ANXIETY AND DEPRESSION MODELS

The following animal models of anxiety and depression are available in the laboratories of NDI neuroscientists for testing potential therapeutics.

ANXIETY TESTS

Maternal separation
Rapidly becoming the “gold standard” for evaluating anxiolytics, this test measures the number of “squeaks” made by guinea pig pups when temporarily separated from their mother. A reduction in the number of “squeaks” over a five-minute separation time has been predictive of clinical efficacy in reducing anxiety.

Elevated plus-maze
Rodents prefer to explore the enclosed 2 arms of a plus-maze elevated above floor level, compared to the un-enclosed 2 arms. Reduction in this preference by a test compound is considered predictive of an anxiolytic effect. This test is typically conducted using adult rodents and is considered supplemental to the maternal separation test (above).

Ultrasonic Vocalization
Ultrasonic vocalization (USV) by rodents can occur under anxiety producing conditions. It is regarded as a “universal” anxiety measure, i.e. is sensitive to both typical and atypical anxiolytics, and occurs in response to aversive stimuli (e.g., following air puff or shock) and in classical conditioning paradigms.

Light-enhanced startle
The startle response exhibited by rodents to a loud sound is increased by the simultaneous presence of a light of higher than normal intensity. This enhanced response is thought to reflect heightened anxiety, as it is blocked by antagonists directed to discrete brain areas implicated in anxiety.

Light/Dark preference
Activity in light and dark portions of a divided box is recorded. Increased avoidance of the lighted portion reflects elevated anxiety, while little or no preference for either the lighted or dark halves of the box reflects decreased anxiety.
DEPRESSION TESTS

**Forced swim test (Porsolt test)**
This is the most frequently used test of learned helplessness, a classical model of depression. The test measures the time an animal remains immobile when immersed in a water-filled cylinder from which escape is not possible. Antidepressant drugs increase swim time and reduce the length of time spent immobile. Strain of animal must be considered as the behavioral effects of some drugs have been shown to vary between in-bred and out-bred mice. For example, imipramine has been shown to be effective in CD1 mice while failing to show effects in NMRI mice.

**Tail suspension test**
The tail suspension test is a model of depression that measures the length of time an animal, usually a mouse, will struggle to escape while being suspended by its tail. Thus, the tail suspension test is sometimes referred to as a “dry land” version of the forced swim test. As with the forced swim test, antidepressant drugs increase the amount of time the animal spends struggling, and reduce the length of time spent immobile. Strain of animal may also be important in this test, as, for example, imipramine has been shown to be effective with NMRI mice but not with CD1 mice. The tricyclic antidepressants, such as amitriptyline and phenelzine, as well as SSRIs, are generally ineffective in this model.

**Chronic mild stress**
Chronic mild stress involves long term exposure to mild aversive stimuli or environmental conditions, such as food or water deprivation, soiled cage, tilted cage, or altered light/dark conditions. This model may represent anhedonia in depression. Sucrose consumption is often used as an index of anhedonia after the chronic stress treatment. All classes of antidepressants have been shown to be effective in this model, while anxiolytic and neuroleptic drugs have not. However, the results of chronic mild stress studies are sometimes difficult to replicate across laboratories.

**Schedule induced polydipsia**
This paradigm is conducted in a test chamber equipped with a food pellet dispenser and a water bottle. In a typical setup, the food pellets are delivered every 60 seconds, while a sipper tube for the water bottle is always available. Rats are food deprived before testing, but not water deprived. Thus, rats readily consume the food pellets when they are delivered. During the 60 seconds in
Schedule induced polydipsia (continued)
between food pellet deliveries, rats tend to drink water from the water bottle. Over time, water consumption in this task develops into polydipsia. Once a baseline level of polydipsia is achieved, antidepressant effects are indicated by a reduction in water intake. A particular advantage of this task is that repeated administration of antidepressant drugs is necessary to lower water intake. Thus, this model can be used to assess response time to antidepressant drugs, a major clinical concern.

Differential reinforcement of low rate responding (DRL)
In this task rats are trained to press a lever in order to receive a food pellet. However, this occurs only under certain conditions. First, only a lever press that is made after 72 seconds have elapsed since the last lever press will result in the delivery of a food pellet. If a rat presses the lever too soon (i.e., before 72 sec have elapsed since the last lever press), then the 72 second counter resets, and the rat will have to wait 72 seconds more before a food pellet becomes available.

In a DRL-72 procedure, antidepressant drugs produce an increase in the reinforcement rate (the number of food pellets earned over a period of time), meaning that rats are more likely to wait to press the lever until after 72 seconds have elapsed. In addition, response rates either increase or remain unchanged after acute administration of an antidepressant drug.

Example data are shown below. In this study, male Sprague Dawley rats were trained over the course of two months, to perform the DRL-72 procedure. Training criteria consisted of less than a 10% change in the number of responses over the course of 6 consecutive sessions. In the results shown below, the tricyclic antidepressant drug imipramine produced a decrease in responses and an increase in the number of reinforcers, which is an antidepressant response.
IN VIVO MICRODIALYSIS

Most modern antidepressant agents target the serotinergic and noradrenergic systems, but most of them also act on the dopaminergic system. Using in vivo microdialysis to measure drug-induced release of noradrenaline, serotonin and dopamine in target brain regions provides an index of the spectrum of action of a test antidepressant compound.

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