Antioxidative factors in milk

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Lipid auto-oxidation in milk is affected by a complex interplay of pro- and antioxidants. Several of these compounds are also important nutrients in the human diet and may have other physiological effects in the gastrointestinal tract and other tissues. Among antioxidative enzymes superoxide dismutase catalyses the dismutation of superoxide anion to hydrogen peroxide. The degradation of hydrogen peroxide can be catalysed by catalase and the selenoprotein glutathione peroxidase. The latter enzyme can also degrade lipid peroxides. Lactoferrin may have an important role by binding pro-oxidative iron ions. The occurrence of different forms of these antioxidative proteins in milk and available data on their functional role are reviewed. More remains to be learnt of individual compounds and as an example the potential role of seleno compounds in milk is virtually unknown. Antioxidative vitamins in milk can provide an important contribution to the daily dietary intake. Moreover vitamin E and carotenoids act as fat-soluble antioxidants, e.g. in the milk fat globule membrane, which is regarded as a major site of auto-oxidation. Vitamin C is an important water-soluble antioxidant and interacts in a complex manner with iron and fat-soluble antioxidants. The concentrations of these compounds in milk are affected by cow feeding rations and milk storage conditions. Since milk contains a number of antioxidants many reactions are possible and the specific function of each antioxidant cannot easily be defined. There are indications that other compounds may have antioxidative function and measurement of total antioxidative capacity should be a useful tool in evaluating their relative roles.

Antioxidative enzymes: Milk: Vitamins: Lactoferrin

Introduction

Milk lipids can undergo auto-oxidation, which may lead to changes in food quality. The mechanisms involved include a complex interplay of pro- and antioxidants consisting both of low-molecular-weight compounds, such as vitamins, and proteins. Moreover, the physical–chemical orientation of different compounds in milk has a pronounced influence.

Antioxidants can function by a number of mechanisms. Different enzymes can prevent the formation of radicals or scavenge radicals or hydrogen peroxide and other peroxides. Other enzymes catalyse the synthesis or regeneration of non-enzymatic antioxidants. Among antioxidant enzymes, superoxide dismutase and catalase have been demonstrated in milk. Another enzyme family with antioxidant functions are the selenium-containing glutathione peroxidases (GSHPx). They catalyse the reduction of different peroxides aided by glutathione or other reducing substrates.

Non-enzymatic antioxidants can be formed in the animal body or need to be supplied in the feed as essential nutrients. The iron-binding protein lactoferrin can act as an antioxidant, and vitamin C (ascorbic acid) and vitamin E (tocopherols and tocotrienols) are antioxidant vitamins. Some carotenoids have provitamin A action but they also have antioxidant functions. Several non-enzymatic antioxidants act as radical scavengers in the lipid phase, such as vitamin E, carotenoids and ubiquinol whereas vitamin C acts in the water phase. Others can react in both the lipid and the water phase, such as some flavonoids, which operate both as radical scavengers and metal ion binders. Moreover, the dietary supply of some B vitamins and trace elements is important for optimal activity of several antioxidant enzymes. Foods can also contain added natural or synthetic antioxidants. This report reviews the occurrence and functional aspects of antioxidants in bovine milk.

Antioxidative enzymes in milk

Superoxide dismutase (SOD)

SOD catalyses the dismutation of superoxide anion to hydrogen peroxide:

$$2O_2^- + 2H^+ \rightarrow H_2O_2 + O_2$$

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There are three types of SOD which contain either manganese, copper/zinc or iron (Fridovich, 1986) and in bovine milk Cu/Zn-SOD is found (Asada, 1976; Korycka-Dahl et al. 1979). The enzyme in milk serum has the same specific activity, molecular weight and electrophoretic properties as bovine erythrocyte Cu/Zn-SOD (Hill, 1975; Asada, 1976; Korycka-Dahl et al. 1979). In human milk the activity of Mn-SOD has also been identified (Kiyosawa et al. 1993).

Bovine SOD is a dimer with a molecular weight of 31–33 kDa (Evans et al. 1974). Each monomer contains approximately 153 amino acids, one disulphide bridge, one Cu²⁺ ion and one Zn²⁺ ion. The Cu²⁺ is located in the active site and is responsible for electron transfer during action of the enzyme (Fridovich, 1974). The Cu²⁺ is partially exposed to the solvent, but the Zn²⁺ is wholly buried within the protein structure and is thought to play a structural role, aiding protein stability (Rottilio et al. 1972).

SOD has been partly purified from skim milk after precipitation of casein with rennet (Hill, 1975). The whey was concentrated, the solution was treated with ethanol and chloroform and the precipitated proteins were removed by centrifugation. SOD was purified from the supernatant by gel chromatography and ion exchange chromatography (Hill, 1975). SOD is absent in cream and only found in skim milk (Asada, 1976). The activity of SOD detected in milk corresponds to between 0.15 mg (Hill, 1975) and 2.4 mg of enzyme per litre milk when compared to the activity of purified bovine erythrocyte SOD (Korycka-Dahl et al. 1979). The concentration of SOD in milk varies between cows and breeds. The average activity of SOD in milk serum from Holsteins was 0.92 U/ml and from Jerseys 1.27 U/ml (Hoolbrook & Hicks, 1978) and in milk from Ayrshire cows the activity was 0.89 U/ml (Kankare & Antila, 1982).

The concentration of SOD in cow’s milk is probably not affected by stage of lactation or age of the cow, and does not vary between morning and night milking. Neither is it affected by high somatic cell counts (Hicks, 1980). The concentration of SOD in milk is approximately 100 times lower than that in bovine blood (Hoolbrook & Hicks, 1978). The SOD concentration in human milk is 2.0 to 2.3 times higher than that of bovine milk (Kiyosawa et al. 1993).

Cu/Zn-SOD is very resistant to various types of denaturing stress including heating. In fact commercial pasteurised milk retains the enzymatic activity at a similar level to unpasteurised milk (Asada, 1976). A minimal pasteurisation (71-7°C for 15 s) does not cause heat inactivation of SOD but heat treatments at >75°C have been shown to inactivate more than 20 % of the SOD. Purified SOD fractionated from bovine milk is more sensitive to thermal processing than the SOD in bovine milk serum (Hicks et al. 1979; Korycka-Dahl et al. 1979).

The most commonly used assay of SOD in milk is based on measuring the inhibition of the reduction of cytochrome c by the superoxide anion, produced enzymatically in the xanthine–xanthine oxidase (XO) reaction (Asada, 1976; Korycka-Dahl et al. 1979). Since endogenous XO occurs in the milk and may interfere with the SOD determination, an improved method has been developed in which XO is reduced by ultrafiltration in the samples prior to analysis (Granelli et al. 1994).

**Catalase**

Catalase catalyses the decomposition of hydrogen peroxide:

\[ 2\text{H}_2\text{O}_2 \rightarrow 2\text{H}_2\text{O} + \text{O}_2 \]

Catalase is a large enzyme, containing haem-bound iron in its active site. Catalase has a very high capacity to destroy \( \text{H}_2\text{O}_2 \) and in terms of molecules of \( \text{H}_2\text{O}_2 \) degraded per minute per molecule of enzyme, it is one of the most active enzymes known (Gutteridge & Halliwell, 1994). The catalase molecule is usually composed of four identical 60 kDa subunits, with a total molecular weight in the region of 240 kDa, containing four moles of protohaem (Shonbaum & Chance, 1976; Ito & Akuzawa, 1983a; Robinson, 1991). Catalase has been purified from bovine milk by several purification steps, including n-butanol extraction, ammonium sulphate treatment, ethanol–chloroform fractionation, DEAE-Sephacel column chromatography and Sephacryl S-300 gel filtration (Ito & Akuzawa, 1983a). Optimum pH and temperature for enzyme activity are pH 8.0 and 20°C (Ito & Akuzawa, 1983a).

Cream contains about 60 % and skim milk about 40 % of the milk catalase. Most of the catalase in the cream is bound firmly to the membranes of fat globules and cannot be released readily into skim milk. Milk contains three types of catalase that are different from one another with respect to chromatographic and electrophoretic mobility whereas the two types in cream have similar immunological characteristics (Ito & Akuzawa, 1983b; Ito et al. 1984). Other data by Kitchen et al. (1970) have indicated that skim milk contains 73 % of the whole milk activity and cream 24 %. The average catalase activity in raw milk, determined by a polarographic method, was 1.95 U/ml. Catalase concentrations in raw milk samples have been found to be higher in April than in November, indicating a seasonal variation (Hirvi & Griffiths, 1998). The activity of catalase in normal milk is generally lower than in mastitic bulk milk (Kitchen et al. 1970), which agrees with the findings that catalase activity correlated positively to somatic cell count (Read et al. 1969).

Catalase is one of the most heat-labile enzymes occurring in milk, with most of the activity being destroyed by treatment at 72°C for 15 s (Ito & Akuzawa, 1983a; Griffiths, 1986; Hirvi & Griffiths, 1998). However, reactivation with a greater than 20-fold increase in activity, was observed in milk which had been first heated at 80°C for 15 s and then stored at 4°C for 24 h. Since catalase is also present within leucocytes and bacteria this can possibly be explained by its protection from denaturation by heat within cells and its subsequent release during storage of the pasteurised product causing an apparent reactivation (Griffiths, 1986).

A method for determining catalase by a disk-flotation procedure has been described. This method is based on the liberation of oxygen due to the action of catalase on hydrogen peroxide (Gagnon et al. 1959). Catalase activity
can also be assayed by the polarographic method in which oxygen released from \( \text{H}_2\text{O}_2 \) is quantitated with an oxygen electrode (Hirvi et al. 1996).

**Glutathione peroxidase**

Glutathione peroxidase removes \( \text{H}_2\text{O}_2 \) and other peroxides at a high rate. Its seleno group is oxidised by the peroxide and then reduced by glutathione (GSH), which is converted into oxidised glutathione (GSSG) (Gutteridge & Halliwell, 1994).

\[
2\text{GSH} + \text{H}_2\text{O}_2 \rightarrow \text{GSSG} + 2\text{H}_2\text{O}
\]

The first report of glutathione peroxidase activity was made by Mills (1957) who demonstrated glutathione-dependent catabolism of \( \text{H}_2\text{O}_2 \) by bovine red cell lysate. Selenium was discovered to be an essential component of glutathione peroxidase (GSHPx) in 1973 (Flohé et al. 1973). Four gene-products are classified as members of the seleno-dependent GSHPx family. The most well known are the cellular or classical GSHPx and the plasma or extracellular GSHPx, which are homotetrameres. Selenium is specifically incorporated into proteins as selenocysteine in response to the opal codon UGA. It can also be incorporated as selenomethionine. Selenocysteine is part of the active site of glutathione peroxidases (Ladenstein et al. 1979; Ren et al. 1997).

The bovine plasma GSHPx has been cloned and contains 226 amino acids with a calculated subunit weight of 24 860 Da. The corresponding amino acid sequence showed an overall identity of 88 % with the human plasma GSHPx, 88-5 % with the rat plasma GSHPx, but only 46-4 % with the cellular bovine glutathione peroxidase (Martón-Alonso et al. 1993).

GSHPx activity has been detected in raw cow’s milk, at levels between 12 and 32 U/ml and its activity correlates significantly with selenium concentration. This suggests that GSHPx is one of the biologically active forms of selenium in cow’s milk (Hojo, 1982; Debski et al. 1987). Both GSHPx activity and selenium contents of human milk have been shown to decrease with time of lactation and reach a plateau at one month postpartum (Hojo, 1986).

The percentage of total peroxidase activity associated with glutathione peroxidase has been reported as 29 %, 27 % and 65 % for human, cow’s and goat milk, respectively. The GSHPx activity in human milk is at the same level as in bovine milk, 31–39 U/ml (Hojo, 1986; Debski et al. 1987; Bhattacharya et al. 1988). Most of the glutathione peroxidase activity has been found in the fractions corresponding to 150–170 and 92–96 kDa in milk from all species examined. These results suggest that a substantial portion of the GSHPx in milk exists in a complex form attached to high-molecular-weight proteins within the casein fraction (Debski et al. 1987; Bhattacharya et al. 1988). For human milk most of the activity corresponds to fractions containing proteins with molecular weights of approximately 92 kDa. The monomer weight was estimated to be 23 kDa indicating that the native enzyme consists of four identical subunits. The selenium content of the purified enzyme was 0.31 % (Bhattacharya et al. 1988). The GSHPx purified from human plasma showed the same characteristics (Takahashi et al. 1987).

Human milk glutathione peroxidase has been purified 4500-fold using acetone precipitation and purification by repetitive ion-exchange and gel filtration chromatography (Bhattacharya et al. 1988). Of the GSHPx activity in human milk 90 % could be precipitated by anti-plasma-GSHPx immunoglobulin G. Thus, most if not all GSHPx activity in human milk is due to the plasma form of the enzyme (Aivissar et al. 1991). In two human milks 4 % and 13 % of total selenium was calculated to be bound to GSHPx (Aivissar et al. 1991).

Only a few studies have been made on the heat stability of GSHPx. When heating milk at 80°C for 10 min no activity remained and no activity of GSHPx could be detected in market-pasteurised milk (Hojo, 1982). The effect of milder heat treatment is not known.

GSHPx activity is usually measured by the coupled assay of Paglia & Valentine (1967) or modifications thereof. It is an indirect assay, which requires a peroxide source and a coupled reaction maintaining the concentration of the initial GSH substrate. The GSHPx activity is quantified indirectly in a sample by its ability to cause an increased rate of loss of NADPH compared to the rate of loss seen in a reagent blank.

Although GSHPx is an important seleno compound in milk, other such compounds may also be antioxidative. Recently, it has been found that several seleno compounds may scavenge peroxynitrite radicals (Sies et al. 1997), but the role of this reaction in milk has not been studied. The mean selenium concentration in bovine milk varied in European studies between 3-9 and 23-2 mg/l (Aiajeos & Romero, 1995) and in Swedish bulk milk 18 mg/kg was found (Lindmark-Månsson, unpublished results). In milks from the United States and Australia extreme values of 64 and 0-2 mg/l, respectively, have been reported. The importance of these wide variations in selenium for the antioxidative properties of milk remains to be elucidated.

**Non-enzymatic antioxidants in milk**

**Lactoferrin**

Lactoferrin is an 80 kDa iron-binding glycoprotein occurring in whey. It is composed of a single polypeptide chain with two binding sites for ferric ions. Also the plasma iron-binding protein transferrin occurs in milk of some species. There is marked species variation and an inverse relationship between the concentrations of lactoferrin and transferrin. In bovine milk their concentration is in the range 20–200 mg/l each. In bovine milk lactoferrin can bind approximately five times more iron than is naturally present. Several functions have been proposed for lactoferrin, including: iron-binding, a role in iron absorption, bacteriostatic or bacteriocidal action, and a role as a growth factor (Renner et al. 1989; Hambraeus & Lönnerdal, 1994). Binding of iron to lactoferrin may decrease the conversion of hydrogen peroxide into hydroxyl radical via the Fenton type of reaction. In model experiments bovine lactoferrin was found to inhibit the oxidation of ascorbic acid and tryptophan (Bihel & Birlouez-Aragon, 1998). Antioxidant
activity of lactoferrin has been observed in other model systems (Gutteridge et al. 1981; Shinmoto et al. 1992). The role of lactoferrin is reviewed elsewhere in this volume (Steijns & van Hooijdonk).

**Vitamin C**

Ascorbic acid can easily be oxidised to dehydroascorbic acid, especially at alkaline pH. Dehydroascorbic acid can either be further oxidised or reconverted into ascorbic acid by an enzyme catalysed reaction. Under some conditions ascorbic acid can act as a pro-oxidant by regenerating the perfr eryl radical at initiation of lipid peroxidation.

Andersson & Öste (1994) found that in unpasteurised milk sampled in March or August the concentration of ascorbic acid was higher (20–27 mg/l) than in samples collected in October (12 mg/l). Lindmark-Månsén (unpublished results) found a mean of 11.6 mg vitamin C/kg bulk milk in Sweden, 10.2 mg/kg consisting of ascorbic acid and 1.4 mg/kg of dehydroascorbic acid and also found higher values of vitamin C in July and September than in January and March.

Several groups have analysed vitamin C in milk after heat treatment. Bilić (1991) measured both ascorbic and dehydroascorbic acid and found a concentration of ascorbic acid in raw milk of 5.9 mg/l and in pasteurised milk 4.0 mg/l, in ultra heat treated (UHT) milk by indirect heating 0.1–1.1 mg/l and UHT milk by direct heating 3.9–4.2 mg/l. The corresponding concentrations of dehydroascorbic acid in raw milk was 1.5 mg/l and in pasteurised milk 0.4 mg/l, in UHT milk by indirect heating 0.1–0.2 mg/l and UHT milk by direct heating 0.3 mg/l. Pizzoferrato (1992) found that the retention of ascorbic acid was higher after pasteurisation at 80–85°C than at 72–75°C and 90°C, and that UHT treatment at 140°C caused lower values than pasteurisation. Vahcic et al. (1992) found that in raw bulk milk the ascorbic acid concentration was 14 mg/l, in pasteurised milk 9 mg/l and in sterilised milk 6 mg/l.

With respect to storage effects Andersson & Öste (1994) found no notable loss in ascorbic acid during storage of pasteurised milk in a domestic type refrigerator for one week. The same group (Andersson & Öste, 1992a,b) also followed the ascorbic acid concentration during storage at different temperatures of UHT milk prepared with different oxygen content. In milk with the lowest oxygen content (0.6 p.p.m.) ascorbic acid decreased to half of its original concentration after 1–2 weeks with somewhat higher stability at 7°C than at 23°C and 35°C. At oxygen contents of 3.5–5.4 p.p.m. the loss was much more rapid.

Thus, the vitamin C level in milk can be markedly influenced both by storage and heating conditions. Also Korhonen & Korpela (1994) concluded that vitamin C and several other water-soluble vitamins are lost during storage and heating of milk whereas fat-soluble vitamins are more stable.

**Vitamin E**

Vitamin E consists of eight vitamers and α-tocopherol is the major one in bovine milk. α-Tocopherol is an important lipid-soluble antioxidant and acts as a radical scavenger. The tocopheroxy radical formed is relatively stable and can be reconverted into tocopherol by reduction with ascorbic acid. Different authors have reported concentrations of α-tocopherol between 0.2 and 0.7 mg/l in bovine milk (Jensen, 1995). γ-Tocopherol has also been demonstrated and trace amounts of some other vitamers. Barreiros et al. (1995) reported α-tocopherol levels of 7.4–10.0 mg/g lipid for different herds and also demonstrated low levels of α-tocotrienol. Lindmark-Månsén (unpublished results) found 1.0 mg/kg of α-tocopherol in Swedish bulk milk with a mean fat content of 4.3 %. Colostrum was shown to contain 1.9 mg/l of α-tocopherol decreasing in approximately four days to the level in fresh milk, 0.3 mg/l (Hidiroglou, 1989).

Several reports on the effect of α-tocopherol supplementation on milk oxidative stability have emerged. Supplementation of cows after one month of lactation with 5 g DL-α-tocopherol intraperitoneally was shown to increase milk α-tocopherol five times and the values remained higher than the original ones for six days (Hidiroglou, 1989). Also intravenously administrated DL-α-tocopherol was found to increase both milk and plasma α-tocopherol levels. St-Laurent et al. (1990) supplemented Holstein cows for five weeks with 0, 700 or 3000 IU per day of α-tocopherol given in a grain mix. In the group given 3000 IU/day milk α-tocopherol increased from 0.55 to 0.8 mg/l but the level declined to presupplementation levels by two weeks after treatment. In that study, the effects on milk flavour of α-tocopherol supplementation to a feed consisting of grain mix, hay and pasture were also investigated in herds with a chronic spontaneous oxidised flavour milk problem. α-Tocopherol supplementation improved milk flavour but there was no relationship between milk α-tocopherol levels and degree of flavour improvement. After the supplementation period all cows got access to spring pasture and then the flavour problem decreased markedly. The same research group also made a study on supplementation with α-tocopherol together with inorganic selenium mixed in the concentrate ration or alfalfa enriched by spraying with selenium before ensiling (Nicholson et al. 1991). There was no significant effect of α-tocopherol or selenium supplementation on spontaneous oxidation flavour but α-tocopherol inhibited the generation of copper-catalysed oxidised flavour. The data also suggested that selenium supplementation improved the transfer of dietary α-tocopherol to milk. Use of higher levels of α-tocopherol supplementation or parenteral injection modes gave more clear improvement of milk oxidative stability (Charmley & Nicholson, 1993; Charmley et al. 1993). Nicholson & St-Laurent (1991) also found that supplementation with α-tocopherol to a corn silage feed was more effective in improving milk oxidative stability than the supplementation to an alfalfa silage feed.

The demands on milk oxidative stability and vitamin E levels increase when cows are fed unsaturated fats to modify milk fatty acid composition. Feeding of rations containing rapeseed were found to increase the proportion of monoenoic fatty acids in milk fat and also its vitamin E content which could explain the prolonged induction time for milk fat oxidation (Flachowsky et al. 1997). Use of
Differences in tocopherol levels were stable for two months but decreased by 3.14% at one month and by 9.30% at two months. The losses were marginally higher after storage at 8°C. With respect to storage Vidal-Valverde et al. (1993) found that addition of carbon dioxide to milk followed by storage at 4°C for one week after pasteurisation had little influence on the concentrations of β-carotene and α-tocopherol.

**Carotenoids**

Like tocopherols, carotenoids are fat-soluble compounds and their concentration is influenced by the total fat concentration in dairy products. They function as singlet oxygen scavengers and may also react with other reactive oxygen species. Lindmark-Månsson (unpublished results) has found 0.20 mg/kg of β-carotene in bulk milk with a mean fat content of 4.3%. Ollilainen et al. (1989) found 0.17 mg/kg of β-carotene in whole milk (3.9% fat) and 0.10 mg/kg in milk with 1.9% fat. Only traces of lutein and other carotenoids were demonstrated. Khachik et al. (1997) in a detailed study identified thirty-four carotenoids and their concentration is influenced by the total fat content of 4.3%. Ollilainen (1989) found that β-carotene levels of 3.5–4.9 mg/g fat in milk with and without off-flavour in different herds. For some herds β-carotene and/or α-tocopherol levels were lower in milks with off-flavour but this finding was not consistent. In a supplementation study (Schweigert & Eisele, 1990) a single intravenous or intramuscular injection of 100 or 500 mg β-carotene to cows was shown to increase milk β-carotene from 0.02–0.05 mg/l to 0.13–0.16 mg/l and the maximum was attained approximately one week after injection. With respect to carotenoid stability during storage Ruas-Madiedo et al. (1998) found that addition of carbon dioxide to milk followed by storage at 4°C for one week after pasteurisation had little influence on the concentrations of β-carotene and α-tocopherol.

**Interaction among antioxidants**

Free radicals can react with unsaturated fatty acids, proteins, DNA and other compounds and cause different types of damage, but some free radicals have positive biological functions. Free radicals vary much with respect to reactivity and a hierarchy or a pecking order can be predicted (Buettner, 1993). The interaction of vitamin E and vitamin C is compatible with this hierarchy. However, many reactions are possible, and the specific function of each antioxidant cannot easily be defined. The occurrence of many antioxidants has led to the development of methods to measure total antioxidant capacity. Korpela et al. (1995) found that cow’s milk had both peroxyl radical trapping capacity and superoxide radical trapping capacity. More work is necessary to identify the most active antioxidant compounds in milk.

Still another way of comparing oxidation in milk-related solutions has been used by Champagne et al. (1990) who investigated the formation of ascorbate free radicals in ascorbic acid-fortified milk and related products. They found that the formation of radicals increased in the following order: ascorbic acid-fortified milk < ascorbic acid solution < milk– or soy-based infant formula < ascorbic acid solution containing iron and copper <...
infant formula containing iron and copper. In all solutions radical formation increased with increasing pH. Several compounds in cow’s milk may thus inhibit the production of ascorbic acid radicals.

Furthermore, the susceptibility to lipid oxidation may be affected after fortification of milk by vitamin C and iron since both ascorbic acid at high concentration and iron may be pro-oxidative. Rosenthal et al. (1993) found no evidence for this since addition of ascorbic acid (1 g/l) with or without iron lactate (50 mg/l) or iron lactate alone to raw milk had no significant effect on lipid oxidation, as assessed by formation of free fatty acid or thiobarbituric acid-reactive substances.

An attempt has been made to integrate some available information on pro- and antioxidant proteins in milk (Fig. 1). The reduction of oxygen to oxygen reactive species is indicated on the left. Hydrogen peroxide can be produced by sulphhydryl oxidase and xanthine oxidase, and xanthine oxidase may also give rise to superoxide. Superoxide dismutase catalyses the formation of hydrogen peroxide from superoxide. The conversion of hydrogen peroxide into the hydroxyl radical can be controlled by the availability of iron ions, influenced by lactoferrin and transferrin. Moreover, hydrogen peroxide can be consumed in reactions catalysed by lactoperoxidase, catalase and glutathione peroxidase.

The oxidation of an unsaturated lipid is summarised to the right. By different reactions lipoperoxides (LOOH) can be formed, such as fatty acid hydroperoxides and phospholipid hydroperoxides, which can be reduced by different glutathione peroxidases. At present we do not have a good overview on the relative importance of all these reactions and more studies are necessary to achieve this goal. One regulatory factor is the availability of different enzyme substrates, and this may vary in milks of different origins and after different treatments.

Conclusions

Oxidative reactions in milk are affected by a complex interplay of pro- and antioxidants. Since many antioxidants can be found in milk several reactions are possible and the specific function of each antioxidant cannot easily be defined at present. A deeper understanding of these mechanisms may lead to optimised milk handling and processing procedures as well as a better assessment of the importance of milk antioxidants in human health.

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