It has become increasingly clear that males and females differ even more dramatically than we previously thought. Not only do they exhibit differing responses to stress and environmental experience, but they can also respond in opposite directions. In rats, it has been shown that exposure to an acute stressful event can enhance subsequent learning in males while dramatically impairing learning in females. These opposite effects of stress on memory formation are accompanied by similarly opposite effects on neuroanatomical measures, such as dendritic spines in the hippocampal formation. Moreover, these opposite effects of stress are mediated by different hormonal systems between the sexes. These unique responses to stressful experience in male versus female rats may be used to model sex differences in mental illness, such as those that exist for depression and posttraumatic stress disorder.

It is obvious that there are many differences between the sexes, and our external differences only mask those beneath. However, for various reasons, some cultural, it is often assumed that male and female response systems differ only as a matter of degree and not of direction. Indeed, it is often assumed that differences in our experiences or response to external events stem from differences in habits or belief systems that are malleable and could change by adopting a perspective more like the other sex. In this review, I will present data from a series of studies that indicate that males and females not only differ in the degree of their response, but often in direction too. To illustrate this phenomenon, I will focus on behavioral and neuronal responses to stressful experience and learning opportunities. These examples arise from studies conducted in the white albino laboratory rat. This approach eliminates some of the cultural and sociological considerations inherent to many discussions about sex differences in behavior. In addition to behavioral measures, I will present data indicating that anatomical measures of plasticity in the male and female brain can respond in opposite directions to the same environmental event. These behavioral and neuronal differences are dependent on the presence of sex and stress hormones, but differing ones for males versus females. Finally, I will discuss how these sexually dimorphic and diergic responses to life experience may be used to model sex differences in mental disorders, such as depression and posttraumatic stress disorder.

**Sex differences in learning and memory**

There are numerous reports of sex differences in basic learning processes. However, they vary greatly depending on the task used and species involved. In general, men tend to outperform women on tasks that require mental and spatial rotation, whereas women tend to...
outperform men when tested for spatial location in a static environment. Also, men are much more accurate at aiming an object at a target, whereas women often excel at tasks that require fine motor skills. Some of these sex differences in performance, such as those for targeting, also exist in nonhuman primates. With respect to the most common laboratory animal, the rat, sex differences in performance are influenced by natural differences in activity levels. Female rats, who are generally more active than male rats, perform best on tasks that require activity, such as active avoidance, and do quite poorly on those that require immobility, such as during fear conditioning or passive avoidance (for a review, see reference 4). Because sex differences in activity may confound differences in learning, we have adopted a task that does not depend on voluntary activity: classical conditioning of the eyelid response. During this task, the animal is exposed to an aversive stimulation of the eyelid, which causes it to blink. During training, the stimulation is preceded by a tone, which predicts the onset of the stimulation. After repeated exposure to these paired stimuli, the animal “learns” that the tone predicts the eyelid stimulation and blinks in response to it. This task has a number of advantages for studying sex differences in learning. First, the eyelink is a discrete response that can be accurately measured and quantified. Second, performance of this task is not dependent on overt activity or exploration. The animal must emit an unconditioned response to the eyelid stimulation and only upon training elicits a conditioned response to the tone. As an additional advantage, the anatomi cal substrates that underlie learning the basic response have been identified. Finally and perhaps most importantly, the task can be and has been conducted in virtually all animals, from mice to rats to monkeys to humans. Since results from animal studies often generate novel hypotheses about human behavior, this paradigm affords the possibility of testing them directly in normal and patient populations. Using this task of classical eyelid conditioning, we have observed that female rats acquire the learned response faster and emit more learned responses during training than do males. This sex difference in conditioning is even more prevalent if one takes into account the stages of estrus, the cyclic behavior of hormones associated with ovulation. Female rats have a 4- to 5-day cycle over which estrogen and progesterone levels change fairly dramatically. Proestrus is a stage prior to ovulation when estrogen levels are relatively high. When trained during this stage, females learn faster and condition more than females in other stages. These data suggest that estrogen is positively related to performance of this associative learning task.

How do these results compare to others in the literature? Certainly, there are numerous reports that learning (or performance) is related to the presence of sex hormones, although these effects vary depending on task and species. Women tested during the phase of the menstrual cycle associated with high levels of estrogen score better on tests of verbal fluency and fine motor skills—tests that women already perform well relative to men. In rats, females tested during proestrus perform poorly during a spatial memory task that is dependent on an intact hippocampal formation, but perform optimally when the task is not dependent on the structure. Some report that females tested during estrus have deficient spatial performance relative to males and females in other stages, whereas others report no effect of estrous cycle on learning, though performance variables were affected. Some of these effects can be ameliorated by previous familiarization with the task demands, suggesting that the stressful nature of some of these tasks contribute to the seemingly variable responses. Given the variation in the task demands, the brain structures involved, as well as the cyclic nature of endogenous hormone levels, it should come as no surprise that the relationship between absolute levels of hormones and learning is inconsistent. Moreover, since hormone levels do vary so frequently over time and experience, their effect on learning could not be absolute. Rather, hormones modulate learning to varying degrees via numerous mechanisms and presumably for numerous adaptive reasons.

**Sex-specific responses to stress and memory formation**

As with learning, there are sex differences in the stress response and these effects are often a matter of degree, not direction. The most robust sex difference occurs with endogenous levels of glucocorticoids. In many species, glucocorticoid levels are higher in females than males. This sex difference is apparent under unstressed and stressed conditions and in rats, glucocorticoid levels are elevated in females during proestrus relative to other stages of estrus. Stressful experience can also elicit very different behavioral responses in males versus females.
For example, we have shown that female rats exposed to an acute stressful event are severely handicapped in their ability to learn an associative response.2,10 Oddly enough, males respond in the opposite direction to females and thus exhibit enhanced performance after exposure to the same stressful event.23,24 The stressful event consists of either brief exposure to intermittent tailshocks or brief swim stress (20 min), both of which are common methods for inducing behavioral depression in laboratory animals. As a measure of learning, we again used the classically conditioned eyeblink response. These opposite responses to stress are not limited to simple associative learning as occurs during classical conditioning with overlapping stimuli. As illustrated in Figure 1, they are also evident during trace conditioning, a more difficult task in which the conditioning stimuli are separated in time. This task critically involves the hippocampal formation, and some have even suggested that it involves conscious awareness.25-27

If these effects of uncontrollable stressful experience on learning in rats are relevant to the human condition, they should possess some characteristics of mental illness, particularly those associated with stressful experience. One that comes to mind is posttraumatic stress disorder (PTSD). After experiencing a traumatic stressful event, some humans develop PTSD, more than twice as many are women.29 Often-times, they reexperience frightening aspects of the traumatic event, particularly if presented with cues that are associated with the event. To determine whether the effects of stress on learning in rats were sensitive to these factors, we exposed rats to cues associated with the stressful event days after it had ceased and at a time when the effects of stress would have dissipated. Indeed, days after the stressor, males reintroduced to the stress context were further enhanced in their performance, whereas females were further impaired.10,30 Minimally, these results suggest that the effects of acute stress on learning are not entirely dependent on sensory stimulation, but rather can be stimulated by associations that were established during stressful environment. More generally, they suggest that the effects of acute stress on later learning in rats may model some disrupting effects of trauma on cognitive processes in humans such as occurs during PTSD.

Stress hormones and stress effects on memory formation

There are numerous examples of sex differences in behavior, but few demonstrating an opposite response to the same stimulus between sexes. What could be responsible for inducing these opposite responses? When exposed to a stressor, the organism responds by activating a complex series of physiological and behavioral responses that are mediated by the sympathetic branch of the autonomic nervous system (ANS) and the hypothalamic-pituitary-adrenal (HPA) axis. The release of glucocorticoids (corticosterone [CORT] in rats) by the adrenal glands is an important part of the organism’s ability to deal with stress.31 Among other effects, increased levels of corticosterone potentiate the release of adrenaline, increase cardiovascular tone, and mobilize the energy needed for flight and flight responses. In a series of experiments, we directly evaluated the potential role of glucocorticoids in the sex and stress effects on conditioning. After removing endogenous glucocorticoids via adrenalectomy, male and female rats were stressed and trained on the classically conditioned eyeblink response. Somewhat surprisingly, adrenalectomy prevented the enhancing effect of stress on learning in males, but did not alter the female response to stress (Figure 2).10,32 Thus, exposure to the
acute stressful event not only has opposite effects on this measure of performance in males and females, but these effects are mediated by different hormonal systems. How do these results compare to others in the literature? This is a difficult question since there are many different types and effects of stress; they are enhancing or disruptive depending on the task, training conditions, and sex of the animal.10,33-36 Despite the differences in response, many are assumed to occur via glucocorticoid activity and most often by activity within the hippocampal formation. The hippocampus has an abundance of glucocorticoid receptors, particularly the type I or mineralocorticoid receptor,37 and the structure is implicated in feedback of the HPA axis.38 Thus, our results regarding the male response to stress are generally consistent with much of the literature. That the female response is not dependent on the presence of glucocorticoids may be an aberration or simply reflect the fact that so few studies have been conducted in the female.

Since glucocorticoids are not critically involved in the stress effect in females, we considered other potential modulators, the first being ovarian hormones. As shown in Figure 3a, their removal via ovariectomy prevented the stress effect on conditioning, suggesting that their presence is necessary for observing an impairment after stress. Of the two primary ovarian hormones, we evaluated a specific role for estrogen. Figure 3 shows that treatment with the estrogen antagonist tamoxifen prevented the stress effect on conditioning. Together these data suggest that estrogen is critically involved in the stress effect on conditioning in females. We have also determined that the detrimental effect of stress on learning is dependent on the stage of estrus in which the learning occurs. Of the stages, females that were trained during proestrus (stressed 24 hours earlier in diestrus) were most impaired by stressor exposure.11 Since this stage is associated with elevated levels of estrogen, the hormone is again implicated in these stress effects on conditioning.

Recall that females under normal unstressed conditions learn faster in proestrus than in other stages. How might estrogen contribute to both enhanced learning under unstressed conditions and impaired learning after stress? It may be useful to consider the effect of stress on learning from a slightly different perspective in which stress does not impair conditioning directly, but rather prevents the enhancement that normally occurs when estrogen levels are elevated.

**Figure 2.** Contribution of adrenal hormones to the opposite effects of stress on learning in males versus females. A. Males adrenalectomized (ADX) prior to stressor exposure were not affected by stress, while those exposed to a sham surgery showed an enhanced response rate. B. In contrast, females adrenalectomized (ADX) prior to the stressor exposure showed impaired response after stress, as did the females exposed to a sham surgery.

**Neuroanatomical correlates of stress and sex differences in learning**

These opposite effects of stress in males and females pose some interesting questions, one being whether there is a neuronal or anatomical substrate that can account for these opposite responses to stress. First, we considered a potential role for dendritic spines, tiny protrusions on many dendrites in the brain, which are a source of excitatory input.39 Because they enable connections and associations to be made between adjacent neurons, it has been hypothesized that they are involved in the formation of associative memories. Despite the
pervasiveness of the hypothesis in the literature, there are minimal data in support of this. In fact, the most potent modulator of dendritic spines so far established is estrogen. Acute exposure to estradiol enhances spine density in the hippocampus of ovariectomized females; moreover, females in proestrus have a greater spine density than females in other stages. As discussed, it has long been assumed that dendritic spines participate in learning processes. So does this change in spine density across the estrous cycle and between the sexes relate to learning ability? At least as measured with classical eyeblink conditioning, there is a positive relationship between spine density and performance in females: females in proestrus outperform females in other stages and thus the variation in spine density correlates with their ability to acquire the learned response.11

If spine density is positively related to learning ability (of this task), then manipulations other than estrogen that modulate this type of learning may be expected to have effects on spine density. Initially, we considered the effects of stress. As discussed, exposure to an acute stressful event enhances later performance in males, but impairs performance in females. In a series of experiments, we tested whether exposure to one of

![Figure 3](image3.png)

**Figure 3.** Contribution of ovarian hormones to the stress effect on learning in females. A. Females that were ovariectomized (OVX) prior to stressor exposure and training were not impaired by stress and exhibited a similar response to those exposed to a sham surgery. B. Treatment with the estrogen antagonist tamoxifen prevented the stress effect on conditioning in females.

![Figure 4](image4.png)

**Figure 4.** A. Photograph of a hippocampal pyramidal cell impregnated with Golgi. Original magnification: 400x. B. A higher magnification (10000x) of the dendrite illustrates the spines. C. Effects of exposure to an acute stressful event on density of dendritic spines in area CA1 of the hippocampus showing increased density in males, but a decreased density in females during proestrus.
these stressors would affect spine density in the hippocampus and whether the effect would be sex-dependent. As illustrated in Figure 4, males exposed to the acute stressful event of intermittent tailshocks possess a greater density of spines than their unstressed male controls. Conversely, proestrous females who normally possess a high density of spines exhibit a decrease after exposure to the stressful event. Thus, spine density is positively related to performance under these specific conditions. To review, females in proestrus have a greater density than females in other stages and males, and they condition more. In response to stress, males have a greater density of spines than unstressed males and they condition more. In response to stress, females have a reduced density of spines and they condition poorly. These data do not indicate that spines are necessary for learning or that their presence mandates that learning will occur. Rather, they suggest that the presence of spines may enhance the potential for learning—should the opportunity arise.

**Sex differences in depression**

What do these dramatically different behavioral and neuronal responses in male and female rats tell us about human behavior and adaptation to stressful experience? Minimally, they indicate that we must be very careful in generalizing results obtained from males to females. A relevant example of this problem concerns the phenomenon of “learned helplessness.” In the 1960s, a number of influential behavioral scientists came upon an interesting observation. They had been using inescapable and escapable shocks in dogs to study the processes of Pavlovian (or classical) conditioning. During their experiments, they noticed that the dogs that were previously exposed to inescapable shock were less likely to learn a later task in which escape was then possible. These animals, as well as the many other species tested in this paradigm, displayed a number of features characteristic of depression. They did not eat as much, had sleeping problems, and were generally inactive. In essence, it appeared as if they had “given up” and no longer had the motivation to learn. A number of psychologists picked up on these similarities and thereafter promoted this “learned helplessness” phenomenon in animals as a model of depression in humans. This model had such wide appeal that it is included in nearly every general and abnormal psychology textbook and was eventually developed into a more sophisticated model of depression known as learned hopelessness.

The incidence and prevalence of depression is higher in women than in men. It would thus be interesting to test for learned helplessness behaviors in females. Unfortunately, only a few studies have done so. In most of these studies, rats were tested in a shuttle-box avoidance paradigm, in which the animal must “learn” to escape from a footshock on one side of the cage. In order to terminate the shock, the animal must escape through an opening to the other side of the cage and back to the initial side. After exposure to inescapable shocks, male rats were impaired in their performance, whereas the females were not affected. Although these results suggest that females are not learning impaired, it is difficult to prove this conclusively. This is in part because females are generally more active than males, thus the sex difference may simply reflect differences in behavior not relevant to learning, per se. Nonetheless, this paradigm is a commonly accepted animal model for depression in humans. That it may not adequately model female behavior suggests that alternative models may be warranted.

Although women are more likely than men to experience major depression in their lifetime, the course of that depression may not differ. There is no sex difference in duration of the first episode, time to recovery, time to first recurrence, and severity of symptoms. These data contrast with those observed for manic-depressive illness, with no apparent difference in prevalence, but rather one of course. It is reported that women cycle from mania to depression more rapidly than do men and they may have more depressive episodes and dysphoria. The increased prevalence of unipolar and course of bipolar depression as well as general changes in personality are often associated with or exacerbated by changes in ovarian hormones levels such as occur prior to ovulation, after pregnancy, and during menopause. It is in this context that we again present our findings regarding the effects of stress on learning in the females, this time highlighting its relationship to changing levels of estrogen. In a typical experiment, female rats are exposed to an inescapable stressor such as intermittent tailshocks or swimming, and we then measure learning 24 hours later. As discussed, exposure to these stressors dramatically impairs sub-
sequent learning in the female rat.9–11 This effect most pronounced when females are stressed during diestrus and trained in proestrus, a time period over which estradiol levels are changing. Thus, the effect of stress is dependent on the stage of estrus and potentially on changing levels of ovarian hormones.11 Initially, we hypothesized that exposure to the stressful event altered the cycle, perhaps by decreasing the release of estrogen. However, experiments to test this hypothesis indicated that acute stress did not disrupt the cycle. We did observe an increase in estrogen levels after its cessation.53 However, injection of stress levels of estradiol did not impair learning as did the stressor. Thus, the effect of stress on memory formation in the female depends less on absolute levels of estrogen and more on their fluctuation during and shortly after the traumatic event. Consistent with some of the emotional disturbances that can occur during menstruation, postpartum, and menopause, these data suggest that females are particularly susceptible to the deleterious consequences of stress when ovarian hormones are fluctuating.

Conclusion: sex differences in mental illness

That females are different from males may come as no surprise. Nor that their brains are different. What might be unexpected is that they would respond in opposite directions to the same environmental event and that their brains would follow in course. In the face of such divergence, perhaps we should reconsider sex differences in mental illness (Table I). Females are not only more likely to experience depression, but also phobias, generalized anxiety disorder, and posttraumatic stress disorder. They are more often diagnosed with eating disorders, as well as borderline and histrionic personality disorders. Males, on the other hand, are more likely to experience autism and antisocial and narcissistic personality disorders, as well as attention deficit disorder and mental retardation. It may be instructive that the mental disorders more common in women are related to affect whereas those more common in men are related to cognition. Exactly how information about sex differences in emotional and cognitive responses in rats can be used to understand or promote mental health in humans is unclear, but a greater appreciation of our differences can only enhance our ability to treat our common afflictions. 

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**Basic research**

**Efectos opuestos de la experiencia estresante en la formación de la memoria en hombres versus mujeres**

Cada vez se ha demostrado con mayor claridad que los hombres y las mujeres difieren aun más dramáticamente de lo que previamente se pensaba. Ellos no sólo muestran diferencias en las respuestas al estrés y a las experiencias ambientales, sino que también pueden responder en sentidos opuestos. En ratas se ha observado que la exposición a un acontecimiento estresante agudo puede favorecer en los machos en forma consecutiva un aprendizaje; en cambio, en las hembras el aprendizaje se deteriora en forma dramática. Estos efectos opuestos del estrés sobre la formación de la memoria se acompañan del mismo modo de efectos opuestos sobre ciertas mediciones neuroanatómicas como la formación de espinas dendríticas en el hipocampo. Además, estos efectos opuestos del estrés están mediados por sistemas hormonales diferentes en cada sexo. Estas respuestas distintivas para la experiencia estresante en hombres versus mujeres pueden ser utilizadas para modelar diferencias por sexo en las enfermedades mentales, como aquéllas que existen para la depresión y el trastorno por estrés postraumático.

**REFERENCES**


**Effets opposés du stress sur l’organisation de la mémoire chez les hommes comparés aux femmes**

Que les hommes et les femmes diffèrent bien plus que nous ne le pensions auparavant est une réalité de plus en plus manifeste. Non seulement leurs réponses au stress et aux changements d’environnement diffèrent mais elles sont parfois opposées. Il a été montré chez le rat que l’exposition à un événement aigu stressant pouvait améliorer l’apprentissage ultérieur chez le mâle alors que celui-ci était fortement freiné chez la femelle. Parallèlement, ces effets opposés du stress sur la mémoire sont des résultats opposés en termes de critères neuroanatomiques, tel le nombre des épines dendritiques dans l’hippocampe. Par ailleurs, ces effets opposés du stress sont médiais par des systèmes hormonaux qui diffèrent selon le sexe. Ces réponses caractéristiques aux expériences de stress chez les rats mâles comparés aux femelles peuvent servir à modéliser les différences selon le sexe qui existent dans les maladies mentales, par exemple la dépression et le syndrome de stress posttraumatique.

**REFERENCES**