37, p < 0.0001$. 1 BS mouse was excluded showing unusual circling behavior	
	C57Bl/6	12	0.29 ± 0.19*		
	CBA/J	12	0.03 ± 0.03*		
	A/J	12	0.30 ± 0.06*		
Forced swim test: (Immobility time – sec)	BS	12	1.20 ± 0.44	Strain: $F(3,43) = 137.32, p < 0.0001$. 1 CBA/J mouse was excluded as an outlier (more than 2xSTD from mean)	
	C57Bl/6	12	125.57 ± 10.58*		
	CBA/J	11	33.44 ± 6.30*		
	A/J	12	167.99 ± 5.02*		
Light/Dark box: (Time in light Compartment – sec)	BS	11	152.82 ± 5.89	Strain: $F(3,43) = 5.17, p < 0.01$. 1 BS mouse was excluded because of camera malfunction	
	C57Bl/6	12	111.25 ± 9.82*		
	CBA/J	12	143.66 ± 26.17		
	A/J	12	87.66 ± 22.32*		
Amphetamine-Induced hyperactivity: (Total ambulatory activity)			Saline	Amph	Strain: $F(3,36) = 14.98, p < 0.0001$. Stimulant: $F(1,36) = 35.98, p < 0.001$. Interaction: $F(3,36) = 6.16, p < 0.01$. 1 C57Bl/6-vehicle mouse was excluded because of computer malfunction, 1 mouse from BS, C57Bl/6 and A/J groups were excluded as statistical outliers;
	BS	5,6	3697.20 ± 481.96	15953.67 ± 2871.14 [#]	
	C57Bl/6	5,5	2774.60 ± 346.87	8500.80 ± 826.44 ^{#, *}	
	CBA/J	6,6	1173.67 ± 171.51	6552.50 ± 1910.41 ^{#, *}	
	A/J	6,5	526.33 ± 58.32	836.00 ± 430.14*	
Spontaneous activity: (Total ambulatory activity)	BS	9	4347.33 ± 622.57	Strain: $F(3,33) = 15.27, p < 0.0001$. Three BS, CBA/J and A/J mice and two C57Bl/6 mice were excluded because of computer malfunction	
	C57Bl/6	10	4235.90 ± 400.87		
	CBA/J	9	1426.55 ± 109.64*		
	A/J	9	1486.77 ± 376.43*		

*Represents significant difference from BS mice ($p < 0.05$). [#]Represents significant difference between amphetamine- and vehicle-treated group within each strain ($p < 0.05$).

$F(1,44) = 0.97, p = 0.32$, Dose × Day, $F(2,44) = 0.28, p = 0.75$, see Figure for results of *post hoc* comparisons). No lithium effect on total liquid consumption was found in this test (data not shown, Dose, $F(2,44) = 0.42, p = 0.65$, Day, $F(1,44) = 76.77, p < 0.0001$, Dose × Day, $F(2,44) = 2.13, p = 0.13$). Lithium had no effect on spontaneous activity across the entire session (Dose, $F(2,45) = 0.24, p = 0.78, N = 16/\text{group}$) and across the first 10-min session (Dose, $F(2,45) = 0.08, p = 0.92$) but it attenuated amphetamine-induced hyperactivity at the 200 mg/kg dose without affecting the activity of non-amphetamine treated mice (Figure 2C, Dose, $F(2,42) = 12.02, p < 0.0001$, Stimulant, $F(1,42) = 52.20, p < 0.00001$, Dose × Stimulant interaction, $F(2,42) = 4.05, p < 0.05$, see Figure for results of *post hoc* comparisons). Lithium had no effects in the FST (Figure 2B, Dose, $F(2,21) = 0.06, p = 0.93$), the light/dark box (Dose, $F(2,45) = 0.60, p = 0.55, N = 16/\text{group}$) or the resident-intruder test (Dose, $F(2,43) = 1.44, p = 0.24, N = 16/\text{group}$). Vehicle-treated mice demonstrated a relatively low aggression rate in this last test (mean = 0.46, SE = 0.06).

Experiment 3: The effects of imipramine in a battery of models for mania

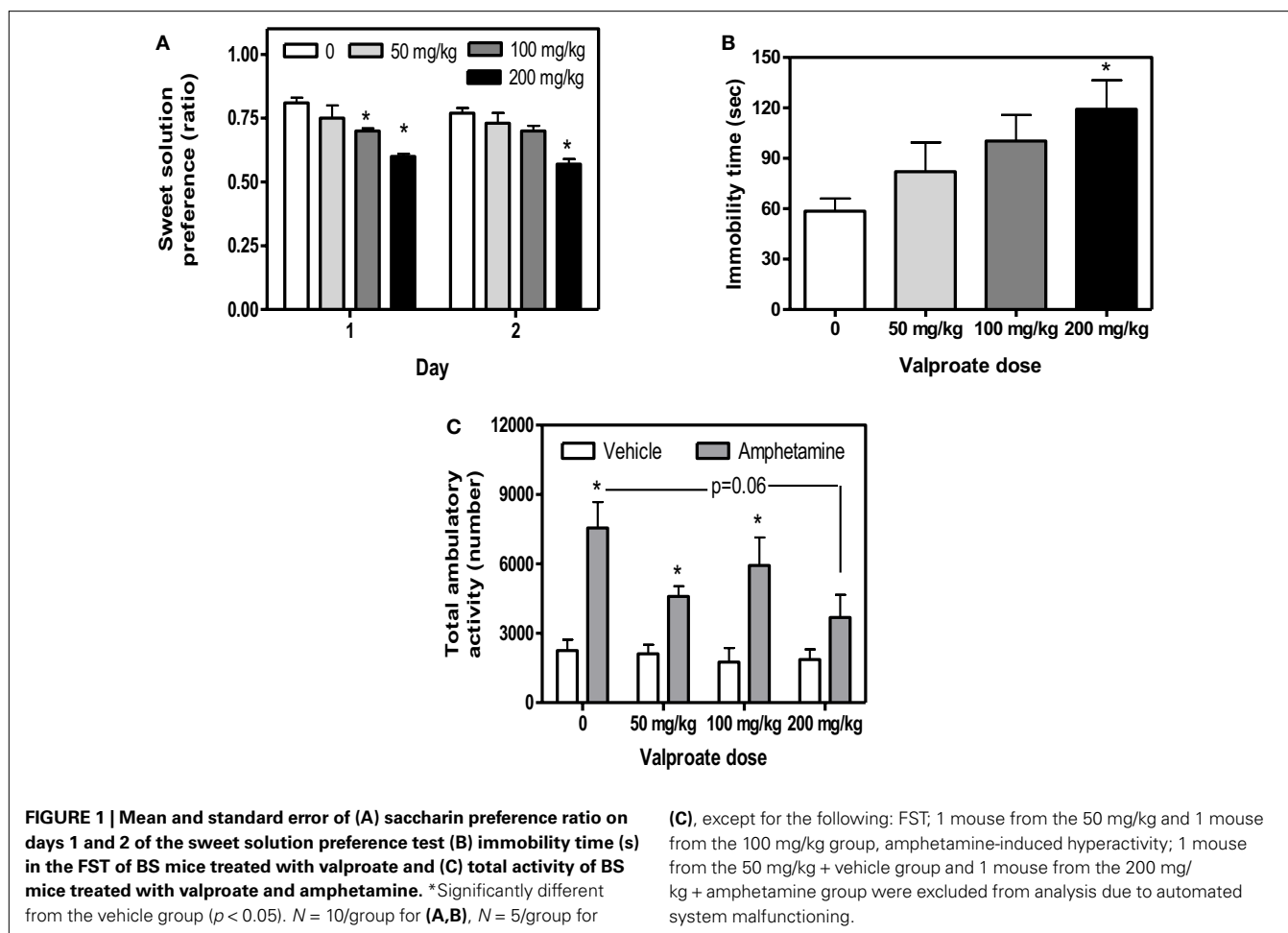
Imipramine administration did not induce any behavioral changes in the test battery for BS mice (Table 2). No imipramine effect on total liquid consumption in the sweet solution preference test was found (data not shown). As mentioned in the Section “Materials and Methods”, due to technical problems the FST results in this

experiment were scored manually. Previous pilots conducted in our laboratory demonstrated no differences between manual and automated scores (data not shown). Moreover, the scores of the control-treated group in this experiment are comparable with the scores of the control-treated groups in Experiments 1 (see Figure 1B) and 2 (see Figure 2B) of this stage.

DISCUSSION

The need to develop more efficient and predictive methods of modeling BPD had been repeatedly emphasized. The present study supports the utility of Black Swiss mice as an appropriate strain to model a number of behavioral domains of the manic pole of BPD. Specifically, the results demonstrate that BS mice show a unique manic-like baseline behavioral pattern and a unique response to mood stabilizing drugs compared with other strains. BS mice show a number of ‘manic-like’ behaviors including increased reward-seeking, risk-taking, aggressive, vigor and goal directed behaviors, as well as intense response to amphetamine. Moreover, in BS mice, most of these behaviors are ameliorated by the prototypic mood stabilizers valproate and lithium but are not affected by the tricyclic antidepressant imipramine (Table 3), further supporting the relevance of the model to the human syndrome.

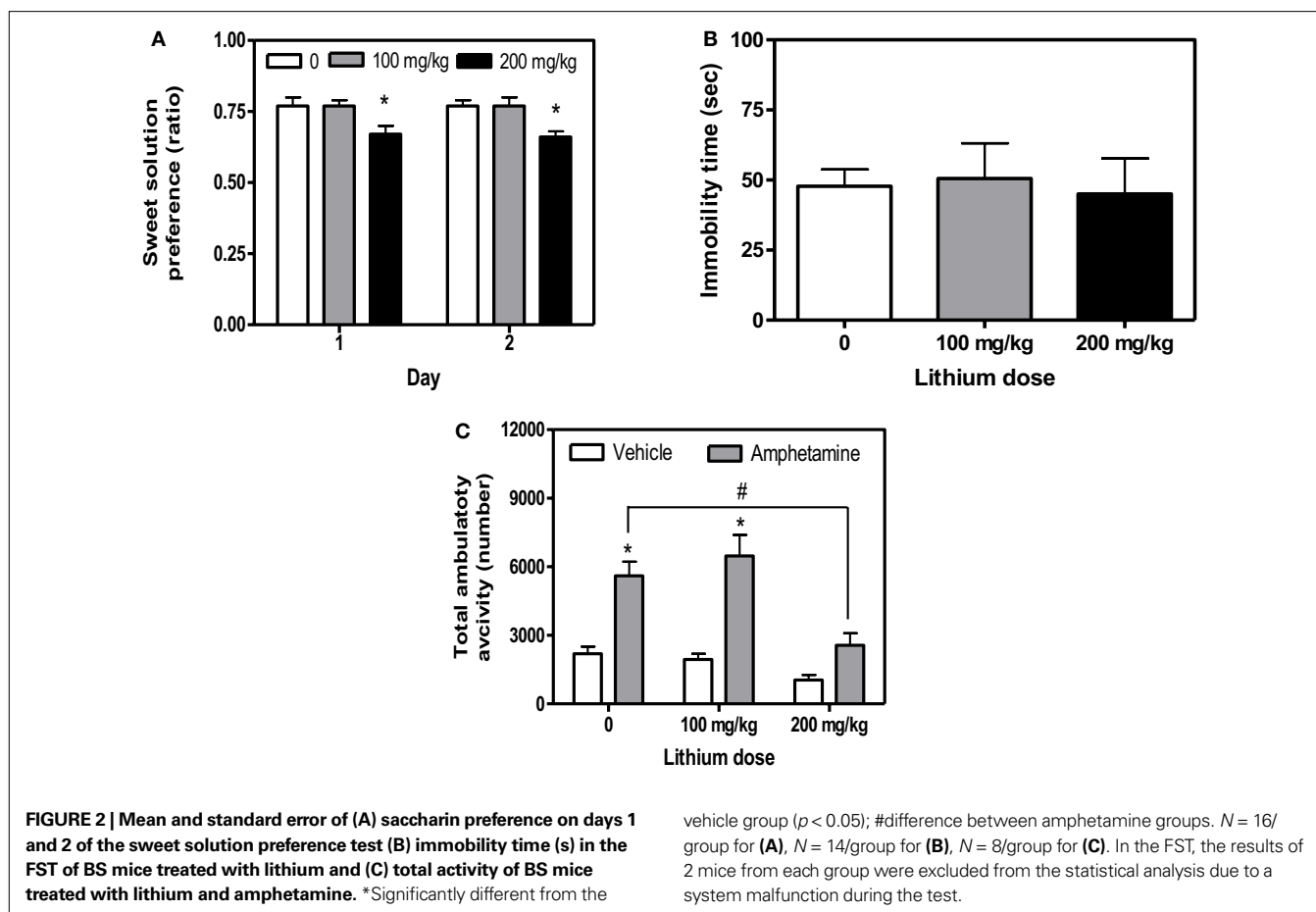
Identifying an appropriate model animal, species or strain for the purposes of modeling a specific disorder is an acknowledged and successful method of research. For instance, the Flinders Sensitive



rats line has been used as a model for depression (Overstreet, 1986) and the Alcohol Preferring P rats have been used as a model for alcoholism (Penn et al., 1978). Lewis rats have been used to study the biological mechanisms underlying obsessive compulsive disorder (Brimberg et al., 2007) and Fischer 344 and Lewis rats have been used to study susceptibility to psychoactive drugs and to stress (Suzuki et al., 1988; Cohen et al., 2006). Strain differences have been suggested as a method to define predictive responses to current medications (Porsolt et al., 1978; Gould et al., 2007) and to select appropriate animals for behavioral models of specific disorders and as background for the development of animal with targeted mutations (Crowley, 1999; Gould et al., 2007). In the field of BPD and mania modeling, recent studies identified a number of mice strains with targeted mutations that demonstrate a variety of ‘manic-like’ behaviors, including mice with targeted mutations of the Clock gene (Roybal et al., 2007), dopamine transporter (Ralph-Williams et al., 2003), and glutamate receptors (Shaltiel et al., 2008) (see Malkesman et al., 2009 for review). The development of model animals through targeted mutations has a number of advantages as they directly indicate possible mechanisms related to the underlying biology of the disorder (Malkesman et al., 2009). However, considering the conceptual and practical problems associated with models based on targeted mutations, there are also advantages in studying the neural basis of strain-specific behaviors (Gould et al.,

2007; Hiscock et al., 2007; Ashkenazy et al., 2008; Flaisher-Grinberg and Einat, 2009b; Flaisher-Grinberg et al., 2009) as well as species-specific behaviors of non-traditional model animals (Smale et al., 2005; Einat, 2007a). Comparison between different strains or different species can provide a strong tool to explore the underlying mechanism of behavior (Smale et al., 2005).

The results of Stage 1 of this study, demonstrating that BS mice show ‘manic-like’ behaviors compared with other strains, combined with the results of Stage 2 showing that these behaviors are attenuated by mood stabilizers but not by antidepressant treatment, provide face and pharmacological validity to the use of this strain in a specific battery of tests for mania. Moreover, the drug response profile demonstrated by BS mice in the current study is unique as it demonstrate ‘anti-manic’ effects in tests traditionally used to assess the effects of depressogenic manipulation and antidepressant drugs (see discussion below). The differences between the strains and the differences in drug effects can be utilized to (1) select an appropriate strain for modeling purposes (2) explore the relevance of these behavioral phenotypes to the underlying biology of the different strains and (3) screen for new and improved mood stabilizing/anti-manic drugs. It is important to note, however, that the findings demonstrating ‘manic-like’ behavioral phenotype in BS mice do not imply that these mice are “manic” as there is no evidence at this time that their behavior has identical causative



factors or general symptomology with the disorder. Moreover, these findings do not contradict the possibility that other mice strains might also demonstrate ‘manic-like’ behaviors in these and other relevant tests. Importantly, an investigation of the biochemical and molecular underpinning of the unique set of behaviors demonstrated by BS mice in comparison to other strains is currently performed in our laboratory. A specific emphasis is being placed on molecular pathways which were previously implicated in mania and/or BPD or its treatment [PKC (Einat et al., 2004, 2007), GSK-3 (Gould et al., 2006), BCL-2 (Lien et al., 2008; Chang et al., 2009) and BDNF (Frey et al., 2006a)].

In the current study it was also demonstrated that the profiles of the effects of lithium and valproate were not identical (see Table 3). Whereas this may seem to question the relevance of the results to the human syndrome, it is suggested that the varied effects in the model are not different than the variability of effects of mood stabilizers in patients (Cousins and Young, 2007). In fact, the differences between the specific effects of lithium and valproate across the test battery suggests a way to distinguish the effects of treatment on separate domains of the disorder (Einat, 2007a). It is however also possible that some of the differences in effects are related to drug levels in the brain. In the present study, drug doses were chosen based on their effects in animal models of affective disorders. For example, 100 and 200 mg/kg lithium increased the threshold for rewarding intracranial self stimulation (Mavrikaki et al., 2009), 50 mg/kg of

lithium reduced amphetamine hyperactivity (Borison et al., 1978). Administration of 200 mg/kg lithium reduced aggression in mice (Nakao et al., 1985) and lithium at 100 mg/kg and valproate at 200 mg/kg attenuated neuropeptide S-induced hyperactivity (Castro et al., 2009). Because specific drug levels in plasma or brain were not assessed in the present study the possibility that other doses and dosing regimens might have different effect profiles is left open. However, the similarities between drug doses in this and in previous studies support the relevance of the current findings. Moreover, it is also possible that even when plasma levels are monitored, the information is not enough to make clear extrapolations to the human conditions. First, the metabolism of mice is different than that of humans and therefore the kinetics of drugs is also different. For example, whereas the standard dose of imipramine for the treatment of depression is 1–4 mg/kg per day (e.g. Leucht et al., 2000), doses used in animal experimentation range between 10 and 30 mg/kg per day (e.g. Einat et al., 1999). Additionally, the presence of the drug in the plasma is not always a representation of its presence in the brain (e.g. Kovacsics and Gould, 2009).

Some of the drug effects in the present study might seem to be in contrast with prior reports. All three drugs were previously found to increase sweet solution preference [lithium (Orsetti et al., 2006) valproate (Monleon et al., 1995; Papp et al., 1996) imipramine (Papp et al., 1996)]. However, these ‘antidepressant-like’ effects were found in animals which were previously exposed

Table 2 | Behavioral effects of imipramine administration in tests for domains of mania.

Test measures collected	Imipramine dose	N	Mean ± SE	Statistical analysis		
Spontaneous activity (Total ambulatory activity)	Control	12	3806 ± 439	$F(3,44) = 0.32, p = 0.81$		
	10 mg/kg	12	3458 ± 532			
	20 mg/kg	12	3216 ± 446			
	30 mg/kg	11	3439 ± 266			
Sweet solution preference (ratio of saccharin consumption to total liquid consumption)	Control	12	Day 1: 0.61 ± 0.03 Day 2: 0.56 ± 0.02	Imipramine: $F(3,43) = 1.96, p = 0.14$; Day: $F(1,43) = 1.83, p = 0.18$; Interaction: $F(3,43) = 0.85, p = 0.48$		
	10 mg/kg	12	Day 1: 0.59 ± 0.03 Day 2: 0.59 ± 0.02			
	20 mg/kg	12	Day 1: 0.59 ± 0.03 Day 2: 0.52 ± 0.02			
	30 mg/kg	11	Day 1: 0.55 ± 0.03 Day 2: 0.54 ± 0.02			
	Forced swim test (immobility time – sec)	Control	11		60.4 ± 10.6	$F(3,42) = 1.03, p = 0.39$. 1 mouse from control and 1 from 30 mg group were excluded because of computer malfunction
		10 mg/kg	12		67.9 ± 12.4	
20 mg/kg		12	54.2 ± 10.9			
30 mg/kg		10	41.5 ± 9.3			
Light/dark box (Time in light compartment – sec)	Control	12	163 ± 2.7	$F(3,43) = 0.38, p = 0.77$. 1 mouse from the 20 mg group was excluded as a statistical outlier (2xSTD away from mean)		
	10 mg/kg	12	160 ± 5.0			
	20 mg/kg	11	154 ± 9.5			
	30 mg/kg	11	155 ± 5.9			
Resident-intruder test (Ratio of aggressive/total interactions)	Control	11	0.43 ± 0.07	$F(3,40) = 0.2, p = 0.89$. 1 mouse from control group, 1 from 10 mg group and 1 from 30 mg group were excluded due to computer malfunction		
	10 mg/kg	11	0.36 ± 0.09			
	20 mg/kg	12	0.37 ± 0.09			
	30 mg/kg	10	0.35 ± 0.08			
Amphetamine-induced hyperactivity (Total ambulatory activity)	Control		Vehicle	Amph	Imipramine: $F(3,38) = 1.85, p = 0.15$. Amphetamine: $F(1,38) = 51.99, p < 0.001$. Interaction: $F(3,38) = 1.96, p = 0.14$. 1 mouse from the control + vehicle group was excluded because of computer malfunction	
	10 mg/kg	5,6	2254 ± 254	5308 ± 1092		
	20 mg/kg	6,6	2658 ± 354	9251 ± 1822		
	30 mg/kg	6,6	1802 ± 363	9374 ± 1318		
	30 mg/kg	6,5	2294 ± 281	6517 ± 1435		

Table 3 | The domains of ‘manic-like’ behavior modeled in the battery, the expected response and actual response to mood stabilizers in BS mice.

Test	Domain of manic-like behavior	Expected response to mood stabilizers	Actual response to lithium	Actual response to valproate
Sweet solution preference	Reward seeking behavior	Reduction in preference	Reduction in preference	Reduction in preference
Forced swim test	Resistance to despair/ increased vigor	Increase in immobility	No effect	Increase in immobility
Resident-intruder test	Aggression	Reduced aggression	No effect	No effect
Dark/light box	Risk taking behavior	Reduced time in light Compartment	No effect	No effect
Amphetamine-induced hyperactivity	Manic-like increased activity or sensitivity to psychostimulants	Reduced hyperactivity	Reduced hyperactivity	Reduced hyperactivity (strong trend)

to depressogenic stimuli while in the present study BS mice were found to demonstrate excessive *baseline* sweet solution preference which was suggested to represent strain-specific, ‘manic-like’ high reward-seeking behavior (Flaisher-Grinberg et al., 2009). Therefore, the reduction in sweet solution preference induced by valproate

and lithium in the current study are in line with the hypothesis and are assumed to represent an anti-manic, rather than an anti-depressant effect (see Table 3). A similar idea can be demonstrated regarding drugs effect in the FST. In the present study, valproate was shown to increase immobility time in the FST whereas previous

studies found that all the drugs used in the present study decrease immobility in the test [lithium (Bersudsky et al., 2007; Gould et al., 2008), valproate (Bourin et al., 1996; Kulkarni and Dhir, 2007), imipramine (Porsolt et al., 1978)]. Importantly, these 'antidepressant-like' effects of both mood stabilizers and antidepressant drugs were demonstrated in studies in which rodent strains with high baseline immobility were used [C57Bl/6 and NIH-Swiss (Bourin et al., 1996), Swiss mice (Porsolt et al., 1978), CD, NMRI and OF-1 (Porsolt et al., 1978; Swiergiel et al., 2007)]. In the present study, BS mice were found to show low baseline immobility in the FST, a behavior which was suggested to represent 'manic-like' excessive vigor and goal directed activity (Roybal et al., 2007; Flaisher-Grinberg and Einat, 2009b). It is therefore reasonable to expect that in these mice the effects of mood stabilizers would be to act as anti-manic, i.e. increase immobility time [valproate (Flaisher-Grinberg and Einat, 2009b); current study; lithium, and a number of mutant mice models (Roybal et al., 2007; Engel et al., 2008)]. In contrast, while imipramine is known to reduce immobility time in many strains, it is suggested that its lack of effect in BS mice in the current study reflect a floor effect, due to the low immobility levels demonstrated by BS mice. Although it is possible that different imipramine doses would have produced an effect, this possibility seems less likely given that the doses used in the study were previously demonstrated to have effects in different mice strains.

In the context of the current results, it is important to note that the spontaneous locomotor activity of BS mice was not affected by any of the drugs and therefore it is reasonable to suggest that the effects of the mood stabilizers were not related to a generalized change in activity but are specific to the tests. This is in contrast to strains such as FVB/NJ mice or some mice strains with targeted mutations (such as the ERK1 heterozygote mice), which were found to show low immobility in the FST combined with generalized hyperactivity (Engel et al., 2008; Milner and Crabbe, 2008), suggesting a generalized, non specific behavioral change.

In the present study it was hypothesized that valproate and lithium would induce an anti-manic reduction in the risk-taking and aggressiveness demonstrated by BS mice. The lack of drug effect in the light/dark box and the resident-intruder test are in contrast with these hypothesis as well as with previous reports demonstrating anxiolytic effects [imipramine (de Angelis, 1992, 1995); valproate (Yoshimura and Ogawa, 1984; Ibarguen-Vargas et al., 2009)] and anti-aggressive effects [imipramine (Pinna et al., 2004); valproate (Einat, 2007b); lithium (Wei et al., 2007; Uchida et al., 2009; Veenema, 2009; Veenema and Neumann, 2009)] in these tests. The lack of effects in the resident-intruder test could be at least partially explained by the low baseline aggression found in Stage 2 of the present study, which could have created a floor effect masking possible drug effects. It is also possible that the previous exposure of mice to a variety of stressors across the test battery had a significant influence on their behavior, as stress was shown to alter aggressive behavior in mice (e.g., Uchida et al., 2009). In the context of the current study, saccharin deprivation following the establishment of saccharin preference (a possible consequence of the sweet solution preference test, conducted before the resident-intruder test) has been found to influence rodents' aggressive behaviors (Belozertseva et al., 2004). The lack of effect of imipramine in the light/dark box might be explained by the fact that in this test BS mice show no

preference for one compartment over the other (baseline behavior is at the range of 150–160 s out of 300-s session, showing lack of preference for either of the compartments and hence, no anxiety-like behavior). Since imipramine is expected to reduce anxiety (de Angelis, 1996), the lack of anxiety demonstrated by BS mice in this test might account for its lack of effect.

In the current study, several rodent tests have been proposed as models for some of the domains of mania and the effects of two mood stabilizers and one antidepressant drug in these tests were assessed. However, the spectrum of behaviors demonstrated in the present study is limited compared with the spectrum of mania and additional tests should be devised that might model further domains of the disorder such as reduced sleep, increased sexual drive and abuse of drugs (Einat, 2007a).

The testing of mice in a continuous set of tests demands a number of important considerations. First, although in the current study tests were arranged from less to more intrusive, it is possible that some tests are more susceptible to previous stress than others (see above for the resident-intruder and light/dark box). Another consideration deals with continuous drug administration. Given that tests are conducted sequentially, this factor introduces variability in the time course of drug treatment at the time of the different tests. Although the effects of acute treatment are not always comparable with the effects of sub-chronic or chronic treatments, this factor might be overcome by starting treatments a few days prior to the behavioral testing. Importantly, most mood stabilizers are expected to have more pronounced effects after chronic treatment compared with acute treatment (Kovacsics and Gould, 2009), whereas other drugs may have effects that tolerate across time. Finally, the inclusion of the amphetamine-induced hyperactivity test in the battery imposes the use of a smaller number of mice in each group in this test. Specifically, in the amphetamine-induced hyperactivity test, half the mice assigned to each treatment group in the other tests are treated with amphetamine and half are treated with vehicle. Although in the current study the number of animal per group in the amphetamine-induced hyperactivity tests ranges from 5 to 8, the findings that lithium and valproate attenuate amphetamine-induced hyperactivity, while imipramine has no effect, are in line with previous reports in rodents [e.g., lithium: (Frey et al., 2006b; Gould et al., 2007; Kim et al., 2008) imipramine: (Dziedzicka-Wasylewska et al., 1997)]. Future experiments will validate these results using a higher number of mice per group.

A different consideration involves timing of testing within the light/dark cycle of the animals. It may seem appropriate to test nocturnal animals during their dark (and not light) phase in a way that will have higher homology with the active phase of diurnal humans. However, the dark phase of nocturnal animals is not the exact reversal of the light phase in diurnal animals as the biology of circadian rhythms is more complicated. For example, although the main circadian hormone melatonin is secreted during the dark in both diurnal and nocturnal mammals, nocturnal mammals are active when melatonin levels are high, while diurnal mammals are active when melatonin levels are low (Challet, 2007). As such, it is possible that regardless of the time of testing, a full homology of the circadian phase between nocturnal mice and diurnal humans is not attainable. Given that the current study did not evaluate the possible effects of light/dark phase on the demonstrated behavioral

profile, additional studies are now designed to explore the behavioral responses of BS mice in the same battery of tests conducted during the dark phase. Moreover, in an additional project, attempts are currently made to explore the possible utilization of diurnal rodents to model affective disorders (Einat and Kronfeld-Schor, 2009).

To summarize, the current study identified BS mice as a potentially preferred strain for modeling a number of the domains of mania. The unique behavioral phenotype demonstrated by BS mice might also be useful in future attempts to elucidate the molecular, cellular, and genetic factors differentiating BS from the other strains.

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