Total and Cancer Mortality After Supplementation With Vitamins and Minerals: Follow-up of the Linxian General Population Nutrition Intervention Trial

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Background
The General Population Nutrition Intervention Trial was a randomized primary esophageal and gastric cancer prevention trial conducted from 1985 to 1991, in which 29,584 adult participants in Linxian, China, were given daily vitamin and mineral supplements. Treatment with “factor D,” a combination of 50 µg selenium, 30 mg vitamin E, and 15 mg beta-carotene, led to decreased mortality from all causes, cancer overall, and gastric cancer. Here, we present 10-year follow-up after the end of active intervention.

Methods
Participants were assessed by periodic data collection, monthly visits by village health workers, and quarterly review of the Linxian Cancer Registry. Hazard ratios (HRs) and 95% confidence intervals (CIs) for the cumulative effects of four vitamin and mineral supplementation regimens were calculated using adjusted proportional hazards models.

Results
Through May 31, 2001, 276 participants were lost to follow-up; 9,727 died, including 3,242 from cancer (1,515 from esophageal cancer and 1,199 from gastric cancer). Participants who received factor D had lower overall mortality (HR = 0.95, 95% CI = 0.91 to 0.99; P = .009; reduction in cumulative mortality from 33.62% to 32.19%) and gastric cancer mortality (HR = 0.89, 95% CI = 0.79 to 1.00; P = .043; reduction in cumulative gastric cancer mortality from 4.28% to 3.84%) than subjects who did not receive factor D. Reductions were mostly attributable to benefits to subjects younger than 55 years. Esophageal cancer deaths between those who did and did not receive factor D were not different overall; however, decreased 17% among participants younger than 55 (HR = 0.83, 95% CI = 0.71 to 0.98; P = .025) but increased 14% among those aged 55 years or older (HR = 1.14, 95% CI = 1.00 to 1.30; P = .47). Vitamin A and zinc supplementation was associated with increased total and stroke mortality; vitamin C and molybdenum supplementation, with decreased stroke mortality.

Conclusion
The beneficial effects of selenium, vitamin E, and beta-carotene on mortality were still evident up to 10 years after the cessation of supplementation and were consistently greater in younger participants. Late effects of other supplementation regimens were also observed.


With incidence rates exceeding 100 per 10,000 person-years, the people of Linxian, China, have some of the highest rates of esophageal squamous cell carcinoma and gastric cardia adenocarcinoma in the world (1,2). Several studies in the early 1980s showed that nutritional deficiencies were common in this area, suggesting a link between these deficiencies and the high cancer rates (3). The Linxian General Population Nutrition Intervention Trial (NIT), a large-scale, randomized, double-blind, primary prevention trial, was designed to test the efficacy of four combinations of vitamins and minerals in reducing esophageal and gastric cardia cancer incidence and mortality in Linxian (4–6). The results of this trial, which started March 1, 1986, and concluded May 31, 1991, showed that supplementation with the antioxidant combination of selenium, vitamin E, and beta-carotene statistically significantly reduced total mortality, total cancer mortality, and gastric cancer mortality (5). The identification of statistically significant effects raises questions about the durability of these effects and also potential good or bad late effects related to supplementation.
Subjects and Methods

**Design of the Trial and Posttrial Follow-up**

The design and conduct of the Linxian General Population NIT and its extended follow-up have been described elsewhere (4–6). In brief, participants were recruited in 1985 from four northern communes in Linxian, a rural county in Henan Province. Residents 40–69 years of age with no history of cancer or debilitating disease were eligible for this trial and were asked to enroll. In all, 29,584 subjects, 60% of those invited, were randomly assigned in the trial. These individuals were interviewed for medical history, family history of cancer, diet, and alcohol and tobacco consumption; were given a brief medical exam; and were asked to donate 10-mL blood sample.

The nine nutrients studied in this trial and their daily doses were retinol (5000 IU, as retinol palmitate), zinc (22.5 mg, as zinc oxide), riboflavin (3.2 mg), niacin (40 mg), ascorbic acid (120 mg), molybdenum (30 µg, as molybdenum yeast complex), selenium (50 µg, as selenium yeast), alpha-tocopherol (30 mg), and beta-carotene (15 mg). Doses ranged from one to two times US Recommended Daily Allowances. An independent study of each of these nutrients and vitamins, although desirable, was not practical. Therefore, these nine nutrients were combined into four regimens or factors: retinol and zinc (factor A); riboflavin and niacin (factor B); vitamin C and molybdenum (factor C); and selenium, vitamin E, and beta-carotene (factor D).

Ten years after the end of the trial, participants who took factor D still had a 5% reduction in total mortality and 11% reduction in gastric cancer; these effects were concentrated among participants younger than 55 years. Esophageal cancer decreased 17% in participants younger than 55 years, but increased 14% in those older than 55 years.

**Contribution**

Sustained benefits were associated with a combination of selenium, vitamin E, and beta-carotene supplementation. More subtle long-term effects were also observed for other vitamin supplements.

**Implications**

Sustained benefits were associated with a combination of selenium, vitamin E, and beta-carotene supplementation. More subtle long-term effects were also observed for other vitamin supplements.

Since the conclusion of active trial treatment in 1991, follow-up has continued on all participants to collect data on cancer incidence and all-cause mortality. Here we report mortality results for the total 15.25 years of the trial and posttrial follow-up (through May 31, 2001) for the original a priori trial endpoints, that is, the effects of the intervention agents on esophageal and gastric cardia cancer mortality, and for all-cause mortality.

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The conduct of the Linxian General Population NIT was approved by the institutional review boards of the Cancer Institute of Chinese Academy of Medical Sciences and the US National Cancer Institute, and written informed consent was obtained from all participants for participation. The trial was registered as ClinicalTrials.gov number, NCT00342654.

**Statistical Methods**

The main outcomes of this study were esophageal and gastric cardia cancer mortality and total mortality. Participants were censored at their last known follow-up date, date of death, or the administrative closure of follow-up for the study (May 31, 2001), whichever came first. The 15.25-year follow-up period was analyzed as a whole and in three 5-year periods: the trial period (March 1, 1986, to May 31, 1991) when the intervention agents were given and two other periods (June 1, 1991, to May 31, 1996, and June 1, 1996, to May 31, 2001) when active follow-up continued but no additional intervention was performed. Among the cancers, both esophageal squamous cell carcinoma and gastric cardia adenocarcinoma occur at epidemic rates in this population, share some etiologic risk factors, and before widespread use of endoscopy and biopsy, were diagnosed as a single disease referred to as “esophageal cancer” or “hard of swallowing disease” (7). Through 2001, esophageal adenocarcinoma was nonexistent in Linxian. We present data for the effects of the supplements on mortality of esophageal squamous cell carcinoma, gastric cardia adenocarcinoma, gastric noncardia adenocarcinoma, and the combination of esophageal and gastric cardia cancer (the original hard of swallowing disease).

We tabulated baseline frequencies and percentages of participants by demographics. We compared risk between those who received each factor and those who did not. In the analysis of factorial trials, this kind of analysis is known as “at the margins analysis” and has the most power to examine the effect of each factor (8). Cox proportional hazards models were used to calculate hazard ratios (HRs) and 95% confidence intervals (95% CIs) for each factor, adjusting for the other three treatment factors, sex, age (continuous), and commune (four communes). These analyses were conducted on 29,559 of 29,584 initial study participants; 25 were lost to follow-up before the intervention began (Figure 1). Interactions between each of the factors and age group or sex were examined by including appropriate terms in the Cox models. This testing of interactions between each of the treatments (factors) and age and sex was planned a priori (9,10) The 55-year age cut point was chosen because it is the midpoint in the 40- to 69-year age distribution of the trial population. Kaplan–Meier estimates of cumulative event rates were plotted to compare time to death for each factor, for all subjects, and by sex and age group (11). In addition to “pure” risks determined from Cox models and Kaplan–Meier estimates, we also estimated selected “crude” risks without adjustment for covariates (ie, cumulative incidence), analogous to the approach of Fine and Gray (12). The pure cumulative risks slightly exceeded the crude cumulative risks, as expected (data not shown). Indeed, the estimated pure risks exceeded the crude risk by no more than 0.01 for total mortality, total cancer mortality, gastric cancer mortality, and esophageal cancer mortality. To estimate the calendar time–specific hazard ratios for the figures, we calculated smoothed (moving) hazard ratio estimates using a generalized additive model with a local regression (loess) smoother with span 0.4 and a test-based confidence band.

The assumption of a constant treatment hazard ratio across the three study periods was verified for each of the analyses by examining treatment by time period interactions. All P values are two-sided and P values less than .05 were considered statistically significant. Analyses were conducted using SAS version 9.1.3 service pack 4 (SAS Institute, Inc, Cary, NC); figures were produced

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**Figure 1.** CONSORT flow diagram of the Linxian General Population Trial.
using S-Plus 6.2 for Windows and Sigma Plot 8.0 (Systat Software, Inc, San Jose, CA).

Results

Through May 31, 2001, there were 381,954 total person-years of follow-up over a median of 15.25 years of observation. The baseline demographic characteristics, smoking and alcohol use, and family history of esophageal cancer for all subjects are shown in Table 1. As expected, there were no statistically significant differences between any of these baseline characteristics by treatment group assignments. Smoking prevalence was comparable and alcohol consumption was much less common than in the United States. A family history of esophageal cancer was substantially more common than in the West.

Total and cause-specific numbers of deaths for the entire study period and for the three 5-year periods are shown in Table 2. A total of 9727 deaths were reported during the 15.25 years of follow-up. Deaths due to cancer and cerebrovascular diseases each accounted for approximately one-third of the total deaths. The most common specific causes of death were cerebrovascular events (n = 2984), esophageal cancer (n = 1515), and gastric cancer (n = 1199). Total numbers of deaths were 2528 in the trial period (follow-up period 1), 3555 in follow-up period 2, and 3644 in follow-up period 3.

Adjusted hazard ratios (95% confidence intervals) for the associations between each intervention factor and total and cause-specific deaths for the entire study period and for each of the 5-year follow-up periods are also shown in Table 2. Subjects given factor A (retinol and zinc) had a marginally increased risk of total mortality for the total follow-up period (HR = 1.04, 95% CI = 1.00 to 1.09, P = .035) compared with subjects not given factor A. In the factor A group, the hazard ratio point estimate for total deaths remained consistently higher than unity in all three follow-up periods, with period-specific estimates of 1.03, 1.05, and 1.05, respectively, although none of the period-specific estimates was statistically significant. This increased risk was mainly due to the effect of factor A on noncancer deaths: The hazard ratio for cerebrovascular deaths was 1.08 (95% CI = 1.00 to 1.16; P = .045) and for other causes of death was 1.06 (95% CI = 0.99 to 1.13; P = .088). The hazard ratio point estimates associated with these causes of death remained higher than unity (although not statistically significant) for all three follow-up periods. Neither sex (interaction P = .372) nor age group (interaction P = .757) statistically significantly modified the effect of factor A on total mortality. Factor A was also associated with period-specific protective effects for other cancer deaths in period 2 (HR = 0.76, 95% CI = 0.58 to 1.00; P = .050) and for cardia cancer deaths in period 3 (HR = 0.73, 95% CI = 0.57 to 0.93; P = .012), although neither effect was consistent in the other time periods.

Intervention with factor B (riboflavin and niacin) was not associated with total deaths in the overall (HR = 0.98, 95% CI = 0.94 to 1.02; P = .318) or period-specific analyses when compared with lack of factor B, and hazard ratio point estimates remained close to 1 in all three follow-up periods (Table 2). This factor was not associated with cause-specific deaths overall or in any follow-up period, except for a marginally statistically significantly protective effect on total cancer deaths in follow-up period 2 (HR = 0.89, 95% CI = 0.80 to 1.00; P = .043). The effect of factor B on total deaths was not statistically significantly modified by sex (interaction P = .177) or age group (interaction P = .109).

Factor C (vitamin C and molybdenum) was not associated with total deaths in overall (HR = 0.97, 95% CI = 0.94 to 1.01; P = .177) or period-specific analyses when compared with lack of factor C (Table 2). However, this factor was inversely associated with overall risk of cerebrovascular deaths (HR = 0.92, 95% CI = 0.86 to 0.99; P = .023), an effect not seen during the trial period but which became apparent during later follow-up. There were no statistically significant associations between factor C and total cancer mortality, esophageal or gastric cancer mortality, or other causes of mortality in the overall or period-specific analyses, although a marginally statistically insignificant increase in the combined esophageal and cardia cancer mortality endpoint was noted in the overall analysis (HR = 1.08, 95% CI = 1.00 to 1.17; P = .052). Sex did not modify the effect of factor C on total mortality (interaction P = .290), but there was a statistically significant interaction with age (interaction P = .003) such that younger (<55 years at random assignment) participants had reduced risk (HR = 0.89, 95% CI = 0.83 to 0.96; P = .001) and older (≥55 years at random assignment) participants were unaffected (HR = 1.01, 95% CI = 0.97 to 1.06; P = .607). Statistically significant age group interactions were also seen for factor C with total cancer mortality and esophageal cancer mortality; younger patients were unaffected but older participants had elevated hazard ratios (Table 3).

Factor D (selenium, vitamin E, and beta-carotene) reduced total mortality (HR = 0.95, 95% CI = 0.91 to 0.99; P = .009; reduction in cumulative mortality from 33.62% in the no-factor D group to 32.19% in the factor D group). Moving hazard ratio curves (Figure 2)

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Table 1. Baseline demographic characteristics, smoking and alcohol consumption, and family history of cancer

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>All participants (% of total)</th>
<th>Range for the eight treatment arms</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. of participants</td>
<td>29 659</td>
<td>3688–3708</td>
</tr>
<tr>
<td>Age group (y)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;50</td>
<td>12 368 (42%)</td>
<td>42%</td>
</tr>
<tr>
<td>50–59</td>
<td>10 258 (35%)</td>
<td>34%–35%</td>
</tr>
<tr>
<td>≥60</td>
<td>6936 (23%)</td>
<td>23%–24%</td>
</tr>
<tr>
<td>Sex</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>16 378 (55%)</td>
<td>55%–56%</td>
</tr>
<tr>
<td>Male</td>
<td>13 181 (45%)</td>
<td>44%–45%</td>
</tr>
<tr>
<td>Smoking*,†</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nonsmoker</td>
<td>20 613 (70%)</td>
<td>70%–71%</td>
</tr>
<tr>
<td>Smoker</td>
<td>8842 (30%)</td>
<td>29%–30%</td>
</tr>
<tr>
<td>Alcohol drinking†,‡</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nondrinker</td>
<td>22 539 (77%)</td>
<td>76%–77%</td>
</tr>
<tr>
<td>Drinker</td>
<td>6915 (23%)</td>
<td>23%–24%</td>
</tr>
<tr>
<td>Family history of esophageal cancer†</td>
<td></td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>20 800 (71%)</td>
<td>70%–72%</td>
</tr>
<tr>
<td>Yes</td>
<td>8651 (29%)</td>
<td>28%–30%</td>
</tr>
</tbody>
</table>

* Ever smoking cigarettes for 6 or more months.
† Data on smoking (n = 104), drinking (n = 105), and family history (n = 108) were not available for some subjects.
‡ Ever drinking any alcoholic beverage in the last 12 months.
show the smoothed hazard ratios remained less than 1.0 for most of the 15.25 years of observation, indicating a protective effect during most of this period. The estimated hazard ratios for the three follow-up periods were 0.91 (95% CI = 0.84 to 0.99; P = .023), 0.99 (95% CI = 0.93 to 1.06; P = .723), and 0.94 (95% CI = 0.88 to 1.00; P = .460), respectively (Table 2). The hazard ratios for the three major group causes of mortality for the full 15.25 years of follow-up were also less than 1.0 but were not uniformly statistically significant: 0.95 (95% CI = 0.89 to 1.02; P = .148) for cancer, 0.98 (95% CI = 0.98 to 1.05; P = .613) for cerebrovascular events, and 0.92 (95% CI = 0.86 to 0.98; P = .013) for other causes. The majority of the overall effect was attributed to a reduced risk of death from gastric cancer and causes of death other than cancer or cerebrovascular diseases.
The effect of factor D on total mortality was not modified by sex (interaction $P = .843$), but was modified by age group (interaction $P = .024$). Total and age-specific cumulative event and moving hazard ratio curves presented in Figure 2 show cumulative mortality by factor D. The hazard ratios were $0.95$ (95% CI = 0.91 to 0.99; $P = .009$) for all study subjects, $0.88$ (95% CI = 0.82 to 0.95; $P < .001$) for subjects younger than 55 years old at study entry, and $0.98$ (95% CI = 0.94 to 1.03; $P = .367$) for subjects 55 or older at study entry (Table 3). Therefore, virtually the entire effect of factor D on total mortality was due to effects in individuals younger than 55 years.

Similar cumulative event rate curves and moving hazard ratio curves for total cancer mortality (Figure 3), total gastric cancer...
mortality (Figure 4), and esophageal cancer mortality (Figure 5) all show that the effects of factor D were predominantly or exclusively in subjects younger than 55 years. The hazard ratios for total cancer mortality associated with factor D were 0.95 (95% CI = 0.89 to 1.02; \( P = .047 \)) for all subjects, 0.85 (95% CI = 0.76 to 0.95; \( P = .046 \)) for those younger than 55, and 1.102 (95% CI = 0.94 to 1.12; \( P = .976 \)) for those 55 or older (Table 3). Corresponding hazard ratios for gastric cancer mortality were 0.89 (95% CI = 0.79 to 1.00; \( P = .043 \)), 0.83 (95% CI = 0.69 to 1.00; \( P = .046 \)), and 0.93 (95% CI = 0.80 to 1.07; \( P = .307 \)). Cumulative crude gastric cancer mortality for all subjects was 4.2% in the no-factor D group compared with 3.8% in the factor D group, an overall reduction of 0.4%. For esophageal cancer mortality, effect modification by age was even more pronounced. There was no overall association between factor D and esophageal cancer mortality for all subjects (HR = 1.101, 95% CI = 0.91 to 1.11; \( P = .905 \)); however, in subjects younger than 55 years, factor D esophageal cancer mortality decreased (HR = 0.83, 95% CI = 0.71 to 0.98; \( P = .025 \)), whereas in individuals aged 55 years or older, it increased (HR = 1.14, 95% CI = 1.00 to 1.30; \( P = .047 \)) (Table 3).

## Discussion

The initial results of the Linxian General Population NIT, published in 1993, showed no association between factors A, B, or C and overall mortality, total cancer mortality, or mortality from esophageal or gastric cancers (5). However, factor D, which included selenium, vitamin E, and beta-carotene, statistically significantly reduced total mortality, total cancer mortality, and mortality from gastric cancer (5). An important question remained, however, whether the preventive effects of factor D would last beyond the trial period. The results of the continued follow-up show that hazard ratios, as indicated by moving hazard ratio curves, remained less than 1.0 for each of these endpoints for the majority of the follow-up period; 10 years after completion of the trial, the group that received factor D still showed a 5% reduction in total mortality and an 11% reduction in gastric cancer mortality. Overall, one in 70 people who took factor D was spared death from all causes, and one in 227 was spared death from gastric cancer.

Stratification of results by sex and age was planned a priori. There were no statistically significant interactions with sex. However, when stratified by age, factor D had a strong protective effect in individuals younger than 55 years but almost no effect on subjects aged 55 years or older. This pattern was seen consistently for total mortality, total cancer mortality, gastric cancer mortality, and esophageal cancer mortality. Indeed, the effect of factor D on esophageal cancer was reversed by age, showing a protective effect for younger but a harmful effect for older individuals.

Because this trial provided selenium, vitamin E, and beta-carotene as one factor, it was not possible to disentangle the effects of these three supplements. However, observational case-cohort studies using subjects in this cohort and patients with upper...
gastrointestinal tract cancers that developed during the intervention period showed inverse associations between risk of esophageal cancer and baseline serum levels of selenium and alpha-tocopherol, but not beta-carotene (13–15). Higher baseline serum selenium also was associated with reduced risk of gastric cardia cancer (13). These results suggest that the protective effects seen in the randomized trial were due to the selenium and vitamin E components. In a subcohort of 1103 subjects from this trial followed through May 31, 2001, higher baseline serum selenium levels also were associated with statistically significant reductions in esophageal and gastric cardia cancer mortality (16). A separate randomized controlled trial in Linxian (17) gave further support for a preventive effect of selenium in subjects with preexisting esophageal squamous dysplasia, the precursor lesion of esophageal squamous cell carcinoma. Compared with control subjects, those with mild dysplasia who received 10 months of daily supplementation with 200 µg of selenomethionine were more likely to have regression and less likely to have progression of their esophageal squamous dysplasia.

In addition to evaluating the durability of the beneficial effects observed during the trial period for factor D, we also evaluated other postintervention events in this trial to look for late effects, and several were noted. When the full 15.25 years of follow-up were considered, nutritional supplementation with factor A (vitamin A and zinc) was associated with increased total mortality, mainly due to an increase in stroke deaths among subjects given factor A compared with those who were not given factor A, whereas supplementation with factor C (vitamin C and molybdenum) was associated with a decrease in stroke deaths and with a slight increase in esophageal/cardia cancer deaths.

Increased mortality among factor A recipients was not expected, given the low retinol levels of the population at the start of the trial, the modest doses of both vitamin A and zinc, and the generally null or protective results from previous observational studies in this cohort relating high serum retinol and tissue zinc concentrations to various cancer endpoints (15,18). In fact, a previous analysis of stroke (the main contributor to the observed increase in total mortality) in this trial showed a protective effect for persons who took the combination of factor A and factor D (HR = 0.71, 95% CI = 0.50 to 1.00, for group AD vs placebo) (19).

We were surprised that factor C, a combination of vitamin C and molybdenum, appeared to be associated with increased risk for the combined esophageal and cardia cancer endpoint, given the well-known role of vitamin C as an antioxidant and inhibitor of carcinogenic N-nitroso compound production in the stomach (20). Furthermore, many epidemiological studies have shown reduced risk of these cancers in persons with high consumption of fruits and vegetables and rich sources of vitamin C, and the only prospective study of plasma vitamin C and gastric cancer showed a protective effect for high concentrations (21). No similar prospective studies of plasma vitamin C values in esophageal cancer are known.
The decreased risk of stroke among trial participants who received factor C was not wholly unexpected. Higher plasma vitamin C levels have been associated with reduced risk of stroke in prospective epidemiological studies (22–24), although randomized trials that have included vitamin C as part of an antioxidant vitamin treatment arm have not shown any effect (25, 26). There are no data on molybdenum and stroke.

The effects of vitamin and mineral supplements on reduction of total mortality and cancer mortality have been heavily debated over the past 25 years. The results of this study need to be interpreted in the context of other trials of vitamin and mineral supplementation. By the 1980s, it was established that antioxidants could quench free oxygen radicals and potentially reduce the risk of cancer by preventing DNA damage by these radicals. Observational epidemiological studies showed inverse associations between cancer incidence and dietary intake of several vitamins and minerals, but more definitive evidence awaited the completion of randomized trials (27). It was generally assumed that prescribing pills would be a more convenient and acceptable way to prevent cancer than proscribing carcinogens. These facts and assumptions motivated the design and conduct of the first generation of randomized controlled cancer prevention trials, including the Linxian General Population NIT, to reduce cancer risk using vitamins and minerals. The results of the largest and most informative of these trials (i.e., those with more than 10,000 participants) were often contrary to the initial expectations. Beta-carotene supplementation increased total mortality in both the Alpha-Tocopherol, Beta-Carotene (ATBC) Cancer Prevention Study (28) and the beta-Carotene and Retinol Efficacy Trial (CARET) Study (29), whereas no mortality benefit for beta-carotene was seen in either The Physicians’ Health Study (PHS) (30) or the Women’s Health Study (31). The most likely explanation for the unexpected findings from these four large trials conducted in the West is a mismatch of the design of the trials with the population attributes: each of these trials tested supplements that were associated with increased mortality risk in pharmacological doses of micronutrients in already well-nourished populations (32).

A recent meta-analysis of randomized trials of antioxidant supplements for prevention of cancer and other diseases (33) combined the results of these large-scale studies with many smaller studies and concluded that treatment with beta-carotene, vitamin A, or vitamin E likely increased total mortality, and the effect of vitamin C or selenium on total mortality needed further study. The results of this study agree in part with the meta-analysis conclusions. In this trial, factor A (which included vitamin A) increased mortality, factor C (which included vitamin C) was not associated with overall mortality, and factor D (which included selenium) decreased mortality. However, beta-carotene and vitamin E, two supplements that were associated with increased mortality risk in the meta-analysis, were in factor D, which reduced mortality in this trial. Several potential explanations exist for these apparently discrepant results. First, the protective effect of selenium may have been stronger than the possible deleterious effects of beta-carotene.
and vitamin E in this trial, so the overall effect of factor D was beneficial. This hypothesis is supported by medium-sized trials that have shown beneficial effects for selenium in reducing mortality and cancer risk (34–36).

A second possible explanation for these discrepancies is that baseline nutritional status of the populations studied influenced the supplementation effects. The people of Linxian are nutritionally deficient (3,37), so vitamin and mineral supplements may be more beneficial to them than to other populations that have been studied. The results of the Dysplasia NIT (38), a medium-sized randomized nutrition intervention trial that was conducted among subjects with cytologically diagnosed esophageal squamous dysplasia in Linxian at the same time as the General Population NIT, showed that supplementation with 26 minerals and vitamins was associated with a non–statistically significant 7% reduction in mortality risk. Results from the Nutritional Prevention of Cancer Trial showed that the benefits for selenium supplementation on total cancer mortality (39) and the development of prostate cancer (36) were essentially limited to participants with lower selenium levels at the start of the trial. Further support for this hypothesis comes from the data of the meta-analysis of antioxidant supplement trials and total mortality itself (33). We classified the 68 study populations included in this analysis as Western (n = 58), East Asian (n = 8), or other (n = 2) and found statistically significant heterogeneity between the results of the studies performed in Western and East Asian populations. The Western studies had a combined odds ratio (OR) of 1.04 (95% CI = 1.01 to 1.06) and the East Asian studies had a combined OR of 0.92 (95% CI = 0.84 to 1.02) (χ² for heterogeneity P = .02). Because nearly all of the events in the East Asian group came from Linxian, which we know has borderline or deficient nutrition, this difference in meta-analysis results may well reflect differences in the baseline nutritional status of the populations evaluated.

A third possible explanation for the heterogeneity of results observed among studies that have evaluated the association of vitamin supplements and total mortality or gastrointestinal cancer risk is effect modification by the stage of disease at study entry. Our results show that only individuals younger than 55 years benefited from factor D. This result may indicate greater benefit earlier in the course of carcinogenesis and is consistent with a “point of no return,” beyond which supplementation with vitamins is not useful and may be harmful, preferentially benefiting the developing tumor more than the host. This hypothesis may help explain why observational studies, which reflect long-term intake of vitamins and vitamin-containing fruits and vegetables, have usually shown beneficial associations, whereas trials, which have largely been conducted in older patients, have sometimes shown harmful effects from vitamin interventions.

Participants in the ATBC and CARET studies, in addition to being older than those in the Linxian general population trial (ages 50–69 years for ATBC and 45–74 years for CARET), were heavy smokers and some were exposed to asbestos, both powerful carcinogenic exposures that may have put them beyond the point in the disease process that they could benefit from supplements. Detailed analyses of both of these studies have shown that the increased risk associated with vitamin use was almost exclusively seen in current (as opposed to former) smokers (40) and in those who smoked most (41). In contrast, the PHS study, which included fewer than 10% smokers, showed no adverse effect of beta-carotene (30).

In addition to this report, three other cancer prevention nutrition intervention trials have reported results from continued follow-up after the termination of intervention. Follow-up of the participants of the ATBC Study for up to 8 years after the end of the intervention showed that both the harmful effects of beta-carotene (ie, increased total mortality and lung cancer incidence) and the beneficial effect of vitamin E (ie, decreased prostate cancer incidence) disappeared, albeit slowly (42). However, analyses of cerebral infarcts (80% of all strokes) among vitamin E recipients in the ATBC Study showed reversed effects during the trial (relative risk [RR] = 0.86, 95% CI = 0.75 to 0.99) (43) and the 6 years posttrial (RR = 1.13, 95% CI = 1.00 to 1.27) (44). After 6 years of postintervention follow-up in the CARET study, the relative risk of total mortality remained greater than 1.0, but this elevated risk diminished and was no longer statistically significant (45). Lung cancer mortality, however, was still statistically significantly increased. The Calcium Polyp Prevention Trial reported that the protective effect of calcium supplementation on colorectal adenoma recurrence found during the trial period (RR = 0.81, 95% CI = 0.74 to 0.98) (46) continued up to 5 years after supplementation ended and was, if anything, stronger after than during the intervention itself (RR = 0.63, 95% CI = 0.46 to 0.87) (47).

Durability of cancer prevention effects after cessation of intervention has also been observed with nonnutritional agents. In fact, the most consistent example of a sustained cancer prevention effect reported to date from any cancer prevention agent tested in trials is for tamoxifen in the primary prevention of breast cancer. Posttrial follow-up from three tamoxifen trials (48–50) consistently found benefit after the conclusion of active treatment, and in one trial (50), statistically significant reduction in risk (among patients with estrogen receptor-positive tumors) was seen only after treatment had ceased.

This study has several strengths. It was a randomized double-blind design and had excellent compliance and long-term follow-up with virtually complete ascertainment of cases in a well-defined population.

This study also has limitations. Interventions with factors containing multiple agents do not allow evaluation of the effects of individual agents alone, nor were we able to evaluate more than one dose for each of the agents supplemented. The people of Linxian are deficient in many micronutrients, which may limit the generalizability of these results to well-nourished populations. If the protective effects of this study are due to replacement of essential nutrients in a nutritionally deprived population, then similar interventions might be useful in similarly deprived populations in the West, including the United States, although populations with low rates of esophageal and gastric cancer mortality are unlikely to avert as many deaths as high-rate populations such as that in Linxian. Finally, the smoothed hazard ratios that we presented were intended to provide an alternative visual representation of the effects at specific points in time and to complement the cumulative view offered by the Kaplan–Meier curves. These smoothed hazard ratios should be interpreted with caution, however, because the confidence intervals around these curves nearly always include 1.0. Thus, such curves are affected by the play of chance and may be
biased by choice of smoothing parameters, edge effects, and other factors.

It should be noted that the follow-up period occurred during a time of dramatic economic progress in China as a whole. Although documented improvements in dietary intakes in Linxian during follow-up were modest (37), more substantial undocumented changes almost certainly occurred. Effects of dietary improvements should have been evenly distributed across all participants in the various randomized treatment groups in the trial. Thus, if the effects of the supplements and the dietary micronutrient intake are additive, any dietary changes that might have occurred should not bias the treatment group effects in the postintervention period. If instead the benefit from supplementation was to correct a deficiency state to exceed some minimum required threshold, then, if all people started to become less deficient because of dietary improvements over time, the observed treatment effects would be expected to weaken. It is all the more remarkable then that benefits persisted despite this likely improved nutrition and its attendant attenuation of treatment effects. It is also possible that improved diet may have modified effects in the postintervention period, including the enhanced benefit widely observed in younger participants and the emergence of late effects, most notably the benefit for factor C on cerebrovascular deaths.

In summary, 10 years of postintervention follow-up of participants in this cancer prevention trial demonstrated the durability of previously observed beneficial effects on mortality from supplementation with selenium, vitamin E, and beta-carotene. The persistence of risk reduction for up to 10 years after treatment in this trial reinforces the validity of the original trial findings and is consistent with an emerging new paradigm in cancer prevention, namely, that prevention may be achievable with short-term as opposed to life-long treatment. Striking age interactions were seen, suggesting that supplements may be more beneficial in younger age groups. Late beneficial and harmful effects on mortality were not observed during the trial period of supplementation were also seen for other supplementation groups. Durability and late effects should be examined in other prevention trials.

References

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**Notes**

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