Original paper

Beneficial effects of Camelina sativa oil on behavioural (memory, anxiety, depression and social-related) manifestations and oxidative stress parameters in a mice model of irritable bowel syndrome

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Abstract

Irritable bowel syndrome (IBS) is a common chronic functional gastrointestinal impairment, which includes also a variety of psychological and psychiatric comorbidities. The oxidative stress was earlier reported as presenting an important part of the IBS complex pathophysiology. In the present study, we determined the fatty acids profile and thereafter the effects of Camelina sativa Madalina variety cold pressed oil in a zymosan-induced model of IBS. We are showing here some facilitatory effects of the Camelina sativa oil in this context, both on the behavioral and oxidative stress-related actions. Thus, all these results are suggesting some beneficial effects for the present Camelina sativa oil used in this study on behavioural (memory, anxiety, depression and social-related) manifestations and oxidative stress parameters in a zymosan-induced mice model of irritable bowel syndrome. This could be relevant for the connections between the physiological manifestations in IBS and its depressive-like or anxiolytic-like manifestations. In conclusion, Camelina sativa oil exerted facilitatory effects on both the behavioral (short memory, as tested in the Y maze task, anxiety-tested in the elevated plus maze, depression, as observed in the forced swimming test and social-dominance test) and oxidative stress-related actions (mainly manifested through a general increase of the two main antioxidant enzymes determined here: superoxide dismutase and glutathione peroxidase, as well as a decrease in the lipid peroxidation marker malondialdehyde).

Keywords

Camelina sativa oil, behavioural, memory, anxiety, depression, oxidative stress, irritable bowel syndrome.

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Introduction

Irritable bowel syndrome (IBS) is a common chronic functional gastrointestinal impairment. IBS symptomatology includes a high degree of variability mainly due to individual differences. Generally characterized by abdominal bloating and bowel habit changes such as diarrhoea, constipation, abdominal bloating, incomplete defecation accompanied by pain or burning sensation, IBS diagnosis is also based on other symptoms derived or caused by the primary symptoms such as flatulence and gas emission, mucous stools. Despite the burden and sometimes the severity of these symptoms, IBS is characterized by the absence of any structural damage which makes this disorder hard to diagnose through radiological or endoscopic changes, and laboratory tests are deficient (LONGSTRETH & al, 2006 [1]).

In 1892, Osler invented the term mucosal colitis when describing a mucosal and abdominal disorder with a high incidence in psychopathology patients. Although the psychopathology of IBS is not yet fully known, the hypothesis of the involvement of the gut-brain axis is accepted, however deficiency or impairment of brain-gut axis’s at any level could complement the picture of a condition based on both the gastrointestinal tract and the central nervous system (MAYER, 2011 [2]). In this way, IBS was recently described, by ROME IV, as a complex disorder involving gastrointestinal symptomatology as well as neurological impairments in the enteric nervous system, hypothalamic-pituitary-adrenal (HPA) axis, and also central nervous system (COJOCAŘIU & al, 2019 [3]). Previous research showed that important stress-coping mechanisms are implicated in IBS as well as several cognitive changes regarding affective spectrum behaviours (depressive and anxious behaviours) and social deficiencies (LEFTER & al, 2018 [4]).

Since any impairment in the body caused or could be caused by a signalling misuse, and oxidative stress occurs on the biggest signalling pathway (through reactive species transport and metabolism) (COJOCAŘIU & al, 2019 [3], LEFTER & al, 2018 [4]), any imbalance in the normal redox state leads to toxicity by producing free radicals and peroxides with deleterious effect on proteins, lipids and cellular DNA. In this way, the vicious circle between the production of reactive oxygen species and the inability of the body to detoxify the biological system of these species reverberate in the deficiency to repair the resulting damage. It is a fundamentally known pattern that the nervous system is susceptible to lesions caused by oxidative stress due to lipid structures and low antioxidant content.

However, oxidative stress was described as causes, components or effects of many disorders including gastrointestinal impairments (ACHITEI & al, 2013 [5], ILIE & al, 2020 [6]). In this way, previous research reported the implication of oxidative stress in several IBS animal models and also in human IBS patients (COJOCAŘIU & al, 2019 [3], LEFTER & al, 2018 [4], CHAUDHARI & al, 2014 [7], BÁTRINĂ & al, 2020 [8]).

Therapeutic approaches involving treatments with effect on symptomatology were used, given the fact that there is no specific IBS cure so far (BASNAYAKE, 2018 [9]). In this regard, besides the pharmaceutical compounds most of which are designed for other targets, the tendency is to resort to cognitive behavioural therapy, multi-component psychological therapy, dynamic psychotherapy, hypnotherapy and herbal therapy for relieving the symptoms (FORD & al, 2014 [10]).

The Camelina oil obtained from plants harvested in Romania was previously obtained, analysed (DOMIL & PIRVULESCU, 2015 [11], POPĂ & al, 2019 [12]) and it was recently demonstrated to exhibit potential in different applications in pharmaceutical and cosmetic industry due to the fact that it can stimulate the keratinocytes differentiation and turn-over, collagen synthesis, and the fibroblasts cellular division (CRACIUN & al, 2019 [13]). Thus, in this study, we determined the fatty acids profile and the effects of Camelina sativa Madalina variety cold pressed oil in a zymosan- induced model of IBS. Zymosan is a structural component, a known polysaccharide prepared from the cell wall of Saccharomyces cerevisiae (PILLEMER & ECKER, 1941 [14]), which served as a model for recognition of microbes by the innate immune system in one hand, and induces inflammatory signals in macrophages through Toll-like receptors TLR2 and TLR6, on the other hand (UNDERHILL, 2003 [15]). Moreover, zymosan interacts with dectin-1 receptors on macrophages to stimulate the release of pro-inflammatory mediators (TAGHAVI & al, 2018 [16]).

Given the fact that our group previously demonstrated the antioxidative capacity and neuropharmacological active profile of Chrysanthellum americanum extracts in a rat model of IBS (COJOCAŘIU & al, 2019 [3]) and described the alleviating potential on behavioural and oxidative stress in a complex nice model of IBS (data not published yet), our aim was to reveal Camelina sativa Madalina variety cold pressed oil effect in a zymosan- induced model of IBS on both behavioral and oxidative stress-related actions. In this way, we determined fatty acid profile of the oil by gas chromatography, then we evaluated short-term memory following behavioral testing using Y Maze test (YM), the anxious behavior using Elevated Plus Maze test (EPM) and the antidepressant effect after using Forced Swimming Test (FST). We determined the oxidative status evaluating superoxide dismutase (SOD) and glutathione peroxidase (GPx) enzymes activity and malondialdehyde (MDA) levels. Moreover, we were interested in revealing correlations between the behavioural parameters we determined and the levels of the oxidative stress markers.

Materials and Methods

1. Determination of fatty acids profile

Seeds of Camelina sativa, Madalina variety were obtained from plants cultivated without using fertilisers at Moara Domnească Teaching Farm of University of Agromonic Sciences and Veterinary Medicine Bucharest, in 2017. The pressed cold oil was obtained and analysed by using a modified earlier published method (COPOLÔVICI & al, 2017 [17], CSAKVAR & al, 2019 [18], RABA & al, 2018 [19]). The methyl esters of fatty acids contained in the samples were obtained and chromatographed as follows: 0.3 mL of oil samples together with 0.6 mL of methanol/ toluene/sulphuric acid (88/10/2, v/v/v) solution mixture were pipetted into Eppendorf tubes, vigorously vortexed and then kept at 80°C for one hour in a block heater (Stuart
block heater, Staffordshire, UK). The resulting methyl esters were extracted twice with 0.5 ml of n-heptane and analysed by gas chromatography coupled with mass spectrometry (GC-MS 2010 Plus-TQ8040 equipment, Shimadzu, Kyoto, Japan). An Optima DB 5 chromatographic capillary column (30 m length; 0.25 mm i.d.; 0.25 µm film thickness, MACHEREY-NAGEL GmbH & Co. KG, Düren, Germany) and helium as the carrier gas at a flow rate of 0.93 L min⁻¹ have been used. The injector temperature and MS source were maintained at a temperature of 250°C and 200°C, respectively. Identification of fatty acid methyl esters has been performed based on their mass spectra using commercial mass spectra libraries: NIST 14 library and Willy 09 library, and real standards. All the chemical reagents were of chromatographic grade and were bought from Sigma-Aldrich, Germany. All measurements were performed in triplicates.

2. Animals and Treatments

Male Swiss mice at an initial body mass of 30-40 g were used according to Helsinki Declaration and in accordance with the legislation of Romania and the European Union on the use of animals in biomedical experiments. The animals (n=24) were randomly assigned to 4 groups (n=6 per group): (1) control group; (2) IBS group (with zymosan administration); (3) IBS – Camelina oil group which received both zymosan and Camelina oil treatments; (4) Camelina oil group which received Camelina cold pressed oil. Zymosan (Sigma) was administered for 3 consecutive days under permanent observation. Camelina sativa oil was administered in a dose of 0.6 ml, for 5 consecutive days. In the case of combined administration group, the Camelina oil was administered 5 day after the end of zymosan treatment. Saline solution (NaCl 0.9%) was administered to control group at a volume of 2 ml / kg body weight, equivalent to other treatments volumes.

3. Behavioral testing

The animals were subjected to behavioral tests in the following order: Y maze test, Elevated-plus maze, Forced swim test, Social dominance test and were performed from the following day after all treatments were completed.

3.1. Y Maze Test

Y maze test is generally used for short-term memory assessment by evaluating the successive and alternating exploration behaviour of the three arms of the Y-shaped apparatus. The maze used in the present study consisted of three arms and an equilateral triangular central area.

3.2. Elevated plus maze

Elevated maze test is a common behavioural test used to assess anxiety-like behaviours. The test is based on the conflict that arises between the mouse's tendency to explore the labyrinth and the tendency to avoid a new stimulus that can be adversely. For assessing anxiety, the time spent in the open arms, head-dipping behaviour, time of grooming and periods of freezing were recorded, in a mice dimension specific maze.

3.3. Forced swimming test

The forced swim test is used to observe the depressive behaviour and behavioural despair. A transparent glass water-safe-filled container was used. The protocol consists in placing individuals in a transparent cylinder (30 cm in diameter, height 59 cm) filled with water (25 cm, 26°C), being forced to swim. The animals were exposed to the experimental conditions for 15 minutes. In the following say, the animals were re-exposed in a shorter period, during which the following behavioral parameters are recorded: immobility (floating), struggle (vigorous upward movements) and swimming.

3.4. Social dominance test

The social dominance test is useful for identifying deficits in social interactions and to assess animal aggressivity. In a transparent plexiglas tube measuring 45 cm in length and 4 cm in (inside) diameter, the subjects are observed regarding the tendency to cross the tube to the other end. The animals interact in the middle and the more dominant animal will show greater aggression and force its opponent out of the tube.

4. Biochemical determinations

4.1. Superoxide dismutase (SOD) activity determination

To determine the enzymatic activity of SOD, the “19160 SOD determination kit” determination kit (Sigma-Aldrich, USA) was used. This kit uses an indirect method of determination using nitroblue-tetrazolium, being the most common and efficient method. The principle of the method is based on the use of WST tetrazolium salt, highly water soluble, which produces a colored substance by reducing the superoxide anion. The rate of molecular oxygen reduction is linearly correlated with xanthine oxidase activity and is inhibited by SOD. Since the absorbance at 450 nm is proportional to the concentration of the superoxide anion, the SOD activity can be quantified by measuring the reaction medium colour intensity at 450 nm and expressed as inhibition rate (%). The specific activity of SOD was then reported considering the total soluble proteins content (determined using Bradford assay technique). The SOD activity was therefore expressed as enzyme units/ milligram (U/mg) (CIÖBICA & al, 2012 [20]).

4.2. Glutathione peroxidase (GPx) activity determination

To determine the enzymatic activity of GPx, the “Glutathione Peroxidase Cellular Activity Assay Kit” (Sigma-Aldrich, United States of America) was used. The kit uses an indirect measurement method based on the oxidation of glutathione in oxidized glutathione, a catalysed reaction of the glutathione peroxidase enzyme and coupled with the reduced glutathione recycle, the glutathione reductase-catalyzed reaction. Throughout this series of reactions, the concentration changes of the NADPH enzyme cofactor, the reduced form, are measured. Decrease in absorbance measured at 340 nm during NADPH oxidation to NADP + is directly proportional to GPx activity, since the enzyme is the modulating torque of the reaction. The reaction is carried out at 25 °C and pH 8.0 and is initiated by the addition of an organic peroxide, tert-butyl hydroperoxide (t-Bu-OH) to the reaction medium. Preparation of reagents Glutathione Peroxidase Assay Buffer, NADPH Assay Reagent, tert-butyl hydroperoxide 30 mM were performed according to the manufacturer instructions. The results were reported to the total protein samples content, and GPx activity was expressed as GPx enzyme units/ mg (U/mg) (CIÖBICA & al, 2012 [20]).

4.3. Malondialdehyde concentration determination

Malondialdehyde (MDA) concentrations were determined using thiobarbituric acid reactive substances (TBARs) assay method, previously described (CIÖBICA &
a, 2011 [21]). A 0.1 ml brain and colon extracts samples were pipetted into a centrifuge tube, 1 ml of 50% trichloroacetic acid solution and 1.1 ml of thiobarbituric acid were added. The tubes covered with a small glass funnel were kept for 20 minutes in a boiling water bath. After cooling the tubes under the jet of water, they were centrifuged for 10 minutes at 3,000 rpm and spectrophotometrically read (λ = 532 nm) versus control consisting of 0.1 ml of distilled water, 1 ml of 50% trichloroacetic acid and 1.1 ml of thiobarbituric acid (Beckman Coulter, Canada). The signal was read against an MDA standard curve and the results were expressed as nmol/mg protein.

5. Statistical analysis

The raw data was statistically analysed using one-way analysis of variance (ANOVA) using a statistical analysis software (Minitab 17). F values for which p < 0.05 were regarded as statistically significant. Post hoc analyses were performed using Tukey’s honestly significant difference test in order to compare groups.

Results and Discussion

The fatty acids profile of Camelina sativa, Madalina variety cold pressed oil has is presented in Table 1 and the main components are similar with the composition of Camelina oil of different varieties earlier published (DOMIL & PİRVELESCU, 2015 [11], POPA & al, 2019 [12], AHMED & al, 2019 [22], KANCLERZ & al, 2019 [23], KICZOROWSKA & al, 2019 [24], KURSAJ-POPOWSKA & al, 2019 [25], TEJERA & al, 2016 [26], TERPINC & al, 2012 [27]). The chromatographic analyses revealed that the oil contained the highest amount of polynsaturated fatty acids (PUFA: 54.84%), followed by monounsaturated fatty acids (MUFA 35.87%) and a low content of saturated fatty acids (SFA: 9.29%). The main components determined in our sample were: 37.77% α-linolenic acid (C18:3, ω-3), 17.07% linoleic acid (C18:2, ω-6), 16.88% cis-11-eicosenoic acid (C20:1, ω-9), 14.05% oleic acid (C18:1, ω-9), 5.52% palmitic acid (C16:0), 4.03% erucic acid (C22:1, ω-9), 2.11% stearic acid (C18:0), 1.86% arachidic acid (C20:0).

The components presented in concentration smaller than 1% were: cis-13-eicosenoic acid (C20:1, ω-9), behenic acid (C22:0), myristic acid (C14:0), palmitoleic acid (C16:1, ω-7). The fatty acid profile and the value of the ratio ω-6: ω-3 (0.45) obtained in our present study is very similar with that found by Erkkila et al. (ERKKILA & al, 2019 [28]). The values of the ratio ω-6: ω-3 was determined to be found in the range 0.42-0.58, and this is dependent on the climate conditions, quality of land, (POPA & al, 2019 [12]) presence of the heavy metals (PARK & al, 2015 [29]), etc.

In this report we tested the effects of Camelina sativa pressed oil in a zymosan-induced model of IBS. In fact, the connections between the physiological manifestations in IBS and its depressive-like or anxyolitic-like manifestations was previously suggested (COJOCARIU & al, 2019 [3], BALMUS & al, 2019 [31]) and we are showing here some facilitating effects of the Camelina sativa pressed oil in this context, both on the behavioral and oxidative stress-related actions. Thus, regarding the memory, as tested in the Y maze, we could see in Figure 1 a significant improvement (p< 0.05) in the short-term immediate spatial memory processes, as demonstrated by the increase in the spontaneous alternation percentage in IBS+ Camelina oil group, when compared to IBS group alone.

| Fatty Acid, Omega-n | Percentage of Fatty Acids (%) |
|---------------------|-----------------------------|
|                     | Present study, Madalina variety, 2017, Romania | Madalina variety, 2018, Romania (POPA & al, 2019 [12]) | Banat region, Romania (DOMIL & PİRVELESCU, 2015 [11]) | Poland (KANCLERZ & al, 2019 [23]) | UK (O’DWYER & al, 2013 [30]) | Finland (ERKKILA & al, 2019 [28]) |
| Myristic acid (C14:0) | 0.08 ± 0.03 | 0.037 | ND | ND | LL | 0.1 |
| Palmitoleic acid (C16:1, ω-7) | 0.04 ± 0.02 | 0.070 | ND | ND | LL | 0.1 |
| Palmitic acid (C16:0) | 5.52 ± 0.07 | 4.240 | 5.17 | 5.23 | 5.61 | 5.7 |
| Linoleic acid (C18:2, ω-6) | 17.07 ± 0.31 | 15.81 | 17.32 | 18.07 | 17.95 | 16.4 |
| α-Linolenic acid (C18:3, ω-3) | 37.77 ± 0.51 | 27.05 | 30.46 | 34.73 | 33.8 | 38.4 |
| Oleic acid (C18:1, ω-9) | 14.05 ± 0.57 | 10.96 | 22.06 | 15.58 | 16.37 | 13.2 |
| Stearic acid (C18:0) | 2.11 ± 0.05 | 1.86 | 3.03 | 2.33 | 2.49 | 2.5 |
| cis-13-Eicosenoic acid (C20:1, ω-9) | 0.87 ± 0.04 | 1.52 | ND | ND | ND | ND |
| cis-11-Eicosenoic acid (C20:1, ω-9) | 16.88 ± 0.30 | 14.04 | 17.16 | 14.31 | 14.24 | 14.7 |
| Arachidic acid (C20:0) | 1.36 ± 0.10 | ND | 1.37 | ND | LL | 1.5 |
| Erucic acid (C22:1, ω-9) | 4.03 ± 0.06 | ND | 3.12 | ND | LL | 3.4 |
| Behenic acid (C22:0) | 0.23 ± 0.03 | 0.28 | ND | ND | LL | 0.3 |
| Nervonic acid (C24:1, ω-9) | ND | ND | 1.21 | ND | LL | 0.6 |
| SFA | 9.29 | 6.42 | 9.84 | 7.56 | 11.78 | 10.2 |
| MUFA | 35.87 | 26.59 | 40.97 | 34.22 | 31.97 | 32.7 |
| PUFAs | 54.84 | 42.86 | 47.78 | 58.8 | 55.38 | 57.1 |
| ω-6: ω-3 | 0.45 | 0.58 | 0.56 | 0.52 | 0.56 | 0.42 |

ND, not detected, LL: low levels (not show)
Figure 1. The effects of Camelina oil administration in a mice model of IBS on short term memory observed in Y Test. The results are expressed as means ± S.E.M. The columns that do not share a letter are significantly different (a,b – Tukey HSD test).

In addition, in regards to the depressive-like behaviour, there are significant decreases (p < 0.05) in the floating time in the IBS+ Camelina oil group, when compared to IBS group alone (Figure 3a), as well as in the struggle time, as it can be seen in the Figure 3b below:

Figure 2. The effects of Camelina oil administration in a mice model of IBS on anxious behaviour observed in Elevated Plus maze: a. open arms time; b. open arms entries; c. closed arms entries; d. grooming behaviour durations. The results are expressed as means ± S.E.M. The columns that do not share a letter are significantly different (a,b – Tukey HSD test).

Figure 3. The effects of Camelina oil administration in a mice model of IBS on anxious behaviour observed in Forced Swim Test: a. struggle behaviour duration; b. floating behaviour duration. The results are expressed as means ± S.E.M. The columns that do not share a letter are significantly different (a,b – Tukey HSD test).
Beneficial effects of Camelina sativa oil on behavioural (memory, anxiety, depression and social-related) manifestations

Also, there were significant ameliorations of the social manifestations, as it can be seen below in the IBS + Camelina oil group, as compared only with the rats which had the experimental model of IBS (Figure 4).

![Figure 4](image)

**Figure 4.** The effects of Camelina oil administration in a mice model of IBS on social domination and aggressive behaviour observed in Social Dominance Test. The results are expressed as means ± S.E.M. The columns that do not share a letter are significantly different (a,b – Tukey HSD test).

Moreover, when it comes to the effects of this extract on the oxidative stress status, as it can be seen in the Figures 5, 6, 7, and 8, our results are suggesting an antioxidant effect. Significant differences between the studied groups were observed regarding the SOD activity (Overall ANOVA p<0.01) (Figure 6) and MDA levels (Overall ANOVA p<0.01, p<0.01) (Figure 8) in both brain and bowel tissues extracts. Relevant, but not statistically significant overall changes were observed for GPx activity (Figure 7) and total soluble proteins (Figure 5).

![Figure 5](image)

**Figure 5.** The effects of Camelina oil administration in a mice model of IBS on the total soluble proteins levels evaluated from brain and bowel tissues. The results are expressed as means ± S.E.M. The columns that do not share a letter are significantly different (a,b,c,d – Tukey HSD test).

![Figure 6](image)

**Figure 6.** The effects of Camelina oil administration in a mice model of IBS on the SOD specific activity evaluated from brain and bowel tissues. The results are expressed as means ± S.E.M. The columns that do not share a letter are significantly different (a,b,c – Tukey HSD test).
Thus, all these results are suggesting some beneficial effects for the *Camelina sativa* Madalina variety oil used in this study on behavioural (memory, anxiety, depression and social-related) manifestations and oxidative stress parameters in a zymosan-mice model of irritable bowel syndrome, even though some studies regarding cognitive deficiencies in IBS patients turned out to be contradictory, some authors considering the data as being not entirely clear on this matter (BERRILL & al, 2013 [32], FARUP & HESTAD, 2015 [33]). This could be relevant for the connections between the physiological manifestations in IBS and its depressive-like or anxiolyitic-like manifestations, in accordance with specialized literature in what concerns the human studies (FARUP & HESTAD 2015 [33], CRYAN & al, 2002 [34]), like we previously discussed and published (COJOCARIU & al, 2019 [3], LEFTER & al, 2018 [4], BALMUS & al, 2019 [31]), since we are showing here some facilitating effects of this specific *Camelina* oil in this context, on both the behavioral and oxidative stress-related actions.

Also, this could be connected with the fact that this oil was previously suggested to have laxative effects (KANCLERZ & al, 2019 [23], FRASER & al, 2017 [35]), as this correlates for example with increased MDA concentration in the oil group (e.g. as in increased membrane permeability), when compared for example with the controls (Figure 8). Regarding *Camelina sativa*, this specific plant is known for being a rich source of omega-3 fatty acids, and one possible mechanism regarding its protective effect could be linked to decreased inflammation, especially when recent studies revealed beneficial effect in immunoregulatory function, with a decrease in mRNA expression levels of IFN γ after a 12-week diet enriched with *Camelina sativa* oil. Moreover, there are documented benefits in terms of cardiovascular health, given the reduced levels of ICAM1 expression induced by a lean fish diet, when compared to a fatty fish or a control diet (DE MELLO & al, 2019 [36]).

Recent studies indicated that polyphenols are implicated in reversing oxidative stress, alleviating some cognitive and mood disorders (SAJILATA & al, 2008 [37], SANMUKHANI & al, 2014 [38]). Even though *Camelina sativa* oil did not cause obvious changes in oxidative stress status in subjects with impaired glucose metabolism (ERKKILA & al, 2019 [28]), this plant is considered to be more stable toward oxidation, with an antioxidant activity high enough to be considered a potential source of bioactive antimicrobial preservative used in pharmaceutical applications (KUMAR & al, 2015 [39]).

Oxidation stability is also given by the increased level of gamma-tocopherol, chlorogenic acid, which slows the release of glucose into the bloodstream (ABRAMOVIC & al, 2007 [40], HRASTAR & al, 2011 [41], and it is known for increasing serum concentrations of eicosapentaenoic acid (EPA) and docosahexaenoic acid (DHA) (CUTULI, 2017 [42]), with beneficial effects on cardiovascular disease (MOZAFFARIAN, 2008 [43]), autoimmune (CALDER, 2006 [44]), mental disorders (McCANN & AMES, 2005 [45]) and fetal development (TEJERA & al, 2016 [26]). Even though two strains of *Camelina sativa* crop have been genetically modified to produce oil containing EPA or DHA, to compensate for the low synthesis capacity in humans, these seed oils could be a valid replacement of fish oil to provide a sustainable solution for EPA and DHA demands in human consumption (WEST & al, 2019 [46]).

Also, there is an obvious link between PUfAs consumption and modulatory effects on expression and secretion of cytokines and chemokines (IL-1, IL-6 and TNF alfa) in both humans and pigs (TARANU & al, 2014 [47]), while there is a stimulatory or an inhibitory effect of ω-3 fatty acids on the pro-inflammatory cytokines in mice (CALDER, 1996 [48]). *Camelina sativa* oil cakes consumption in pigs indicated an obvious modulation between Th1/Th2 cytokine balance, and induction or suppression on one of these two cytokines, could be easily be used in nutritional treatment strategies and immunological approaches (TARANU & al, 2014 [47]). In addition, the aminoacid of the *Camelina* oil content could also play a role in this context (BÂTRÎNA & al, 2020 [8]).
Conclusions

Our data presented here is suggesting some beneficial effects of the Camelina oil on behavioural (short term memory – tested in the Y maze task, anxiety – tested in the elevated plus maze, depression – as observed in the forced swimming test and social behaviour observed in social-dominance test) manifestations and oxidative stress parameters (the two main antioxidant enzymes determined here: superoxide dismutase and glutathione peroxidase, as well as a reduction in the lipid peroxidation marker malondialdehyde) in a zymosan-induced mice model of irritable bowel syndrome.

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