**Elevated plus maze test combined with video tracking software to investigate anxiolytic effect of exogenous ketogenic supplements**

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**KEYWORDS:**

Elevated plus maze test, Exogenous ketone supplements, Gavage, Ketogenic diet, Glucose, Anxiety

**SUMMARY:**

Here, we present a protocol to investigate changes in anxiety level of rodent animal models. The elevated plus maze (EPM) test used together with a video tracking software provides a reliable method to document the effect of various potential anxiolytic treatments in pre-clinical laboratory scenarios.

**ABSTRACT:**

The overall goal of this study was to describe the methodology of the elevated plus maze (EPM) test in combination with a video tracking software. The purpose of the method is to document the effect of various potentially anxiolytic treatments on laboratory rodent models. The EPM test is based on the rodents’ proclivity toward protected, enclosed dark spaces and unconditioned fear of open spaces and heights, and their innate intense motivation to explore novel environments. EPM is a widely used behavioral test for investigating the anxiolytic or anxiogenic responses of rodents given drugs that are known to effect behavior. Observation demonstrating a decreased proportion of time spent on closed arms, increased proportion of time spent on open arms, reduced number of entries to closed arms and elevated number of entries to open arms measured by EPM may reflect reduced anxiety level. The effect of exogenous ketone supplements on anxiety-related behavior was tested in Sprague Dawley (SPD) rats using this method. Exogenous ketone supplements were chronically fed for 83 days or sub-chronically orally gavaged daily for 7 days acutely before conducting the EPM test. Behavioral data collection was performed using the SMART video-tracking system by a blinded observer at the end of treatments. The main findings indicated that EPM was an effective method to detect the ketone supplement-induced anxiolytic effect and can be considered a sensitive measure to assess changes in anxiety behavior associated with drug or metabolic-based therapies.

**INTRODUCTION:**

The goal of this article is to describe the methodology of the elevated plus maze (EPM) test in combination with a video tracking software in order to monitor changes in anxiety related behavior and novel treatments in laboratory rodent models. The EPM test is a relatively simple behavioral assessment method, which was developed for investigation of quantifying anxiety behavior levels and anxiety responses of rats after application of drug treatments1. Indeed, it has been demonstrated that EPM test is a widely used and effective behavioral assay for investigation of the changes in anxiety level of rodents1,2. Applicability of EPM test in rodents (mainly rats and mice) is based on their proclivity toward enclosed, dark spaces (approach), an unconditioned fear of open spaces/heights (avoidance) and their high level of innate motivation to explore novel environments. Consequently, the EPM test is a well-established methodology based on an approach-avoidance conflict2,3.

EPM is a plus-shaped apparatus consisting four elevated arms, which was described by Handley and Mithani4 (**Figure 1**), and consists of two opposite arms that are open to the surroundings (open arms), whereas the two closed opposite arms (closed arms) are equipped with walls. After treatment, if increased time spent on the open arms and/or increased number of open arm entries compared to control (untreated) animals is detected on the EPM, this indicates an anxiolytic effect2,3. The most robust avoidance response was demonstrated in the first 5 min after the start (placement of rats in the intersection of four arms of EPM) of the EPM assay 5; therefore, behavior after a treatment is commonly recorded for 5 min on EPM. As additional measures of anxiety level, number of head dips, rears (vertical standing of rodent on two hindlegs), fecal boli, as well as total arm entries (spontaneous motor activity) and different postures (stretching or freezing) can also be recorded on EPM2. Thus, multiple behavioral parameters can be compiled to provide a comprehensive assessment of anxiety-related behavior.

In order to increase validity of results, 2-3 behavioral assays are commonly used together, such as the light–dark choice test, social interaction test and EPM test to measure anxiety level in different animal models 6. The EPM assay performed alone on rodents is also a suitable method to investigate the anxiolytic or anxiogenic effect of different drugs 7. The EPM is sensitive not only to benzodiazepine-type anxiolytics (*e.g.,* diazepam) 8, but also, among others, to amino acid, monoamine, peptidergic and nucleosidergic compounds (*e.g.,* NMDA antagonist AP7, AMPA antagonist CNQX, µ-opioid receptor agonist morphine, NPY1 antagonist BIBP3226, Substance P, Ghrelin, Oxytocin, serotonin receptor agonists and antagonists such as 8-OH-DPAT and WAY-100635, β1-adrenergic antagonist Betaxolol) 9-12. Consequently, EPM assay on rodents is a suitable and sensitive method to investigate the influence of different treatments that influence brain areas involved in the anxiolytic effect (*e.g.,* amygdala, hippocampus, and limbic areas) and mechanisms of action (*e.g.,* serotonergic, GABAergic and adenosinergic system) implicated in anxiety 2. The agents tested in these EPM studies include exogenous ketone supplements that alter brain signaling in subtle ways that may require a sensitive method to detect behavioral changes.

In this article we describe the EPM test used in combination with a video tracking software which helps eliminate experimental bias and facilitates the collection and analysis of behavioral alterations in response to novel anxiolytic treatments.

**PROTOCOL:**

Animal treatment and measuring procedures were performed in accordance with the University of South Florida Institutional Animal Care and Use Committee (IACUC) guidelines (Protocol #0006R). All efforts were made to reduce the number of animals used.

**1. Preparations**

Note: The protocol typically requires laboratory-bred rats or mouse for EPM testing. However, other animals, such as guinea pig have also been tested on EPM 13. It is important to consider the color contrast between the animals in the maze and the maze color when using video-tracking. The contrast is less important for researchers watching animals live or *via* video. The settings of the video tracking software needs to be configured to document that the animals are black or white on either a black or white maze. Problems with configuration settings can occur with a clear acrylic maze, but a matte grey maze can be optimal for both rodent colors.

1.1. Select animals for the experiment considering the potentially influencing factors, such as strain, sex, estrus cycle and age, as well as body weight 2.

1.2. Determine the number of animals per group for the test based on the individual experiment.

Note: The group size will be dependent on effect size that is expected with the test treatment. Power analyses are generally done before the experiment is initiated to determine the minimum number of subjects to be included given the variability in animal's responses in any given task, as well as the number of experimental groups/conditions.

1.3. Design the experiment (in which a battery of different behavioral test, such as open field test, EPM test, hole-board test and forced swimming test will be used) carefully.

Note: Pre-exposure of rodents to a novel test environment (such an open field test) immediately before EPM tests may change behavior of animals on EPM 1,2.

1.4. Handle all animals in a similar way before EPM test.

Note: It has been demonstrated that different stress factors, application of drugs (*e.g.,* injections), shipping stress and handling can change behavior and behavioral responses of rodents on EPM 16. Thus, habituation of animals to animal house (*e.g.,* after shipment, 1-2 weeks before EPM test), experimental conditions, and treatment procedures (*e.g.,* gavaging) are necessary. It is also important that handling of rodents and any experience with prior stressors, particularly immediately before testing, is consistent across animals and treatment groups.

1.5. Conduct the behavioral studies in nocturnal animals, such as rats and mice, using a reverse light cycle, so that the behavioral assessment can be performed when the animals are in their dark, active phase.

NOTE: The effects of different housing conditions and light cycle/circadian rhythm on behavior and their influence on EPM results were demonstrated previously17, since the animals' hormones are regulated by light cycle.

1.6. Use the same experimenters during the procedures and ask them to avoid perfume or soaps with strong odor.

1.7. Ask the experimenters not to talk near the animal during the experiment or move objects near the EPM environment.

Note: It is critically important that the observer makes minimal movements and no noise when collecting behavioral data.

1.8. Clean the entire EPM after each trial to erase smells of previous animals which might interfere with exploration of the test animal.

1.9. (Recommended) Handle the animals for several days before EPM (picking up gently by the torso and holding for a minute or two) to acclimate them to the experimenter.

1.10. When placing the animals on the EPM make sure to handle all animals in a consistent manner and place each rodent in the EPM in the same position facing the same arm (*e.g.,* in center facing toward open arm away from the experimenter).

**2.** **Application of exogenous ketone supplements**

2.1. Measure the body weight of animals before starting treatments to determine dosage calculation for treatment (*e.g.,* intragastric gavage).

2.2. Familiarize the animals to the intragastric gavage method (adaptation period) using water by gavage for 5 days before ketone supplementation (standard rodent chow/SD + water gavage; *e.g.,* 2.5 g/kg body weight water/day). Exclude the use of any animal that does not adapt to the intragastric gavage method.

2.3. Following the adaptation period feed the animals chronically for 83 days and sub-chronically for 7 days with SD and gavage daily with either water (*e.g.,* 5 g/kg body weight/day; control group: n = 8), ketone supplements such as ketone ester (KE; 1,3-butanediol-acetoacetate diester; *e.g.,* 5 g/kg body weight/day; n = 8), ketone salt (KS; Na+/K+ - beta-hydroxybutyrate/βHB mineral salt; *e.g.,* 5 g/kg body weight/day; n = 8) or KS+ medium chain triglyceride (1:1 ratio, KSMCT; n = 8) 18-20.

Note: The animals that received intragastric gavage were tested on EPM one hour after treatment. Rats fed with standard rodent chow and gavaged with water (excluding ketone supplementation) served as control groups.

**3. Anxiety assay**

**3.1. EPM apparatus**

3.1.1. Use the same apparatus across a study to standardize results. EPM is a plus-shaped apparatus, which consists of four arms (*e.g.,* arms may be 10 cm wide and 50 cm long): Two opposite arms are opened and two closed opposite arms are equipped with high (*e.g.,* 30 cm) walls. The apparatus is elevated above the floor (*e.g.,* by 55 cm) 2.

Note: The most commonly used parameters are accumulated time in the open arms and the number of entries to open arms, however time spent in closed arms and center, and number of entries to closed arms and center are measured, as well as distance travelled in each areas.

3.1.2. Light up the EPM by using indirect lighting (*i.e.,* direct the light source toward the ceiling instead of directly illuminating the EPM apparatus) and ensure all four arms are similarly illuminated (without shadows, **Figure 2**).

Note: Changes in level of light alter behavior of rodents on EPM. Therefore, similar illumination is needed in consecutive experimental animals and days (*e.g.,* 2800 lumens in the room) 2.

**3.2. Video-tracking system**

Note: Use a video-tracking system with computer interface and video camera for data collection, which will automatically collect behavioral data in rats (**Figure 3**). For the video tracking system a wide variety of standard analog cameras or user-defined image sources (Infrared cameras, camcorder, WIA-compliant USB camera, webcams, *etc.*) can be used. When analyzing the recorded video, the movement tracking software supports all common video formats, such as .avi, .vob, .wmv,.asf, .mov, .qt, .mpg, .mpeg, .mp4, .3gp, .mkv. If the video does not playback correctly it might require a specific codec, additional video formats are supported if the corresponding codec is installed in the system. The movement tracking software can also be used to analyze previously acquired video and process the images from different sources, such as DVD/HD recorders, digital video files (AVI, DIVX, MPEG, *etc.*), webcams, DV cameras and WIA-compatible imaging devices.

**3.2.1. System setup**

3.2.1.1. Plug the installation key of the movement-tracking software into a USB 2.0 port and launch the installation tool.

3.2.1.2. Fix the camera above the experimental area and ensure that it will stay immobile along the whole experiment.

3.2.1.3. Set up a new experiment in the movement tracking software system by using the instruction manual. Select the **New task**. Double click on the icon of the protocol which the new experiment should follow (**Figure 3, Supplementary File 1**).

3.2.1.4. Enter details to label/describe the experiment in the **Experiment Info** dialog.

3.2.1.5. Specify the source of the video sequences to process.

3.2.1.6. Define the transformation rule for a correct distances measurement. The calibration process enables the movement-tracking software to be informed of the actual dimensions of the experimental area in order to obtain reliable values for distances and speeds.

3.2.1.7. Determine the regions of interest (zones) in the working area.

3.2.1.8. Adjust the parameters of the detection process.

3.2.1.8.1. In order for the movement-tracking software to precisely detect the position of the animal in the image, some detection adjustments must be set.

3.2.1.8.2. The tracking process requires a clear and well-contrasted image by using fine adjustment of general brightness and contrast parameters in the **Brightness & Contrast** section of the **Detection Settings** panel. As needed adjust these settings for the whole image or for user-defined zones.

3.2.1.9. Put a rat into each arena to test the detection process.

3.2.1.10. Press the **Start Test** button to verify if the detection process can identify the subject correctly. Confirm the detection is activated by the appearance of a dot on the screen. The Calibration process has to be done before starting the Test.

3.2.1.11. Detection is considered confirmed when the only black dot shown in the player is the animal being tracked. The red tracking line needs to closely follow all the animal displacements. Proper tracking is also confirmed with a white label listing the animal number and corresponding coordinates based on displacement. If such detection is not obtained, adjust the **Threshold** and **Erosions** parameter for optimizing the detection and tracking process.

3.2.1.12. Adjust **Threshold** and **Erosion** parameters to get a sharper and noise-free test image.

3.2.1.13. If the tracking path is correctly detected, press the **Stop Test** button (**Figure 4**). If these adjustments are going to be used for every new experimental file, press the **Save** button as default button. Press the **Accept** button to save the new detection settings.

3.2.1.14. Set the time conditions of the trials.

3.2.1.15. If the experimental protocol requires the track acquisition process to start at the same time the subject is placed into the experimental area it is possible to setup the remote unit that comes with the software or to use a wireless mouse.

NOTE: This option provides the possibility of remote controlling the start and stop, and thus the recording session can be completed without the computer.

**3.2.2. Setup Subjects in the system**

3.2.2.1. Manage the experimentation subject’s database. To create a database of experimental subjects, enter the **Subjects Database** manager by pressing the **Subjects** button in the **Experimentation Assistant** bar.

3.2.2.2. Press the **+** button to add new subjects to the database.

3.2.2.3. With the one subject option already selected, enter the subject’s code.

3.2.2.4. Fill the rest of the subject’s information in the **Subject Properties** section.

3.2.2.5. Press the **Create** button to add the new subject.

3.2.2.6. Define the experimentation plan. Use the **Scheduler** to define the different phases, sessions, trials and subjects planned to be executed within the experimental project. The trial is selected automatically as “the next trial” to be executed. This property is shown as a green tick at the left side of the trial name.

**3.2.3. Data acquisition by simultaneous recording and tracking**

Note: When a live image source is selected, the **Player** panel provides an embedded recording module for easily capturing the video coming from the selected camera.

3.2.3.1. Prepare the movement-tracking software for data acquisition (calibration, zone definition, detection settings, time settings, scheduler).

3.2.3.2. Open the **Data acquisition** panel.

3.2.3.3. Start recording the video of the experiment without the animal by pressing the **Start recording** button available on the software.

3.2.3.4. Place the animal into the experimental area.

3.2.3.5. Start the data acquisition process by pressing the **Start** button of the **Time control** panel.

The tracking process will be carried out simultaneously to the recording process. AS needed, ask the experimenter to note down the behavioral variables manually, such as rears, head dips, and falls (**Figure 5**).

3.2.3.6. Collect the EPM data manually as well as a blinded observer (separate the observer from the EPM by a curtain) in the testing room.

3.2.3.7. Wait until the end of the tracking process recording, or press the **Stop** button on the **Time control** panel.

3.2.3.8. Remove the animal from the experimental area. Stop the video recording process by pressing the **stop** button available on the movement-tracking software player.

3.2.3.10. Prepare the experimental area for the next animal by washing and drying it. Repeat the cycle again.

**3.2.4. Data analysis**

3.2.4.1. To access the **Analysis** tool, press the **Analysis** button in the **Experimentation Assistant** bar.

3.2.4.2. To generate analysis reports of the finished trials: Select the trials to analyze. Configure and select the analysis report. Set the time intervals to be analyzed. Generate and review the reports. Export the results to a spreadsheet or to image formats (**Figure 6**).

**3.3. EPM for measurement of anxiety level**

3.3.1. Perform the EPM experiments under non-stress conditions (in dimly lit and quiet room) after oral gavage.

NOTE: Make sure that the experiments are run in close time interval (*e.g.,* between 12.00 and 14.00) because the circadian rhythm can influence the behavior on EPM 15,17. Avoid unnecessary movements and noise during the experiment.

3.3.2. Before start of the test make sure that the EPM is cleaned and dried and the video-tracking system is ready to use.

3.3.3. Transfer the rats in their home cage to the experimental room 30 min prior to beginning the experiment.

3.3.4. Place the rat at the intersection of the four arms of the EPM, facing the open arm opposite to where the experimenter is.

3.3.5. Start the video tracking software as well as manually record the behavior of the animal for 5 minutes.

3.3.6. If the animal falls off from EPM, pick it up and place it back to the same point of the EPM where it fell off. Exclude the behavioral data of this animal from the analysis.

Note: A loud noise or movement may immobilize/freeze animals on open arms. If loud noise happens during the experiment, exclude behavioral data of these animals from analysis.

3.3.8. At the end of the 5 minute test stop the video tracking software and remove the animal from the EPM. Place it back into its home cage.

3.3.9. Before the next experiment/animal, clean the EPM with a disinfecting detergent (*e.g.,* Quatricide) followed by tap water. Dry the apparatus by paper towel.

**4. Analyses of data collected by the video tracking system**

4.1. Based on recorded data, analyze the amount of time spent in the open arms and in closed arms; number of entries made to the open arms, closed arms and to the center zone; latency to entry into the closed arms; distance travelled in open arms, closed arms and the center zone.

**Note:** the animal is considered to be in an area when all four paws are in the area.

4.2. Determine the effects of treatments on behavior by using analysis of variance (ANOVA) with Fisher’s LSD test/Tukey’s multiple comparisons test.

**REPRESENTATIVE RESULTS:**

The current experiment investigates the hypothesis that exogenous ketone supplementation administered either chronically (fed for 83 days) or sub-chronically (orally gavaged for 7 days) has anxiolytic effect on two months old male Sprague-Dawley (SPD) rats (250-350g). Chronic administration consisted of the following ketone supplements: Low-dose ketone ester (LKE; 1,3- butanediol-acetoacetate diester, ~10 g/kg/day, LKE), high dose ketone ester (HKE; ~25 g/kg/day, HKE), beta-hydroxybutyrate-mineral salt (bHB-S; ~25 g/kg/day, KS) and bHB-S + medium chain triglyceride (MCT; ~25 g/kg/day, KSMCT). For sub-chronic experiments the following treatment groups were used: KE, KS and KSMCT (5 g/kg/day). The control groups included standard diet (SD) or SD with water gavage (Control). All data was represented as the mean ± standard error of the mean (SEM). The results were considered significant when *p* < 0.05. Significance was determined by one-way ANOVA with Fisher’s LSD test.

Rats in the KSMCT group spent significantly more time in the open arms (*p* = 0.0094), compared to the control group after chronic feeding. The time spent in the closed arms was significantly less in LKE, KS and KSMCT groups (*p* = 0.0389, 0.0077 and 0.0019, respectively), while the KS group spent significantly more time in the center (*p* = 0.0239) compared to the control (SD) group (**Figure** 7A; 18).

Rats in the KS and KSMCT groups traveled significantly longer distance in the open arms (*p* = 0.036 and 0.0165), while the rats in the LKE, KS and KSMCT groups showed significantly less distance traveled in the closed arms (*p*=0.0252, 0.00041 and 0.0032, respectively), compared to control (SD) (**Figure** 7B). The KS and KSMCT groups had greater distance traveled in the center area (*p*=0.0206 and 0.0482, respectively), when compared to control, while in KSMCT group the latency to first entrance to closed arms was significantly greater after chronic feeding (*p* = 0.0038; 18)(**Figure** 7C).

The time spent in the open arms was greater in the KE group (*p*= 0.0281) after 7 days of oral gavage, while in KE, KS and KSMCT groups the time spent in the center decreased (*p*= 0.0005, <0.0001 and 0.023, respectively), compared to control (**Figure** 8A; 18). In the KE and KS groups the number of entries to closed arms was significantly lower (*p*= 0.0436, 0.0234, respectively) after 7 days of administration (**Figure** 8B), while the rats in the KS group also entered the center less frequently (*p* = 0.0193), compared to the control (SD).

**FIGURE AND TABLE LEGENDS:**

**Figure 1**: **Elevated plus maze (EPM) used for testing rats.** Each arm is 10 cm wide and 50 cm long with two opposite arms opened with a raised edge. The two closed opposite arms are equipped with 30 cm high walls. The runway height from the floor is 55 cm.

**Figure 2**: **Examples of direct and indirect lightings.** Ensure the light source is pointed toward the ceiling, while the direct light above the experimental area should be blocked. It is important to use indirect light during EPM experiments in order to similarly illuminate all four arms without shadows.

**Figure 3: The experimentation assistant bar of the movement-tracking software**. It is designed to provide access to the main operations. The buttons correspond to the task within the typical experimentation process, while only the currently allowed tasks are active.

**Figure 4**: **The subject track is marked with a red line following the animal’s movement**. By adjusting the threshold, the background can be decreased until only the animal is detected and tracked by the red line. The track is following the center of mass of the subject and the current position coordinates are indicated.

**Figure 5: Elevated plus maze with a Sprague Dawley rat in the open arm.** An example of the experimental setup is demonstrated.

**Figure 6**: **Accumulated movement track of the animal during a trial.** As part of the data analysis the collected trajectory trace of the subject in the tracking area can be displayed.

**Figure 7: Behavioral response of SPD rats in the EPM after 83 days of chronic feeding of exogenous ketone supplementation**. Representative results collected by the EPM and the movement-tracking system 18. (**A)** The KSMCT group spent a greater percentage of time in the open arms, while the LKE, KS and KSMCT groups spent less time in closed arms, compared to control (SD)group; (**B**) TheKS and KSMCT groups traveled more distance in the open arms, while LKE, KS and KSMCT groups traveled less distance in closed arms showing reduced anxiety compared to control (SD) group; **(C)** The KSMCT group entered the closed arms later, indicating reduced anxiety compared to the control (SD) group. Abbreviations: SD, standard rodent chow+water (25 g/kg body weight (b.w.) water/day); LKE, SD+LKE (1,3-butanediol-acetoacetate diester, 10 g/kgb.w./day); HKE, SD+HKE (25 g/kg b.w./day); KS, SD+beta-hydroxybutyrate-mineral salt (bHB-S; 25 g/kg b.w./day); KSMCT, SD+ bHB-S+medium chain triglyceride (MCT; 25 g/kg b.w./day); SPD: Sprague-Dawley rat; EPM: elevated plus maze; (\**p* < 0.05; \*\**p* < 0.01; \*\*\**p* < 0.001; \*\*\*\**p* < 0.0001). This figure has been modified from Ari *et al*.18.

**Figure 8**: **Behavioral response of SPD rats after 7 days of oral gavage of exogenous ketone supplementation**. Representative results were collected by the EPM using movement-tracking software system 18. (**A**) The KE group spent more percentage of time in the open arms, while KE, KS and KSMCT groups spent less time in the center, compared to the control (SD) group, indicating reduced anxiety; (**B**) Less entries were detected in the closed arms by rats in the KE and KS groups, compared to control (SD) group; Abbreviations: SD, standard rodent chow+water (5 g/kg b.w. water/day); KE, SD + ketone ester (1,3-butanediol-acetoacetate diester, 5 g/kg b.w./day); KS, SD+beta-hydroxybutyrate-mineral salt (bHB-S; 5 g/kg b.w./day); KSMCT, SD+ bHB-S +MCT (5 g/kgb.w./day); SPD: Sprague-Dawley rat; EPM: elevated plus maze; (\**p* < 0.05; \*\*\**p* < 0.001; \*\*\*\**p* < 0.0001). This figure has been modified from Ari *et al*.18.

**DISCUSSION:**

In general, there are several commonly used tests, such as the light–dark choice test, social interaction test and EPM test are used to measure anxiety level in different animal models. However, EPM assay alone is a suitable method to investigate for example the effect of exogenous ketone supplements on anxiety level 18, 20.

The main advantage of the EPM method is thatit relies on the rodents’ instinctive proclivity toward dark, enclosed spaces in addition to the unconditioned fear of heights and avoidance of open spaces.On the other hand, other methods used to study anxiety-like behavior are based on the behavioral responses to certain noxious stimuli, such as electric shock, food/water deprivation, loud noises, and exposure to predator odor3. These tests usually result in a conditioned response, while the EPM also represents a more humane alternative. Furthermore, the EPM can be a useful tool to study the involvement of different brain regions (*e.g.,* limbic regions, hippocampus) and underlying mechanisms (*e.g.,* GABA, glutamate, serotonin, adenosine *etc.*) of anxiety behavior 2.

When applying treatments that are quite stressful for the animals (*e.g.,* the oral gavage) it is important that all animals are handled the same way and by the same person, especially when assessing potentially subtle anxiolytic effects. If possible, introduction of the drug/compound in drinking water or *via* a palatable 'treat' may be a preferred method. To ensure that the same amount is administered to each animal an oral gavage can be used. Based on the pharmacokinetic properties of the compound it is usually advisable to test the animals on EPM within one hour after gavaging. When selecting experimental subjects it is important to consider their strain, sex, estrus cycle and age as well as body weight, according to the objectives and test substances 2. In regards to age, when designing EPM studies and interpreting data it is important to consider that the percentage of open arm entries linearly increases with age 21 and the aging-related changes in EPM behavior are strain-specific 22.

When conducting EPM test there are potential problems that need to be addressed. Sometimes animals need to be excluded from analysis due to outlier tendencies (*e.g.,* never leaves the area where it was placed, almost falls off the apparatus, it is distracted by a noise of event outside of the apparatus). Further complications with EPM testing may include treatments which cause sedation or hyperactivity, because these types of effects need to be assessed *via* EPM parameters.

It is important to expose animals to EPM test once, because decreased activity on the open arms and decreased total time spent on the central platform were demonstrated on the second (repeated) exposure of rodents compared to the first exposure on the EPM 14,15. Therefore, single exposure of rodents to the EPM test is strongly recommended. However, if there is minimum 3 weeks between first and second exposure to EPM and the EPM setup is moved to another room (different environment) the animals may be investigated by EPM test more than once 2.

EPM is available in different materials, sizes (*e.g.,* for mouse or rat) and colors which needs to be considered when choosing our study subjects. It is important to keep in mind that the odors left by the previous animal on the apparatus may change the behavior of the subsequent animal. Therefore we recommend using EPM made of a material that is easy to clean, such as acrylic glass (not transparent), which does not retain odors after washing. Avoid EPM apparatus made of wood. Preferably use matte color that is different from the color of animals tested on EPM (*e.g.,* black color if white animals are tested). The better the contrast between the animal and the enclosure, the better the detection of the animal and the higher the reliability and precision of the results obtained (distance covered, speed, tracking). EPM apparatus made from matte gray material is useful with white, black and white and black animals.

Further advantage of the video tracking system is that in addition to EPM, it offers a flexible and easy way to set it up with a wide variety of behavioral tests, such as Water Maze, Open Field, Plus/Radial Arm/T-Y Mazes, Place Preference, Forced Swimming and Tail Suspension tests.

In summary the goal of this article was to describe the EPM test used in combination with a video tracking software to collect and analyze behavioral alterations in response to novel anxiolytic treatments. The possible applications of the EPM include prescreening of newly developed pharmacological agents for treatment of anxiety-related disorders. In addition to the anxiolytic and anxiogenic agents, the behavioral effect of different hormones and drugs of abuse can also be investigated. The influence of aging and exposure to various stressors can also be assessed. This study has concluded that when proper steps are taken, the use of EPM has proven to be a sensitive method to assess behavioral changes associated with ketone supplementation 18, 20.

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**DISCLOSURES:**

International Patent # PCT/US2014/031237, University of South Florida, D.P. D’Agostino, S. Kesl, P. Arnold, “Compositions and Methods for Producing Elevated and Sustained Ketosis”. Provisional patent #62289749, University of South Florida, C. Ari, D.P. D`Agostino, “Exogenous ketone supplements for reducing anxiety-related behavior”. D.P. D`Agostino and C. Ari are co-owners of the company Ketone Technologies LLC. These interests have been reviewed and managed by the University in accordance with its Institutional and Individual Conflict of Interest policies. All authors declare that there are no additional conflicts of interest.

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