Therapeutic effects of an alpha-casozepine and L-tryptophan supplemented diet on fear and anxiety in the cat

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Abstract

Objectives This study assessed the anxiolytic effectiveness of a test diet (Royal Canin Feline Calm diet) supplemented with L-tryptophan and alpha-casozepine.

Methods Subjects were 24 cats that were classified as mildly or markedly fearful based on the presence of a person in their home room. Three different protocols were used to assess anxiety: (1) evaluation of the response to a human in the cat’s home room (home room test); (2) analysis of the response to placement in an empty test room (open-field test); and (3) analysis of the response to an unfamiliar human (human interaction test). All three protocols were first run at baseline, and the results were used to assign the animals to control and test diet groups that showed equivalent fear and anxiety. Both groups were retested on the three protocols after 2 weeks (test 1) and again after 4 weeks (test 2).

Results The diet groups differed for two behavioral measures in the open-field test: inactivity duration and inactivity frequency. The control group showed statistically significant increases in inactivity duration between baseline and test 1 and baseline and test 2, while the group fed the test diet showed a marginally not significant decrease in inactivity duration between baseline and test 1 and a not significant decrease for test 2. There was also a significant increase in inactivity frequency between baseline and test 1 in the test diet group and marginally not significant decrease in the control group. There were no differences between groups in the approach of the cats toward people for the home room test and the human interaction test.

Conclusions and relevance These results suggest that the test diet reduced the anxiety response to placement in an unfamiliar location, but that fear in the presence of an unfamiliar person was not counteracted by the diet.

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and lack of control that might be expressed as hyper-
vigilance, restlessness, escape or avoidance, circling, yawn-
ing, lip licking and distress vocalization.\textsuperscript{1,2,4}

Pharmacological therapy, in combination with envi-
ronmental and behavioral modification, is often indi-
cated to deal with the problem and address the wellbeing
of the pet. To date, however, no pharmacologic therape-
utics have been approved for the treatment of fear and
anxiety in cats. While a variety of natural products claim
to be effective in reducing anxiety, these claims have not
been comprehensively assessed.

Royal Canin Feline Calm is an alpha(\(\alpha\))-casozepine
and L-tryptophan supplemented balanced diet. Both
nutrients have been reported to have anxiolytic effects.\textsuperscript{7-9}
Tryptophan is an essential amino acid and the metabolic
precursor to melatonin and serotonin (5HT) that has been
implicated in the regulation of many behavioral pro-
cesses, such as mood, aggression and susceptibility to
stress.\textsuperscript{10-13} L-tryptophan has been evaluated in the treat-
ment of behavior disorders in cats including signs of
repetitive behavior, vocalization and agonistic behaviors.\textsuperscript{8}
However, supplementation with L-tryptophan alone may
not be effective, because of limitations in its ability to pass
through the blood–brain barrier as it competes with other
large neutral amino acids for a common transporter.
Therefore a lowered level of large neutral amino acids in
relationship to tryptophan may increase tryptophan avail-
ability, leading to increased 5HT synthesis in the brain.\textsuperscript{13}
\(\alpha\)-casozepine is a bioactive peptide originating from \(\alpha\)
S1 casein, a protein in cow’s milk, which has an affinity
for gamma-aminobutyric acid (GABA) receptors in the
brain. It is reported to have an anxiolytic effect similar to
benzodiazepines, with none of the side effects such as
incoordination and disinhibition of aggression.\textsuperscript{9,14,15}
Beata et al reported that fear of strangers, contact with
familiars, general fears, fear-related aggressions and
autonomic disorders were all significantly improved.\textsuperscript{9}
In a recent study, the ‘Calm’ diet was found to reduce uri-
nary cortisol and increase the plasma tryptophan/large
neutral amino acid ratio; however, no behavioral changes
were observed. In that study, all enrolled cats were per-
ceived to be normal, which limited the possibility of
observing a reduction in stress.\textsuperscript{16}

The purpose of the present study was to evaluate the
effectiveness of the Royal Canin Feline Calm diet (test
diet) in reducing fear and anxiety in cats in an open-field
test (OFT) and a human interaction test (HIT). We have
previously demonstrated that cats that exhibit fear of
people show greater inactivity in an OFT than non-
fearful cats, and that inactivity in cats with mild fear of
people is intermediate.\textsuperscript{6} In the HIT, non-fearful cats had
significantly shorter and more frequent bouts of inactiv-
ity than either mildly fearful or highly fearful cats.
Treatment with diazepam caused a reduction in inactiv-
ity in both the OFT and HIT.\textsuperscript{6} Therefore, in this study we
selected cats that had been categorized as either fearful
or mildly fearful.

Materials and methods
Experimental design
A parallel group design was used to evaluate the anxio-
lytic effectiveness of the test diet. At baseline, physical
examination, complete blood count, biochemical profile
and urinalysis were all within normal limits. Cats were
fed twice daily in separate feeding cages. Each cat was
given a measured amount of the prescribed food calcu-
lated for its optimum weight. All cats were initially
placed on the control diet (Purina Cat Chow) for 2 weeks
of adaptation, after which they were tested in the home
room, OFT and HIT over three consecutive days. These
baseline data were used to place subjects into two groups
that showed equivalent fear and anxiety. One group was
then switched to test diet and the other maintained on
control diet for the following 4 weeks. Subjects were
retested on the three tasks after 2 and 4 weeks.

All procedures were approved by the animal use and
care committee and were conducted in accordance with
the guidelines set forth by the Canadian Council on
Animal Care. The facility is a licensed animal research
facility under the Ontario Ministry of Agriculture, Farms
and Rural Affairs.

Subjects
The subjects were 24 domestic shorthair cats (five males;
19 females) from the Vivocore colony, ranging in age
from 4–12 years. Cats were selected from the population
of fearful and mildly fearful cats as described in our ini-
tial study\textsuperscript{6} (Table 1).

Test protocols
Three different test protocols were conducted to assess
measures of fear and anxiety. The first involved charac-
terizing the cat’s response to an unfamiliar person enter-
ing their home room (home room test).\textsuperscript{6} The second
(OFT) examined the cat’s response to placement in an
empty test room. The final protocol examined the cat’s
response to placement in the test room with an unfami-
 liar person (HIT). Both the OFT and HIT have been previ-
ously described.\textsuperscript{6}

Home room test
All cats had been categorized using a home room test as
described in our model development study.\textsuperscript{6} Home room
assessment was performed by the first author (GL), a vet-
erinary behaviorist with whom the cats had minimal
familiarity. Briefly, the cats classified as fearful had been
obtained approximately 6 months earlier from a research
facility in which they had been identified as fearful. These
cats were housed together and continued to actively
avoid their caregivers. In the home room test, they
actively avoided and displayed threats if approached. The other cats were intermixed in various housing rooms. The cats were assessed as non-fearful if they would voluntarily approach or did not withdraw when approached and would solicit or accept physical contact (touch, stroke) without signs of fear or avoidance. Those cats that would allow approach but would withdraw when contact was attempted were considered mildly fearful, and those cats that actively avoided approach or displayed signs of threats or aggression were considered fearful. Non-fearful cats were excluded from the study.

**Open-field test**

The OFT was conducted over 10 mins in a dedicated room (2.74 m × 3.66 m and 2.98 m in height) that included three cameras for recording movement, with one of the cameras set up to also record audio. The behavioral measures were recorded with the use of a behavioral tracking system, EthoVision XT (Noldus). Behavioral measures included: distance traveled; duration and frequency of inactivity, defined as when an animal was sitting or lying down or not exhibiting any overt behavior; and vocalization frequency.

**Human interaction test**

In the HIT, a person unfamiliar to the cat was seated on the floor in the center of the room before entry of the cat. For each test a different person was used but the same person was used for all cats at each test. The person was instructed not to wear any perfume or cologne. During the test the person was instructed not to move, look directly at the animals or make any attempt to elicit attention. The human interaction was conducted over 10 mins and we recorded the same behavioral measures that were assessed in the OFT. In addition, we quantified frequency and time spent in proximity to the person (within 0.5 m) and in contact with the person.

**Statistical analyses**

All statistical analyses were conducted using Statistica version 11.0 software package with significance set to \( P < 0.05 \). Analysis of data obtained during the treatment phase for each measure for measures that were common to both the OFT and HIT was based on a repeated measures analysis of variance (RMANOVA) with time of test (baseline, test 1 and test 2) and task (open field and human interaction) as within-subject variables and both group (control, treatment) and level of fear (mildly fearful, fearful) as the between-subject variables. For all analyses in which the tasks were compared, task (open field vs human interaction) was added as a second within-subject variable. For the home room test data, task was not included as a main effect in the RMANOVA.

| Identification | Test group | Age   | Sex (neuter status) | Anxiety status |
|----------------|------------|-------|---------------------|---------------|
| Herman         | Control    | 5.45  | M (n)               | Fearful       |
| Katniss        | Control    | 5.63  | F (n)               | Fearful       |
| Larisa         | Control    | 4.56  | F                   | Mildly fearful|
| Lily           | Control    | 9.70  | F (n)               | Mildly fearful|
| Maddox         | Control    | 4.33  | M (n)               | Mildly fearful|
| Melanie        | Control    | 4.25  | F                   | Mildly fearful|
| Mimi           | Control    | 4.42  | F                   | Mildly fearful|
| Mina           | Control    | 5.86  | F (n)               | Fearful       |
| Moon           | Control    | 4.25  | F                   | Mildly fearful|
| Onca           | Control    | 11.61 | F (n)               | Mildly fearful|
| Poppins        | Control    | 6.18  | F                   | Mildly fearful|
| Slash          | Control    | 5.51  | M (n)               | Fearful       |
| Andrew         | Treatment  | 4.38  | M (n)               | Mildly fearful|
| Aya            | Treatment  | 5.51  | F (n)               | Fearful       |
| Delta          | Treatment  | 4.30  | F (n)               | Mildly fearful|
| Gael           | Treatment  | 5.45  | M (n)               | Fearful       |
| Iris           | Treatment  | 5.53  | F (n)               | Fearful       |
| Jackie         | Treatment  | 6.18  | F (n)               | Mildly fearful|
| Lacey          | Treatment  | 6.17  | F                   | Mildly fearful|
| Marissa        | Treatment  | 4.42  | F                   | Mildly fearful|
| Masha          | Treatment  | 6.17  | F                   | Mildly fearful|
| Milly          | Treatment  | 4.42  | F                   | Mildly fearful|
| Prudence       | Treatment  | 5.51  | F (n)               | Fearful       |
| Ruby           | Treatment  | 4.42  | F                   | Mildly fearful|

\( n = \) neutered
Post-hoc analyses were carried out using Fisher’s least significant difference test. For the HIT the Mann–Whitney U-test was used for group comparisons due to the skewed nature of score distributions which violated the assumptions necessary for use of parametric statistics. As guidelines for statistical interpretation, we used the 0.05 level of significance to evaluate whether or not we could reject the null hypothesis. In cases where the level of significance was >0.05 and <0.10, we a priori characterized the result as marginally not significant, which justified further examination.

Results

Home room test

The group means for the home room test are summarized in Table 2. The home room scores for each group were compared using an RMANOVA with time of test (baseline, 2 weeks and 4 weeks) as a within-subject variable and group as the between-subject variable. There were no differences between groups, but there was a change with time of test \( F(2,44) = 6.18705, P < 0.005 \). A post-hoc Fisher’s test revealed that home room scores for both test 1 (\( M = 2.6042; \ SEM = 0.2369; P < 0.005 \)) and test 2 (\( M = 2.6875; \ SEM = 0.2655; P < 0.01 \)) were significantly lower than baseline (\( M = 3.2917; \ SEM = 0.279 \)).

| Group       | Time of testing |          |          |          |
|-------------|----------------|----------|----------|----------|
|             | Baseline       | Test 1   | Test 2   |
| Control     | Mean 3.17      | 2.42     | 2.54     |
|             | SEM 0.41       | 0.36     | 0.33     |
| Treatment   | Mean 3.42      | 2.79     | 2.83     |
|             | SEM 0.39       | 0.31     | 0.43     |

Behavioral measures of anxiety

Table 3 summarizes the grouped data for the OFT.

Distance traveled

The RMANOVA revealed a statistically significant main effect of level of fear \( F[1,21] = 10.5974, P < 0.005 \), which reflected less activity in the fearful animals \( [M = 6.809, \ SEM = 5.623] \) than in the mildly fearful \( [M = 29.229; \ SEM = 3.98] \). The analysis also revealed a marginally significant effect by treatment and group \( F[2,42] = 2.737, P < 0.08 \). To explore this further, we next looked at each group separately for both the OFT and HIT, using an RMANOVA on distance traveled, with time of test as a within-subject variable.

For the OFT (Table 3, raw data for which can be found in the supplementary material), the analysis for the control group revealed a significant effect of treatment day \( F[2,22] = 3.504; P < 0.05 \). For the treatment group, by contrast, the effect was non-significant \( (P > 0.1) \). We then used the Dunnett test to compare baseline with test sessions for both the control and treatment group. The analysis revealed a statistically significant decrease between baseline and test 1 \( (P = 0.05) \) and a marginally not significant decrease between baseline and test 2 \( (P < 0.09) \) for the animals on the control diet, and a non-significant increase for the animals on the test diet. Figure 1 shows that these results reflect reduced activity with repeated testing in control group, but not in the group with the test diet. In the HIT there were no significant changes.

Inactivity duration

There were two significant effects. The first was a significant main effect of level of fear \( F[1,21] = 8.325143, P < 0.01 \) and the second was a significant three-way interaction between time of test, task and group \( F[2,42] = 5.349922, P < 0.01 \). The level of fear effect was due to a significantly \( (P < 0.01) \) lower overall inactivity

Table 3 Open-field measures

| Time of testing | Group   | Distance | Inactivity | Vocalization |
|-----------------|---------|----------|------------|--------------|
|                 |         | (m)      | Duration (s) | Frequency | Frequency |
| Baseline        | Control | Mean 30.16 | 464.39 | 18.58 | 40.92 |
|                 |         | SEM 5.91  | 27.24  | 3.14   | 13.86 |
|                 | Treatment | Mean 24.56 | 501.76 | 13.58 | 42.92 |
|                 |         | SEM 10.62 | 27.56  | 3.09   | 12.63 |
| Test 1          | Control | Mean 16.62 | 514.96 | 13.83 | 35 |
|                 |         | SEM 4.87  | 24.30  | 3.23   | 13.02 |
|                 | Treatment | Mean 29.14 | 456.44 | 20.08 | 58.5 |
|                 |         | SEM 8.57  | 32.63  | 3.81   | 15.82 |
| Test 2          | Control | Mean 18.68 | 525.43 | 14.42 | 30.92 |
|                 |         | SEM 5.05  | 16.68  | 3.05   | 10.56 |
|                 | Treatment | Mean 25.17 | 490.51 | 17.17 | 40.58 |
|                 |         | SEM 7.03  | 22.45  | 3.07   | 10.01 |
duration in the mildly fearful animals ($M = 474.11$, SEM = 16.06) than in the fearful animals ($M = 554.36$, SEM = 22.707). The origins of the three-way interaction were due to open-field inactivity decreasing on the first test session in the test diet group, but increasing in the control group. In the HIT, inactivity went down in both groups on test session 1.

To further analyze the data we then looked at the OFT separately using an RMANOVA comparing the groups at each of the three time points. There was a significant level of fear effect ($F[1,21] = 9.486518$, $P < 0.05$) and a significant time of test by group interaction ($F[2,42] = 4.28005$, $P < 0.05$). For the fear effect, a post-hoc Dunnett test revealed that the anxiety effect in the open field was due to fearful cats having a greater inactivity duration compared with mildly fearful ($P < 0.05$).

To assess the origins of the interaction we first looked at the Tukey HSD test and compared each score with every other score. None of the differences achieved statistical significance. In order to assess potential trends, we then examined the data using the Fisher multiple comparison test. This revealed statistically significant increases in inactivity duration between baseline and test 1 for the control animals ($P < 0.05$). By contrast, the test diet group showed a marginally not significant decrease between baseline and test 1 ($P < 0.07$). For test 2 there was a significant increase from baseline in inactivity in the control group ($P < 0.05$) while the test diet group maintained a non-significant decrease in inactivity (Figure 2).

Inactivity frequency

The results of the analysis of inactivity frequency revealed a statistically significant effect of level of fear ($F[1, 21] = 6.812399$, $P < 0.05$) and a statistically significant three-way interaction between time of test, task and group ($F[2,42] = 3.859167$, $P < 0.05$). To determine the origins of the interaction we compared inactivity frequencies separately in the OFT and HIT using an RMANOVA. The results of the ANOVA in the OFT demonstrated a significant effect of level of fear ($F[1,21] = 7.899711$, $P < 0.05$) and a significant interaction between time of test and group ($F[2,42] = 5.053722$, $P < 0.05$). The post-hoc Dunnett test revealed that the anxiety effect in the open field was due to mildly fearful cats having a greater inactivity frequency compared with fearful cats ($P < 0.05$). For post-hoc analysis of group effects, we first compared means for each of the groups at each time point using the Tukey HSD test, which did not reveal any statistically significant differences. Next, to determine whether there were any clear trends we used the Fisher LSD test; the group effect due to an increase in inactivity frequency between baseline and test 1 for the test diet group was $P < 0.05$ and was non-significant for test 2, while the control group showed a marginally not significant decrease at test 1 $P < 0.08$ and non-significant decrease at test 2 (Figure 3). For the HIT, there were no significant differences.

Vocalization frequency

The results revealed a significant main effect of task ($F[1,21] = 11.25315$, $P < 0.005$) and no other significant
main effects or interactions. Figure 4 shows that the task effect reflected greater vocalization in the OFT than in the HIT, which is consistent with the findings in our model development study.⁶
Additional measures for the human interaction test

For each of the HIT sessions, there were four measures related to frequency and duration of approach and contact. For each of the measures, the data were negatively skewed because a large percentage of animals in both groups showed no contact and no approach (4/12 control and 6/12 treatment) (Table 4). Since the score distributions violated the assumptions necessary for use of parametric statistics, the group comparisons were made with the Mann–Whitney U-Test, which did not reveal any statistically significant effects (0.40 \( \leq P \leq 0.887 \)).

Discussion

The purpose of this study was to compare the effectiveness of a diet supplemented with L-tryptophan and α-casozepine (Royal Canin Feline Calm diet) with a control diet in reducing fear and anxiety in cats that showed fearfulness to people. We looked at three different test protocols: (1) response to an unfamiliar environment (OFT); (2) response to an unfamiliar person (HIT); and (3) assessment of fear in the home room (home room test).

Behavioral measures of fear and anxiety were based on activity, inactivity and vocalization, which were quantified using a behavioral tracking system. In the HIT, we also monitored time spent close to and in contact with the person.

For the inactivity measures in the OFT, the group on the control diet showed increases in inactivity compared with baseline while the group on the test diet showed decreases, which resulted in significant interactions between group and time of test. Although the paired comparisons using the more conservative Tukey HSD test did not reveal statistically significant differences from baseline, the Fisher multiple comparison test found a statistically significant increase over baseline on both the first and second treatment test in the control group. By contrast, the group fed the test diet showed a marginally not significant decrease between baseline and the first OFT and a non-significant decrease in test 2. Similarly, for distance traveled there was a significant decrease in distance traveled at test 1 in the control group and a non-significant increase in the test diet group.

The other measure that showed sensitivity to the test diet in the OFT was inactivity frequency. This is a measure of the number of times that an animal stops moving and then restarts; the results parallel the inactivity duration data, with the test diet group showing a significant increase in inactivity frequency over baseline at test 1 and the control group a marginally not significant decrease.

Given the link between inactivity in an unfamiliar environment and anxiety, these results provide potential evidence of the anxiolytic effectiveness of the test diet. These findings are consistent with the effects seen with diazepam treatment in our previous study, in which inactivity duration was significantly decreased and inactivity frequency increased (Figure 5).

There were no significant effects of diet in the HIT. Given the results from the OFT, this absence of significant effects in the HIT may reflect a selective benefit of the test diet in reducing anxiety, but not fear. Similarly,
we have previously shown that in the HIT, the anxiolytic drug diazepam did not promote approach to humans in fearful cats.6

The decline in vocalization frequency from the OFT to the HIT in both groups is consistent with our previous study, in which much higher levels of vocalization were seen in the OFT than the HIT. One possible interpretation is that the presence of the person increased fear, which reduced vocalization.6

We also found differences in fear level with mildly fearful cats when compared with fearful cats, showing a decrease in distance traveled and inactivity duration, and greater inactivity frequency. This result replicates our previous observation in which fearful cats were more inactive than non-fearful cats, with mildly fearful cats intermediate.6

We interpret these results to indicate that the effectiveness of the test diet in reducing overall anxiety is probably limited to moderate anxiety-provoking situations. The absence of an effect on the fear response in the presence of an unfamiliar person may indicate a dissociation between reducing anxiety associated with an unfamiliar location but not in reducing the fear toward an animate stimulus. Another possibility is that the cats used in this study, which were pre-selected based on a fearful response to people, may have been too fearful for the diet to effect any improvement. Similarly, in our previous study fearful cats given diazepam spent the greatest time away from an unfamiliar person. Therefore a future study might be designed to evaluate a greater number of cats using cats that are minimally fearful or non-fearful in the home room.

Table 4 Human interaction test measures

| Time of testing | Group   | Proximity to person | Contact with person |
|-----------------|---------|---------------------|---------------------|
|                 |         | Duration (s) | Frequency | Duration (s) | Frequency |
| Baseline        | Control | Mean       | 28.59  | 1.08        | 3.09  | 1.25        |
|                 |         | SEM        | 16.17  | 0.61        | 1.37  | 0.55        |
|                 | Treatment | Mean    | 47.32  | 1.33        | 19.34 | 1.92        |
|                 |         | SEM        | 28.22  | 0.67        | 12.19 | 1.24        |
| Test 1          | Control | Mean       | 23.67  | 2.92        | 4.59  | 2.00        |
|                 |         | SEM        | 12.13  | 1.42        | 2.42  | 1.05        |
|                 | Treatment | Mean    | 12.29  | 1.50        | 7.59  | 1.58        |
|                 |         | SEM        | 8.28   | 0.97        | 6.18  | 1.11        |
| Test 2          | Control | Mean       | 18.21  | 3.00        | 1.07  | 0.75        |
|                 |         | SEM        | 12.65  | 1.96        | 0.75  | 0.51        |
|                 | Treatment | Mean    | 16.67  | 2.17        | 5.58  | 1.42        |
|                 |         | SEM        | 10.92  | 1.35        | 4.52  | 0.89        |

Figure 5 Comparison of effects of diazepam and Royal Canin Feline Calm diet on open-field inactivity duration. Inactivity duration showed a marginally not significant decrease ($P = 0.07$) after 2-week treatment with test diet while the control group significantly increased ($P < 0.05$) (left). Similarly, inactivity duration decreased significantly after diazepam therapy (right) compared with baseline ($P < 0.001$).
assessment. In addition, as a previous study with α-casozepine found that some cats improved in the second month of treatment, further studies might consider a longer course of therapy.8

As the test diet is supplemented with both α-casozepine and L-tryptophan, the present results do not distinguish whether the beneficial effects are a result of either of these alone, or the two in combination.

Conclusions

These results are suggestive of an anxiolytic effect of the Royal Canin Feline Calm diet to placement in an unfamiliar location based on a reduction in inactivity duration and increased inactivity frequency in the Calm diet compared with an increase in inactivity duration and decreased inactivity frequency in the control diet. Future studies of longer duration might further confirm the effect of the diet on fear and anxiety.

Supplementary material

Raw data for the open-field testing from which Table 3 was compiled.

Conflict of Interest

Gary Landsberg, Dr Milgram, Christina de Rivera and Stephanie Kelly do not have any potential conflicts of interest to declare other than providing the research for this study as contracted by Royal Canin. Isabelle Mougeot is an employee of Royal Canin.

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References

1 Landsberg GM, Hunthausen W and Ackerman L. Behavior problems of the dog and cat. 3rd ed. Edinburgh: Saunders Elsevier, 2013, pp 181–210.
2 Sherman BL and Mills DS. Canine anxieties and phobias: an update on separation anxiety and noise aversions. Vet Clin North Am Small Anim Pract 2008; 38: 1081–1106.
3 Cooper LL. Fears, phobias and anxieties – cats. In: Tilley LP and Smith FWK (eds). Blackwell’s five-minute veterinary consult: canine and feline. 6th ed. Ames, Iowa: Wiley Blackwell, 2016, pp 488–489.
4 Theimer T. The biology of fear and anxiety-related behaviors. Dialog Clin Neurosci 2002; 4: 241–249.
5 Martin KM, Martin D and Shaw JK. Small animal behavioral triage: a guide for practitioners. Vet Clin North Am Small Anim Pract 2014; 44: 379–399.
6 De Rivera C, Ley J, Milgram NW, et al. Development of a laboratory model to assess fear and anxiety in cats. J Feline Med Surg 2017; 19: 586–593.
7 Kato M, Miyaji K, Ohtani N, et al. Effects of prescription diet on dealing with stressful situations and performance of anxiety-related behaviors in privately owned anxious dogs. J Vet Behav 2012; 7: 21–26.
8 Pereira GG, Fragoso S and Pires E. Effect of dietary intake of L-tryptophan supplementation on multi-housed cats presenting stress related behaviours. Proceedings of the British Small Animal Veterinary Association Congress; 2010; Birmingham, UK.
9 Beata C, Beaumont-Graff E, Coll V, et al. Effect of alpha-casozepine (Zylkene) on anxiety in cats. J Vet Behav 2007; 2: 40–46.
10 Leathwood PE. Tryptophan availability and serotonin synthesis. Proc Nutr Soc 1987; 46: 143–156.
11 van Hierden YM, Koolhaas JM and Korte SM. Chronic increase of dietary L-tryptophan decreases gentle feather pecking behavior. Appl Anim Behav Sci 2004; 89: 71–84.
12 Koopmans SJ, Ruis M, Dekker R, et al. Surplus dietary tryptophan reduces plasma cortisol and noradrenaline concentrations and enhances recovery after social stress in pigs. Physiol Behav 2005; 85: 469–478.
13 DeNapoli JS, Dodman NH, Shuster L, et al. Effect of dietary protein content and tryptophan supplementation on dominance aggression, territorial aggression, and hyperactivity in dogs. J Am Vet Med Assoc 2000; 4: 504–508.
14 Palestrini C, Minero M, Cannas S, et al. Efficacy of a diet containing caseinate hydrolysate on signs of stress in dogs. J Vet Behav 2010; 5: 309–317.
15 Miclo L, Perrine E, Driou A, et al. Characterization of alpha-casozepine, a tryptic peptide from alpha-s1 casein with benzodiazepine-like activity. FASEB J 2001; 15: 1780–1782.
16 Miyaji K, Kato M, Ohtani N, et al. Experimental verification of the effects of normal domestic cats by feeding prescription diet for decreasing stress. J Appl Anim Welf Sci 2015; 18: 355–362.