Screening and Routine Supplementation for Iron Deficiency Anemia: A Systematic Review

Marian S. McDonagh, PharmD, Ian Blazina, MPH, Tracy Dana, MLS, Amy Cantor, MD, MPH, Christina Bougatsos, MPH

BACKGROUND AND OBJECTIVES: Supplementation and screening for iron-deficiency anemia (IDA) in young children may improve growth and development outcomes. The goal of this study was to review the evidence regarding the benefits and harms of screening and routine supplementation for IDA for the US Preventive Services Task Force.

METHODS: We searched Medline and Cochrane databases (1996–August 2014), as well as reference lists of relevant systematic reviews. We included trials and controlled observational studies regarding the effectiveness and harms of routine iron supplementation and screening in children ages 6 to 24 months conducted in developed countries. One author extracted data, which were checked for accuracy by a second author. Dual quality assessment was performed.

RESULTS: No studies of iron supplementation in young children reported on the diagnosis of neurodevelopmental delay. Five of 6 trials sparsely reporting various growth outcomes found no clear benefit of supplementation. After 3 to 12 months, Bayley Scales of Infant Development scores were not significantly different in 2 trials. Ten trials assessing iron supplementation in children reported inconsistent findings for hematologic measures. Evidence regarding the harms of supplementation was limited but did not indicate significant differences. No studies assessed the benefits or harms of screening or the association between improvement in impaired iron status and clinical outcomes. Studies may have been underpowered, and control factors varied and could have confounded results.

CONCLUSIONS: Although some evidence on supplementation for IDA in young children indicates improvements in hematologic values, evidence on clinical outcomes is lacking. No randomized controlled screening studies are available.
Iron-deficiency anemia (IDA), defined as iron deficiency (serum ferritin <12 µg/L) with hemoglobin levels <110 g/L,1,2 can present a significant burden of disease in infancy and childhood. Iron is required in the production of hemoglobin, an essential protein found in red blood cells, and is stored in the body for use in hemoglobin production. Iron deficiency occurs when the level of stored iron becomes depleted. IDA occurs when iron levels are sufficiently depleted to produce anemia, characterized by hypochromic microcytic red blood cells.3 Although infants in the United States with iron deficiency are usually asymptomatic, IDA has been associated in some observational studies with cognitive and behavioral delays in children. However, these studies had methodologic flaws4; for example, the outcomes examined were varied and not clearly clinically important. The effect of IDA in infancy and childhood has been reported in few well-designed, long-term controlled studies.

Iron deficiency among infants and toddlers in the United States has a prevalence of ∼8% in the general population5–7; however, only about one-third of children who are iron deficient have associated anemia.5,8,9 The prevalence of IDA in children between the ages of 1 and 5 years is estimated to be ∼1% to 2% in the United States.8,10 The prevalence in children from low-income families is estimated to be slightly higher (ie, ∼3% for boys and 4% for girls based on 1 study of 432 one- to three-year-old children residing in California).7 Current evidence regarding the prevalence of IDA in infants <1 year old in the United States is lacking, although estimates for low-risk infants in other developed countries range from 2% to 4%.11,12 Screening young children for IDA may lead to earlier identification and therefore earlier treatment, which has the potential to prevent negative health outcomes. However, the advent of iron fortification in the United States in many children’s food products may influence our current understanding of IDA.

In 2006, the US Preventive Services Task Force (USPSTF) concluded that the evidence was insufficient to recommend for or against routine screening for IDA or routine iron supplementation for asymptomatic children aged 6 to 12 months who are at average risk for IDA (I Recommendations).13 These recommendations were based on a lack of evidence that screening resulted in improved health outcomes, as well as poor and conflicting evidence regarding the benefit of iron supplementation in children who are not at increased risk of IDA. At that time, the USPSTF recommended routine iron supplementation for asymptomatic children aged 6 to 12 months who are at increased risk for IDA (B Recommendation), based on evidence that iron supplementation may improve neurodevelopmental outcomes in children who are at increased risk of IDA, which outweighs any potential harms.

The present review was commissioned by the USPSTF to update the previous recommendations.13 The scope of this review includes evidence regarding the benefits and harms of routine iron supplementation, screening for IDA in children ages 6 to 24 months, and the association between a change in iron status and improvement in child health outcomes in populations relevant to the United States.

**METHODS**

Detailed methods and data for this review (including search strategies, inclusion criteria, abstraction and quality rating tables, information on risk factors and risk assessment tools, and results related to biochemical and composite intermediate outcomes) are provided in the full report.14 The protocol was developed by using a standardized process15 with input from experts and the public. In consultation with the USPSTF, analytic frameworks and Key Questions were developed for routine supplementation (Supplemental Appendix Figure 2) and for screening for IDA (Supplemental Appendix Figure 3) to show the linkages between Key Questions and bodies of evidence.

A research librarian searched the Cochrane Central Register of Controlled Trials, the Cochrane Database of Systematic Reviews (through the second quarter, 2014), and Medline (1996–August 2014) for relevant studies to update the previous USPSTF reviews.16,17 Because the previous research focused on systematic reviews and key studies of treatments for IDA, we also searched the reference lists of systematic reviews18–20 to identify any additional, relevant studies published before 1996.

Studies were selected on the basis of inclusion and exclusion criteria developed for each Key Question. Articles were selected for full review if they were related to IDA in children who received an intervention (supplementation or screening and related treatment) between the ages of 6 and 24 months. We restricted inclusion to English-language articles and excluded studies published only as abstracts. For all Key Questions, the focus was on studies that involved iron supplementation and treatment regimens commonly used in clinical practice in the United States. We excluded studies conducted in resource-poor populations, including nutritionally deficient populations in developing countries and populations in areas expected to have a high prevalence of hemoparasites, by selecting studies conducted in countries listed as having "high" or “very high” human development based on the international United Nations Human Development Index.21 At least 2 reviewers independently evaluated each study to determine eligibility.
Clinical outcomes of study were morbidity (including growth; cognitive, psychomotor; and neurodevelopmental outcomes; and diagnosis of developmental delay), mortality, and quality of life. Harm outcomes included accidental overdose, study discontinuations, and other harms related to screening, supplementation, or treatment. Included intermediate outcomes were incidence of IDA, iron deficiency, and anemia, as well as hematologic indices such as ferritin levels. Randomized controlled trials, nonrandomized controlled clinical trials, and controlled cohort studies were included for all Key Questions.

Details about the study design, patient population, setting, screening method, interventions, analysis, follow-up, and results were abstracted. A second investigator reviewed the data abstraction for accuracy. Two investigators independently applied criteria developed by the USPSTF15 to rate the internal validity (quality) of each study as good, fair, or poor. Discrepancies were resolved through a consensus process. When otherwise not reported and where possible, relative risks (RRs) and 95% confidence intervals (CIs) or P values were calculated.

The aggregate quality of the body of evidence for each Key Question (ie, good, fair, poor) was assessed by using methods developed by the USPSTF; these assessments were based on the number, quality and size of studies; consistency of results between studies; and directness of evidence.15 Meta-analysis was not attempted due to the limited number of studies for each Key Question and differences among studies in design, population, and outcomes.

RESULTS

Figure 1 shows the results of the literature search and selection process.

Routine Iron Supplementation

Benefits of Routine Iron Supplementation in Children Ages 6 to 24 Months

A 1996 review conducted for the USPSTF17 found adequate evidence that iron prophylaxis resulted in reductions in the incidence of iron deficiency and IDA, but few data focusing on clinical outcomes were reported. The 2006 update16 did not assess the effect of supplementation on intermediate outcomes, and it found mixed evidence regarding the benefit of iron supplementation on neurodevelopmental test scores.

Overall, 10 trials of iron supplementation were included in this update.22–34 One study was rated as good quality,29 7 as fair quality,22–28,32 and 2 as poor quality.30,31 In general, children were enrolled between 6 and 9 months of age. Iron supplementation was administered for durations ranging from 3 to 18 months.

Supplementation was provided as oral iron drops, iron-fortified formula, and as iron-fortified milk, foods, or meat. Controls used in the studies varied, and included a non-iron-containing formula or supplement, a specific diet, cow’s milk, or nothing. Race or ethnicity was poorly reported. Enrolled sample sizes ranged from 24 to 493, except for 1 larger study of 1798 children; many studies analyzed fewer numbers due to loss to follow-up or refusal to undergo venipuncture. Only 1 study analyzed children on an intention-to-treat basis;29 the proportion of the sample available for analysis at the end of the other studies ranged from 53% to 92%. Most studies excluded children born prematurely and those with conditions likely to affect iron absorption, growth, or development, thus ruling out some children at higher than average risk for IDA but not specifically targeting those at average risk.

Methodologic shortcomings included unclear methods of randomization and allocation concealment,25–27,30–32 lack of or unclear methods of blinding,22,23,25,27,30,31 and high or differential loss to follow-up.24,27,28,30,31 In addition, studies may be underpowered; although 7 studies reported some power or sample size calculations, they were limited to certain outcomes and varying differences in effect sizes.24,26–29,31,32 For example, 2 studies were reported to be powered for developmental outcomes, with 1 sufficient to detect a 5-point difference on the Bayley Scales of Infant Development28 and 1 sufficient to detect a 2.5-point difference in “developmental scores” (scale not mentioned).26 We did not pool the results because of the heterogeneity of the studies in terms of supplementation method, dose, duration, timing of initiation and follow-up, and methodologic limitations. In addition, risk factors were largely not reported, and no studies stratified results according to risk groups.

Six fair-quality placebo-controlled trials of routine iron supplementation in young children sparsely reported various growth outcomes; 5 trials found no clear effect of supplementation on weight, length, or head circumference after 3 to 12 months of follow-up (Table 1).22,25,26,28–30 As noted earlier, studies may have been underpowered to detect growth outcomes. Most sample sizes varied from 70 to 428, with 1 study including 1657 children. The only study reporting statistically significant differences in growth parameters found lesser growth values in the iron-supplemented group, possibly due to baseline differences in these growth outcomes.26 The group that received iron began the study with lesser values (weight: 7.98 vs 8.09 kg; length: 66.6 vs 66.9 cm [both P < .01]). Although this study was the largest, it was conducted in Chile, had a high incidence of IDA in the control group (22.6%), and suffered from methodologic flaws. Children were...
initially randomized to receive low- or high-iron supplementation, but the low-iron intervention was replaced with a no-iron intervention partially through the study, partly because the interim analysis suggested that the low-iron condition was sufficient to prevent IDA. For analysis, all children who received any iron supplementation were combined and compared with children who did not receive supplementation, breaking randomization and leading to baseline differences between the groups. Because randomization was broken, we viewed the results as a fair-quality, comparative observational study. The authors found the following significant results for the iron-supplemented versus no-iron groups, respectively, after 12 months: weight, 10.0 vs 10.1 kg ($P < .05$); length, 74.7 vs 75.1 cm ($P < .001$); length for age ($z$ score), $-0.27$ vs $-0.15$ ($P < .01$); and head circumference, 46.7 vs 47.0 cm ($P < .001$); the changes were significant possibly because the iron-supplemented group had lower values at baseline. Other clinical outcomes, such as diagnosis of psychomotor or neurodevelopmental delay or quality of life, were not reported in any trial.

Although not clearly clinical outcomes, developmental test scores after follow-up periods of 3 to 12 months were reported in 3 fair-quality trials. Two trials ($N = 428$ and 1657, respectively), including the Chilean study with methodologic flaws mentioned earlier, found no statistically significant difference between groups on the Bayley Scales of Infant Development (Table 2). Differences between groups were small and ranged from 0.6 to 0.7 on mental development and 0.2 to 0.7 for psychomotor development. One trial of children potentially at higher risk for IDA used the Griffiths scale to measure psychomotor development. Although scores in both groups declined and were within normal limits at 24 months, they declined less in the iron-supplemented group (general quotient score at 24 months: $-9.3$ vs $-14.7$; $P = .04$).

Ten trials of iron supplementation in children reported inconsistent
<table>
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<tr>
<th>Study Year, Country, N</th>
<th>Risk Factors Reported</th>
<th>Interventions and Comparator</th>
<th>Outcomes: Supplementation Versus Control Groups</th>
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<td>IDA (Hb &lt; 110 g/L and Iron Deficiency)</td>
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<tr>
<td>Domellöf et al. 2001, Sweden, N = 70, 3 mo; fair</td>
<td>Race: Not reported; Preterm and low birth weight infants excluded</td>
<td>A. Placebo from 4 to 6 mo and iron supplement from 6 to 9 mo of age (n = 34) B. Placebo from 4 to 9 mo (n = 36)</td>
<td>No effect (numbers not reported)</td>
</tr>
<tr>
<td>Getman et al. 2004, United States, N = 294, 3 mo; fair</td>
<td>Race: 55% vs 48% black; Preterm and low birth weight infants excluded</td>
<td>A. Oral multivitamin drops, 10 mg of iron/d B. Multivitamin drops without iron</td>
<td>8% (11/138) vs 8% (11/144) anemic and had 2 other abnormal hematologic values; RR: 1.04 (95% CI: 0.47–2.33)</td>
</tr>
<tr>
<td>Gill et al. 1997, United Kingdom and Ireland, N = 302, 11 mo; fair</td>
<td>Race: not reported; Preterm and low birth weight infants excluded</td>
<td>A. Iron-fortified formula B. Noniron-fortified formula C. Cow’s milk</td>
<td>3.1% (54/1114) vs 22.9% (156/681); RR: 0.14 (95% CI: 0.09–0.20)</td>
</tr>
<tr>
<td>Lucoff et al. 2003, Chile, N = 1557, 6–12 mo; fair</td>
<td>Race: not reported; Preterm and low birth weight infants excluded</td>
<td>A. Multiple interventions with varying iron concentrations B. No iron supplementation</td>
<td>3.1% (54/1114) vs 22.9% (156/681); RR: 0.14 (95% CI: 0.09–0.20)</td>
</tr>
<tr>
<td>Makrides et al. 1998, Australia, N = 92, 6 mo; fair</td>
<td>Race: not reported; Preterm and low birth weight infants excluded</td>
<td>A. High-iron weaning diet B. Control weaning diet</td>
<td>0 vs 0</td>
</tr>
<tr>
<td>Morley et al. 1999, United Kingdom, N = 428, 9 mo; fair</td>
<td>Race: not reported; Preterm and low birth weight infants excluded</td>
<td>A. Iron-fortified formula, 1.2 mg iron/L B. Unfortified formula, 0.9 mg iron/L C. Cow’s milk</td>
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findings for incidence of IDA, iron deficiency, and anemia, as well as changes in hemoglobin and serum ferritin (Table 1 [poor-quality studies were omitted from tables]).22,24–32 Iron supplementation was not found to influence the incidence of IDA in 4 good- or fair-quality studies (N values ranged from 62 to 284).24,26,27,29,32 One study, the aforementioned larger Chilean study (N = 1657) with a high incidence of IDA in the control group (22.6%), reported a significant benefit (RR: 0.14 [95% CI: 0.09–0.20]).26 Overall, the studies reported ranges of IDA from 0% to 8% for those in the supplementation group and 0% to 22.6% in the placebo group.

For incidence of iron deficiency, 2 fair-quality studies with high incidences in the control groups suggest a significant benefit from supplementation. These findings included 1 study (N = 302) that compared iron-fortified formula with non-iron-fortified formula and cow’s milk (6% vs 22% vs 43%)24 and the large (N = 1657) Chilean study (26.5% vs 51.3%; RR: 0.52 [95% CI: 0.45–0.59]).26 Three other studies found no difference in rates of iron deficiency,24,27,30 including the only study conducted in the United States (RR: 0.94 [95% CI: 0.74–1.20]).24 Overall, the studies reported rates of iron deficiency ranging from 3.9% to 78% for those in the supplementation group and from 7.7% to 84% in the placebo group.

Six trials (3 fair-quality and 1 poor-quality) reported ranges of IDA from 0% to 8% for those in the supplementation group and 0% to 22.6% in the placebo group.

TABLE 1

<table>
<thead>
<tr>
<th>Study, Year, Country, N</th>
<th>Duration, Quality</th>
<th>Risk Factors Reported</th>
<th>Interventions and Comparator</th>
<th>Outcomes: Supplementation Versus Control Groups</th>
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<tr>
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<td></td>
<td>IDA (Hb &lt; 110 g/L and Iron Deficiency)</td>
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<tr>
<td>Szmyt et al, 2009, N = 225</td>
<td>5 mo, good</td>
<td>Race: 84% vs 78% white</td>
<td>No difference between groups (data only reported in a figure)</td>
<td>118.8 vs 121.5</td>
</tr>
<tr>
<td>Williams et al, 1999, United Kingdom, N = 92, 10–12 mo; fair</td>
<td>Race: 74% white, 24% black, 2% Asian; Receiving income support: 59%; Preterm and low birth weight infants excluded</td>
<td>—</td>
<td>At 18 mo of age: 2% (1/46) vs 33% (15/46); RR: 0.07 (95% CI: 0.01 to 0.48)</td>
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Two trials were omitted from the table due to poor-quality rating (Yalçin et al, 2000 and Yeung and Zlotkin, 2000). Hb, hemoglobin; NS, not significant.

Iron deficiency was defined variably as serum ferritin <10 μg/L,25, 27, 30 <15 μg/L,24 or <12 μg/L in addition to 1 of 2 other hematologic indicators.26

RR not calculable based on data reported.

Groups differed significantly at baseline.
reported significant benefits, with RRs ranging from 0.07 (95% CI: 0.01–0.48)\(^{26}\) to 0.14 (95% CI: 0.09–0.20).\(^{23}\) However, variability in the definitions of anemia, the unknown mix of baseline risk of children enrolled, and the variation in control group rates across these studies (from 13% to 33%) limit the interpretability of the findings for US populations. Overall, the studies reported rates of anemia ranging from 0% to 22% for those in the supplementation group and from 13% to 33% in the placebo group.

In addition, hemoglobin results were reported in 8 studies,\(^{24,30,32}\) with small differences between groups; 3 were significant.\(^{25,26,28}\) Nine studies reported ferritin concentrations, with conflicting results.\(^{22–30,32}\)

**Harms of Routine Iron Supplementation in Children Ages 6 to 24 Months**

None of the studies of iron supplementation reported serious harms, including accidental overdose or withdrawals due to adverse events. Five studies reported on adherence to the assigned regimen and found no impact based on iron content. In some cases, however, the control group was preferred (e.g., cow’s milk over fortified or unfortified formula).\(^{24,28–31}\)

In 1 fair-quality trial, no clinically significant adverse events thought to be related to study interventions were reported.\(^{25}\) No differences in rate of gastrointestinal adverse events in toddlers consuming iron-fortified milk and those consuming unfortified milk were found in a good-quality trial (2% vs 2%; RR: 1.0 [95% CI: 0.9–11]).\(^{29}\) No other studies reported the incidence of gastrointestinal adverse events.

**Screening for IDA**

**Benefits and Harms of Screening Asymptomatic Children Ages 6 to 24 Months for IDA**

As in previous reviews,\(^{16,17}\) no studies evaluating the benefits or harms of screening programs for asymptomatic children ages 6 to 24 months for IDA were found.

**Benefits and Harms of Treatment of IDA in Children Ages 6 to 24 Months**

No new studies of oral iron treatment of IDA in infants and children 6 to 24 months of age were found. Of the studies included in the previous reviews, only 1 study (\(N = 110\)) met our current criteria.\(^{35}\) This study was rated as poor quality due to baseline differences in age and unclear reporting of methods. Improved growth velocity and hemoglobin and ferritin levels were found, but no differences were reported in Denver Developmental Screening Test psychomotor development outcomes compared with control subjects.

No newly published studies reporting harms of iron treatment in children ages 6 to 24 months were found. One older randomized controlled trial (\(N = 334\)), published in 1991\(^{36}\) and not included in the previous reviews, reported no differences between children receiving iron treatment and those receiving placebo in overall incidence or incidence of specific adverse events, including gastrointestinal events.

**Association Between a Change in Iron Status and Improvement in Child Health Outcomes in Populations Relevant to the United States**

No studies met the established criteria to evaluate an association between improvement in impaired iron status and child health outcomes in populations relevant to the United States.

One poor-quality and 2 fair-quality supplementation studies provided evidence regarding changes in iron status and measures of growth or development scale scores in children with normal iron status at baseline.\(^{22,25,30}\) Two fair-quality studies (Table 3) found statistically significant changes in iron status measures but no differences between groups in measures of weight or height after 9 and 18 months.\(^{22,25}\) A poor-quality trial found large numerical (but nonsignificant) differences in hemoglobin and serum ferritin levels and a significant difference in transferrin saturation, but no differences in weight, height, or head circumference outcomes or in neurodevelopmental outcomes based on the Bayley Scales of Infant Development.\(^{30}\)

**DISCUSSION**

As in the previous USPSTF reviews,\(^{16,17}\) we found no evidence regarding the effects of routine iron supplementation in young children on
diagnosis of psychomotor or neurodevelopmental delay or quality of life, and the evidence regarding developmental test scores after 3- to 12-month follow-up periods (although not clearly clinical outcomes) does not indicate important differences.\textsuperscript{23,26,28,30}

We found no evidence of important or clear benefit in growth outcomes, which is consistent with the findings of a recent meta-analysis of 21 randomized controlled trials that included studies from any country.\textsuperscript{18}

Although the study findings are not consistent, the evidence from 10 trials of iron supplementation in children indicates no benefit in terms of incidence of IDA, anemia, or hemoglobin; the findings were inconsistent regarding incidence of iron deficiency and serum ferritin concentrations.\textsuperscript{22–32}

Some of the variation in findings may have been due to inadequate sample sizes for specific outcomes. This situation is often ideal for pooling studies to gain statistical power; in this case, however, we found both clinical and methodologic heterogeneity and did not combine the studies. For example, there was important variability in the definitions of IDA, anemia, and iron deficiency (mostly unknown baseline risk of children enrolled) and wide variation in control group rates across these studies. Although the prevalence rates of iron deficiency in the United States are currently estimated at 8%\textsuperscript{5–7}, control group rates in these studies ranged from 13% to 33%, such that generalizability of these findings to the US population is unclear. Studies that did find a benefit generally had higher rates of iron deficiency in the control groups compared with studies which found no benefit, suggesting that baseline risk is important in determining who will benefit from supplementation, in terms of preventing iron deficiency. An additional factor potentially contributing to variability was the use of cow’s milk as a control in several studies; use of cow’s milk is considered a risk factor for IDA and is advised against before 12 months of age by the American Academy of Pediatrics, and the Centers for Disease Control and Prevention.\textsuperscript{2,37,38}

In addition, 1 of the largest studies included in the review that consistently found statistically significant results supporting supplementation for most hematologic values was conducted in Chile and had a high incidence of hematologic values in the control groups. This study, which initially randomized children to receive low- or high-iron supplementation, broke randomization, leading to baseline differences between the groups.\textsuperscript{26}

Harms of routine iron supplementation in children were rarely reported, and supplementation did not result in higher rates in studies reporting harms. Although an older meta-analysis of 28 studies (randomized controlled trials and cohort studies) found a slightly increased risk of diarrhea with iron supplementation (RR: 1.1 [95% CI: 1.0–1.2]),\textsuperscript{39} the majority of studies were conducted in developing countries, and the age of the populations ranged from 2 days to 14 years.
As in the previous reports, evidence regarding the benefits and harms of screening for IDA in children ages 6 to 24 months is absent. Similarly, we found only very limited evidence regarding the benefits and harms of IDA treatment that is generalizable to children ages 6 to 24 months in the United States. Based on this evidence, benefits were shown only for some iron status measures in the short term. Previous USPSTF reports concluded that there was no evidence regarding the relative harms of treatment. This update identified only 1 additional study, which indicated no differences between children receiving iron supplementation and placebo in the incidence of overall or specific adverse events, including gastrointestinal events.

The potential for long-term benefit of preventing IDA in young children presumes that improvement in iron status is associated with good long-term clinical outcomes, such as normal growth and neurodevelopment. Evidence capable of showing this specific association was extremely limited and did not support a clear association between change in iron status and differences in growth or neurodevelopment. Limitations of our report include restricting inclusion of studies published in English and studies conducted in developed countries or studies in developing countries where the population enrolled was similar to the population of the United States, particularly in terms of rates of malnutrition, hemoparasite burden, and general socioeconomic status. A number of studies of iron supplementation and treatment that were conducted in developing countries were excluded. Malnourishment, very low socioeconomic status, and/or presence of parasitic endemic diseases were common in the included populations in these studies. Also excluded were studies of iron supplementation that enrolled children aged <6 months; this population was outside the scope of the review. Good-quality, randomized controlled trials of routine supplementation, screening programs, and treatment of IDA in children 6 to 24 months of age, with adequate sample sizes for key iron status and clinical health outcomes, are needed. Such trials should clearly report prognostic baseline characteristics of enrolled children, details of interventions, longer term benefits (particularly developmental outcomes using appropriate neurodevelopmental tests), and harms, and the studies should use appropriate controls (ie, not cow’s milk). In addition, these studies should report neurodevelopmental diagnoses rather than test scores.

CONCLUSIONS

Expanded and better research is needed to assess the benefits and harms of routine iron supplementation and screening to prevent IDA in young children in developed countries. At present, the limited evidence indicates no benefits in growth and neurodevelopmental test scores with supplementation, and hematologic outcomes are variably affected. The benefits and harms of treatment are largely unclear, as is the association between improvement in IDA or iron deficiency and clinical outcomes.

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