ABSTRACT: Background: Iron deficiency anemia (IDA) has been implicated in the etiology of transient ischemic attack and ischemic stroke. This study aimed to: 1) document IDA prevalence in patients ≥65 years of age admitted to hospital with transient ischemic attack or first ischemic stroke, and 2) investigate dietary intake as a predictor of iron status. Methods: Ninety-four patients were enrolled. An algorithm containing values for hemoglobin, ferritin, total iron binding capacity, transferrin saturation, and serum transferrin receptor measured at admission was used to identify IDA. Usual dietary intake was assessed with the Clue II food frequency questionnaire. Results: Prevalence estimates were 6.4% for IDA, 2.1% for iron deficiency without anemia, and 6.4% for anemia from other causes. IDA prevalence was significantly higher than published National Health and Nutrition Examination Survey III (NHANES III) estimates for gender-specific age groups ≥ 70 years (One-Sample Proportion Test; males p = 0.038 [n= 37]; females p = 0.002 [n=44]). A comparison of IDA prevalence against selected controls from the NHANES III database yielded an odds ratio (OR) of 6.3, 95% confidence interval (CI) 0.8 to 53.7, which was not statistically significant (Fisher’s Exact Test; n=94; p = 0.118). Multivariate linear regression analysis of dietary intake with indicators of iron status (n=58) revealed only iron supplements (p=0.013) and heme iron intake (p = 0.038) as negative predictors of total iron binding capacity (p<0.05). Conclusions: These findings support the initiation of a prospective case control study to investigate IDA as a risk factor for ischemic stroke in elderly patients.

Iron deficiency anemia (IDA) has been proposed as an etiological factor for ischemic stroke. There is considerable evidence for this for pediatric transient ischemic attack (TIA) and ischemic stroke1-7, and IDA has emerged in a recent case control study as a significant risk factor for stroke in otherwise healthy young children7. In adult ischemic stroke, however, it is less well-established that IDA confers risk. This hypothesis was first supported by a series of case reports linking IDA to thrombocytosis and stroke, implicating reactive thrombocytosis as a contributing mechanism8 10. A follow-up study to the First National Health and Nutrition Examination Survey (NHANES I) identified a significant U-shaped relationship between transferrin saturation and stroke in white women 45-74 years of age11, suggesting an increased risk for stroke at both low and high levels of circulating iron. Lower serum iron concentration, an indicator of reduced iron transport and negative iron balance, has been reported as a significant predictor of stroke in elderly individuals.12-16
The diagnosis of IDA in elderly stroke patients is difficult because it requires differentiating between IDA and the anemia of chronic disease (ACD). Bone marrow aspirate, the gold standard for determining iron status, is invasive and often unreliable when performed in a routine clinical laboratory. Serum ferritin, although commonly used to assess iron stores, is unreliable when performed in a routine clinical laboratory. Serum transferrin receptor (sTfR) concentration rises with chronic infection, inflammation, liver disease, and malignancy. Serum transferrin receptor values continue to rise as iron deficiency progresses to the anemia stage (reviewed in) and can both distinguish between IDA and ACD and identify iron deficiency coexisting with ACD. Values for sTfR remain normal in inflammatory conditions and ACD, offering a particular advantage for assessing iron (Fe) status during the inflammatory response to stroke.

The role of diet as an independent contributor to iron status in the elderly is also uncertain. Nutritional status is often compromised by multiple mechanisms that impact on dietary intake, such as impaired functional and cognitive status, multiple medical conditions and medications, poor oral health, poverty, changes in taste and smell, and dysphagia. Gastrointestinal blood loss, non-steroidal anti-inflammatory drugs, anti-coagulants, genitourinary disease, or frequent blood drawings can also contribute to IDA in aged individuals. Positive associations have been reported between iron status and dietary factors such as iron supplements, heme iron (found in meat, poultry and fish), vitamin C, protein, and alcohol. However, the strength of these associations is limited by varying accuracy among the different indicators of iron status that have been employed. The relationship between dietary intake and iron status has not been studied in stroke patients.

Despite the intriguing evidence that IDA may increase risk for ischemic stroke and TIA, there are no reported prevalence estimates for IDA in elderly stroke patients. Thus, our objective was to employ a rigorous algorithm that included measurement of sTfR to determine the prevalence of IDA in a series of patients 65 years-of-age or older admitted to hospital with TIA or first presentation of ischemic stroke. Prevalence data for IDA were compared with existing published figures for free-living elderly. We also investigated whether previous dietary intake was a predictor of iron status for patients newly admitted with stroke or TIA.

**METHODS**

**Subjects**

Data were obtained prospectively from elderly patients ≥ 65 years of age (n=94) admitted sequentially to the Neurology service at Royal University Hospital, Saskatoon, Saskatchewan between May 23 and December 23, 1999. These patients were diagnosed with first ischemic stroke or TIA clinically by a neurologist. Exclusion criteria were intracerebral or subarachnoid hemorrhage, stroke due to a brain tumour, or previous known history of stroke. Consent was obtained from the patient or next-of-kin. Ethics approval was obtained from the University of Saskatchewan Advisory Committee on Ethics in Human Experimentation.

Medical charts were reviewed in order to document major risk factors for stroke (age, gender, race, hypertension, smoking status, carotid stenosis, atrial fibrillation, previous TIA, diabetes mellitus, family history, dyslipidemia, and cardiac disease) and potential medical causes of IDA (gastrointestinal blood loss, non-steroidal anti-inflammatory drugs or anticoagulant use, and genitourinary disease).

**Laboratory analysis**

Blood samples were obtained within 24 hours of admission. Serum samples were stored at -70°C until analysis for sTfR using an enzyme immunoassay kit and manufacturer reference range (Ramco Laboratories, Houston, Texas). The remaining laboratory tests (serum ferritin, serum iron, transferrin saturation, total iron binding capacity (TIBC), white blood cell count, red blood cell count, hemoglobin, hematocrit, mean corpuscular volume, mean corpuscular hemoglobin, mean corpuscular hemoglobin concentration, red blood cell distribution width, platelet count, and mean platelet volume) were conducted using the methodology and reference ranges of the Department of Laboratory Medicine, Saskatoon Health Region.

**Classification of iron status**

Based on the algorithm shown in the Figure, patients were classified into the following groups: a) IDA, b) occult iron deficiency, c) ACD, d) IDA coexisting with ACD, and e) other anemia.

**Comparison of IDA prevalence with published estimates**

We undertook a preliminary comparison of our IDA prevalence figures against a published population estimate. Because of the lack of Canadian data for comparison, the IDA prevalence in our study group was compared with results from the NHANES III Study, which was selected as the largest comprehensive population-based study of Fe status relevant to our data collection timeline and subject age. This was a six year study (1988-1994) conducted by the National Center for Health Statistics, Centers for Disease Control and Prevention to assess the health and nutritional status of the civilian non-institutionalized population of the United States. In the Third National Health and Nutrition Examination Survey (NHANES III), IDA was identified by two or more abnormal values for free erythrocyte protoporphyrin, transferrin saturation, or ferritin.
Two methods were used for comparison: 1) First, our prevalence estimates for males and females were compared with the published prevalence figures from NHANES III for free-living elderly subjects (n=3,067)\textsuperscript{19}. As the latter data were only specified for the age groups of 50-69 years and ≥ 70 years, IDA prevalence was calculated for those subjects from our study who were ≥ 70 years (n=81). 2) Secondly, we made a small refinement in the comparison by using a limited case control approach to compare our IDA prevalence figure with that from 94 age- and gender-matched controls (with black and Hispanic individuals excluded) from the NHANES III database\textsuperscript{38} who resided in areas in the Midwestern U.S.A. with populations <1,000,000 and did not have a previous history of stroke. Although it was not possible to conduct a comprehensive case control study that matched subjects for stroke risk factors, this secondary analysis did enable selection of NHANES III individuals who were better controls with respect to age, gender, ethnicity, geographical location, and environment (rural or smaller city size and altitude).

Dietary intake

Usual dietary intake was assessed by interviewing the patient and/or surrogate source during the hospital stay in order to complete a CLUE II Health Habits and History (modified Block) Questionnaire, a 63-item food frequency questionnaire. This questionnaire records usual diet over the past year and has been directly validated against antioxidant status\textsuperscript{39} and indirectly validated for iron\textsuperscript{40}. The questionnaires were produced, scanned, and analyzed by Block Dietary Data Systems (Berkeley, California).

Statistical analysis

Statistics were performed using SPSS Statistics 17.0 software, with all results considered significant at p<0.05.

For the first comparison of IDA prevalence estimates, we utilized the One-Sample Proportion Test with continuity correction\textsuperscript{41}. For the second comparison of IDA prevalence between the two groups, an odds ratio and 95% confidence interval were calculated, and Fisher’s Exact Test was used to test the hypothesis that the odds ratio=1. Fisher’s Exact Test was chosen since two cells had an expected count less than \textsuperscript{541}. Similarly, Fisher’s Exact Test was chosen to determine whether other potential causes of IDA, that is, non-steroidal anti-inflammatory drugs, anticoagulants, gastrointestinal disease, or genitourinary disease, were associated with the presence of IDA. Chi-square analysis was used to test whether there were differences in the proportion of cases diagnosed with stroke versus TIA by gender.

Multivariate analysis was used to examine associations between dietary variables that would be expected to influence outcome measures of iron status. Initial bivariate analysis was used to examine the association between single predictor variables (age, body mass index, gender, kilocalories, protein, dietary iron, supplemental iron, total iron, dietary vitamin C, supplemental vitamin C, total vitamin C, fiber, heme iron, and alcohol), and outcome measures of iron status (sTfR, total iron binding capacity, hemoglobin, and presence or absence of IDA). Those variables found to be significant predictors at p<0.20 were considered in subsequent model building. Forward stepwise selection procedure was used and stepwise criteria for entry were set as follows: probability of F to enter ≤ 0.20, probability of F to remove ≥ 0.30.

RESULTS

Sample demographics

The overall consent rate was 92%. Demographic characteristics are shown in Table 1 and demonstrate an equal number of females and males with 62.8% of the sample ≥ 75 years of age. Eighty-four percent (n=79) of subjects presented with stroke, and 16% presented with TIA (n=15). These diagnoses were not significantly different by gender (X\textsuperscript{2}=0.079, p > 0.05). The presence of risk factors for stroke as identified by chart review is also shown in Table 1.

Prevalence of IDA

The number of cases identified for IDA, iron deficiency without anemia, IDA co-existing with ACD, ACD, and other anemias is found in the Figure. The overall prevalence of IDA in our sample of stroke and TIA patients was 6.4%, accounting for 40% of all cases of anemia identified. Only one of these six individuals had IDA co-existing with ACD. Earlier stage iron deficiency without anemia was present in 2.1% of the sample. Anemia from other causes was identified in 6.4% of the patients. Table 2 contains descriptive statistics for key hematological data.
Table 1: Demographic and risk factor characteristics of subjects at study enrolment

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Frequency</th>
<th>Total</th>
<th>Percent</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Age (years)</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>65-74</td>
<td>35</td>
<td>94</td>
<td>37.2</td>
</tr>
<tr>
<td>75+</td>
<td>59</td>
<td>94</td>
<td>62.8</td>
</tr>
<tr>
<td><strong>Gender</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>47</td>
<td>94</td>
<td>50.0</td>
</tr>
<tr>
<td>Female</td>
<td>47</td>
<td>94</td>
<td>50.0</td>
</tr>
<tr>
<td><strong>Other risk factors</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hypertension</td>
<td>61</td>
<td>94</td>
<td>64.9</td>
</tr>
<tr>
<td>Smoking</td>
<td>18</td>
<td>94</td>
<td>19.1</td>
</tr>
<tr>
<td>Carotid Stenosis</td>
<td>26</td>
<td>71</td>
<td>36.6</td>
</tr>
<tr>
<td>Atrial Fibrillation</td>
<td>18</td>
<td>94</td>
<td>19.1</td>
</tr>
<tr>
<td>Previous TIA</td>
<td>20</td>
<td>94</td>
<td>21.3</td>
</tr>
<tr>
<td>Diabetes Mellitus</td>
<td>28</td>
<td>94</td>
<td>29.8</td>
</tr>
<tr>
<td>Family History of Stroke</td>
<td>4</td>
<td>94</td>
<td>4.3</td>
</tr>
<tr>
<td>Elevated Cholesterol</td>
<td>33</td>
<td>60</td>
<td>55.0</td>
</tr>
<tr>
<td>Cardiac Disease</td>
<td>48</td>
<td>94</td>
<td>51.1</td>
</tr>
</tbody>
</table>

1For whom data was available by chart review. 2History of hypertension, hypertension medication use or diagnosis at admission. 3Serum total cholesterol >5.2 mmol/L, history of dyslipidemia or lipid lowering medication. 4History of coronary artery disease, angina, myocardial infarction, cardiomyopathy, congestive heart failure, mitral stenosis/calci-fication, or left ventricular hypertrophy.

Table 2: Summary of hematological parameters used to identify IDA

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Gender</th>
<th>n</th>
<th>Mean (SD)</th>
<th>Median</th>
</tr>
</thead>
<tbody>
<tr>
<td>Serum Transferrin Receptor (µg/ml)^1</td>
<td>F</td>
<td>45</td>
<td>4.8 (2.6)</td>
<td>4.0</td>
</tr>
<tr>
<td></td>
<td>M</td>
<td>46</td>
<td>4.9 (3.6)</td>
<td>3.9</td>
</tr>
<tr>
<td>Serum Ferritin (µg/L)^2</td>
<td>F</td>
<td>47</td>
<td>90 (71)</td>
<td>83</td>
</tr>
<tr>
<td></td>
<td>M</td>
<td>47</td>
<td>210 (402)</td>
<td>97</td>
</tr>
<tr>
<td>Total Iron Binding Capacity (µmol/L)^3</td>
<td>F</td>
<td>47</td>
<td>56 (11)</td>
<td>57</td>
</tr>
<tr>
<td></td>
<td>M</td>
<td>47</td>
<td>50 (11)</td>
<td>52</td>
</tr>
<tr>
<td>Transferrin Saturation (%)^4</td>
<td>F</td>
<td>47</td>
<td>21 (12)</td>
<td>19</td>
</tr>
<tr>
<td></td>
<td>M</td>
<td>47</td>
<td>25 (13)</td>
<td>23</td>
</tr>
<tr>
<td>Hemoglobin (g/L)^5</td>
<td>F</td>
<td>47</td>
<td>132 (15)</td>
<td>133</td>
</tr>
<tr>
<td></td>
<td>M</td>
<td>47</td>
<td>147 (14)</td>
<td>147</td>
</tr>
</tbody>
</table>

Reference Ranges: ^12.9-8.3 µg/ml; ^220-120 µg/L (female), 20-400 µg/L (male); ^334-51 µmol/L (female), 41-63 µmol/L (male); ^414-51% (female), 25-56% (male); ^5110-160 g/L (female), 135-180 g/L (male).
Table 3 shows comparison of IDA prevalence in the stroke and TIA patients (n=81) with NHANES III prevalence figures for gender-specific age groups ≥ 70 years. The prevalence of IDA in our study was significantly higher than that found in NHANES III for both men and women (males p = 0.038; females p = 0.002). The second limited case-control comparison (n=94) yielded a similar difference in IDA prevalence between our subjects and the reference group to that obtained with the first analysis (Table 4). This secondary analysis yielded an odds ratio of 6.3, which was not statistically significant (p = 0.118).

Predictors of IDA

Nonsteroidal anti-inflammatory drug use (p=0.414), anticoagulant use (p=0.681), gastrointestinal disease (p=1.00), and genitourinary disease (p=0.501) were not significantly associated with the presence of IDA.

Dietary information was available for only 58 patients (62% of the sample). Results from multivariate linear regression analysis of dietary intake with indicators of iron status revealed few significant associations. Gender emerged as a highly significant predictor of hemoglobin (p<0.01), and supplemental iron (p = 0.013) and heme iron intake (p = 0.038) were negative predictors of total iron binding capacity (p<0.05).

DISCUSSION

This prospective study is the first to document the prevalence of IDA in elderly patients at hospital admission for TIA or first stroke. Due to the lack of consensus criteria for investigating IDA in the elderly, we developed a powerful classification scheme that included the specificity of sTfR values for identifying tissue iron deficiency while limiting the influence of any single variable. We describe a prevalence of IDA of 6.4%, with IDA accounting for 40% of all cases of anemia. Our patient sample appears to be relatively representative of the larger population, given the high consent rate (92%) and the similarity of our subjects’ baseline characteristics to those of the Registry of the Canadian Stroke Network42. Specifically, profiles were similar for the risk factors of hypertension, smoking and male gender, whereas the prevalence of diabetes and dyslipidemia was higher in our study, possibly due to the older subject age. A smaller proportion of our cases (15%) presented with TIA compared to 33% for the Stroke Registry, and it is likely that TIA was underrepresented in our study because such patients are not always admitted to hospital.

Our preliminary analyses suggest that IDA may be more prevalent in elderly patients newly admitted for TIA or first stroke than in the general free-living elderly population. The first approach, a gender-specific comparison of our prevalence figures with those published for free-living subjects 70 years of age and older from the NHANES III Study19, demonstrated a higher prevalence of IDA in the stroke and TIA patients for both genders. This comparison has limitations since an unknown number of the elderly reference group would have had a previous stroke. The NHANES III sample also included individuals from across the United States, so factors such as altitude, ethnicity, and lifestyle, including diet, may have differed from our Canadian prairie province-based sample. The second comparison, in which we selected more similar control subjects from this database38, yielded a similar absolute difference in IDA prevalence between our sample and the reference group. An odds ratio calculation revealed that our first-time stroke or TIA patients were 6.3 times more likely to have IDA. Although not statistically significant, this odds ratio suggests that the finding may be clinically important; the wide confidence interval suggests that the study was underpowered to

Table 4: Prevalence of IDA in stroke and TIA study subjects compared with NHANES III controls

<table>
<thead>
<tr>
<th>Iron and Stroke/TIA study</th>
<th>IDA present Frequency (%)</th>
<th>IDA absent Frequency (%)</th>
<th>Odds ratio (95 % CI)</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male</td>
<td>8.1%</td>
<td>2%</td>
<td>2.07</td>
<td>0.038</td>
</tr>
<tr>
<td>Female</td>
<td>6.8%</td>
<td>1%</td>
<td>3.11</td>
<td>0.002</td>
</tr>
</tbody>
</table>

1Includes subjects ≥70 years (n=37 males, n=44 females). 2One-Sample Proportion Test with continuity correction.

1Age and gender-matched control subjects (with black and Hispanic individuals excluded) without history of previous stroke who resided in areas in the Midwestern U.S.A. with populations <1,000,000 (n=94). 2Fisher’s Exact Test.
make this comparison. Finally, these conclusions are guarded for both comparisons, since our algorithm for detecting IDA was more rigorous than the methodology used in the NHANES III Study. In particular, sTfR measurements have superior predictive value in the presence of the anemia of chronic disease.

The experimental design utilized was not intended to establish IDA as a cause of stroke. However, the intriguing results do suggest that a comprehensive prospective case control study matching subjects for all major stroke risk factors should be undertaken to further investigate whether IDA increases risk for stroke in elderly individuals. Our prevalence estimate would provide valuable baseline data for calculating the sample size required for adequate power. This undertaking is especially compelling given the recent finding that healthy young children presenting with vaso-occlusive stroke are ten times more likely to have IDA than those without stroke. While it is acknowledged that the etiology of pediatric stroke differs from that of the adult, IDA has now been accepted as an etiological factor for stroke in children.

We also found a prevalence of 2.1% sub-clinical iron deficiency, which is characterized by iron storage depletion and tissue iron deficiency prior to the onset of anemia. While most studies on stroke risk have addressed IDA, there is insufficient evidence on underlying mechanisms to exclude sub-clinical iron deficiency as a possible risk factor for stroke. In fact, two epidemiological studies associated stroke risk with biochemical measures of early stage iron deficiency. Several mechanisms have been proposed by which IDA could promote the development of stroke, including thrombocytosis, altered erythrocyte deformability and blood flow, anemic hypoxia and endothelial dysfunction associated with inflammation. Some of these mechanisms are altered by early stage iron deficiency, but unfortunately, none have been carefully investigated in the setting of stroke.

A secondary objective of our study was to examine the role of dietary intake as a predictor of iron status. Iron supplements and heme iron (the more readily absorbed form) emerged as significant negative predictors of total iron binding capacity, which increases with iron deficiency and reflects the iron binding potential of transferrin. This is in agreement with previous reports of an inverse association of TIBC with iron supplements. The lack of association with total dietary or non-heme iron is not surprising, since the latter is less well absorbed and more influenced by dietary promoters and inhibitors of iron absorption. Nonheme iron absorption can also be impaired by atrophic gastritis, which is common in elderly persons.

No dietary variable emerged from our study as a significant predictor of hemoglobin, the indicator of late-stage iron deficiency anemia and stroke in elderly individuals. While dietary inadequacies may not have been prolonged enough to affect hemoglobin levels, in contrast to others who documented inverse associations between sTfR and iron intake in younger subjects, we found no associations between dietary variables and sTfR. Although these data suggest that dietary intake was not a major contributor to IDA in our patients, this conclusion is based on only 58 dietary analyses. Several medical conditions and medications are also known contributors to IDA in the elderly. These factors did not appear to be critical in our patients since we found no associations between IDA and nonsteroidal anti-inflammatory drug or anticoagulant use, gastrointestinal disease or genitourinary disease. However, this conclusion is limited by the relatively small number of IDA cases.

An additional 6.4% of our patients presented with anemia that could not be explained by IDA or the anemia of chronic disease. Although our algorithm was not intended to identify the type of these anemias, such cases deserve further study since anemia of all causes has also been implicated as a precipitant of stroke and related mortality. This does not preclude the possibility that subclinical iron deficiency or IDA confer added risk through specific mechanisms beyond anemic hypoxia.

In summary, the 6.4% prevalence of IDA documented in an elderly stroke and TIA group at hospital admission provides intriguing evidence to support a future case control study in elderly stroke patients. Further investigation should clarify whether IDA is a preventable cause of stroke in the elderly. The study of possible mechanisms for increased stroke risk in the setting of IDA is also warranted. Although our study has focused on IDA, other forms of anemia as well as early stage sub-clinical iron deficiency, should be further studied.

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REFERENCES


