Anxiety Is Correlated With Running in Adolescent Female Mice Undergoing Activity-Based Anorexia

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Activity-based anorexia (ABA) is a widely used animal model for identifying the biological basis of excessive exercise and starvation, 2 hallmarks of anorexia nervosa (AN). Anxiety is correlated with exercise in AN. Yet the anxiety level of animals in ABA has not been reported. We asked: Does food restriction as part of ABA induction change the anxiety level of animals? If so, is the degree of anxiety correlated with degree of hyperactivity? We used the open field test before food restriction and the elevated plus maze test (EPM) during food restriction to quantify anxiety among singly housed adolescent female mice and determined whether food restriction alone or combined with exercise (i.e., ABA induction) abates or increases anxiety. We show that food restriction, with or without exercise, reduced anxiety significantly, as measured by the proportion of entries into the open arms of EPM (35.73%, p = .04). Moreover, ABA-induced individuals varied in their open arm time measure of anxiety and this value was highly and negatively correlated to the individuals’ food restriction-evoked wheel activity during the 24 hr following the anxiety test (R = -.75, p = .004, N = 12). This correlation was absent among the exercise-only controls. In addition, mice with higher increase in anxiety ran more following food restriction. Our data suggest that food restriction-evoked wheel running hyperactivity can be used as a reliable and continuous measure of anxiety in ABA. The parallel relationship between anxiety level and activity in AN and ABA-induced female mice strengthens the animal model.

Keywords: activity-based anorexia, anxiety, anorexia nervosa, food restriction, exercise

Supplemental materials: http://dx.doi.org/10.1037/bne0000040.supp

Anorexia nervosa (AN) is a psychiatric disorder with a mortality rate of 10%–15% (Birmingham, Su, Hlynsky, Goldner, & Gao, 2005; Bulik, Slos-Op’t Landt, van Furth, & Sullivan, 2007; Sullivan, 1995). There is no accepted pharmacological treatment for this disorder, and the underlying pathology in the illness is not well understood. This necessitates the development and use of animal model. The latter may yield insight into the emergence of disease, a preclinical phase that is difficult to study in the human population. An animal model also makes it possible to conduct post mortem analysis of the brain to study neurochemical changes underlying the biobehaviors of disease.

Currently, the most widely accepted animal model for AN is activity-based anorexia (ABA). In this model, access to a running wheel is combined with a restricted feeding schedule. Several species of rodents, including rats and mice, respond to this environment by increasing their voluntary wheel activity several-fold (Chowdhury, Wable, Sabaiauskas, & Aoki, 2013; Epling, Pierce, & Stefan, 1983; Gutierrez, 2013; Hall & Hanford, 1954; Klenotich & Dulawa, 2012; Routtenberg & Kuznesof, 1967). This combination of food restriction with greatly increased wheel activity leads to exaggerated weight loss and eventual death, unless wheel access and food restriction are removed from the environment. It is important that the ABA model captures two hallmarks of AN—excessive voluntary exercise as well as starvation. Because many of the ABA animals run on the wheel during the limited hours of food access, the food restriction imposed by the experimenter turns into self-starvation.

AN is commonly comorbid with an anxiety disorder (Kaye, Bulik, Thornton, Barbarich, & Masters, 2004). A high tendency to control anxiety is found in AN, and individuals suffering from AN may get a sense of control over emotional distress through dieting and exercise (Fiore, Ruggiero, & Sassaroli, 2014). Anxiety control may also mediate the drive for thinness, leading to overexercise (Fiore et al., 2014). Indeed, no less than 40% and as many as 80% of the individuals with AN exhibit excessive exercise (Davis, Katzman, & Kirsh, 1999; Hebebrand et al., 2003), and this often...
precedes the formal diagnosis. The levels of anxiety and severity of exercise are correlated in patients diagnosed with AN (Holtkamp, Hebebrand, & Herpertz-Dahlmann, 2004; Peñas-Lledó, Vaz Leal, & Waller, 2002; Shroff et al., 2006). It is possible that anxiety, either preceding or secondary to food restriction, leads to hyperactivity.

An animal model of the disease might be used to investigate the important role that anxiety may play in AN. Yet the anxiety level and its relationship with wheel running in ABA has never been reported. Outside of the ABA model, the relationship between wheel running and anxiety is fairly complicated, because it is affected by age, sex, and social conditions of the rodents (Scioliño & Holmes, 2012). Wheel running was found to be anxiolytic in singly housed female wild-type mice, which is the group most comparable to our experimental animals (Pietropaolo et al., 2008), although to our knowledge, we are the first to examine the effect of a short duration of wheel access. In this study, our goal was to examine the relationship between the severity of response to ABA and anxiety. We sought to achieve this by looking at the measures of each—wheel activity in ABA and behavioral tests for anxiety. Our first aim was to determine whether food restriction alone or wheel access alone evoke a measurable change in anxiety and whether food restriction combined with access to a running wheel abates or increases anxiety measures. This question was answered by using behavior tests to quantify anxiety of ABA animals just before and during food restriction and comparing these values to the anxiety measures of age-matched animals that were only food-restricted, only given access to a running wheel, or exposed to neither treatments. Specifically, we employed the widely accepted test, namely the elevated plus maze test (EPM) to measure anxiety levels during food restriction of the ABA paradigm and compared this with the open field (OF) test conducted before food restriction to determine whether individual differences in anxiety existed prior to food restriction and were altered by it.

We have previously shown that running activity of individual animals correlates negatively with their level of α4-containing GABA receptors in the hippocampus (Aoki et al., 2014) and to the extent of GABAergic innervation of the hippocampal pyramidal cells (Chowdhury et al., 2013). Thus, individual differences in the ABA model are a meaningful way of exploring the underlying pathology. In this study, our second aim was to determine whether individual differences in the severity of response to ABA could be explained by individual differences in anxiety measured before food restriction and/or in the state anxiety measured during food restriction.

Materials and Methods

Animals

C57BL6 mice were purchased as breeders from Charles River Laboratories (Wilmington, MA) at age P35. Following a week of acclimation, they were paired. The breeders were never assigned to any experimental condition. Only their litters were used for the experiments. Litters from these breeders were weaned at P25, and animals were group-housed with same-sex littermates in a 12-hr light/dark cycle (lights on at 0700 h). Female mice were used in this study because of the high prevalence of AN in females (Fairburn & Harrison, 2003). On P37, female mice were assigned to one of four experimental groups (Supplemental Table 1, available online as supplemental material). Several litters were divided into two of the four experimental groups, and each group had animals from multiple litters to ensure that the group differences were not confounded by differences in litters. All procedures relating to the use of animals were in accordance with the Institutional Animal Care and Use Committee of New York University (A3317-01).

ABA Induction and Behavioral Controls

Control (CON) animals were housed in standard home cages with ad libitum access to food (dry chow (PMI Mouse Diet 5001 from LabDiet, St. Louis, MO; 336 kcal per 100 g, 28.507% protein, 57.986% carbohydrates, 13.496% fat) and wet gel (Diet-Gel from Clear H2O, Portland, ME; 76A; 99.8 kcal per 100 g, 4.7% protein, 17.9% carbohydrates, 1.5% fat, 73.4% moisture)) and no running wheel access for the duration of the study. At noon on P37 (start of Day 1 of experiment), animals in the ABA and EX (exercise control) group were placed in standard home cages with running wheels attached (low-profile mouse wheel, Med Associates, Inc., St. Albans, VT) and ad libitum access to food to record baseline wheel activity. Starting on P40 at noon (start of Day 4 of experiment) until noon on P43 (end of Day 6), animals in the ABA and FR (food restriction control) group were given unlimited access to food for the first 2 hr of the dark cycle; food was not available for the remaining 22 hr per day (Figure 1A). EX animals continued to have unrestricted access to food. Body weight, food intake, and wheel running activity (where applicable) were measured daily within 20 min prior to the start of the dark cycle.

Loss of wheel data: Because of technical failure, continuous wheel data were not collected for 4 animals of the ABA group from 8 a.m. to 9 p.m. on P42. This interval comprised of the final 4 hr of FR2 (8 a.m. to noon) and the initial 9 hr of FR3 (noon to 9 p.m.). Wherever we report data from FR2 and FR3, wheel counts for these periods of loss were omitted for all animals.

Behavioral Testing

OF. On P39 before noon (end of Day 2), corresponding to the end of 2 days of single housing with ad lib food access, mice were tested for 10 min in the OF test in a room separate from the one in which they were housed. The field dimensions were 70 × 70 × 30 cm. The tests were conducted between 1000 and 1300 h, following habituation (30 min in the homecage) to the light and noise level of the room where the open field apparatus was located. The mice were tracked by infrared beam breakages recorded by MDBActivity Monitor. Both the apparatus and the software were manufactured by Med Associates. Each animal was placed in the center of the open field at the beginning of the test. The open field was cleaned with 30% ethanol between animals. The raw data were imported into Matlab (Version 2010b, MathWorks, Natick, MA) and analyzed for the position of the mouse once every second. The field was divided into 16 equal squares, and the central 4 squares were considered to be the center of the field.

EPM. EPM tests were conducted on P41 (end of Day 4/FR1), 48 hr after recovery from the open field, and 21 hr after onset of food restriction as applicable to ABA and FR. Habituation to the testing room was provided for 30 min in the homecage. The maze,
Figure 1. The schedule of activity-based anorexia (ABA) and experimental controls and their wheel activity. (A) Control (CON) animals ($N = 9$) were housed singly in standard home cages with ad libitum access to food (and no running wheel access) for the duration of the study. At noon on Postnatal Day (P) 37 (beginning of Day 1 of experiment), animals in the ABA ($N = 12$) and exercise control (EX; $N = 7$) group were placed in standard home cages with running wheels attached and ad libitum access to food to record baseline running activity. Starting at noon on P40 (beginning of Day 4) until noon on P43 (end of Day 6), animals in the ABA and food restriction control (FR; $N = 8$) group were given unlimited access to food for the first 2 hr of the dark cycle; food was not available for the remaining 22 hr per day. For the ABA and FR animals, Days 4, 5, and 6 of the experiment are referred to as FR1 (food restriction Day 1), FR2 (food restriction Day 2), and FR3 (food restriction Day 3), respectively. EX animals continued to have unrestricted access to food. Body weight, food intake, and wheel running activity (where applicable) were measured daily within 20 min prior to the start of the dark cycle. Open field (OF) and elevated plus maze (EPM) testing was conducted on days 2 and 4 of the experiment respectively. (B) Averaged wheel activity of ABA animals and age-matched controls. Food restriction greatly increased wheel running activity. The white bars represent average wheel activity measured during the food restriction phase for ABA animals and corresponding days for EX animals (fed ad lib during the phase; Days 4–6). **** indicates $p < .001$, ns indicates no difference, comparing the black bars with respective white bars of the same group. (C) Wheel activity of ABA and EX. Daily activity of 12 ABA (on the left) and seven EX (on the right) animals was measured as the distance run on the wheel. The dotted line parallel to the x-axis indicates the days of food restriction for the ABA group. During food restriction, the ABA animals increased their running much more than the EX animals which were fed ad lib.
custom made with white polypropylene, was placed on a platform in a room illuminated with dim ambient lighting. Each of the four arms was 30 cm long and 5 cm wide on the inside. The walls of the two closed arms were 15 cm high and 1 cm thick. The floor of the maze was elevated 39 cm above the platform. Each animal was placed in the center of the elevated plus maze facing an open arm. A video camera (Panasonic model no. WV-BP334, Panasonic Corporation of North America, Newark, NJ) was positioned about 433 cm above the mouse to record behavior during the 10-min test. At the end of the test, the animal was weighed and returned to its cage. The maze was cleaned with 30% ethanol between animals.

The videos were recorded using EthoVision software version 4.1.106 (Noldus Information Technology, Wageningen, The Netherlands). Subsequently, the videos were scored by an observer, blind to the experimental condition of the animal, to note the location of the animal at every second. The locations possible were as follows: open arm, closed arm, and center. The mouse was classified to be present in a particular arm when all four paws had crossed into that arm.

**Methodological Considerations**

The EPM was scheduled at the end of 21 hr of food restriction to study the emergence of anxiety in the mice. The timing allowed us to examine whether the change in anxiety preceded the maximal rise in wheel activity, which for some mice exposed to ABA is on the second day of food restriction (Chowdhury et al., 2013). Some but not all animals decrease their activity on the third day of food restriction (unpublished data and this study). Why this may occur only in some animals is unknown. The timing of the EPM ensured that none of the mice would be inactive, a state that would render the EPM measure uninterpretable.

There is considerable evidence that repeating the OF or EPM raises issues with interpretation because of habituation to the environment of the test (Almeida, Garcia, & de Oliveira, 1993; Bond & Giusto, 1977; Bronstein, 1972, 1973; Bronstein, Wolkoff, & Levine, 1975; File, 1993; File, Mabbutt, & Hitchcott, 1990; Kiemiesky, Sick, & Kruppenbacher, 1977; Lee & Rodgers, 1990; Rodgers, Lee, & Shepherd, 1992; Rodgers & Shepherd, 1993; Russell & Williams, 1973; Treit, Menard, & Royan, 1993; Valle, 1971). Therefore, we have chosen two different tests in the baseline and food restriction periods in the experiment.

**Statistical Analysis**

Normality of the distribution of measures was tested using the D’Agostino & Pearson omnibus normality test and Shapiro-Wilk normality test. Two-way analysis of variance (ANOVA) was used to evaluate differences between the four groups, using wheel access and food access as the two factors. When two-way ANOVA was performed to evaluate the effect of experimental days on wheel activity, the day of experiment was used a repeated measure. Pearson’s correlation was computed between normally distributed measures of anxiety and ABA. Spearman correlation was used if the measures were not distributed normally. Statistical software used was GraphPad Prism Version 6.

Exclusion criteria: ABA and EX groups each had one animal that did not use the running wheel (<20 wheel revolutions on 3 consecutive days during the experiment). These two animals were excluded from analysis.

**Results**

**Wheel Running Reveals Group Differences and Individual Variability**

Previously, we had shown that food restriction evokes a robust (66%) increase of voluntary wheel running of adolescent female mice (Chowdhury et al., 2013). However, it remained untested whether animals without food restriction would also increase wheel running, simply from exposure to the wheel over a period of 8 days. To determine whether the increased wheel running was evoked specifically by food restriction, we analyzed the wheel running pattern of ABA (food restricted) and EX (fed ad libitum) mice, matched for age and genetic background.

Baseline running was computed as average running wheel activity on Days 2 and 3 of the experiment. The first day of wheel activity was not included in the analysis because the mice were getting acclimated to the novelty of the wheel and being singly housed. Wheel activity during the food restriction period for the ABA animals was computed as the average daily running wheel activity on Days 4, 5, and 6 of the experiment. As was shown previously, ABA animals increased their running after the onset of food restriction (mean increase = 7.9 ± 1.3 km per 24 hr, N = 12). This increase was significantly higher, t(17) = 3.38, p = .003, than the increase shown by EX animals (1.6 ± 0.8, N = 7) over the same number of days but in the absence of food restriction (mean difference between the groups = 6.3 ± 1.8; Figure 1B). ABA animals had a higher mean baseline running than EX animals, but this difference across the treatment groups was far less than the difference that emerged after food restriction for the ABA group, relative to the age-matched EX group (Figure 1B). We will refer to the increased running following food restriction as hyperactivity. The hyperactivity of ABA animals changed over the experimental days (Figure 1C), suggesting an evolving impact of the food restriction. This prompted us to look at the evolution of the relationship between running activity of individual animals on different days with anxiety measures.

**EPM Open Arm Entries Are Increased by Food Restriction but Not by Wheel Access**

The number of entries into the open arms is a measure influenced by anxiety (Walf & Frye, 2007). Entries into the open arm is also influenced by total activity levels, as is indicated by the positive correlation of total entries with entries into the open arms (R = .7 and .82; p = .010 and .003 for ABA and CON groups, respectively). To normalize for individual differences in overall activity level, the number of entries into the open arms was divided by the total entries (Lister, 1987). This ratio was 35.73% higher for the FR and ABA groups (Mfood restricted = 0.118 ± 0.01) compared with the EX and CON groups (Mfood lib = 0.087 ± 0.008), main effect of food in a two-way ANOVA, F(1, 32) = 4.578, p = .040; no effect of the wheel, F(1, 32) = 0.24, p = .62, no interaction F(1, 32) = 0.15, p = .7 (Figure 2A). This effect of food restriction on open arm entries is consistent with the demonstrated anxiolytic effect of mild-to-moderate calorie restriction (Riddle et al., 2013; Yamamoto et al., 2009).

The time spent in the open arms is widely used to measure anxiety in mice, with greater time spent in the open arm being
Considered as less anxious behavior (Carola, D’Olimpio, Brunamonti, Mangia, & Renzi, 2002; Walf & Frye, 2007). We evaluated whether access to the wheel (i.e., ABA and EX groups of mice) or food restriction (i.e., the ABA and FR groups of mice) for a day (as EPM was conducted at the end of FR1) increased or decreased anxiety of the animals as a group. Two-way ANOVA revealed no significant differences among the four groups in the time spent in the open arms (Figure 2B). There was no main effect of wheel access, $F(1, 32) = .29$, $p = .59$, or interaction, $F(1, 32) = .01$, $p = .93$. There was, however, a trend to an effect of food access, with the food restricted groups (i.e., with wheel = ABA; without wheel = FR, $M_{FR \text{ food restricted}} = 86.05 \pm 12.49$ s) spending more time in the open arms than the ad lib fed groups (i.e., with wheel = EX; without wheel = CON, $M_{ad \text{ lib}} = 57.13 \pm 7.96$ s), $F(1, 32) = 2.82$, $p = .10$.

EPM Open Arm Time and Proportion of Entries Into Open Arms Are Correlated With Wheel Running Following Food Restriction in ABA Animals

Two-way ANOVA did not reveal an effect of 4 days of wheel activity upon anxiety measured by time spent in the open arms or entries made into the open arms as a proportion of total entries. However, because individual animals exhibited great variability in wheel activity, we surmised that individuals’ differences in wheel activity might be related to individuals’ state anxiety. Indeed, within the ABA group, there was a highly significant negative correlation between the time spent on the open arm and the average wheel activity during the 3 postfood restriction days ($R = -.79$, $p = .002$, $N = 12$; data not shown). Further analysis of the individual days of food restriction revealed that there was a negative correlation between the time spent on the open arm and the daily running activity on FR3 (i.e., day following the EPM; Table 1, Figure 3A, left) but not FR1.

In the EX group, there was no correlation between the time spent on the open arm, and the average wheel activity during the Days 4, 5, and 6, corresponding to ABA group’s FR1 through FR3 ($R = .31$, $p = .49$; data not shown). Analysis of wheel running during Days 4 and 5, separately, which correspond to ABA group’s FR1 and 2, also did not reveal correlation to the open arm time in the EPM (Table 1, Figure 3A, right). The contrasting results between the ABA and EX groups suggest that wheel running per se is not a manifestation of anxiety, but becomes so in the presence of food restriction.

The fraction of entries made into the open arm by ABA animals correlated negatively with the wheel activity on the first two days of food restriction FR1 and FR2 (Table 1, Figure 3B, left). These findings are consistent with the idea that wheel activity is an expression of the animal’s anxiety level that emerges by FR1.

This was not so in the EX group. The lack of correlation between the fraction of entries into the open arm (Table 1, Figure 3B, right) and the daily running activity on Days 4 and 5 supports the idea expressed above that wheel running in the absence of food restriction is not a manifestation of anxiety.

The Severity of Response to ABA as Measured by Wheel Activity Is Correlated With the Change in Anxiety Level Produced by Food Restriction

The anxiolytic effect of food restriction, as indicated by EPM and two-way ANOVA (Figure 2), would predict that if wheel activity in ABA is positively correlated with anxiety level, then a reduction in anxiety level, such as by food restriction, would lead to a decrease in wheel activity by the ABA group. This contradicts our observation. Instead, we observed that wheel running by the ABA group increased after a supposedly anxiolytic treatment of food restriction (Figure 1B) and the extent of running correlated tightly and negatively with anxiety level as measured in the EPM (Figure 3B). This contradiction was resolved by observing that the ABA as well as the FR mice underwent differential changes in anxiety level evoked by food restriction, while the EX and CON mice did not undergo any change in anxiety level (Figure 4A). We measured the change in anxiety of individual animals by using the time spent in the center of the OF (Carola et al., 2002) to measure their level of anxiety prior to food...
restriction, relative to EPM anxiety measurements after food restriction (Figure 4A). Carola et al. have shown a correlation between the measurements of anxiety in OF and EPM. We also observed that the time spent in the center of the OF was correlated positively to the EPM open arm time for the EX and CON animals ($R/H_11005^{.94, .73, p/H_11005^{.001, .014}}$). These correlations indicate that both OF and EPM were reliable, stable tests for measuring anxiety of the EX and CON groups, whereas the anxiety measures changed much for the FR and ABA groups, as a result of food restriction. Correspondingly, the ABA and FR groups showed a lack of correlation between the EPM open arm time and time in center of the OF ($R/H_11005^{.04, .585, p/H_11005^{.899, .128}}$, respectively).

Within the ABA group, the medians of the two measures, OF center time and EPM open arm time, were quite different at 39 s and 73.5 s, respectively. To make a meaningful comparison, we computed a modified $z$-score for the anxiety level measured in the OF as follows: modified $z$-score = (time spent in center of minus its median value)/interquartile range. Similarly, we calculated the modified $z$-score for the anxiety level measured by the EPM test as follows: modified $z$-score = (time spent in the open arms minus its median value)/ interquartile range. We chose to use the modified $z$-score instead of the standard $z$-score, because the OF center time measure of the ABA group was not normally distributed ($D^{/}D_{/}A^{/}g_{/}o_{/}s_{/}t_{/}i_{/}n_{/}o_{/}s_{/}i_{/}u_{/}b_{/}u_{/}s_{/}t_{/}e_{/}t_{/}e_{/}n_{/}t_{/}a_{/}$. $K_2 = 13.88, p = .001$, Shapiro-Wilk normality Test $W = 0.81, p = .013$).

Using these modified $z$-scores, the ABA mice could be categorized as belonging to one of the following three subtypes:

1. Those for whom food restriction was anxiolytic, identified as exhibiting higher modified $z$-score for the EPM than the OF measure.
2. Those for whom food restriction was anxiogenic, identified as exhibiting lower modified $z$-score for the EPM than the OF measure.
3. Those that underwent no change in anxiety level, identified as exhibiting very similar or the same modified $z$-score for OF and EPM.

We considered whether the severity of response to ABA, as measured by the increase in running, following the onset of food restriction relative to the baseline running, was related to the change in anxiety level, computed as a difference between anxiety measurement by EPM after and by OF before food restriction. Indeed, changes in wheel running (and thus of severity of response to ABA) correlated negatively and strongly with changes in anxiety ($R/H_11002^{.612, p/H_11005^{.034}}$; Figure 4B, left). The change in modified $z$-scores for the EX group were not correlated to the change in wheel activity ($R = 0, p = .97$, Figure 4B, right).

**Wheel Activity Is Not Correlated to the Total Entries in the EPM**

The total entries in the EPM are a measure of general activity (Lister, 1987; Walf & Frye, 2007). To examine whether the difference in wheel activity between the ABA and EX groups was a generalized
change in activity or specific to wheel running, we compared the total entries between groups. There was a significant effect of wheel in a two-way ANOVA on total entries, $F(1, 32) = 20, p < .001, M_{wheel} = 58.79 \pm 3.78$ is less than $M_{nonwheel} = 82.24 \pm 3.97$, but no effect of food access, $F(1, 32) = 2.38, p = .13$, or interaction, $F(1, 32) = 3.8, p = .06$ (data not shown). However, the mean values of total entries for the ABA ($M = 58 \pm 4.81$) and the EX ($M = 60.14 \pm 6.56$) groups were not different from one another (Tukey’s post hoc test, $p = .99$). From this post hoc test, we conclude that the difference in activity induced by food restriction is only in the domain of wheel running (Figure 1B), and not generalized to all types of activity, when the wheel access is available. Furthermore, the wheel activity is uncorrelated with the total entries in the EPM (Table 1), indicating that wheel activity is a specific type of activity and is unlike general walking or exploratory activity.

The FR group ($M = 92 \pm 4.3$) exhibited greater number of total entries than the ABA group ($q = 6.811, p < .001$). This could be because food restriction rendered the FR animals hyperactive.

### Figure 4.

The overall effect of food restriction on anxiety levels is anxiolytic, but on individual animals, it is qualitatively and quantitatively different. Activity-based anorexia (ABA) and food restricted (FR) in (A) demonstrate the different changes in anxiety level of individual animals. The change induced in the anxiety level in ABA mice is correlated with the change induced in the wheel activity from baseline levels by food restriction (B). (A) The $x$-axis shows the time spent in the center of the open field (OF) prior to food restriction, and the $y$-axis shows the time spent in the open arms of the elevated plus maze (EPM) after 21 hr of food restriction. The ABA and FR mice do not show any correlation between these two measures, whereas exercise control (EX) and control (CON) mice show a positive correlation. (B) The $x$-axis shows the change in the modified z-scores (based on the median) of the EPM open arm time relative to the OF center time for the ABA (on the left) and EX animals (on the right). Positive values indicate an increase in the z-score, hence a relative increase in nonanxious behavior in the EPM than in the OF, and hence an anxiolytic effect of food restriction. Negative values correspondingly indicate an anxiogenic effect. The $y$-axis is the increase in the average daily wheel activity during the food restriction phase (post food restriction) relative to baseline (pre food restriction). The change in the modified z-scores of anxiety level is negatively correlated with the change in activity from the pre- to the postfood restriction phase in the ABA animals, but not in the EX animals.
(Gelegen et al., 2007; Klenotich & Dulawa, 2012) but without a wheel as an outlet for this hyperactivity.

**Time Spent in the Center of the OF Before Food Restriction Does Not Correlate to the Wheel Activity Preceding or Following Food Restriction**

To ensure that there were no preexisting differences in anxiety among the experimental groups, we tested ABA, FR, EX, and CON animals in the OF before food restriction was imposed upon the FR and ABA groups, but after acclimation of the EX and ABA groups to the wheel. Comparison of the time spent in the center of the OF revealed no group differences, two-way ANOVA, \( F(1, 32) = 0.72, \ p = .40 \), for main factor food; \( F(1, 32) = 0.03, \ p = .84 \), for main factor wheel; \( F(1, 32) = .35, \ p = .55 \), for interaction (data not shown). The time in center of the OF was not correlated with the average wheel activity in baseline (\( R = .08, \ p = .74 \)) or daily wheel activity on experimental Days 2 or 3 (Table 1) for the ABA and EX groups combined. The ABA and EX groups were combined for these correlational analyses because they were identical during this period preceding the onset of food restriction.

To test whether preexisting individual differences in the level of anxiety could predict the severity of response to ABA in terms of wheel activity, we examined the relationship between the prefood restriction OF behavior and wheel running hyperactivity following food restriction. There was no correlation between the time spent in the center of the OF and measures of wheel activity on FR1 or FR2 of the ABA group or on Days 4 or 5 of the EX group (Table 1). It is thus evident that the response to ABA is not predicted by the anxiety level measured in the absence of food restriction.

**Weight Lost by the Beginning of Third Day of Food Restriction Is Predicted by the Total Entries and Entries Into the Open Arms in the EPM**

Weight loss is a cardinal feature of ABA and AN (American Psychiatric Association, 2013; Epling & Pierce, 1996). Because we identified a strong negative correlation between wheel activity of ABA animals during FR2 and anxiety measured by the EPM (Figure 3), we asked whether the weight loss on FR3, resulting from wheel running during FR2, might be related to anxiety.

By FR3, ABA animals weighed 76.25% (\( SEM = 0.62, \ N = 12 \)) of the baseline, as measured the day before food restriction started. FR animals weighed 76.84% (\( SEM = 1.4, \ N = 8 \)). Weighed over the same period, EX animals were at 101.5% of baseline (\( SEM = 0.89, \ N = 7 \)) while CON animals were at 101.1% (\( SEM = 0.59, \ N = 9 \)). Two-way ANOVA revealed a strong main effect of food restriction, \( F(1, 31) = 711.7, \ p < .001 \) and no additional effect of wheel access, \( F(1, 31) = 0.01, \ p = .91 \) or interaction, \( F(1, 31) = .28, \ p = .6 \). However, for the ABA group, the weight on the last day (end of FR3) as a proportion of weight before food restriction correlated positively with the entries into the open arms (\( R = .58, \ p = .046, \ N = 12 \)). The higher the number of entries made by the animal, the higher was its proportional weight and hence, the lower its weight loss. These correlations were absent in the EX and CON groups but also notably absent in the FR group (\( R = .43, .07, .49, \ p = .32, .85, .38 \), respectively). There was also a positive correlation between weight at the end of FR3 and the total number of entries (\( R = .64, \ p = .022, \ N = 12 \)) uniquely for the ABA group, and not for the EX, FR, or CON (\( R = .43, .41, .66, \ p = .32, .48, .67 \), respectively) groups. These data indicate that the weight loss in the ABA group is related to anxiety (total and open arm entries), but that relation does not exist for the FR group. Thus, the nature of weight loss in the two groups is differentiated based on access to the running wheel.

**Distance Traveled in the EPM Is Not Affected by Food Restriction**

The distance traveled in the EPM is an indicator of exploration by the animal (Lister, 1987). There were no differences among the four groups in the distance traveled in the EPM, two-way ANOVA, \( F(1, 32) = 1.3, \ p = .25 \), for main factor food; \( F(1, 32) = 2.15, \ p = .15 \), for main factor wheel; \( F(1, 32) = 3.03, \ p = .091 \), for interaction. This supports the idea that change in overall exploratory drive does not underlie the effects on anxiety-like behavior.

**Wheel Activity Preceding Food Restriction Does Not Correlate to Anxiety Level**

Wheel activity in the baseline period indicates the preference of the animal for the running wheel, but might the individual differences during this baseline period reflect preexisting differences in anxiety? To answer this question, we measured the correlation of wheel running during the baseline period to anxiety. There was no correlation between the baseline running and the time spent in the center of the OF for the ABA and EX group combined (\( R = -.05, \ p = .80 \); data not shown), indicating that the preference for wheel running was unrelated to anxiety level.

**Discussion**

Animal models are crucial to understanding the underlying circuitry of the disease. Establishing the strength of a model in reproducing the disease symptomatology and biochemistry is essential. The ABA rodent model of AN has been used since the 1960s. It is well established that ABA mimics the core phenotype of AN, including severe starvation, rapid weight loss, voluntary overexercise, and loss of estrous cycle function (Aoki et al., 2012; Golden & Shenker, 1992; Gutierrez, 2013; Hall & Hanford, 1954; Klenotich & Dulawa, 2012). Metabolic and neurobiological similarities include hypothermia and hypothalamic hypothyroidism (Bannai et al., 1988; Gelegen et al., 2008; Golden & Shenker, 1994; Hebebrand et al., 2003; Mantzoros, Flier, Lesem, Brewerton, & Jamerson, 1997; Nakazato, Hashimoto, Shimizu, Niitsu, & Iyo, 2012). However, it has not been examined whether ABA can reproduce the correlation between levels of anxiety and exercise in AN. This study set out to determine whether the ABA model can be used to understand the shared etiology and brain circuitry that underlies the comorbidity of AN with anxiety and the relationship between anxiety and exercise in AN.

We examined whether food restriction alone or wheel access alone evokes a measurable change in anxiety and whether food restriction in combination with wheel access changed anxiety level in the ABA model. The change in anxiety was measured by conducting behavior tests just before and during food restriction. We also sought to determine whether individual differences in...
severely of response to ABA were related to individual differences in anxiety measured before food restriction and/or in the anxiety measured during food restriction.

The Open Field Measurement Provided a Stable Baseline Against Which to Measure the Change Induced by Food Restriction

The OF measurement was performed at the end of 2 days of social isolation (for all groups) and wheel access (for ABA and EX animals). The nonfood restricted groups, CON and EX, showed a correlation between the time spent in the center of the OF and time spent in the open arms of the EPM, which are the measures indicating anxiety-related behavior. The correlation suggested that even if social isolation, wheel access, and other unidentified factors affected the anxiety level of the mice, these changes were already captured by the measurement in the OF. There was no further change in anxiety level in the absence of food restriction. So we conclude that the OF measurement was a stable baseline to compare with the EPM measurement, in order to obtain the effect of food restriction on anxiety behavior.

Food Restriction Changes the Anxiety Level in Mice

We have found that food restriction alone as well as in combination with wheel access do evoke changes in the anxiety level of mice (two-way ANOVA; Figure 2). This is an effect of food restriction and not of wheel access, because we see a change in anxiety levels from the prefood restriction phase in food restricted animals (both FR and ABA) but no change in anxiety level in the EX group. Social isolation in adolescence is another known stressor (Evans, Sun, McGregor, & Connor, 2012; Gan, Bowline, Lourenco, & Pickel, 2014). However, our data rule out the contribution of social isolation upon change in anxiety exhibited by the FR and ABA group, because no change was detected among the EX or CON group, which received identical durations of social isolation. Thus, we have been able to isolate the effect of the food restriction on changes in the anxiety levels in the ABA model of AN.

The Nature of Food Restriction Is Modified by Wheel Access

The FR group of mice also becomes hyperactive, as reflected by their higher number of entries in the EPM. However, their weight loss was not predicted by the open or total entries in the EPM, whereas this was the case of the ABA animals. This suggests that the weight loss in ABA animals was related to activity, whereas for the FR group, it was not. Thus, access to the wheel modifies the nature of the weight loss in the ABA model, further establishing the importance of excessive exercise as a key component contributing to the weight-loss aspect of severity of response to ABA.

The Nature of Wheel Activity Is Modified by Food Restriction

The anxiety level during ABA is tightly correlated to the level of hyperactivity of the mice with higher levels of anxiety in the EPM associated with higher values of wheel activity, most notably on the day following the EPM, and somewhat to a lesser extent with running on the day preceding EPM (Figure 3). The anxiety level measured in the EX mice was not related to their wheel activity. This difference suggests that the nature of wheel running is different under the two environmental conditions. In the presence of food restriction, wheel activity and anxiety are coregulated but not so in absence of food restriction. The strong correlation between anxiety measurements based on EPM and wheel running indicated that we could use wheel running in the ABA mice to monitor changes in anxiety across multiple days of food restriction.

Individual Variability in the Change in Anxiety and Hyperactivity Following Food Restriction

A strength of the ABA model is the individual variability in the hyperactivity response to the food restriction, which can be used to identify the mechanisms underlying the severity of response to ABA, as well as resilience (Aoki et al., 2014; Chowdhury et al., 2013). In this study, we show a parallel individual variation in the change in the anxiety level induced by food restriction, both in terms of the direction and magnitude of change. Moreover, we have shown a strong correlation between the change in the level of anxiety and change in wheel activity. The strong correlation between EPM open arm time as well as fraction of entries into the open arms and wheel activity suggests three possibilities: one, anxiety and hyperactivity are coregulated by a common neural pathway (Path 1 of Figure 5); two, anxiety is causal to the hyperactivity in ABA (Path 2 of Figure 5); three, a combination of the above two possibilities exists (Path 3 of Figure 5). Finally, it must be noted that all ABA animals in this study exhibited some increase in wheel activity, including those that showed a decrease in the level of anxiety. This suggests the presence of a component

Figure 5. Schematic of possible pathways of regulation of anxiety and hyperactivity in the presence of food restriction. Path 1 suggests that the release of corticosterone (cort) following the activation of the hypothalamic-pituitary-adrenal axis might lead to changes in anxiety as well as hyperactivity. Path 2 suggests that cort release might lead to changes in anxiety, which generates the changes in level of wheel activity. Path 3, a combination of Paths 1 and 2, suggests that the initial rise in wheel activity following food restriction may be because of cort. Wheel activity might counter any rise in anxiety (as shown with the gray dashed line) and this feedback mechanism might increase the wheel activity further.
mediating an increase in wheel activity following food restriction that is separable from the mechanism related to anxiogenesis. An example of such a component may be the food anticipatory activity, which is hypothesized to be regulated by food-entrainable circadian oscillators in the brain and periphery (reviewed in Escobar, Cailotto, Angeles-Castellanos, Delgado, & Buijs, 2009; Paton & Mistlberger, 2013).

**Potential Common Neural Pathways Regulating Anxiety and Hyperactivity**

We propose the following neural pathway that may coregulate anxiety and hyperactivity. The hypothalamic-pituitary-adrenal (HPA) axis is activated following food restriction (Duclos, Gatti, Bessièrè, & Mormède, 2005; Duclos, Gatti, Bessiere, & Mormede, 2009), resulting in the release of corticosterone (cort). Cort rapidly increases mEPSCs (miniature excitatory postsynaptic currents) in the amygdala and thus the excitability of amygdalar neurons, which might itself result in anxiogenesis (Joels, Sarabdjitsingh, & Karst, 2012). Both the glucocorticoid and mineralocorticoid receptors in the amygdala are involved in anxiogenesis in response to cort (Myers & Greenwood-Van Meerveld, 2007). It is conceivable that not all animals respond to the food restriction with similar level of HPA axis response and thus have different anxiety responses. For example, signaling of the cannabinoid CB1 receptor dampens the response to stress, as well as the level of anxiety (Hillard, 2014). A different level of expression of this receptor may account for the individual variability in the response to stress.

Animals in the ABA paradigm show a fourfold higher level of cort relative to the EX animals, and the cort levels are tied to the level of hyperactivity (Duclos et al., 2009). Cort release in relation to feeding time may be associated with its metabolic roles in the storage of glucose or in the prevention of protein breakdown (Dallman et al., 1999). In addition, cort is known to increase locomotor activity via increasing the dopamine levels in the nucleus accumbens (Piazza et al., 1996) in the dark period, but not in the absence of food restriction (ABA vs. EX)? Food deprivation increases the activity of amygdalar cells (Moscarello, Ben-Shahar, & Ettenberg, 2009). The amygdala has robust projections to the ventral and medial striatum, which influence behavior based on stimulus-reinforcement associations (Cador, Robbins, & Everitt, 1989; McDonald, 1998). Increased activity of these direct projections may increase the reward associated with wheel running, such that the mice increase their wheel activity.

Our previous data showed that there was a change in the GABAergic receptor expression in the amygdala following ABA, in a direction consistent with an increase in outflow from the amygdala (Wable et al., 2014). An increase in signaling to the output regions, such as the striatum, could explain the different behavior of the network under the two conditions - wheel access, with and without food restriction.

Finally, it is possible that the initial rise of wheel activity, that is, on the first day of food restriction, is because of the direct effect of cort on the dopamine levels in the nucleus accumbens (Piazza et al., 1996). Subsequently, the anxiolytic effect of exercise may act to further increase its reward value and hence, the running on the second and third day of food restriction, until the animals are too exhausted to exhibit this increased wheel activity.

**Molecular Markers Possibly Common to an Individual’s Response to ABA and to Their Level of Anxiety**

The possible role of the benzodiazepine (BZD)-insensitive, extrasynaptic type of GABAergic receptors in increasing hyperactivity in the ABA model (Aoki et al., 2014; Wable et al., 2014) suggest that non-BZD anxiolytics should be investigated in reducing severity of response to ABA. The lack of literature on the use of benzodiazepines in the treatment of AN suggests that they are not effective and this might be related to the role of BZD-insensitive GABAergic signaling in the brain of AN individuals. Our data offer an alternative to be explored for treating ABA, and possibly AN, using agents that up-regulate the α4-containing GABARs.

Current literature suggests that low levels of leptin could underlie the hyperactivity exhibited by individuals suffering from AN as well as rodents in ABA (Hebebrand et al., 2003). Leptin could also be tied to anxiety states (Finger, Dinan, & Cryan, 2010). Leptin receptor signaling in the midbrain dopamine neurons modulates amygdala function and expression of anxiety (Liu, Perez, Zhang, Lodge, & Lu, 2011). Thus, leptin and its signaling could be a marker common to anxiety and severity of response to ABA.

The correlation between the severity of response to ABA as measured by wheel activity and the change in anxiety level induced by food restriction further provides support to the idea that anxiety and wheel activity are regulated by a common neural pathway that is activated by the stress of food restriction.

**A Difference Between the Anxiety in ABA and AN**

There is one characteristic in which the model departed from the human disease. A high proportion of the individuals with a dual diagnosis of AN and anxiety develop a childhood anxiety disorder preceding the onset of AN (Kaye et al., 2004). We did not see any correlation between preexisting anxiety level measured using open
field and the extent of hyperactivity in ABA animals. While it is possible that the open field is not the most suitable test for measuring anxiety in pubertal female mice, the lack of predictability from preexisting anxiety level is likely related to the fact that ABA can be induced in wild-type rodents without any particular predisposition to wheel running or any specific anxious phenotype. Gelegen et al. showed that more anxious mouse strains are more hyperactive in ABA than the less anxious strains (Gelegen et al., 2007). There are two differences between the Gelegen et al. study and our study: One, we analyzed individual differences by using correlation analyses between levels of activity and anxiety, rather than relying only on group comparisons of the mean values as in the Gelegen study. We measured activity and anxiety in the same mice. This was not so in the Gelegen study; the anxiety measurements were performed for the most part in male mice in separate studies (Singer, Hill, Nadeau, & Lander, 2005; van Gaalen & Steckler, 2000) except for Ponder, Munoz, Gilliam, and Palmer, 2007, who included female mice in a study on fear conditioning, a measure related to anxiety). Two, our study was conducted on mice in the adolescent stage of development (<2 months old), whereas the subjects in the Gelegen et al. studies were 3–5 months old. It is possible that younger female mice are more vulnerable to hyperactivity induced by ABA, which may explain why the C57BL/6 mice in their study were reported to show no hyperactivity in response to ABA induction, while the C57BL/6 mice in our study showed robust hyperactivity that was of different levels across individuals. This highlights the importance of approximating the disease as closely as possible in the animal model.

Conclusion

The correlational relationship between anxiety level and hyperactivity observed in the ABA animals parallels the observations reported for individuals diagnosed with AN (Breus & O’Connor, 1998; Holtkamp et al., 2004; Peñas-Lledó et al., 2002; Shroff et al., 2006). These parallelisms strengthen the animal model and indicate that the ABA model can be used to understand the neural circuitry that underlies the comorbidity of anxiety and AN.

References


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