High Phosphatidylcholine Hydroperoxide Level in Plasma of Guinea Pigs with Low and Excess Supplementation of Ascorbic Acid

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Summary Graded amounts (0, 50, 500 and 5,000 mg/liter) of ascorbic acid (AsA) were given in drinking water to guinea pigs for 21 days to prepare AsA-deficient, low-AsA, moderate-AsA and excess-AsA animals, and the plasma phospholipid hydroperoxide level and lipid concentration were quantitatively determined to investigate the antioxidant effect of AsA in vivo. Phosphatidylcholine hydroperoxide (PCOOH) was a predominant phospholipid hydroperoxide present in the plasma, and the PCOOH concentration was significantly higher in AsA-deficient, low-AsA and excess-AsA animals (80.4 nM, 54.8 nM and 42.2 nM, respectively) as compared with that in moderate-AsA animals (27.2 nM). Hyperlipidemic plasma characterized as high cholesterol and high triacylglycerol concentrations was confirmed in AsA-deficient animals. Molar ratios of plasma AsA and \( \alpha \)-tocopherol against \( 10^4 \) moles of phospholipids were significantly lower in AsA-deficient and low-AsA animals (0.6–2.1 and 5.5–8.5, respectively) than in moderate-AsA and excess-AsA animals (14.2–18.0 and 11.2–11.9, respectively). In plasma, a high correlation coefficient (\( r = 0.979 \)) was observed between PCOOH and AsA for which there was optimum AsA level to keep the low PCOOH and such correlation was stronger than that (\( r = 0.558 \)) observed with \( \alpha \)-tocopherol. The results indicated that AsA has an important function to control the phospholipid hydroperoxide level in plasma and that moderate supplementation of AsA is required to reveal its optimal antioxidant effect in vivo. The present study also showed that AsA-deficiency especially invites an increase in plasma PCOOH together with a hyperlipidemic state which are risk factors in developing atherogenesis.

Key Words ascorbic acid, \( \alpha \)-tocopherol, phosphatidylcholine hydroperoxide, plasma lipids, hyperlipidemia, guinea pig

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Ascorbic acid (AsA) is known to act as antioxidant scavenging singlet molecular oxygen (1), superoxide and hydroxyl radical (2, 3) which are reactive oxygen species causing membrane lipid peroxidation. In *in vitro* systems, regeneration with AsA of a lipophilic antioxidant α-tocopherol from tocopheryl radical has also been suggested with respect to its antioxidative mechanism (4). In the animal experiments, feeding of AsA-deficient diet (below 0.1 mg AsA/kg diet) causes an increase in the level of thiobarbituric acid-reactive substances (TBARS) as a tentative indication of lipid peroxides in the plasma of guinea pigs (5) and in the livers and brains of rats (6). With AsA-deficiency, the plasma α-tocopherol level is decreased in rats (6). On the other hand, AsA has been known to act as prooxidant when AsA chelates with iron reportedly led to the production of peroxidative Fe(II) (7). In rats, supplementation of excess AsA (1,500 mg/kg diet) brings about a high incidence of erythrocyte hemolysis accompanied by an increase in liver TBARS and a decrease in plasma tocopherol (8). Therefore, it is expected that there is some optimum level for dietary AsA, by which AsA can act as an effective antioxidant but not as a prooxidant in tissue organelles *in vivo*. In human plasma, high accumulation of phosphatidylcholine hydroperoxide as a primary oxidation product of phosphatidylcholine located at the surface of lipoprotein particles has been found in patients with hyperlipidemia (9). The enhanced phosphatidylcholine hydroperoxide level in plasma should directly reflect the formation of oxidized lipoprotein particles that are a possible causative agent for developing atherogenesis (10). As the hyperlipidemic state has been observed in the plasma of AsA-deficient guinea pigs (11), its contribution in modifying the plasma lipid hydroperoxide level is also interesting. In the present study, graded amounts of AsA were given to guinea pigs, and the plasma phosphatidylcholine hydroperoxide level and lipid profile were investigated to estimate the optimum amount of AsA which can prevent both the lipid hydroperoxide accumulation and normalize the hyperlipidemic state of the plasma.

**MATERIALS AND METHODS**

*Animals and diets.* Hartley strain male guinea pigs (3 weeks old, *n* = 20) were purchased from Agricultural Cooperative Association for Laboratory Animals (Hamamatsu). The animals were submitted to prefeeding to a body weight of 410 g with a commercial pellet rations (guinea pigs diet GM-1, containing 1,500 mg vitamin C/kg and 50 mg vitamin E/kg, Funabashi Farm Co., Funabashi). Each animal was kept in a wire-bottomed stainless-steel cage, and water was supplied *ad libitum*. In the feeding experiment, the AsA intake was adjusted by drinking water to prepare AsA-deficient (0 mg/liter, group A), low-AsA (50 mg/liter, group B), moderate-AsA (500 mg/liter, group C) and excess-AsA (5,000 mg/liter, group D) animals, respectively. The animals of all groups were maintained on an AsA-free diet (contained 50 mg vitamin E/kg; Oriental Yeast Co., Tokyo) for 22 days of the feeding period. On day 22 of feeding, the animals were fasted overnight and

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anesthetized with diethylether. Heparinized blood was withdrawn by heart puncture and was centrifuged at low speed (1,000 $\times$ g) at 4°C for 10 min to separate the plasma.

**Plasma lipid analyses.** Plasma total-cholesterol (TC), free-cholesterol (FC), triacylglycerol (TG) and phospholipids (PL) were determined using the cholesterol-E-test, free cholesterol-E-test, triglyceride-E-test and phospholipid-C-test (Wako Pure Chemical Co., Osaka), respectively. Plasma AsA was measured by the dinitrophenylhydrazine method (12). Plasma $\alpha$-tocopherol was determined by high-performance liquid chromatography (13). Fatty acid composition of plasma lipids was analyzed as methylesters (14) using a Shimadzu GC-8A gas chromatograph.

**Lipid hydroperoxide assay.** Phosphatidylcholine hydroperoxide (PCOOH) present in plasma was determined by chemiluminescence-high performance liquid chromatography (CL-HPLC) as has been reported by Miyazawa et al. (15,16).

**Statistical analysis.** The data are expressed as mean values and standard deviations (SD). All data were analyzed using ANOVA and Student’s t-test. The coefficient of correlation between PCOOH and AsA levels in plasma was compared with the regression curve of secondary degree.

## RESULTS

### Body weight gain

Figure 1 shows the weight gain of guinea pigs during the experimental period (22 days) given graded doses of AsA. AsA-deficient animals (group A) showed the smallest weight gain among the four groups, and apparent weight loss was observed

![Graph showing weight gain of guinea pigs](image)
in group A animals after the 18th day due to the lack of appetite, and at the 22nd
day AsA-deficient animals (group A) revealed typical scurvy symptoms. Among
the AsA-supplemented animals (groups B, C and D), the weight gain of excess-AsA
animals (group D) was smaller than that of moderate-AsA animals (group B) and
of low-AsA animals (group C). No significant difference was observed between the
moderated-AsA animals and the low-AsA animals.

**Plasma AsA and α-tocopherol**

The AsA concentration in the plasma of AsA-deficient animals (group A) and
of low-AsA animals (groups B) were significantly lower than that of moderated-
AsA animals (group C) and of excess-AsA animals (group D) (Table 1). The
lowest AsA concentration was observed in the plasma of AsA-deficient animals
(group A). No difference was from 0.6 to 2.1 observed between the moderate-AsA
animals and the excess-AsA animals. The molar ratio of AsA against 10⁴ moles of
plasma phospholipids was in AsA-deficient and low-AsA animals while in moderat-
ed-AsA and excess-AsA animals the molar ratio was 14.2–18.0.

The α-tocopherol concentration, expressed as μg/ml plasma, in AsA-deficient
animals (group A) and in low-AsA animals (group B) was somewhat higher (0.4–
0.5 μg/ml) than that in moderate-AsA animals (group C) and excess-AsA animals
(group D) (0.3 μg/ml) (Table 1), while the molar ratio of α-tocopherol against 10⁴
moles of plasma phospholipids was significantly lower in AsA-deficient animals and
low-AsA animals (5.5–8.5) than that in moderate-AsA animals and excess-AsA
animals (11.2–11.9). The plasma α-tocopherol level of excess-AsA animals and of
moderate-AsA animals was almost the same.

As shown in Fig. 2, a biphasic relationship was seen between plasma AsA and
α-tocopherol levels; these levels were inversely proportional to each other when
compared by units of μg/ml plasma (Fig. 2-A), but were rather proportional when
compared by the units of mol/10⁴ mol plasma phospholipids (Fig. 2-B).

**Table 1. Ascorbic acid and α-tocopherol levels in the plasma of guinea pigs given
graded doses of ascorbic acid.**

| Group            | Ascorbic acid (μg/ml) | Molar/10⁴ mol PL | α-Tocopherol (μg/ml) | Molar/10⁴ mol PL |
|------------------|-----------------------|------------------|----------------------|------------------|
| A (AsA-deficient)| 1.1±0.3a              | 0.6±0.2a         | 0.5±0.1b             | 5.5±1.3a         |
| B (low-AsA)      | 1.9±0.4b              | 2.1±0.4b         | 0.4±0.1a,b           | 8.5±1.7a,b       |
| C (moderate-AsA)| 8.4±1.8c              | 14.2±3.9c        | 0.3±0.0a             | 11.2±1.5b        |
| D (excess-AsA)   | 8.9±1.0c              | 18.0±2.0c        | 0.3±0.0a             | 11.9±2.7b        |

Values are M±SD (n=5). a,b,c Values with different superscript letters in a column
are significantly different (p<0.05). The animals fed ascorbic acid (AsA)-deficient
diet were supplemented with AsA by drinking water in the following dose: A, AsA 0
mg/liter; B, AsA 50 mg/liter; C, AsA 500 mg/liter; and D, AsA 5,000 mg/liter. PL, phospholipids.
Fig. 2. Relationship between ascorbic acid (AsA) and \( \alpha \)-tocopherol levels in the plasma of guinea pigs received graded amounts of AsA. A, relationship in \( \mu \text{g/ml} \) plasma; B, relationship in mol/10\(^4\) mol PL (phospholipids). Abbreviations are the same as those given in Fig. 1. Bars represent SD of means.

Fig. 3. Plasma lipid profile of guinea pigs received graded doses of ascorbic acid (AsA). The animals were maintained on AsA-deficient diet and supplemented with AsA by drinking water in the following doses for 22 days. A, AsA 0 mg/liter (AsA-deficient); B, AsA 50 mg/liter (low-AsA); C, AsA 500 mg/liter (moderate-AsA); D, AsA 5,000 mg/liter (excess-AsA). TC, total-cholesterol; FC, free cholesterol; TG, triacylglycerol; PL, phospholipids. Values are M±SD (n=5). Values with different superscript letters are significantly different at \( p<0.05 \).

**Plasma lipid profile**

Figure 3 shows the plasma lipid concentration of guinea pigs given graded amounts of AsA for 22 days. In the plasma of AsA-deficient animals (group A), the total cholesterol, free cholesterol, triacylglycerol and phospholipid concentrations were higher than in any of the other three groups. The plasma cholesterol and
Table 2. Fatty acid compositions of plasma total lipids of guinea pigs given graded doses of ascorbic acid (AsA).

| Fatty acid | AsA 0 mg/liter (AsA-deficient) | AsA 50 mg/liter (low-AsA) | AsA 500 mg/liter (moderate-AsA) | AsA 5,000 mg/liter (excess-AsA) |
|------------|--------------------------------|---------------------------|-------------------------------|--------------------------------|
| 14:0       | 1.4±0.3                        | 0.8±0.1                   | 0.7±0.1                       | 0.9±0.2                       |
| 16:0       | 18.3±0.3                       | 20.0±0.5                  | 17.6±0.8                      | 22.6±1.3                      |
| 16:1 n7    | 1.6±0.5                        | 1.1±0.1                   | 1.1±0.1                       | 0.6±0.4                       |
| 18:0       | 10.1±0.3                       | 10.4±0.4                  | 11.2±0.3                      | 11.1±0.1                      |
| 18:1 n9    | 16.1±1.2                       | 19.1±0.3                  | 16.4±0.4                      | 18.3±0.3                      |
| 18:2 n6    | 42.1±3.0                       | 39.4±0.9                  | 41.7±0.2                      | 39.4±0.9                      |
| 20:4 n6    | 4.9±2.1                        | 6.1±0.4                   | 5.8±0.5                       | 3.3±1.4                       |
| 22:5 n3    | 0.5±0.1a                       | 0.2±0.1a                  | 0.1±0.0a                      | 2.9±0.7b                      |
| 22:6 n3    | 2.1±0.3b                       | 1.0±0.1a                  | 2.8±0.5b                      | 0.8±0.6a                      |

Values are M±SD (n=5). a,b Values with different superscript letters of each fatty acid are significantly different (p<0.05) among the animal groups.

Polar lipids

The major fatty acids of the plasma total lipids of guinea pigs were linoleic (39.4–42.1%), palmitic (17.6–22.6%), oleic (16.1–19.1%) and stearic (10.1–11.2%) acids (Table 2). The compositional proportion of such major fatty acids was not changed even after the change of AsA intake. A little change was observed in the proportions of 22:5 n-3 (docosapentaenoic acid, DPA) and 22:6 n-3 (docosahexaenoic acid, DHA), while 20:5 n-3 (eicosapentaenoic acid, EPA) was not detected in the plasma. No significant difference was observed in the fatty acid composition of plasma phosphatidylcholine among the four animal groups (Table 3).

**Plasma PCOOH**

In the plasma of guinea pigs, PCOOH was a predominant phospholipid hydroperoxide. Among the four animal groups that received graded doses of AsA, the highest PCOOH concentration (80.4 nM) was observed in the plasma of AsA-deficient animals (group A) (Table 4). The plasma PCOOH level of the moderate-AsA animals (group C; 27.2 nM) was lowest among the four groups, and that of excess-AsA animals (group D; 42.2 nM) and low-AsA animals (group B; 54.8 nM) was significantly higher than the moderate-AsA animals. The molar ratio
Table 3. Fatty acid composition of plasma phosphatidylcholine of guinea pigs given graded doses of ascorbic acid (AsA).

| Fatty acid | Group                  | Weight % |
|------------|------------------------|----------|
|            | AsA 0 mg/liter (AsA-deficient) | 2.1±0.5  |
| 14:0       | AsA 50 mg/liter (low-AsA)   | 4.2±1.5  |
| 16:0       | AsA 500 mg/liter (moderate-AsA) | 1.6±0.4  |
| 16:1 n7    | AsA 5,000 mg/liter (excess-AsA) | 4.5±1.1  |
| 18:0       |                        | 18.7±0.4 |
| 18:1 n9    |                        | 1.4±0.1  |
| 18:2 n6    |                        | 36.5±1.3 |
| 20:4 n6    |                        | 9.9±1.2  |
| 22:5 n3    |                        | 28.6±1.0 |
| 22:6 n3    |                        | 1.3±0.1  |

Values are M±SD (n=5).

Table 4. Phospholipid hydroperoxides in plasma and livers of guinea pigs given graded doses of ascorbic acid.

| Group        | Plasma                     | Livers                     |  |
|--------------|----------------------------|----------------------------|---|
|              | PCOOH                      | PCOOH                      | PEOOH                      |
|              | nM mol/10^5 mol PL         | nmol/g mol/10^5 mol        | nmol/g mol/10^4 mol        |
| A (AsA-deficient) | 80.4±20.1^c     1.4±0.2^b  | 4.2±1.0^a 225±54^a        | 1.9±0.3^a 102±16^a         |
| B (low-AsA) | 54.8±21.9^b     1.3±0.3^b  | 3.8±1.2^a 168±53^a        | 1.8±0.2^a 80±11^a          |
| C (moderate-AsA) | 27.2±5.5^a     0.9±0.1^a  | 3.6±0.5^a 208±29^a        | 1.7±0.3^a 98±18^a          |
| D (excess-AsA) | 42.2±8.6^b     1.2±0.1^b  | 3.6±0.4^a 180±25^a        | 1.8±0.2^a 90±11^a          |

Values are M±SD (n=5). ^a,b,cValues with different superscript letters in a column are significantly different (p<0.05). The animals fed ascorbic acid (AsA)-deficient diet received AsA by drinking water in the following doses for 22 days: A, AsA 0 mg/liter; B, AsA 50 mg/liter; C, AsA 500 mg/liter; D, AsA 5,000 mg/liter. PCOOH, phosphatidylcholine hydroperoxide, PEOOH, phosphatidylethanolamine hydroperoxide; PL, phospholipids.

of PCOOH calculated against 10^5 moles of plasma phospholipids in the AsA-deficient animals and in the low-AsA animals showed significantly higher values (1.3–1.5) than that of the moderate-AsA animals (0.9) (Table 4). Liver phospholipid hydroperoxide concentrations showed no change even after the change of AsA intake (Table 4).

The relationship between the PCOOH concentration and the AsA level in the plasma (Fig. 4-A) showed a high correlation coefficient (r=0.979) with the regression curve of secondary degree, and it revealed that moderate concentration...
Fig. 4. Relationships of phosphatidylcholine hydroperoxide (PCOOH) against ascorbic acid (A) and α-tocopherol (B) in the plasma of guinea pigs given graded doses of ascorbic acid. Abbreviations are the same as those given in Fig. 1.

of AsA is required in plasma to prevent the increase in PCOOH. To keep the plasma PCOOH as low as 1 mole per 10⁵ moles of phospholipids, 50–150 moles of AsA were required per 10⁵ moles of plasma phospholipids. The plasma PCOOH concentration of excess-AsA animals was significantly higher than that of moderate-AsA animals. A weak correlation ($r=0.558$) was observed between the molar ratios of PCOOH and α-tocopherol when compared by the units of the number of phospholipid molecules in the plasma (Fig. 4-B).

**DISCUSSION**

In the present study, four groups of guinea pigs were given graded doses of AsA were designed to study the effect of AsA supplementation in the diet. Guinea pigs require at least 0.5 mg of AsA/100 g body weight/day to prevent scurvy (17). The low-AsA animals (group B; 50 mg of AsA/liter drinking water) prepared in the present study was comparable to 2 mg AsA/100 g body weight/day. The amount of

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AsA ingested by the moderate-AsA animals (group C; 500 mg AsA/liter drinking water; corresponding to 20 mg AsA/100 g body weight/day) was considered to be optimum for maintaining general health because it was in the range employed in standard commercial diets (1,000–1,500 mg AsA/kg; 10–30 mg AsA/100 g body weight/day) of guinea pigs, which are designed to achieve optimum growth and longevity of this animal species. The excess-AsA animals (group D; 5,000 mg AsA/liter drinking water; corresponding to 200 mg AsA/100 g body weight/day) was selected in order to examine the effect of AsA overloading (10 times AsA of moderate-AsA animals). No significant difference was observed in the total volume of drinking water consumed which was supplemented with different amounts of AsA.

Supplementation of AsA in the drinking water was effective to increase the plasma AsA level, and the plasma AsA concentration of moderate-AsA animals and of excess-AsA animals was 4–8 times higher than that of AsA-deficient animals and of low-AsA animals (Table 1). The plasma AsA level of the excess-AsA animals was almost the same as that of the moderate-AsA animals. This may be ascribed to the excretion of excess AsA into the urine. In humans, excess AsA is not absorbed from intestine but is excreted into the urine (18).

Hyperlipidemic plasma (characterized as increases of total cholesterol, free cholesterol, triacylglycerol and phospholipids) was most apparent in AsA-deficient animals (group A) (Fig. 2). Marginal intake of AsA also invited increases of plasma free-cholesterol and phospholipids, as shown in the low-AsA animals (group B). Such influence on plasma lipid concentrations might reflect the disturbance of lipid metabolism in AsA-deficient guinea pigs, although the AsA-deficiency did not affect the fatty acid composition of plasma total lipids and of plasma phosphatidylcholine (Tables 1 and 2). Ginter et al. (19) have reported that AsA deficiency in the guinea pig causes an increase in cholesterol in the plasma and in the tissue organelles along with a decrease in the activity of cholesterol-7α-hydroxylase. Ha et al. (20) showed that AsA deficiency results in a decrease in the tissue carnitine level in the guinea pig which blocks the transport of fatty acids into mitochondria, thus shunting triacylglycerol synthesis in the liver. Kotze et al. (21) have suggested that hypertriglyceridemia caused by a deficiency of AsA is associated with the inhibitory effect of AsA on lipoprotein lipase activity.

In the present study, lower AsA intakes as observed in AsA-deficient animals and in low-AsA animals resulted in a decrease in the plasma level of α-tocopherol (Table 1). This decrease became more clear when the molecular number of α-tocopherol was compared against the number of plasma phospholipid molecules. Hrubá et al. (22) demonstrated that the α-tocopherol contents in plasma and in other tissues are low in AsA-deficient guinea pigs. Such lower levels of AsA and α-tocopherol in the plasma could profoundly influence the plasma lipid peroxide level. As a result, the PCOOH concentration was 3 times higher in AsA-deficient animals and 2 times higher in the low-AsA animals as compared with that of the moderate-AsA animals (Table 4). Such an increase in plasma PCOOH should
directly reflect the stimulation of hydroperoxidation of the phospholipid molecules in the plasma lipoproteins. Kimura et al. (23) have reported that osteogenic disorder (ODS) rats lacking the ability to synthesize AsA shows high levels of TBARS in plasma and livers under AsA-deficiency.

The animals which ingested excess AsA (group D) also showed a PCOOH increase in the plasma (Table 4, Fig. 4). This may reflect the prooxiant effect of excess AsA on phospholipids in plasma lipoprotein particles. Inorganic iron reduced by AsA has been suggested to be an active catalyst of phospholipid peroxidation (7). Generally, iron in plasma is bound to transferrin (24), but in a recent study (25) an alternative low molecular mass species of plasma iron, non-transferrin-bound iron, has been shown to occur in significant concentrations under conditions of transferrin saturation. Randell et al. (26) have reported the effect of AsA on translocation of non-transferrin-bound iron into cells, and they suggested that AsA may reductively mobilize the internal iron store into the regulatory pools. Therefore, we presumed that the PCOOH increase in plasma observed under excess-AsA conditions may be due to AsA-catalyzed phospholipid hydroperoxidation which is possibly accelerated by combination with non-transferrin-bound iron. Previously, Miyazawa et al. (27) observed a significant decrease in \( \alpha \)-tocopherol in the tissue organelles of guinea pigs treated with an excess-AsA diet. Therefore, the decrease of antioxidative molecules such as \( \alpha \)-tocopherol under excess AsA conditions, in which \( \alpha \)-tocopherol may be consumed to scavenge ascorbate radical, may also contribute in the increase in the plasma PCOOH level.

As liver PCOOH and phosphatidylethanolamine hydroperoxide (PEOOH) concentrations were not changed even after changing the ingested amount of AsA, hydroperoxidation of phosphatidylcholine in lipoprotein particles should occur in the endothelial cell space and not in the liver.

Recently, oxidative modification of plasma lipoprotein is recognized as one of the most essential reactions in promoting atherogenesis (28). The onset of atherogenesis in aorta has been noted in AsA-deficient guinea pigs (29). Rath and Pauling (30) reported that a high plasma level of lipoprotein(a) is associated with coronary heart disease in humans and that the incidence of cardiovascular disease is decreased by elevating the AsA intake. These studies have suggested that AsA-deficiency especially could increase the incidence of atherogenesis.

The present study clearly demonstrated that the antioxidant function of AsA is important to control the PCOOH level in plasma and that moderate intake of AsA is required to reveal its optimal antioxidant effect in vivo.

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