Vitamin E and selenium: contrasting and interacting nutritional determinants of host resistance to parasitic and viral infections

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Beneficial effects of trace amounts of dietary Se were first observed in vitamin E-deficient rats so that a strong metabolic interaction between these two micronutrients was apparent since the initial discovery of the nutritional value of Se (Schwarz & Foltz, 1957). Later work showed that Se and vitamin E tended to spare one another’s requirement for the prevention of certain nutritional deficiency diseases (e.g. exudative diathesis in chicks, see Thompson & Scott, 1969). However, there were other conditions that appeared to respond specifically to only one of these nutrients (e.g. resorption gestation in rats, see Harris et al. 1958). As our knowledge about Se and vitamin E grew, the complexity of their interactions became clear with some conditions being specific for one or the other nutrient, whereas other syndromes responded equally well to either nutrient (Table 1). Many of the early results could be adequately explained by the

<table>
<thead>
<tr>
<th>Conditions responsive to dietary:</th>
<th>Se</th>
<th>Vitamin E</th>
</tr>
</thead>
<tbody>
<tr>
<td>Liver necrosis</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>In the second generation:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sparse hair coat</td>
<td>+</td>
<td></td>
</tr>
<tr>
<td>Poor growth</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>Poor sperm motility</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>Cataracts</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>Discoloration of body fat</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Discoloration of the uterus</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Depigmentation of incisors</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td><em>In vitro</em> haemolysis (no glucose)</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Impaired reproductive capacity of females</td>
<td>-</td>
<td>+</td>
</tr>
</tbody>
</table>

The rat was selected as the model since this is the mammalian species for which the Se and vitamin E deficiency diseases are best clarified.
antioxidant properties of both these nutrients: Se as a component of the peroxide-destroying enzyme glutathione peroxidase (EC 1.11.1.9) and vitamin E as a lipid-soluble antioxidant (Hoekstra, 1975). More recent work has established metabolic roles for Se that appear to be unrelated to its antioxidant activity (e.g. as a component of iodothyronine 5'-deiodinase (EC 3.8.1.4), see Arthur et al. 1990).

Nutritional antioxidants such as Se and vitamin E have received a great deal of attention lately with regard to their possible protective benefits against chronic degenerative diseases such as cancer and cardiovascular disease (Collins et al. 1994; Riemersma, 1994). On the other hand, our work has shown that Se and vitamin E status can also have a profound impact on the ability of a host to resist acute infectious disease. Here we review effects of Se and/or vitamin E nutriture on two infectious-disease models in mice: malaria and coxsackievirus-induced heart muscle damage (the latter perhaps related to Keshan disease).

**MALARIA**

Our interest in the relationship between nutritionally-induced oxidative stress and malaria was first stimulated by the Chinese traditional antimalarial drug Qinghaosu (QHS; Klayman, 1985; Hien & White, 1993). This compound contains a cyclic endoperoxide group and is thought to destroy the parasite by generating oxy free radicals (Clark et al. 1983). QHS and its derivatives represent a potent new class of antimalarials and synthesis of novel QHS analogues is a highly active research area (Posner et al. 1994).

If indeed QHS acted through the generation of free radicals, dietary deprivation of vitamin E, a free-radical quencher (Packer, 1991), might be expected to enhance the antimalarial activity of the drug. This in fact was shown to be the case since low doses of QHS have a much stronger effect against *Plasmodium yoelii* in vitamin E-deficient mice than in normal mice (Levander et al. 1989a). However, no such enhancing effect of QHS activity was seen under conditions of Se deficiency. This differential effect of vitamin E deficiency and Se deficiency on the antimalarial efficacy of QHS might prove helpful in clarifying the mode of action of the nutrients and/or the drug. For example, the fact that vitamin E status influences the potency of the drug while Se status does not could suggest that the drug exerts its action in a lipid phase of the cell which is inaccessible to the cytoplasmic glutathione peroxidase. Another possible explanation for the results is that the malarial parasite apparently does not possess a Se-dependent glutathione peroxidase (Jung et al. 1989), so perhaps one would not expect the parasite's response to the drug to be affected by host Se status.

The successful demonstration that vitamin E nutriture could have an impact on the pharmacological effectiveness of QHS prompted us to test the effect of dietary fish oil, a known tocopherol antagonist (Dam, 1962), on the potency of the drug. An experiment was designed to study three-way interactive effects of QHS, fish oil, and vitamin E deficiency on malarial infection but the vitamin E-deficient mice consuming fish oil were so strongly protected against the disease that it was not possible to ascertain any additional beneficial effect of the drug (Levander et al. 1989a). A similar protective effect of fish oil against malaria in vitamin E-deficient mice had been noted about 30 years earlier by Godfrey (1957a) working at Mill Hill in London. Subsequent work by us showed that a variety of fish oils and fish-oil concentrates fed in the diet exhibited this
Table 2. **Contrasting effects of selenium and vitamin E deficiency on the ability of mice fed on diets containing fish oil to resist malarial infection** (Adapted from Morris et al. 1990)

(Mean values with their standard errors for ten animals)

<table>
<thead>
<tr>
<th>Dietary supplement (mg/kg)</th>
<th>Plasma vitamin E concentration (µmol/l)</th>
<th>Erythrocyte glutathione peroxidase activity (units/g haemoglobin)</th>
<th>Rate of survival at intervals (d) after inoculation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vitamin E</td>
<td>Se</td>
<td>Mean</td>
<td>SE</td>
</tr>
<tr>
<td>100</td>
<td>0-5</td>
<td>4-1</td>
<td>0-2</td>
</tr>
<tr>
<td>100</td>
<td>0</td>
<td>4-4</td>
<td>0-4</td>
</tr>
<tr>
<td>0</td>
<td>0-5</td>
<td>&lt;1-2</td>
<td></td>
</tr>
</tbody>
</table>

* Weanling male outbred CD-1 Swiss albino mice were fed on a torula yeast-based diet containing 50 g menhaden (fish) oil/kg with or without vitamin E and/or Se as RRR-α-tocopheryl acetate or sodium selenite respectively. After 4 months, all mice were inoculated intraperitoneally with $5 \times 10^4$ erythrocytes parasitized by *Plasmodium yoelii* (17X strain). Survival was monitored for 60 d after infection. Survival in the group fed on the vitamin E-deficient diet containing fish oil after 19 d was not as good as that seen in other experiments (Levander & Ager, 1993), perhaps because these mice were older at inoculation (19 weeks v. 7 weeks).

Antimalarial activity in vitamin E-deficient mice (Levander et al. 1990). A plant oil containing n-3 fatty acids (linseed oil) also had this antimalarial property in vitamin E-deficient mice as did ground flaxseed and chemically purified ethyl linolenate (Levander et al. 1991).

A series of experiments was carried out to determine the practical limitations of this nutritionally-induced antimalarial action. Feeding the pro-oxidant vitamin E-deficient diet containing fish oil for only 2 weeks was sufficient to confer substantial antimalarial protection to the mice, so a severe vitamin E deficiency is apparently not needed to obtain this effect (Levander et al. 1989c). Presumably, if the plasma tocopherol levels could be decreased without affecting tissue stores too much, the dietary treatment would still have its antimalarial effect since the erythrocytic form of the parasite ‘sees’ only the bloodstream. In fact, it was possible to demonstrate antimalarial effects of a fish-oil concentrate fed in diets that resulted in low but finite plasma tocopherol concentrations, although the degree of protection offered by the diet was inversely related to the resultant level of vitamin E in the plasma (Levander & Ager, 1993). Taken together, these observations suggest that human populations marginally deficient in vitamin E might respond favourably to the antimalarial action of fish oil.

As in the case of the experiment with QHS, Se deficiency had little or no effect on the survival of infected mice fed on fish oil, whereas vitamin E deficiency markedly improved the survival of infected mice fed on fish oil (Table 2). This was true despite a prolonged feeding period with the Se-deficient diet (4 months before inoculation) resulting in a severe Se deficiency (erythrocyte glutathione peroxidase activity in the deficient group was less than 3% of that in the control group). Thus, once again host Se deficiency did not have the same deleterious consequences for the parasite as host vitamin E deficiency.

Two difficult problems in dealing with malaria in the field are the emergence of drug-resistant parasites (Bloland et al. 1993) and the occurrence of the cerebral form of
the disease (Miller et al. 1994). Chloroquine-resistant plasmodia are now a fact of life in many parts of the world and the parasite is developing resistance to several drugs simultaneously. Animal studies have shown that resistance can also develop to drugs of the QHS family (Chawira et al. 1986) so that the therapeutic utility of even this class of antimalarials may eventually be lost. In this context, it is of interest to note that feeding the pro-oxidant vitamin E-deficient diet containing menhaden oil protects against chloroquine-resistant lines of three species of murine malaria: *P. yoelii*, *P. berghei*, and *P. vinckei* (Levander et al. 1989b; Ager et al. 1991; Ager & Levander, unpublished results). Such dietary manipulation was also effective against a sulphone-resistant line of the murine parasite, so that the benefits of nutritionally-induced oxidative stress against the disease were not limited to one type of drug-resistant parasite (Ager & Levander, unpublished results). Even if pro-oxidant nutritional intervention never succeeds as a practical tool for human malaria control, it would seem that phenomena such as those discussed previously might provide useful new theoretical insights into the general problem of drug resistance.

Cerebral malaria is a severe complication of the disease which often results in the death of the patient (Miller et al. 1994). Although several factors are thought to determine whether this particular form of the disease will develop (e.g. local endemicity of the disease, age of the patient etc.), often the nutritional status of the infected individual is not considered. Early animal experiments indicated that feeding mice high levels of a synthetic antioxidant (butylated hydroxyanisole; BHA) offered protection against cerebral malaria (Thumwood et al. 1989). This finding gave rise to the concept that excess production of free radicals might be involved in the pathogenesis of this form of the disease. Several years later, the somewhat paradoxical observation was made that dietary fish oil, a pro-oxidant stress, also protected partially against murine cerebral malaria (Blok et al. 1992). In the latter report, indomethacin, a potent cyclooxygenase inhibitor, had no effect on the course of the infection, thereby ruling out any role of prostaglandins in the protective effect of the fish oil. Moreover, *ex vivo* biosynthesis of tumour necrosis factor, a cytokine presumed to play a key role in the pathogenesis of cerebral malaria, was not decreased by the dietary fish oil treatment. We also found a partial protective effect of dietary fish oil against murine cerebral malaria and that protection could be made virtually complete when the fish oil was fed in a vitamin E-deficient diet (Ager et al. 1993). We were also able to show that feeding either of two structurally-unrelated synthetic antioxidants, N,N’-diphenyl-p-phenylenediamine (DPPD) or probucol, blocked the protective action of the pro-oxidant vitamin E-deficient diet containing menhaden oil in a cerebral malaria model (Levander et al. 1995). Both these observations are most compatible with the hypothesis that the protective effect of dietary fish oil against cerebral malaria is due to dietary-induced oxidative stress. Thus, dietary-induced oxidative stress protects not only against the later sequelae of malarial infection (e.g. anaemia) by suppressing parasite growth, but also against the early manifestations of the disease (e.g. central nervous system consequences) in a model of cerebral malaria. Whether oxidative stress status plays any role in the outcome of human cerebral malaria is not known at this time, although increased levels of lipid peroxidation products have been reported in the cerebrospinal fluid of such patients (Das et al. 1991).

The biochemical mechanism by which fish oils exert their beneficial effect against malaria is not certain since these oils are known to have a wide-ranging impact on several
aspects of metabolism (Simopoulos, 1991). Nonetheless, the most reasonable and straightforward explanation for this antimalarial property of fish oil seems to be its ability to exert pro-oxidant stress on the parasite. Indeed, it is difficult to account for many of our experimental observations on any other basis (dependence of fish oil on vitamin E deficiency for full protection, blockage of the protective effect of the fish oil by concurrent consumption of synthetic antioxidants etc.). On the basis of research conducted thus far, one cannot eliminate the possibility that dietary fish oil could have some effect against malaria other than increased oxidative stress. However, it has long been appreciated that plasmodia are remarkably sensitive to oxidative stress and are not able to grow well under normal O2 tensions (Scheibel et al. 1979). Thus, the concept that parasite growth is suppressed by feeding the host a pro-oxidant diet seems eminently plausible. By forcing the parasite to incorporate highly-unsaturated fatty acids into its membranes in the absence of vitamin E (plasmodia cannot synthesize their own fatty acids and must rely on host nutrition to fulfil this need), the already precarious balance between pro-oxidant and antioxidant factors in the parasite is upset and the parasite is killed. Current research does not allow us to conclude whether this chain of events occurs at the level of the parasite or host erythrocyte membrane or both.

One might ask whether the dietary fish oil in the vitamin E-deficient diet acts by stimulating the immune response of the host against the parasite. However, feeding fish oil generally suppresses rather than stimulates innate immune function (Kelley & Daudu, 1993). Acquired immune response, also, does not appear to play a role here either, since the protective effect of the fish oil vitamin E-deficient diet was seen equally well in nude (athymic) or SCID-beige mice (D. W. Taylor, O. A. Levander, V. R. Krishna, C. B. Evans, V. C. Morris and J. R. Barta, unpublished results). Nonetheless, once mice are cured of malaria by the pro-oxidant diet, they remain resistant to subsequent malarial rechallenge no matter which diet is fed to the host (Ager et al. 1992). Thus, short-term nutritional protection against malaria by feeding the vitamin E-deficient diet with fish oil appears to confer long-term immunological protection against the parasite. A similar phenomenon has recently been reported in Mg-deficient mice infected with Babesia hyloymphsci (Maurois et al. 1994). Babesia belong to the same phylum of protozoan parasites as Plasmodium and Mg deficiency is known to exert an increased oxidative stress on erythrocytes in hamsters (Mesocricetus auratus; Freedman et al. 1992). In his pioneering studies, Godfrey (1957b) showed that fish oil in a low-vitamin E diet protected against Babesia rodhaini in mice.

Our success in controlling plasmodial growth by manipulating host oxidative stress status through diet has led us to try this approach with other parasitic diseases. Feeding fish or flaxseed oil in a normal starter ration decreased caecal lesions in chicks infected with the coccidial parasite, Eimeria tenella (Allen et al. 1994). Like Babesia, Eimeria is of the same phylum as Plasmodium, although its life-cycle and host habitats are quite different. Coccidiosis is an economically significant problem for the poultry industry and the parasite has developed resistance to many commonly used anti-coccidial drugs, so any new leads to the control of the parasite could be useful. On the basis of an earlier report that feeding a diet deficient in vitamin E and Se to mice impaired development of Schistosoma mansoni (DeWitt, 1957), we tested the effect of the vitamin E-deficient diet with fish oil on the progression of schistosomiasis in mice (Morris et al. 1994). As shown by others (Fusco et al. 1992), dietary menhaden oil increased the cercarial penetration by S. mansoni. However, we found that feeding menhaden oil in a vitamin E-deficient diet
decreased the liver surface nodule scores and the number of adult worms despite the increased cercarial penetration. Given the fact that schistosomes contain a Se-dependent glutathione peroxidase (Roche et al. 1994), it might be profitable to examine the influence of host Se deficiency on the growth and maturation of S. mansoni.

Will the dietary manipulation of host oxidative stress status ever find application in attacking human malaria? Malaria is now the leading killer of children under 5 years in sub-Saharan Africa and the World Health Organization estimates that globally the number of persons infected with malaria is increasing at a rate of approximately 5%/year (Kolberg, 1994). In vitro addition of n-3 fatty acids inhibits the growth of P. falciparum (Fevang et al. 1992; Kumaratilake et al. 1992) so that the phenomena discussed in the present review are apparently not limited only to murine malaria. On the other hand, serum and erythrocytes collected from an individual supplemented with tocopherol-fortified fish oil had no effect on the in vitro growth of blood stages of P. falciparum (Abu-Zeid et al. 1993). Supplementation of normal (laboratory chow) diets with menhaden oil resulted in substantial (though not complete) protection against murine malaria so that vitamin E deficiency does not seem to be required to obtain at least some beneficial effect of the fish oil (Levander & Ager, 1993). A pilot study conducted in China indicated that patients clinically cured of malaria with QHS recrudesced more slowly when given a supplement of fish oil concentrate not containing vitamin E, although all patients eventually redeveloped the disease (Levander et al. 1994). Perhaps fish oil might find a use in the therapy of human malaria as an adjunct to treatment with pro-oxidant drugs such as those related to QHS. Low doses of artelinic acid, a derivative of QHS, had a more powerful chemosuppressive action in mice fed on menhaden oil than in mice fed on lard even though both groups were supplemented with vitamin E (Levander & Ager, 1993). Additional research is needed to determine what, if any, role dietary-induced oxidative stress may play in the control of human malaria.

KESHAN DISEASE

The discovery that nutritional levels of Se protect against the endemic juvenile cardiomyopathy known in China as Keshan disease has to rank among the greatest triumphs of public health trace element nutrition. Several lines of evidence suggest that low Se status is the basic underlying cause of this condition. For example, Keshan disease has been linked to many indicators of poor Se nutrition including decreased blood and hair Se content and low dietary Se intakes (Keshan Disease Research Group, 1979a). Moreover, the Chinese scientists carried out an extensive Se supplementation trial which provided convincing evidence for the ability of Se to prevent the disease (Keshan Disease Research Group, 1979b).

Despite their obvious success in associating Keshan disease with impaired Se nutrition, the Chinese scientists were always very careful to point out that certain features of the disease could not readily be explained on the basis of poor Se status alone (Yang et al. 1988). In particular, there was an appreciable seasonal and annual variation in the incidence of the disease which suggested an infectious component. Several enteroviruses were isolated from Keshan disease victims. One such virus, coxsackie B4, was found to cause extensive damage to heart muscle when injected into mice fed on a low-Se diet based on grains from a Keshan disease area (Ge et al. 1987). Supplementation of the
Table 3. Effect of vitamin E and dietary fat on the heart damage caused by a myocarditic coxsackievirus (CVB3/20) in mice* (Adapted from Beck et al. 1994a)

(Mean values and standard deviations for five or ten animals)

<table>
<thead>
<tr>
<th>Vitamin E (mg/kg)</th>
<th>Dietary fat (g/kg)</th>
<th>Histopathological score at intervals (d) after inoculation</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Lard</td>
<td>Menhaden</td>
</tr>
<tr>
<td>38.4</td>
<td>40</td>
<td>0</td>
</tr>
<tr>
<td>0</td>
<td>40</td>
<td>0</td>
</tr>
<tr>
<td>38.4</td>
<td>0</td>
<td>40</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>40</td>
</tr>
</tbody>
</table>

a, b, c, d: Means with unlike superscript letters were significantly different (P<0.05).

* Weanling male C3H/HeJ mice were fed on a torula yeast-based diet containing lard, menhaden oil or corn oil and 0.2 µg Se as sodium selenite/g either with or without vitamin E (RRR-α-tocopheryl acetate). After 4 weeks, all mice were inoculated intraperitoneally with 10^5 tissue culture infectious dose-50 of CVB3/20 in 0.1 ml RPMI-1640 medium. Pathological score: 0, no lesions; 1, foci of mononuclear cell inflammation associated with myocardial cell reactive changes without myocardial cell necrosis; 2, inflammatory foci clearly associated with myocardial cell reactive changes; 3, inflammatory foci clearly associated with myocardial cell necrosis and dystrophic calcification; 4, extensive inflammatory infiltration, necrosis and dystrophic calcification.

mice fed on the deficient diet orally or by adding Se to the drinking water before inoculation reduced the severity of heart lesions caused by the virus.

As a follow-up to those studies, Beck et al. (1994a,b,c,d) initiated a series of experiments to determine the effect of Se deficiency on the cardiotoxicity of coxsackievirus B3. In agreement with the Chinese results, Se deficiency increased the histopathological damage caused by a myocarditic strain of the virus, coxsackievirus B3/20 (CVB3/20; Beck et al. 1994c). Likewise, vitamin E deficiency also increased the cardiac damage caused by CVB3/20 and substituting menhaden (fish) oil for the lard in the diet resulted in even greater damage (Table 3; Beck et al. 1994a). Neither Se nor vitamin E deficiency had any effect on neutralizing antibody response or natural-killer-cell function. However, either deficiency resulted in decreased lymphocyte proliferation as stimulated either by mitogen (Concanavalin A) or by antigen (prepared from membranes of HeLa cells infected with coxsackievirus). This impairment of cell-mediated immune function may have allowed the virus to multiply to a greater extent, thereby accounting for the increased viral titres seen in the hearts of the Se- or vitamin E-deficient mice.

Beck et al. (1994a,b) then extended their nutritional studies to a strain of the virus, coxsackievirus B3/0 (CVB3/0), that is amyocarditic (i.e. causes no heart damage under normal conditions). When the host mice were fed on a diet deficient in either vitamin E or Se, the normally benign CVB3/0 was able to cause mild but significant heart damage (Beck et al. 1994a,b). Furthermore, when CVB3/0 was isolated from the hearts of either vitamin E- or Se-deficient mice, passed through HeLa cells, and re-inoculated into normal (i.e. non-deficient) mice, the virus retained its capacity to cause heart damage. This observation suggested the possibility that passage of the benign CVB3/0 through a deficient host had somehow altered its genetic make-up such that now the virus exhibited virulence. Similar results were obtained with the normally myocarditic virus (CVB3/20) in that the increased virulence apparent in a deficient host was retained after passage.

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through HeLa cells and re-inoculation into normal mice. Therefore, inoculation of either a myocarditic or amycarditic form of the virus into either a Se- or vitamin E-deficient host resulted in an increased virulence that was stable to passage through cell culture and manifested itself as greater heart damage upon re-inoculation into a normal host.

The most plausible, and yet controversial, explanation for these results is that the viral genome mutated as a result of the virus having been passed through an oxidatively-stressed host. It should be emphasized that the viral preparations used here (CVB3/0 and CVB3/20) are homogeneous cloned and sequenced viruses, not the heterogeneous viral population that you might expect to find in a wild-type virus. This would rule out the possibility that the growth of a particular sub-population within the original inoculum might be favoured under conditions of host oxidative stress, thereby leading to an apparent genotypic change. Coxsackievirus is a non-enveloped virus, therefore it is difficult to visualize how such a virus could pick up host factors that might influence virulence and retain them during passage through cell culture.

If indeed a genetic change in the virus is the explanation for our results, these experiments would provide a novel insight into our understanding of host–virus interrelationships. To our knowledge, these studies would provide the first example of an influence of host nutritional status on the genetic constitution of a virus. How could such a change be accomplished? Perhaps dietary oxidant stress compromises the host immune system so that the virus replicates to higher titres thereby increasing the chances of mutation. RNA viruses, such as the coxsackievirus, are characterized by relatively high mutation rates, in the range of $10^{-3}$–$10^{-5}$ substitutions per nucleotide copied (Domingo & Holland, 1994). Up to ten genomic changes may account for the difference in cardiovirulence between the CVB3/0 and CVB3/20 described previously (Chapman et al. 1994). On the other hand, only a single amino acid residue accounted for the virulent phenotype of a pancreatropic variant of coxsackievirus B4 (Caggana et al. 1993).

What would be the consequences of any such nutrition-driven mutation in the viral genome? Morse (1993a) has coined the term ‘emerging viruses’ to refer to ‘viruses that either have newly appeared in the population or are rapidly expanding their range, with a corresponding increase in cases of disease.’ Change in host nutritional status was thought to occasionally play a role in the recognition of viruses already widespread in the human population through the precipitation of new or more serious disease. Increased viral traffic (i.e. transfer and dissemination of viruses to new host populations) was considered the major determinant of ‘new’ viral diseases with viral evolution being of lesser relative importance. Our results with dietary-induced oxidative stress, however, suggest that mutation rates in RNA viruses may be accelerated greatly during nutritional deprivation so that evolution of new viruses may assume greater significance under some conditions.

Our dietary antioxidant deficiency–viral infection model has been useful also in exploring other host factors that determine virus pathogenicity. For example, mice are known to develop resistance to the coxsackievirus as they get older so that after about 8 weeks of age they become refractory to viral challenge. If mice were fed on a Se-deficient diet, however, they retained susceptibility to CVB3-induced myocarditis up to 14 weeks of age (Beck et al. 1994d). Susceptibility of mice to CVB3-induced myocarditis is also a function of genetic background of the host since different strains of mice vary widely in their ability to resist myocardial disease. The biochemical basis for these differences in susceptibility to viral disease due to host genetics is not completely understood but may be related to several factors such as the presence of viral receptors, interferon production.
Table 4. **Contrasting effects of selenium and vitamin E deficiency on two mouse models of infectious disease**

<table>
<thead>
<tr>
<th>Test diet</th>
<th>Effect of deficiency</th>
<th>Malaria</th>
<th>Myocarditis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vitamin E-deficient (lard)</td>
<td>++</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>Vitamin E-deficient (fish oil)</td>
<td>++</td>
<td>--</td>
<td></td>
</tr>
<tr>
<td>Se-deficient</td>
<td>0</td>
<td>++</td>
<td>--</td>
</tr>
</tbody>
</table>

+ + , Very beneficial effect; +, beneficial effect; 0, no effect; −, detrimental effect; −−, very detrimental effect.

or major histocompatibility complex haplotypes. We were able to render mice genetically resistant to CVB3-induced myocarditis susceptible by feeding them a diet deficient in both Se and vitamin E (Williams et al. 1994).

Our results also could have important implications regarding the aetiology of Keshan disease. If an enterovirus such as coxsackievirus is involved in Keshan disease, the mouse studies indicate that vitamin E, as well as Se, could have a role in the disease since deficiency of either nutrient increased the heart damage caused by the virus. Indeed, marginal vitamin E nutriture was reported in residents of Keshan-disease areas (Yang et al. 1984; Xia et al. 1989). Moreover, rats fed on diets comprised of cereals from an endemic area developed liver necrosis, indicating that the diet was low in both vitamin E and Se (Ge et al. 1987). On the basis of changes in rat heart muscle mitochondria, Yang et al. (1994) recently concluded that Se deficiency alone is not sufficient to account for the pathogenesis of Keshan disease. Other dietary components, including vitamin E, were suggested as possible complicating factors. Further work is needed to establish the relative role of Se and vitamin E in the aetiology of Keshan disease.

**CONCLUDING REMARKS**

The results reviewed here demonstrate the profound impact of dietary oxidative stress on the outcome of infection in two mouse models of disease (Table 4). In one case (malaria), vitamin E deficiency protected against the disease, whereas in the other (viral myocarditis), vitamin E deficiency made the condition worse. On the other hand, Se deficiency had no effect on the course of murine malaria, but strongly exacerbated viral-induced heart damage. Work by others has indicated that oxidative stress may be involved in a number of other parasitic and viral diseases such as trypanosomiasis (Godfrey, 1958; Docampo & Moreno, 1984), babesiosis (Godfrey, 1957b; Maurois et al. 1994), HIV/AIDS (Droge et al. 1992; Greenspan & Aruoma, 1994) or influenza (Hennet et al. 1992). Should our results linking an increase in the pathogenesis of coxsackievirus with the host diet be explained by alterations in the genome of the virus, then the implications for human health of similar genetic changes possibly occurring in other viruses might be profound indeed.

Our studies are the product of effective collaboration of a nutritional biochemist, a parasitologist, and a viral immunologist. Such collaborations can lead to novel, and sometimes unanticipated, results. In the preface to his recent book, Professor S. S. Morse pointed out that “finding answers to the questions posed by emerging viruses

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requires attacking the problem from several different perspectives, often crossing disciplinary lines' (Morse, 1993b). And yet the difficulties confronting such interdisciplinary collaborations are well known (organizational obstacles, funding impediments, etc.; Kahn & Prager, 1994). Nonetheless, it is hoped that others will join in exploring the fascinating frontier delineated by the interface of oxidative stress status and infection.

Note added in proof: Beck et al. (1995) have in fact recently provided evidence that the genomic sequence of an avirulent Coxsackievirus B3 is altered as a result of being passed through a Se-deficient mouse.

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