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Effect of Calcium Supplementation on Blood Lead Levels in Pregnancy: A Randomized Placebo-Controlled Trial

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BACKGROUND: Prenatal lead exposure is associated with deficits in fetal growth and neurodevelopment. Calcium supplementation may attenuate fetal exposure by inhibiting mobilization of maternal bone lead and/or intestinal absorption of ingested lead.

OBJECTIVE: Our goal was to evaluate the effect of 1,200 mg dietary calcium supplementation on maternal blood lead levels during pregnancy.

METHODS: In a double-blind, randomized, placebo-controlled trial conducted from 2001 through 2003 in Mexico City, we randomly assigned 670 women in their first trimester of pregnancy to ingest calcium (n = 334) or placebo (n = 336). We followed subjects through pregnancy and evaluated the effect of supplementation on maternal blood lead, using an intent-to-treat analysis by a mixed-effects regression model with random intercept, in 557 participants (83%) who completed follow-up. We then conducted as-treated analyses using similar models stratified by treatment compliance.

RESULTS: Adjusting for baseline lead level, age, trimester of pregnancy, and dietary energy and calcium intake, calcium was associated with an average 11% reduction (0.4 µg/dL) in lead blood level relative to placebo (p = 0.004). This reduction was more evident in the second trimester (–14%, p < 0.001) than in the third (–8%, p = 0.107) and was strongest in women who were most compliant (those who consumed ≥ 75% calcium pills; –24%, p < 0.001), had baseline blood lead > 5 µg/dL (–17%, p < 0.01), or reported use of lead-glazed ceramics and high bone lead (–31%, p < 0.01).

CONCLUSION: Calcium supplementation was associated with modest reductions in blood lead when administered during pregnancy and may constitute an important secondary prevention effort to reduce circulating maternal lead and, consequently, fetal exposure.


Despite improvements in environmental policies and significant reductions in average U.S. blood lead levels, lead exposure remains a concern for pregnant and lactating women. This is particularly true among certain population subgroups at increased risk, such as women from developing countries and those with occupational exposures [Centers for Disease Control and Prevention (CDC) 2007; Meyer et al. 2003]. In addition, overall declines in environmental sources highlight maternal bone as a long-lived endogenous source of exposure that poses a potential hazard for the developing fetus and breastfeeding infant (Hu and Hernández-Avila 2002). Redistribution of cumulative maternal bone lead stores into the circulation occurs during periods of increased bone resorption, such as pregnancy and lactation (Gulsion et al. 2003; Manton et al. 2003; Téllez-Rojo et al. 2004). Prenatal lead exposure has adverse influences on infant birth and neurodevelopmental outcomes across a wide range of exposure (Bellinger 2005; Hu et al. 2006), and maternal bone lead has been shown to be an independent risk factor (Gomaa et al. 2002; Gonzalez-Cossío et al. 1997; Hernández-Avila et al. 2002).

The potential role of nutrition in altering susceptibility to lead exposure and toxicity has long been recognized (Aub et al. 1932; Hu et al. 1995; Mahaffey 1974, 1990). Dietary intake concurrent to exposure is known to have an impact on lead dynamics, and nutrients may interact with lead by binding lead in the gut, competing with lead for absorption, altering intestinal cell avidity for lead, and altering affinity of target tissues for lead (Ballew and Bowman 2001).

Inadequate calcium consumption has been shown to increase lead absorption (Heard and Chamberlain 1982) and retention (Six and Goyer 1970). Lead competes with calcium at calcium-binding sites and may subsequently alter protein function and calcium homeostasis (Sauk and Somerman 1991). Evidence indicates that low dietary calcium and vitamin D are risk factors for elevated bone lead levels (Cheng et al. 1998). Higher milk intake during pregnancy also has been associated with lower maternal and umbilical cord lead levels in postpartum women (Hernández-Avila et al. 1997), suggesting that calcium status may be an important factor in the maternal–fetal transfer of lead across the placenta.

Calcium requirements are increased substantially during pregnancy and lactation in order to meet the needs of the developing fetus and nursing infant for skeletal mineralization and growth (Prentice 2000). Maternal calcium homeostasis is maintained by controlling intestinal calcium absorption, renal calcium excretion, and mobilization of skeletal mineral stores (Kovacs and Kronenberg 1997). The role of dietary calcium and mineral adequacy on skeletal changes of pregnancy and lactation is controversial; however, it is recommended that pregnant and breastfeeding women consume 1,000–1,300 mg calcium per day, depending on their age (Institute of Medicine 1997).

In a randomized, double-blind, placebo-controlled trial of 1,200 mg daily calcium supplementation in lactating women, we have previously shown that calcium supplementation reduced maternal lead blood by 15–20% (Hernández-Avila et al. 2003) and breast milk lead by 5–10% (Ettinger et al. 2006) over the course of lactation. Our objective in the present study was to evaluate the effect of 1,200 mg daily calcium supplementation on maternal blood lead levels during pregnancy, the period of greater relevance for maternal–fetal transfer of lead.

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Calcium supplementation and blood lead in pregnancy

Materials and Methods

Study population and design. We recruited pregnant women from 2001 through 2003 at the Mexican Social Security Institute (Instituto Mexicano del Seguro Social) prenatal clinics that serve a low- to moderate-income population in Mexico City. We assessed 3,836 women for eligibility, of whom 1,981 did not meet study eligibility criteria (pregnancy of no more than 14 weeks' gestation; not presenting with a high-risk pregnancy; plans to reside in the metropolitan Mexico City area for ~3 years) or had other reasons not being enrolled (n = 2). Of the remaining 1,855 eligible women, 670 (36%) agreed to participate and signed the informed consent, and were randomly assigned to receive a daily supplement of 1,200 mg calcium [two 600-mg calcium carbonate tablets (Wyeth Consumer Health Care/Lederle Laboratories, Inc., México City, Mexico) at bedtime; n = 334] or placebo (n = 336).

We assessed blood lead levels, dietary calcium intake, and reported use of lead-glazed ceramics (LGC) at three time points: baseline (first trimester), 6 months (second trimester), and 8 months (third trimester). We assessed compliance by pill count at each follow-up visit. We defined women who had at least one blood lead measurement at 6 or 8 months' gestation (n = 565; 84%) as having completed follow-up. Eight women did not have baseline blood lead levels, yielding a total of 557 subjects (83%) available for inclusion in the final analyses (Figure 1).

The research protocol was approved by the Human Subjects Committee of the National Institutes of Public Health, the Mexican Social Security Institute, the Brigham and Women’s Hospital, and the Harvard School of Public Health and complied with both Mexican and U.S. federal guidelines governing the use of human participants. All participating mothers received a detailed explanation of the study intent and procedures and were advised on identifying and avoiding LGC pottery use during pregnancy before signing the approved written informed consent.

Blood lead measurement. Blood lead measurements (1.0 µg/dL = 0.0483 µmol/L) were performed using graphite furnace atomic absorption spectrophotometry (Perkin-Elmer model 3000; Norwalk, CT, USA) at the American British Cowdray (ABC) Hospital Trace Metal Laboratory according to a technique described in Miller et al. (1987). The laboratory participates in the CDC blood lead proficiency testing program administered by the Wisconsin State Laboratory of Hygiene (Madison, WI, USA) and maintained acceptable precision and accuracy over the study period.

Bone lead measurement. At 1 month postpartum (~5 days), maternal bone lead was estimated by a spot-source cadmium-109 K-X-ray fluorescence (K-XRF) instrument at the research facility at the ABC Hospital. We used two 30-min in vivo measurements of each subject’s mid-tibial shaft (representing cortical bone) and patella (trabecular bone). The physical principles, technical specifications, validation, and use of the K-XRF technique have been described in detail elsewhere (Chertle et al. 2003; Hu et al. 1998). For quality control, we excluded bone lead measurements with uncertainty estimates > 10 and 15 µg lead/g mineral bone for tibia and patella, respectively.

Dietary intake. We assessed maternal dietary intake in each trimester of pregnancy using a semiquantitative food frequency questionnaire designed to estimate usual dietary intake over the prior month. We based the questionnaire on the semiquantitative food frequency questionnaires and validation methodology used in the Harvard Nurses’ Health Study and Health Professionals’ Follow-up Study (Willett et al. 1985, 1987). We translated the questionnaire and validated it for use specifically for the Mexican Spanish-speaking adult population (Hernández-Avila et al. 1998).

Statistical analysis. We compared baseline characteristics of participants between the calcium and placebo groups using Wilcoxon rank-sum (Mann–Whitney U-test) two-sample test of equality or Student’s t-test, as appropriate. We performed a similar comparison between those included in the analyses and those lost to follow-up.

We evaluated the effect of calcium supplement on blood lead concentration using an intent-to-treat analysis by means of a mixed-effects regression model with a random intercept for each subject. This approach takes into account the within-subject correlation structure attributed to the repeated measurements, yielding valid standard errors of the effect estimates. Blood lead concentrations in the second and third trimester of pregnancy were the outcome variables; however, we used models featuring natural-log-transformed blood lead because this parameterization provided the best fit. In order not to exclude very low blood lead concentrations from the analysis, we substituted 27 blood lead measurements (1.6% of the total) below the limit of detection (1 µg/dL) with random numbers following a uniform distribution between 0 and 1. We adjusted models for the following baseline variables: first trimester log-transformed blood lead concentration, maternal age (years), treatment group, daily calcium (grams per day) and energy intake (kilo calories per day), and trimester of pregnancy.

To assess the overall intent-to-treat effect of calcium supplementation on blood lead...
concentrations throughout the last two trimesters of pregnancy, we fitted the following model:

$$
\ln(BP_{ij}) = (\alpha + u_i) + \beta_1S_i + \beta_2BP_{ij} + \beta_3C_{it} + \beta_4C_{jt} + \beta_5A_i + \beta_6T_j + \epsilon_{ij}
$$

[1]

where $\ln(BP_{ij})$ is the log-transformed blood lead concentration for subject $i$ at trimester $j$, $\alpha + u_i$ represents the error term associated with the $i$th subject, $S_i$ is the dummy variable that indicates treatment assignment, $\ln(BP_{ij})$ is the log of blood lead concentration for subject $i$ at trimester $j$, $T_j$ is the $j$th trimester of pregnancy, $C_{it}$ is the baseline daily dietary intake, $A_i$ is age, and $\epsilon_{ij}$ denotes the random ratio deviation [$e_{ij} \sim n(0, \sigma^2)$]. The overall treatment effect estimate is the coefficient $\beta_1$.

We fitted a second model to estimate the treatment effect at each trimester:

$$
\ln(BP_{ij}) = (\alpha + u_i) + \beta_1S_i + \beta_2BP_{ij} + \beta_3C_{it} + \beta_4C_{jt} + \beta_5A_i + \beta_6T_j + \epsilon_{ij}
$$

[2]

where $S_i$, $T_j$ denotes the interaction term between blood lead levels and trimester of pregnancy. The treatment effect estimate in the second trimester is the coefficient $\beta_1$, and the effect in the third trimester is $(\beta_1 + \beta_2)$.

We used a secondary dose–response study to further assess the effectiveness of supplementation. We assessed compliance by pill count at each visit and analyzed it as proportion of expected pills used between baseline (first trimester) and end of follow-up (8 months' gestation). We defined treatment compliance group in three ways: $\geq 50\%$ of pills consumed, $\geq 67\%$ of pills consumed, and $\geq 75\%$ of pills consumed. To try to disentangle the effect of calcium supplementation on bone lead mobilization versus gastrointestinal absorption, we developed models with an interaction model for postpartum bone lead concentrations, and thus bone lead mobilization rates, on blood lead would depend on the use of LGC. We fitted the following model:

$$
\ln(Pb_{ij}) = \beta_0 + \beta_1S_i + \beta_2BP_{ij} + \beta_3LGC_{it} + \delta_1S_iBP_{ij} + \delta_2S_iLGC_{jt} + \text{covariates}, + \epsilon_{ij}
$$

[3]

where $Pb_{ij}$ is blood lead concentration for the $i$th subject at the $j$th trimester, $S_i$ is the supplementation group, $BP_{ij}$ is the first available postpartum bone lead measurement, $LGC_{jt}$ is current use of LGC in the $j$th subject at the $i$th trimester, and $\delta_1$ represents the difference in the effect of supplementation between the high and low bone Pb concentration groups, and $\delta_2$ represents the difference in the effect of supplementation between the current use/none of LGC groups. Covariates are baseline blood lead level, baseline daily calcium dietary intake, baseline daily energy intake, age, and trimester of pregnancy. Because we were trying to disentangle biologic mechanisms, we restricted these models to those with $\geq 75\%$ compliance.

Finally, to account for possible heterogeneity of treatment effects according to initial blood lead levels, we also performed analysis by baseline blood lead group ($\leq 5\mu g/dL$ vs. $>5\mu g/dL$) using an intent-to-treat analysis and then among only those women with $\geq 50\%$ compliance.

We performed all statistical analyses using Stata for Windows (version 9.0; StataCorp LP, College Station, TX, USA).
Results

We randomized 670 eligible women to receive calcium supplementation (n = 334) or placebo (n = 336) (Figure 1). Baseline characteristics were largely similar for both the calcium and placebo groups. Mean maternal age was 1 year higher in the control group (26.9 years) than in the calcium group (25.9 years; p = 0.02) (Table 1). Approximately 35% of women reported current use of traditional LGC for storing, preparing, or serving food; however, we found no significant differences by treatment group. Dietary calcium intake, also not significantly different between the two groups, was about 900 mg/day on average. Geometric mean (and geometric standard deviation) prerandomization blood lead levels were 3.8 (2.0) and 4.1 (2.0) µg/dL for the calcium and placebo groups, respectively (p = 0.05).

A total of 565 women (84%) completed follow-up. Comparing the group that completed follow-up (placebo n = 277; calcium n = 288) with those who lost to follow-up [placebo n = 59 (18%); calcium n = 46 (14%)], we found no significant differences by treatment group assignment (p = 0.18). Those women who remained in the study reported higher daily energy intake (p < 0.01) and higher use of LGC (p = 0.04) at baseline. Those women who completed follow-up reported higher current use of LGC (36%) than those who did not complete follow-up (26%); among those completing follow-up, however, we found no significant differences in reported LGC use by treatment group.

In the intent-to-treat analysis (n = 557), calcium supplementation was associated with an overall average reduction of 11% in maternal blood lead concentrations relative to placebo (p = 0.004) (Table 2). In a secondary analysis, this reduction was more evident in the second trimester (14% reduction, p = 0.001) than in the third trimester (8% reduction, p = 0.107). These results did not change when we controlled for hematocrit level (data not shown).

When we assessed the dose–response effect of calcium supplementation for women “as treated” (n = 557) using models stratified by treatment compliance, we saw a clear dose–response effect of calcium on blood lead concentration (Table 3). Among those women who consumed ≥ 50% of pills, calcium was associated on average with a 15% reduction in blood lead levels compared with those taking placebo (p < 0.001). This increased to 19% (p < 0.001) and 24% (p < 0.001) for those who consumed ≥ 67% of pills and ≥ 75% of pills, respectively (p for trend < 0.001). Figure 2 shows the effects of calcium and placebo on maternal blood lead over time among the high-compliance group.

Among the low-compliance group (< 50% of pills consumed), blood lead was higher in the calcium-supplemented group, suggesting that these women were somehow different from the low compliers receiving placebo. In fact, in the group that completed follow-up (n = 565), those with low compliance reported higher current use of LGC in the calcium group (35%) compared with placebo (27%), which might explain the apparent increase in blood lead among the low-compliance group. We found no significant differences in reported LGC use among the high-compliance group.

Figure 3 shows the proportional reduction (95% confidence intervals (CIs)) in blood lead due to calcium supplementation, stratified by use of LGC and patella lead lead, among the high-compliance group. Among women consuming ≥ 75% of pills, those with higher patella bone lead experienced greater reductions than those women with lower bone lead levels, corresponding to a 23% reduction (p = 0.01) for those with no reported use of LGC and a 31% reduction (p < 0.01) for those who reported use of LGC. In this subset of most compliant women with high patella bone lead (> 5 µg/dL) and reported use of LGC, the effect corresponded to an average blood lead reduction of 1.95 µg/dL (95% CI, –0.78 to –2.87).

We repeated the analysis by baseline blood lead group (< 5 µg/dL vs. ≥ 5 µg/dL) using intent-to-treat and as-treated analyses among only those women with compliance ≥ 50% of pills consumed (Table 4). The effects of calcium appeared stronger in the group with higher blood lead at baseline (17% reduction), compared with those with baseline blood lead levels < 5 µg/dL (7% reduction). However, when we restricted the analysis to those women who were more compliant, the reductions were similar between the women with higher (≥ 5 µg/dL) and lower (< 5 µg/dL = 14%) blood lead at baseline. Among those women with low compliance (< 50% of pills; n = 82), those with low baseline blood lead (< 5 µg/dL) appeared to experience a paradoxical effect of calcium on blood lead levels (an increase of 34%). Those who initiated the study with higher blood lead (≥ 5 µg/dL) showed the same average effects of treatment (17% reduction), although not statistically significant. Further analysis revealed, however, that the reported use of LGC in low compliers was higher among the calcium group (35%) than in the placebo group (27%), which may account for the apparent differences in treatment effect (7% vs. 17% reduction) observed in the intent-to-treat analysis by baseline blood lead.

Discussion

In this randomized control trial, calcium supplementation (1,200 mg) was associated with modest reductions in blood lead levels when administered during pregnancy. These effects were clearly stronger with increasing compliance, with a 24% average reduction in the most compliant women, and strongest in those with baseline blood lead level > 5 µg/dL (17% average reduction). In the subset of most compliant women with high patella bone lead (> 5 µg/g) and reported use of LGC, we found the greatest reduction in blood lead of 31%, which corresponds to an average reduction of 1.95 µg/dL (95% CI, –0.78 to –2.87).

These results are consistent with our previously published randomized trial, which showed that dietary calcium supplementation among postpartum women reduced maternal blood lead by 15–20% over the course

Table 4. Effect of calcium supplementation* by baseline blood lead level.

<table>
<thead>
<tr>
<th>Baseline blood lead level</th>
<th>No. (calcium/placebo)</th>
<th>p-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Among all women with follow-up (intent-to-treat analysis)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt; 5 µg/dL</td>
<td>349 (183/166)</td>
<td>0.08</td>
</tr>
<tr>
<td>≥ 5 µg/dL</td>
<td>208 (100/108)</td>
<td>0.003</td>
</tr>
<tr>
<td>Among those women with compliance ≥ 50% (as-treated analysis, among high compliers)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt; 5 µg/dL</td>
<td>292 (162/130)</td>
<td>0.01</td>
</tr>
<tr>
<td>≥ 5 µg/dL</td>
<td>183 (87/96)</td>
<td>0.004</td>
</tr>
<tr>
<td>Among those women with compliance &lt; 50% (as-treated analysis, among low compliers)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt; 5 µg/dL</td>
<td>57 (21/36)</td>
<td>0.02</td>
</tr>
<tr>
<td>≥ 5 µg/dL</td>
<td>25 (13/12)</td>
<td>0.49</td>
</tr>
</tbody>
</table>

*Adjusting for baseline blood lead level (log-transformed), maternal age, dietary calcium intake at baseline, daily energy intake at baseline, treatment group, and trimester of pregnancy. %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Δ %Delta

Figure 3. Blood lead proportional reduction estimates due to calcium supplementation (and 95% CIs), stratified by use of LGC (yes/no) and bone lead level (high/low) among the high-compliance group (> 75% of pills by pill count, adjusting for baseline blood lead, age, dietary calcium intake, daily energy intake, and trimester of pregnancy).
of lactation (Hernández-Avila et al. 2003). In that study, the effect among women who were compliant with supplement use (>50% of pills consumed) and had high bone lead (patella lead >3 μg lead/g bone mineral) was an estimated reduction in mean blood lead of 1.16 μg/dL (95% CI, –2.08 to –0.23).

These results are also consistent with the results of a study by Gulson et al. (2004) of blood lead isotopic ratios during pregnancy among women who had recently immigrated to Australia. The authors found that compared with an earlier group of such women they had studied who had calcium-deficient diets, calcium-replete women had a rise in blood lead levels during pregnancy (with an isotopic fingerprint suggesting the lead came from bone) that occurred later in pregnancy and of a smaller magnitude. Although the use of lead isotopic ratios by Gulson et al. (2004) provided very rigorous and precise methodology to their work, the interpretation with respect to calcium supplementation is limited by the small number of women (<20) in their cases series and thus limited statistical power to detect an association (Altman and Bland 1995) and issues of comparability (e.g., the calcium-deficient women were studied at an earlier time and came from Central Europe, whereas the calcium-replete women were studied at a later time, came from Asia, and were otherwise not matched), making the results of our randomized placebo-controlled trial of particular interest.

The effect of calcium may be exerted, at least in part, by decreasing bone resorption and the consequent mobilization of maternal bone lead stores. In a case–crossover trial of calcium supplementation during the third trimester of pregnancy, we have previously shown that maternal bone resorption, as reflected by urinary cross-linked N-telopeptide, was reduced by an average of 13.6 nM bone collagen equivalents/mM creatinine (14%) compared with placebo (Janakiraman et al. 2003), indicating that calcium supplementation can suppress maternal bone mobilization.

The effects of calcium may also be attributed to decreasing the intestinal absorption of lead and/or increasing the excretion of lead from circulation. In the present study, we did not have pregnancy bone lead levels, and Mexican laws forbidding potential radiation exposure during pregnancy did not allow us to obtain bone lead measurements during pregnancy. However, our observation in the stratum of women with no reported LGC use—that the calcium effect is greater in those with high bone lead—suggests that, in this population, the effect may have been exerted mainly through inhibiting bone resorption.

Average baseline dietary calcium intake for women in our trials of Mexican women was less than the U.S. recommended dietary intake of 1,000–1,300 mg/day for pregnant and lactating women (Institute of Medicine 1997). Levels of dietary calcium intake in our studies were, however, consistent with those reported in the Mexican National Nutrition Survey (Barquera et al. 2003) and in a nationally representative sample of U.S. women of childbearing age (Lee et al. 2005). Hertz-Picciotto et al. (2000) followed 195 women over the course of pregnancy and found a U-shaped pattern of maternal blood lead concentration across pregnancy. The late pregnancy increases were steeper among women with low dietary calcium intake in both the low and high age groups, suggesting that lead redistribution may be more pronounced among pregnant women in calcium-deficient states. It is possible that high amounts of calcium are needed to counterbalance the nutritional needs of the developing fetus (Johnson 2001). Other genetic, hormonal, or lifestyle factors may also be involved (Ettinger et al. 2007).

Nonetheless, dietary calcium intake likely plays a limited, but still important, role in decreasing the mobilization of lead from maternal bone and/or decreasing gastrointestinal absorption of ingested lead, thereby decreasing the risk of fetal and infant exposure. Calcium supplementation during pregnancy may also reduce the risk of hypertensive disorders of pregnancy (Hofmeyr et al. 2007) that may also arise secondary to lead exposure (Rothenberg et al. 2002; Sowers et al. 2002) (and thus conferring additional negative effects of lead for both mother and fetus and a potential benefit of calcium supplementation). The risks posed by calcium supplementation at levels approximating recommended daily intakes in this population are negligible. We therefore conclude that dietary supplementation of calcium intake should be considered as a cost-effective means for lowering transgenerational fetal lead exposure. This is particularly important in populations where dietary calcium intake is low. Because bone lead has a half-life of years to decades, women and their infants will continue to be at risk for exposure long after environmental sources of lead have been abated.

References


