# Conjugated linoleic acid and fatty acid binding protein as antioxidants

PIERGIACOMI, V.A.<sup>1</sup>; PALACIOS, A.

#### SUMMARY

Studies were carried out to determine the effect of conjugated linoleic acid (CLA) and rat liver cytosolic protein enriched in fatty acid binding protein (FABP) on the non-enzymatic lipid peroxidation of rat liver microsomes. The inhibition of lipid peroxidation was more evident when the FABP containing fraction obtained from CLA-group was used with either kind of microsomes (CLA and control). The chemiluminescence and polyunsaturated fatty acid composition of rat liver microsomes changed after CLA treatment. When native and peroxidized microsomes obtained from control group were compared, the most affected polyunsaturated fatty acids were: C18:2, C18:3 and C20:4, while in CLA-group C20:4 was mainly peroxidized The simultaneous analysis of chemiluminescence and fatty acid composition demonstrated that CLA and FABP play a role protecting rat liver microsomes against the harmful effect of lipid peroxidation.

*Key words*: (Conjugated Linoleic Acid), (Fatty Acid Binding Protein), (lipid peroxidation), (liver), (microsomes)

Cátedra de Bioquímica, Facultad de Ciencias Veterinarias, Universidad Nacional de La Plata CC 296, B1900 AVW, La Plata, Argentina. Fax: 54-221-4257980. E-mail: vpier@fcv.unlp.edu.ar

<sup>1</sup> Assistant of the Comisión de Investigaciones Científicas y Técnicas de la Prov. de Buenos Aires (CIC) Argentina.

Recibido: marzo 2006 - Aceptado: agosto 2006 - Publicación on line: agosto 2006.

InVet. 2006, 8(1): 139-148 ISSN(papel): 1514-6634 ISSN (on line) 1668-3498

### Ácido linoleico conjugado y proteína transportadora de ácidos grasos como antioxidantes

#### RESUMEN

El objetivo del trabajo fue determinar el efecto del ácido linoleico conjugado (ALC) y de la proteína citosólica de hígado de rata enriquecida en Proteína Transportadora de Ácidos Grasos (PTAG), sobre la peroxidación no enzimática de lípidos de microsomas hepáticos de rata. Luego de la incubación de éstos en un sistema ascorbato-Fe<sup>++</sup> se observó que el total de cpm/mg de proteina originada por quimioluminiscencia fue menor en los microsomas obtenidos de las ratas del grupo ALC respecto a los del grupo control. Cuando la fracción PTAG obtenida del grupo ALC fue agregada a la peroxidación de microsomas de ambos grupos de animales ALC y control, la inhibición de la lipoperoxidación fue más evidente. Además se encontró que ambas fracciones PTAG, tanto la obtenida de animales del grupo ALC como la obtenida del grupo control, tuvieron mayor efecto como antioxidantes cuando se usaron microsomas ALC respecto a microsomas control. La composición de ácidos grasos de los microsomas cambió luego del tratamiento con ALC. Comparando microsomas nativos y peroxidados obtenidos de grupo control, los ácidos grasos polinosaturados afectados fueron: C18:2 (n-6), C18:3 (n-3) y C20:4 (n-6), y en el grupo ALC fue C20:4 (n-6). El análisis de quimioluminiscencia y composición de ácidos grasos demostró que tanto el ALC como la PTAG juegan un rol en la protección de los microsomas de hígado de rata contra los efectos nocivos de la lipoperoxidación.

*Palabras clave*: (Ácido linoleico conjugado), (proteína transportadora de ácidos grasos), (peroxidación lipídica), (hígado), (microsomas).

#### INTRODUCTION

of The oxidative degradation polyunsaturated fatty acids in bio-membranes can be initiated by free radicals and can proceed in a chain reaction causing considerable membrane damage or producing lipid peroxides<sup>31</sup>. Conjugated linoleic acid (CLA) is a collective term referring generally to a mixture of positional and geometric conjugated dienoic isomers of linoleic acid. Numerous CLA isomers are found in milk-fat, cheese, and beef <sup>9,29</sup>. The cis-9, trans-11, isomer of CLA (the principal dietary form) is produced in the rumen of cattle and other ruminant animals during the

microbial biohydrogenation of linoleic and linolenic acids <sup>13,20,23,36,39,42</sup>. It is evident that CLA significantly reduced tumor incidence as well as tumor cell proliferation<sup>14-16,19</sup>. The mechanisms through which CLA inhibits tumorigenesis are moot.<sup>32</sup>. In different studies it has been demonstrated that CLA is a potent antioxidant and that c-9, t-11 isomer is selectively incorporated into cellular phospholipid, which may, at least, in part, explain the anticarcinogenic activity of CLA as antioxidant mechanism<sup>33</sup>. One might speculate that the inhibition of carcinogenesis by CLA could result from the combined effects of a number of CLA activities, possibly including

> InVet. 2006, 8(1): 139-148 ISSN(papel): 1514-6634 ISSN (on line) 1668-3498

direct effects of one or more CLA isomers/ metabolites on cell differentiation and effects of one or more CLA isomers on prostaglandin metabolism which may also influence cancer development at some sites<sup>5,27,34</sup>. Animal fat, which has been maligned for so long, may actually constitute a potent therapeutic component to our diet11, 24. In rat liver, 60 % of cytosolic long chain fatty acids are associated with a low molecular weight protein 15 KDa named Fatty Acid Binding Protein (FABP) 7. In vitro studies of our laboratory have shown that cytosolic fraction enriched with FABP contains antioxidant properties against lipid peroxidation <sup>8</sup>. For this reason, it was of interest to analyze the effect of this protein on the lipid peroxidation process of liver microsomes isolated from rat treated with CLA. The present study aims to compare the polyunsaturated fatty acid composition and non enzymatic lipid peroxidation of rat liver microsomes obtained from animals supplemented with conjugated linoleic acid with control animals, and to analyze the effect of Fatty Acid Binding Protein on the lipid peroxidation process.

#### MATERIAL AND METHODS

Female Wistar AH/HOK rats were obtained from Laboratory Animal Facility, Facultad de Ciencias Veterinarias, Universidad Nacional de La Plata. Octadecadienoic acid conjugated (CLA) and butylated hydroxytoluene (BHT) were from the Sigma Chemical Co. (St. Louis, MO). BSA (fractionV) was obtained from Wako Pure Chemicals Industries Ltd; Japan. Standards of fatty acids methyl esters were generously supplied by NU Check Prep, Inc, Elysian, MN, USA. L [+] ascorbic acid and boron trifluoridemethanol complex were from Merck. All other reagents and chemicals were of analytical grade from Sigma.

#### 1. Animal treatment

Seven- week- old female Wistar AH/HOK rats, weighing 120-137 g were used. Two groups of five rats were considered, Group A received conjugated linoleic acid (CLA), Group B was used as control. All rats were fed commercial rat chow and water ad-libitum. Group A received a daily oral dose of CLA (30 mg) for 10 days. On day 11, rats were sacrificed by cervical dislocation and the liver was rapidly removed.

## 2. Preparation of microsomes and cytosolic fraction enriched in fatty acid binding protein

Liver was cut into small pieces and washed extensively with 0.15 M NaCl. A homogenate 30% (w/v) was prepared in a solution (0.25 M sucrose, 10 mM Tris-HCl pH 7.4) using the Potter-Elvejhem homogenizer. The homogenate was spun at 10,000 g for 10 min. The supernatant (3 ml) obtained was applied to a Sepharose 4B column (1.6 x 12 cm) equilibrated and eluted with 10 mM Tris-HCl (pH 7.4), 0.01 % NaN<sub>2</sub>. The microsomal fraction appearing in the void volume (8-12 ml) was brought to 0.25 M sucrose by addition of solid sucrose. With respect to concentrations and activities of certain microsomal enzymes, the quality of this microsomal preparation has a similar composition to that obtained by ultracentrifugation<sup>41</sup>.

The soluble fraction (14-22 ml) was concentrated and filtrated on Sephadex G-75 (3x39 cm) eluted with buffer Tris-HCl 0.01 M, pH 7.4, 0.01% NaN3, flow rate: 90 ml/h, to obtain two large fractions F1 and F2. F1 corresponded to proteins with molecular weight greater than 68 KDa and F2 represented proteins in the range 12-16 KDa that are enriched with FABP, this fraction was concentrated by ultrafiltration in an Amicon

InVet. 2006, 8(1): 139-148 ISSN(papel): 1514-6634 ISSN (on line) 1668-3498

cell containing a PM10 membrane. All operations were performed at 4 °C and under dim light.

# **3.** Lipid peroxidation of rat liver microsomes

Chemiluminescence and lipid peroxidation were initiated by adding ascorbate to microsomes <sup>44</sup>. Microsomes (1 mg protein) were incubated at 37 °C with 0.01M phosphate buffer pH 7.4, 0.4 mM ascorbate, final volume 2 ml. Phosphate buffer was contaminated with sufficient iron to provide the necessary ferrous or ferric iron for lipid peroxidation <sup>40</sup>. In each assay, 0.25, 0.50 and 1 mg of protein of FABP obtained from two groups (ALC and control) were used. Organelle preparations, which lacked ascorbate, were carried out simultaneously. Membrane light emission was determined over a 120 min period, chemiluminescence was recorded as cpm every 10 min and the sum of the total chemiluminescence was used to calculate cpm/mg protein. Chemiluminescence was measured as counts per min in a liquid scintillation analyzer Packard 1900 TR equipment with for а program chemiluminescence.

#### 4. Fatty acid analysis

Microsomal lipids were extracted with chloroform/methanol (2:1 v/v) <sup>12</sup> from native or peroxidized membranes. Fatty acids were transmethylated with  $F_3B$  in methanol at 60 °C for 3 h. Fatty acids methyl esters were analyzed with a GC-14 A gas chromatograph (Shimadzu Kyoto, Japan) equipped with a packed column (1.80 m x 4 mm i.d.) GP 10 % DEGS-PS on 80/100 Supelcoport. Nitrogen was used as a carrier gas. The injector and detector temperatures were maintained at 250 °C, the column temperature was held at 200 °C. Fatty acid methyl ester peaks were identified by

comparison of the retention times with those of standard.

#### 5. Other methods

Proteins were determined by Lowry's method <sup>28</sup>.

#### 6. Statistical analysis

Results are expressed as mean  $\pm$  S.D. of five independent determinations. Data were evaluated statistically by one-way analysis of variance (ANOVA) and Tukey test. The statistical criterion for significance was selected at different P values and indicated in each case.

#### RESULTS

The polyunsaturated fatty acid composition of total lipids isolated from rat liver microsomes obtained from CLA and control groups is shown in Table 1. The effect of conjugated linoleic acid on the total fatty acid composition of native microsomes showed a significant decrease in linoleic acid C18:2 (n-6) when compared with the control group. When peroxidized microsomes were compared with natives, the arachidonic acid C20:4 (n-6) decreased in the two groups, the linoleic acid C18:2 (n-6) and the linolenic acid C18:3 (n-3) decreased only in the control group. Statistically significant differences in the content of C18:2 (n-6), C18:3 (n-3) and C20:4 (n-6) with respect to the control group were observed in CLAgroup when peroxidized microsomes were compared.

The incubation of rat liver microsomes in the presence of ascorbate -Fe<sup>++</sup> resulted in the peroxidation of membranes as evidenced by the emission of light (chemiluminescence). The lipid peroxidation process measured as light emission decreased when microsomes isolated from CLA-group were compared with

> InVet. 2006, 8(1): 139-148 ISSN(papel): 1514-6634 ISSN (on line) 1668-3498

Fatty acid	CLA group		Control group	
	Native	Peroxidized	Native	Peroxidized
18:2 n-6	$7.232 \pm 0.751^{\rm h}$	$5.406 \pm 2.316^{e}$	$9.374 \pm 1.064^{\text{b,h}}$	$1.516 \pm 0.128^{\text{b,e}}$
18:3 n-3	$0.715 \pm 0.316$	$0.554 \pm 0.216^{\rm f}$	$1.275\pm0.516^{\text{c}}$	tr <sup>c,f</sup>
20:4 n-6	$19.915 \pm 4.480^{a}$	$10.012 \pm 2.543^{\rm a,g}$	$21.115\pm4.031^{\text{d}}$	$3.796 \pm 1.523^{d,g}$
22:6 n-3	tr	tr	tr	tr

Table 1. Polyunsaturated fatty acid composition (area %) of rat liver microsomes native and peroxidized from CLA and control groups

Data are given as the mean  $\pm$  S.D. of five experiments. Native: without incubation in ascorbate Fe<sup>++</sup> system; peroxidized: with incubation in the presence of 0.4 mM ascorbate. Statistically significant differences between native vs. peroxidized microsomes of the two groups are indicated by p<sup>a</sup> < 0.03, p<sup>b</sup> and p<sup>c</sup> < 0.0001, p<sup>d</sup> < 0.002. Statistically significant differences between peroxidized microsomes are indicated by p<sup>e</sup> < 0.04, p<sup>f</sup> < 0.05 and p<sup>g</sup> < 0.01. Statistically significant differences between native microsomes of the two groups are indicated by p<sup>b</sup> < 0.05. tr: trace.

Table 2. Lipid Peroxidation of rat liver microsomes-control. Effect of FABP containing fraction obtained from control and CLA treated groups.

Microsomes	FABP fraction (mg)	Light emission (cpmx10 <sup>-3</sup> )	Inhibition %
Native	00	2300± 205	00
+FABP control	0.25 0.50 1.00	$\begin{array}{c} 1955 \pm 172 \\ 1655 \pm 102 \\ 1311 \pm 133 \end{array}$	$15 \pm 2$ $43 \pm 5^{b}$ $28 \pm 4^{a}$
+ FABP-CLA	0.25 0.50 1.00	$\begin{array}{c} 1840 \pm 230 \\ 1380 \pm 178 \\ 966 \pm 127 \end{array}$	$\begin{array}{c} 40 \pm 5 \\ 20 \pm 3 \\ 58 \pm 4 \\ \end{array}$

Data are given as the mean  $\pm$  S.D. of five experiments. Statistically significant differences between control and supplemented groups are indicated by  $p^a < 0.03$  and  $p^b < 0.02$ .

microsomes obtained from the control group. Moreover, light emission was inhibited by addition of the FABP containing fraction; significant differences were found when the activity of FABP from CLA and the control group was compared. The inhibition of lipid peroxidation microsomes (obtained from CLA and control groups) as a function of FABP containing fraction is illustrated in Tables 2 and 3 respectively. In both cases inhibition of light emission by the FABP containing fraction was protein concentration dependent. As shown in

InVet. 2006, 8(1): 139-148 ISSN(papel): 1514-6634 ISSN (on line) 1668-3498 Tables 2 and 3, the inhibition of lipid peroxidation was more evident when the FABP containing fraction obtained from CLA treated group was used in either kind of microsomes (CLA and control).

Fig 1 shows the total light emission in the organelles obtained after lipid peroxidation of microsomes from CLA and control groups. Both FABPs (CLA and control) were more effective as antioxidant using CLA rather than control microsomes. These results are illustrated when comparing Fig. 1A with Fig.1B statistically.

Microsomes	FABP fraction (mg)	Light emission (cpmx10 <sup>-3</sup> )	Inhibition %
Natives	00	$1349 \pm 196$	00
+FABP control	0.25	$1052 \pm 210$	$22 \pm 3^{a}$
	0.50	715 ± 92	$47 \pm 5^{b}$
	1.00	499 ± 107	$63 \pm 7$
+ FABP-CLA	0.25	$768 \pm 197$	$43 \pm 4^{a}$
	0.50	$418 \pm 89$	$69 \pm 7^{b}$
	1.00	$378 \pm 72$	$72 \pm 6$

Table 3. Lipid Peroxidation of rat liver microsomes-CLA. Effect of FABP containing fraction obtained from control and CLA treated groups.

Data are given as the mean  $\pm$  S.D. of five experiments. Statistically significant differences between control and supplemented groups are indicated by  $p^a < 0.002$  and  $p^b < 0.01$ .

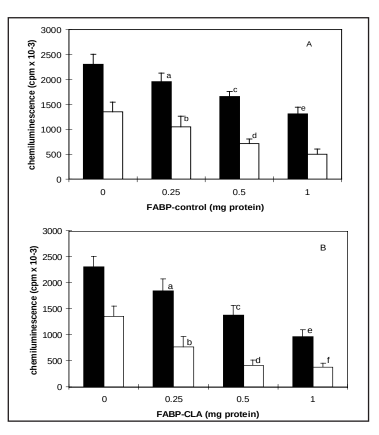


Fig.1. Effect of fatty acid binding protein-containing fraction obtained from control (A) and CLA (B) groups, on ascorbate-Fe<sup>++</sup> lipid peroxidation induced in vitro. Microsomal membranes from control ( $\blacksquare$ ) and CLA ( $\square$ ) supplemented animals. Data are given as the mean  $\pm$  S.D. of five experiments. Statistically significant differences between control and supplemented FABP in lipid peroxidation of microsomes are indicated by p<sup>a</sup>< 0.006, p<sup>b</sup>, p<sup>c</sup> and p<sup>c</sup>< 0.001, p<sup>d</sup>< 0.004 and p<sup>f</sup>< 0.002.

InVet. 2006, 8(1): 139-148 ISSN(papel): 1514-6634 ISSN (on line) 1668-3498

#### DISCUSSION

Although considerable research has already been performed to characterize the changes in structure, composition and physical properties of membranes subjected to oxidation<sup>18, 22,37,38,43</sup>, it is important to know how biological compounds with antioxidant properties contribute to the protection of specialized membranes against deleterious effects produced by reactive oxygen species and other free radicals. Evidence leading to the recognition of the anticarcinogenic activity of the conjugated dienoic derivatives of linoleic acid (CLA) has been reviewed. New data indicated that CLA has a potent antioxidant activity <sup>19,21</sup>. Because the c-9, t-11 CLA isomer is esterified in phospholipid, it may represent a heretofore-unrecognized in situ defense mechanism against membrane attack by oxygen radicals <sup>4,33</sup>. It appears to act as a chain-breaking antioxidant by trapping chain propagating free radicals 6,17. The carcinogenesis inhibition could result from the combined effects of CLA activities <sup>1</sup>. Lipid peroxidation studies in vitro are useful for the elucidation of possible mechanism of peroxide formation in vivo 3,25 since the high concentration of polyunsaturated fatty acids in membranes causes susceptibility to lipid peroxidative degradation <sup>35</sup>. The present study showed that rat liver microsomes obtained from CLA group are protected against lipid peroxidation when compared with similar membranes obtained from control rats, as revealed by the results obtained by chemiluminescence and polyunsaturated fatty acid composition. Moreover, significant differences of fatty acid composition between CLA with respect to the control group were found. Our results showed that CLA decreases polyunsaturated fatty acids, for this reason another author evoke that CLA does not appear

InVet. 2006, 8(1): 139-148 ISSN(papel): 1514-6634 ISSN (on line) 1668-3498 to act as an antioxidant, its ability to decrease polyenoic fatty acid concentration could decrease the formation of highly cytotoxic lipid peroxidation products <sup>2</sup>. In contrast, another investigator suggested an antioxidant mechanism of CLA 24. Nevertheless, the same authors postulate that CLA may induce lipid peroxidation in vivo <sup>26</sup>. It has been recently proposed that CLA increases the concentration of FABP by activation of specific receptors. Conjugated linoleic acid is a ligand and activator of Peroxisome-Proliferator Activated Receptor (alpha) and its effects on lipid metabolism may be attributed to transcriptional events associated with this nuclear receptor. Levels of liver fatty acid-binding protein were induced by CLA in hepatoma cells <sup>10,30</sup>. We agree that perhaps this mechanism is involved in improving the antioxidant activity of FABP, because the inhibition of light emission by FABP was protein concentration dependent. In conclusion, our results are consistent with the hypothesis that formation and action of CLA represents a previously unrecognized in situ defense mechanism against membrane attack by oxygen radicals. Further studies are needed to more adequately evaluate these observations.

#### CONCLUSIONS

The oxidative degradation of polyunsaturated fatty acids in bio-membranes can be initiated by free radicals and can proceed in a chain reaction causing considerable membrane damage or producing lipid peroxides. This article is a contribution to knowledge in the field. Studies were carried out to determine the effect of CLA and rat liver cytosolic protein enriched in FABP on the nonenzymatic lipid peroxidation of rat liver microsomes.

#### ACKNOWLEDGMENTS

We thank Cesar Arcemis for his technical assistance.

#### REFERENCES

- AGATHA, G.; ORLOV, A.; BROSH, A. 2004. Conjugated linoleic acid modulation of cell membrane in leukemia cells. Cancer Lett 209: 87-103.
- 2.BASU, S.; SMEDMAN, A.; VESSBY, B. 2000. Conjugated linoleic acid induces lipid peroxidation in humans. FEBS Lett. 468 : 33-36.
- 3.BENZI, G. 1988. Peroxidation, energy transduction and mitochondria during aging. John Libery Eurotext, London 51-117.
- BERGAMO, P.; LUONGO, D.; ROSSI, M. 2004. Conjugated linoleic acid-mediated apoptosis in Jurkat T cells involves the production of reactive oxygen species. Cell Physiol Biochem 14: 57-64.
- BERNAL-SANTOS, G.; PERFIELD, J.W. 2<sup>ND</sup>; BARBANO, D.M.; BAUMAN, D.E.; OVERTON, T.R. 2003. Production responses of dairy cows to dietary supplementation with conjugated linoleic acid (CLA) during the transition period and early lactation. J Dairy Sci 86 : 3218-3228.
- BROWN, J.M.; BOYSEN, M.S.; CHUNG, S.; FABIYI, O.; MORRISON, R.F.; MANDRUP, S.; MCINTOSH, M.K. 2004. Conjugated linoleic acid induces human adipocyte delipidation- Autocrine/ paracrine regulation of MEK/ERK signaling by adipocytokines. J Biol Chem 279: 26735-26747.
- CATALÁ, A. 1993. Interaction of fatty acids, acyl CoA derivates and retinoids with microsomal membranes: effect of cytosolic proteins. Mo. Cell Biochem 120: 89-94.
- 8.CATALÁ, A.; CERRUTI, A.; ARCEMIS, C. 1995. Inhibition of microsomal chemiluminiscence by cytosolic fractions containing fatty acid binding protein. Arch Physiol Biochem 103: 39-43.
- 9. DORAZIO, N.; FICONERI, C.; RICCIONI, G.; CONTI, P.; THEOHARIDES, T.C.; BOLLEA,

M.R. 2003. Conjugated linoleic acid: a functional food? Int J Immunopathol Pharmacol 16:215-220.

- DUPLUS, E.; GLORIAN, M.; FOREST, C. 2000. Fatty acid regulation of gene transcription. J Biol Chem 275: 30749-30752.
- FLINTOFF-DYE, N.L.; OMAYE, S.T. 2005. Antioxidant effect of conjugated linoleic acid isomers in isolated human low- density lipoproteins. Nutr Res 25 (1): 1-12.
- FOLCH, J.; LEES, N.; SLOANE-STANLEY, GA
  1957. A simple method for the isolation and purification of total lipids from animal tissues. J Biol Chem 226:497-509.
- FUKUDA, S.; FURUYA, H.; SUZUKI, Y.; ASANUMA, N.; HINO, T. 2005. A new strain of butyrivibrio fibrisolvens that has high ability to isomerize linoleic acid to conjugated linoleic acid. J Gen Appl Microbiol 51: 105-113.
- 14. HA, Y.L.; STORKSON, J.; PARIZA, M.W. 1999. Inhibition of benzo(a)-pyrene-induced mouse forestomach neoplasia by conjugated dienoic derivatives of linoleic acid. Cancer Res 50:1097-1101.
- 15. HOFFMANN, K.; BLAUDSZUN, J.; BRUNKEN, C.; HOPKER, W.W.; TAUBER, R.; STEINHART, H. 2005. New application of a subcellular fractionation method to kidney and testis for the determination of conjugated linoleic acid in selected cell organelles of healthy and cancerous human tissues. Anal Bioanal Chem 381: 1138-1144.
- 16. IP, C.; SING, M.; THOMPSON, H.J.; SCIMECA, J.A. 1994. Conjugated linoleic acid suppresses mammary carcinogenesis and proliferative activity of the mammary gland in the rat.Cancer Res 54: 1212-1215.
- IP, M.M.; MASSO-WELCH, P.A.; IP, C. 2003. Prevention of mammary cancer with conjugated linoleic acid: role of the stroma and the epithelium. J Mammary Gland Biol Neoplasia 8:103-118.
- KAGAN, V.E. 1988. Lipid Peroxidation in biomembranes.CRC Press, Boca Raton, FL 55-146.
- 19. KELLY, G. 2001.Conjugated Linoleic Acid: A Review. Alt Med Rev 6:367-382.

InVet. 2006, 8(1): 139-148 ISSN(papel): 1514-6634 ISSN (on line) 1668-3498

- 20. KEPLER, C.R.; HIRONS, K.P.; MCNEILL, J.J.; TOVE, S.B. 1966 Intermediates and products of the biohydrogenation of linoleic acid by Butyrivibrio fibrisolvens. J Biol Chem 241: 1350-1354.
- KHARAZI, P.R.; PEYVAST, G. 2005. Natural antioxidant and anticarcinogenic compounds from two varieties of Iranian garlic. Asian J. Chem. 17: 219-223
- 22. KIM, H.K.; KIM, S.R.; AHN, J.Y.; CHO, I.J.; YOON, C.S.; HA, T.Y. 2005. Dietary conjugated linoleic acid reduces lipid peroxidation by increasing oxidative stability in rats. J. Nutr. Sci. Vitaminol 51: 8-15.
- 23. KRAMER, J.K.G.; FELLNER, V.; DUGAN, M.E.R.; SAUER, F.D; MOSSOBA, M.M.; YURAWECZ, M.P. 1997. Evaluating acid and base catalysts in the methylation of milk and rumen fatty acids with special emphasis on conjugated dienes and total trans fatty acids. Lipids 32: 1219-1228.
- KRITCHEVSKY D. 2000. Antimutagenic and some other effects of conjugated linoleic acid. Brit J Nutrition 83: 459-465.
- 25. LAI, C.H.; YIN, J.D.; LI, D.F.; ZHAO, L.D.; QIAO, S.Y.; XING, J.J. 2005. Conjugated linoleic acid attenuates the production and gene expression of proinflammatory cytokines in weaned pigs challenged with lipopolysaccharide. J Nutr 135: 239-244.
- 26. LEUNG, Y.H.; LIU, R.H. 2005. Trans-10, cis 12 - conjugated linoleic acid isomer exhibits stronger oxyradical scavenging capacity than cis-9,trans-11conjugated linoleic acid isomer. J Agric Food Chem 48: 5469-5475.
- 27. LIEW, C.; SCHUT, H.A.J.; CHIN, S.F.; PARIZA, M.W.; DASHWOOD, R.H. 1995. Protection of conjugated linoleic acids against 2-amino-3methylimidazol (4-5-f) quinoline-induced colon carcinogenesis in the F344 rat: a study of inhibitory mechanism. Carcinogenesis 16:3037-3043.
- LOWRY, O.H.; ROSEBROUGHT, N.J.; FARR, A.L; RANDALL, R.J. 1951. Protein measurement with Folin phenol reagent. J Biol Chem 193: 265-275.
- 29. LUNA, P.; DE LA FUENTE, M.A.; JUAREZ, M. 2005. Conjugated linoleic acid in processed cheeses

InVet. 2006, 8(1): 139-148 ISSN(papel): 1514-6634 ISSN (on line) 1668-3498 during the manufacturating stages. J Agric Food Chem 53: 2690-2695.

- 30. MOYA-CAMARENA, S.Y.; VANDEN HEUVEL, J.P.; BLANCHARD, S.G.; LEESNITZER, L.A.; BELURY, M.A. 1999 Conjugated linoleic acid is a potent naturally occuring ligand and activator of PPARalpha. J Lipid Res 40:1426-33.
- 31. PARADIES, G.; PETROSILLO, G.; PISTOLESE, M.; DI VENOSA, N.; SERENA, D.; RUGGIERO, F.M. 1999. Lipid peroxidation and alterations to oxidative metabolism in mitochondria isolated from rat heart subjected to ischemia and reperfusion. Free Rad Biol. Med. 27 : 42-50.
- PARIZA, M.W.; HA, Y.L. 1990a. Conjugated dienoic derivatives of linoleic acid: a new class of anticarcinogens. Med Oncol Tumor Pharmacother 7:169-171.
- 33. PARIZA, M.W.; HA, Y.L. 1990b. Newly recognized anticarcinogenic fatty acids. Basic Life Sci. 52: 167-170.
- PARIZA, M.W.; PARK, Y.; COOK, M.E. 1999. Conjugated linoleic acid and the control of the cancer and obesity. Toxicol Sci 52: 107-110.
- 35. PARK, Y.; STORKSON, J.M.; ALBRIGHT, J.K.; LIU, W.; PARIZA, M.W. 2005. Biological activities of conjugated fatty acids: conjugated eicosadienoic (conj. 21:2deltac12,t14/c13,t15) acids, and other metabolites of conjugated linoleic acid. Biochem Biophys Acta 1687: 120-129.
- PARODI, P.W. 1997.Cow's milk fat components as potential anticarcinogenic agents. J Nutr 127 : 1055-1060.
- RICE-EVANS, C.; HOCHSTEIN, P. 1981. Alterations in erytrocyte membrane fluidity by phenylhydrazine-induced peroxidation of lipids. Biochem. Biophys. Res. Commun. 100:1537-1542.
- RICHTER, C. 1987. Biophysical consequences of lipid peroxidation in membranes. Chem Phys Lipids 44: 175-189.
- 39. SEHAT, N.; RICKERT, R.R.; MOSSOBA, M.M.; KRAMER, J.K.G; YURAWECZ, M.P.; ROACH, J.A.G.; ADOLF, R.O.; MOREHOUSE, K.M.;

FRITSCHE, J.; EULITZ, K.D.; STENHART, H.; KU, K. 1999. Improved separation of conjugated fatty acid methyl esters by silver ion-high-performance liquid chromatography. Lipids 34: 407-413.

- 40. TADOLINI, B.; CABRINI, L.; MENNA, C.; PINNA, G.G; HAKIM, G. 1997.Iron [III] stimulation of lipid hydroperoxide - dependent lipid peroxidation. Free Rad Res 27 : 563-576.
- 41. TANGEN, O.; JONSSON, J.; ORRENIUS, S. 1973. Isolation of rat liver microsomes by gel filtration chromatography. Anal Biochem 54 : 597-603.
- 42. VAN DE VOSSENBERG, J.L.; JOBLIN, K.N.

2003. Biohydrogenation of C18 unsaturated fatty acids to stearic acid by a strain of Butyrivibrio hungatei from the bovine rumen. Lett Appl Microbiol 37 : 424-428.

- 43. VLADIMIROV, Y.A.; OLENEV, V.I.; SUSLOVA, T.B.; CHEREMISINA, Z.P. 1980 Lipid peroxidation in mitochondrial membrane. Adv Lipid Res 17:173-249.
- 44. WRIGHT, J.R.; RUMBAUGTH, R.C.; COLBY, H.D.; MILES, P.R. 1979. The relationships between chemiluminescence and lipid peroxidation in rat hepatic microsomes. Arch Biochem Biophys 192: 344-351.

InVet. 2006, 8(1): 139-148 ISSN(papel): 1514-6634 ISSN (on line) 1668-3498