THE EFFECT OF RETICULOCYTE HEMOGLOBIN CONTENT ON THE DIAGNOSIS OF IRON DEFICIENCY ANEMIA: A META-ANALYSIS STUDY

EFEKAT SADRŽAJA RETIKULOCITNOG HEMOGLOBINA NA DIJAGNOSTIKOVANJE ANEMIJE USLED NEDOSTATKA GVOŽDA: META-ANALIZA

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Summary

Background: Iron deficiency anemia (IDA) is the most common type of anemia worldwide and has many adverse effects on life quality. This meta-analysis study aims to show that reticulocyte hemoglobin content (CHr) is more effective than routinely used parameters in the diagnosis of IDA.

Methods: Comprehensive and systematic research was done using international databases including PubMed, Web of Science, Cochrane Library, Science Direct, and Google Scholar, which contain all articles published on IDA until December 29, 2020. Seventeen articles were included in the meta-analysis.

Results: The analyses found the Cohen's deffect size (Standardized Mean Difference) values of the parameters. Accordingly, CHr is 2.84 (95% CI 2.36 to 3.31), mean corpus volume (MCV) is 2.46 (95% CI 1.97 to 2.95), ferritin is 2.37 (95% CI 1.63 to 3.11), and transferrin saturation (TSAT) is 3.76 (95% CI 2.14 to 5.38). To diagnose IDA, the sensitivity value of the CHr concentration was found as 83.5% (95% CI 76.1 to 89.8), specificity value to be 91.8% (95% CI 85.5 to 96.4), and mean cut-off value as 28.2 pg.

Conclusions: The results of our study reveal the findings that CHr is a better biomarker than MCV and ferritin used in determining IDA, and its efficacy is lower than TSAT. It is very important to use it routinely for the pre-diagnosis of IDA, which is very important for public health. The groups in the study are heterogeneous but contain bias. Therefore, meta-analyses of studies with less heterogeneity of CHr are needed.

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Kratak sadržaj

Uvod: Anemija usled nedostatka gvožda (IDA) je najčešća vrsta anemije na svetu i ima mnogo negativnih efekata na kvalitet života. Cilj ove meta-analize je da pokaže da je koncentracija retikulocitnog hemoglobina (CHr) efikasnija od rutinski korištenih parametara u dijagnozi IDA.

Metode: Sprovedeno je sveobuhvatno i sistematično istraživanje gde su upotrijebljene međunarodne baze podataka, uključujući PubMed, Web of Science, Cochrane Library, Science Direct i Google Scholar, koje sadrže sve članke objavljene na temu IDA do 29. decembra 2020. Sjeverozapadni metode su upotrijebljene u meta-analizi.

Rezultati: Analizom utvrđena je veličina efekta Koenovog d (standardizovana srednja razlika). Shodno tome, CHr je 2,84 (95% CI 2,36 do 3,31), srednja zapremina tela (MCV) je 2,46 (95% CI 1,97 do 2,95), ferritin je 2,37 (95% CI 1,63 do 3,11), a zasićenje transferinom (TSAT) je 3,76 (95% CI 2,14 do 5,38). Da bi se dijagnostikovala IDA, utvrđena je vrednost osetljivosti koncentracije CHr na 83,5% (95% CI 76,1 do 89,8), vrednost specifičnosti 91,8% (95% CI 85,5 do 96,4), a srednja granična vrednost 28,2 pg.

Zaključak: Rezultati naše studije ukazuju da je CHr bolji biomarker od MCV i ferritina koji se koriste u određivanju IDA, a njegova efikasnost je niža od TSAT. Veoma je važno da se rutinski koristi za preddijagnozu IDA, što je veoma važno za javno zdravlje. Grupe u studiji su bile heterogene, ali imaju elemente pristrasnosti, pa je potrebno sprovedeti meta-analize sa manjom heterogenostim CHr.
Introduction

Iron is an element that has essential functions for human life. While it is found in the structure of hemoglobin (Hb), which provides oxygen transport in the body, it also ensures the fulfillment of iron-related functions by joining the structure of enzyme systems in some tissues (1). Iron deficiency anemia (IDA) occurs when the iron intake in humans is less than its excretion, in other words, when a negative iron balance occurs in the body (increased need for iron, absorption disorders, chronic blood loss) as a result of insufficient iron for Hb synthesis in the stores (1, 2). IDA is the most common type of anemia, and it constitutes the most advanced stage of iron deficiency. The World Health Organization (WHO) describes a hemoglobin value of <130.0 g/L in men, <120.0 g/L in women, and <110.0 g/L in pregnant women as anemia (3, 4). IDA is more common in women than men due to conditions such as menstruation and pregnancy. Adolescence causes an increase in blood pressure due to rapid growth and development and insufficient iron stores. If this condition cannot be compensated, IDA may occur as a result of insufficient intake. The most important reasons for IDA in postmenopausal women and men are the formation and increase of gastrointestinal system (GIS) bleeding (5–7). IDA is characterized by hypochromia and microcytosis in erythrocytes, decreased serum ferritin and serum iron levels, TSAT, and increased total iron-binding capacity (4, 8). Low serum ferritin level in IDA is essential and should not always be associated with IDA. Again, because it is an acute phase reactant, its normal condition does not exclude IDA; the underlying etiology must be defined and regulated (9). In contrast to all these conditions, iron overload reduces the efficiency of iron utilization and induces oxidative stress formation (10). In addition to these, free erythrocyte zinc protoporphyrin (ER-ZPP), soluble transferrin receptor (sTfR), and reticulocyte hemoglobin content (CHR or Ret-He) are among the reliable laboratory test parameters used to describe IDA. Soluble transferrin receptor with increasing erythrocyte ER-ZPP value causes early deterioration of iron condition and emergence of IDA (11–14).

Bone marrow erythropoietic activity and intracellular iron requirement are important criteria in determining sTfR level. Therefore, in conditions associated with iron deficiency and induced erythropoiesis (sickle cell anemia, megaloblastic anemia, thalassemia, polycythemia, etc.), sTfR concentration increases, while aplastic anemia decreases (15, 16). Normal serum sTfR level is 3.5–8.5 mg/L. It is known that a high sTfR (>8.5 mg/L) level is an early and sensitive biomarker for the diagnosis of IDA (11).

The ratio of sTfR concentration to logarithmic ferritin level is also determinant in the differential diagnosis of IDA. A ratio of less than 1 is associated with chronic disease anemia, while the ratio higher than 2 is evaluated in favour of IDA (17).

The decrease in iron concentration increases zinc transport in the intestines, and therefore the increased concentration of ER-ZPP (80 μg/dL) in erythrocytes is associated with iron deficiency. However, routine use of ER-ZPP measurements is difficult and time-consuming due to automation difficulties (18).

CHR, also known as Ret-He, measures the amount of hemoglobin in reticulocytes and is an indicator of cell hemoglobination, reflecting the quality of newly produced reticulocytes. Microcytic, hypochromic red blood cell (RBC) is formed due to ongoing reticulocyte production when there is not enough iron. Thus, RET-He reflects an earlier measure of reduced hemoglobin status compared to hemoglobin and hematocrit (11, 19).

Reticulocytes are separated from the erythroblasts after Hb synthesis, pass into the peripheral blood and turn into mature erythrocytes within a few days. Therefore, CHR is the ideal parameter to be considered for real-time Hb synthesis. Reticulocyte hemoglobin content is affected only by the amount of iron unless there are hematopoietic disorders (10).

Determination of iron status is possible with RET-He measurement. RET-He is determined by automated fluorescence flow cytometry, which measures the mean values of the forward light scattering intensity of mature red blood cells and reticulocytes using a polymethine dye. The values obtained reflect the reticulocyte hemoglobin content (20). Reticulocyte hemoglobin content is more effective in diagnosing iron deficiency, determining early iron deficiency anemia, differentiation of beta-thalassemia feature, and more effective than the other parameters involved in iron metabolism (10, 12). CHR is a less variable parameter that performs better than ferritin in response to intravenous (IV) iron therapy, providing better diagnostic accuracy for iron (15, 16).

In the United States and Europe, CHR has been accepted as a marker in iron deficiency with a diagnostic threshold of 29 pg. However, there is no reference value agreed on the best value for its sensitivity and specificity (17).
This study examined the effectiveness of CHr in addition to routine parameters in determining IDA.

**Methods**

**Literature Search Strategy**

Comprehensive systematic research was carried out using international databases, including PubMed, Web of Science, Cochrane Library, Science Direct, and Google Scholar, to determine all studies on CHr as a biomarker of IDA.

Our study included all articles published until December 29, 2020. The words used as search criteria in this study are as follows: »Reticulocyte hemoglobin AND iron deficiency anemia OR iron deficiency«, »reticulocyte hemoglobin« OR »iron deficiency« OR »iron deficiency anemia«.

This systematic review and meta-analysis were carried out under the guidelines for Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA).

**Selection Criteria**

The full texts of the articles related to the study were examined in terms of exclusion and inclusion criteria. Appropriate studies including clinical features of patients diagnosed with IDA and CHr used as a biomarker in the diagnosis of the disease were included in the meta-analysis.

Prospective/retrospective cohort, case-control, and cross-sectional studies were found suitable for inclusion in our study.

Repeated articles, reviews, case reports, expert opinions, letters, editorials, studies on experimental animals, studies not including control groups, studies where CHr is a biomarker but not related to iron deficiency, studies without available data, studies on results other than mean and standard deviation values, articles published in languages other than English, studies involving chronic patient groups with IDA, and studies only on iron deficiency were excluded (Figure 1).

**Data Extraction and Quality Assessment**

Data extraction, evaluation of literature quality, and evaluation of bias risk were carried out independently by two researchers (Serdar M. and Kılıç M.).

The following features were extracted: Article information (first author, year of publication), country, study designs, gender, device information, study example, patient/control information, the total number of samples, researched parameters. This information is shown in Table I. Microsoft Excel database was used to save the necessary information.

**Statistical Analysis**

The meta-analysis study was performed using MedCalc statistical software, version 19.0.7 (MedCalc Software, Ostend, Belgium). A meta-analysis was performed on Cohen’s d effect size (Standardized Mean Difference) by taking the mean and standard deviation values. In the evaluation, the results obtained with the random-effects model of the studies with low bias risk were taken into consideration. The study aimed to perform group analysis of CHr, MCV, ferritin, and TSAT biomarkers according to the target range in IDA. Using Cochran’s Q statistic and I² statistic respectively, statistical heterogeneity was evaluated. If the I² statistical value was above 50 per cent and the p-value was lower than 0.05, the heterogeneity was considered statistically significant.
| FIRST AUTHOR         | YEAR | COUNTRY      | DEVICE              | STUDY DESIGN | PATIENT GROUP                        | PATIENT/CONTROL                                      | TOTAL NUMBER | FEMALE | MALE | RESEARCHED PARAMETERS               |
|----------------------|------|--------------|---------------------|---------------|--------------------------------------|-------------------------------------------------------|---------------|--------|------|-------------------------------------|
| Chaipokam et al.     | 2017 | Thailand     | Sysmex XE Alpha     | Prospective cohort | Adult-Anemia | Control, IDA, Talasemi, Talasemi trait, AI | 267           | 190    | 77   | MCV, CHr                            |
| Rehu et al.          | 2011 | Finland      | ADVIA 120 and 2120  | Retrospective  | Adult-Anemia                         | Control, IDA, ACD                                       | 250           | 138    | 112  | CHr, MCV                            |
| Brugnara et al.      | 1999 | Amerika      | Technicon H3        |               | Iron Deficiency Children            | Control, IDA, ID                                       | 210           | 90     | 210  | CHr, MCV, Ferritin                  |
| Balci et al.         | 2016 | Turkey       | ADVIA 2120i         | Case-control   | 6–12 Years Old Children             | Control, IDA, B12 vitamin deficiency, mixed anemia     | 112           | 52     | 60   | CHr, MCV, Ferritin                  |
| Ceylan et al.        | 2007 | Turkey       | ADVIA 120           | Case-control   | Adult-Anemia                         | Control, IDA, ID, Talasemi minor, B12 vitamin deficiency | 131           | 91     | 40   | CHr, MCV                            |
| Cai et al.           | 2017 | China        | ADVIA 120           | Case-control   | Adults                               | Control, IDA, NIDA                                      | 140           | 100    | 40   | CHr, MCV, Ferritin                  |
| LUO et al.           | 2007 | China        | ADVIA 120           | Case-control   | Premenopausal Women-Anemia           | Control, IDA, NIDA                                      | 142           | 142    |      | CHr, MCV                            |
| Dinh et al.          | 2020 | Vietnam      | ADVIA 2120i         | Retrospective  | Adult-ESRD                           | Control, IDA, NIDA, ESRD, IDA-ESRD                     | 312           | 188    | 124  | CHr, MCV                            |
| Ageeli et al.        | 2013 | Saudi Arabia | ADVIA 2120i         | Case-control   | Adult-Anemia                         | Control, IDA, ACD, CRF                                  | 320           | 170    | 150  | CHr, MCV, Ferritin, TSAT            |
| Buttarello et al.    | 2016 | Italy        | Sysmex XE-5000      | Case-control   | Adult-Anemia                         | Control, IDA, ID, NIDA, trait β talasemia              | 290           |         |      | CHr, MCV, Ferritin                  |
| Uçar et al.          | 2019 | Turkey       | Sysmex XN 1000      | Case-control   | Adult-Anemia                         | Control, IDA, ID, NIDA                                  | 217           | 171    | 46   | CHr, MCV, Ferritin, TSAT            |
| Toki et al.          | 2017 | Japan        | Sysmex XN 300/XE    | Case-control   | Adult-Anemia                         | Control, IDA, ID, NIDA                                  | 211           | 148    | 63   | CHr, MCV, Ferritin, TSAT            |
| Vázquez-López et al. | 2019 | Spain        | ADVIA 120           | Case-control   | 1–16 Years Old Children              | Control, IDA, ID                                        | 1239          | 620    | 619  | CHr, MCV, Ferritin, TSAT            |
| Malczewska-Lenczowska et al. | 2017 | Poland       | ADVIA 120           | Case-control   | Sports women                         | Control, ID stage I, ID stage II                        | 219           | 219    |      | CHr, MCV, Ferritin                  |
| Chinudomwong et al.  | 2020 | Thailand     | Sysmex XN-10        | Case-control   | Adult-Anemia                         | Control, IDA, ID, NIDA, IDA-inflammation, NIDA-CKD, Talasemi | 958           | 603    | 335  | CHr, MCV, Ferritin                  |
| Urrechaga et al.     | 2011 | Spain        | Sysmex XE 5000      | Case-control   | Anemia and Talasemia                 | Control, talasemia trait, mild IDA, severe IDA          | 473           |         |      | CHr, MCV, Ferritin, TSAT            |
| Rungngu et al.       | 2016 | Indonesia    | Sysmex XE-2100      | Case-control   | 6–12 Years Old Children-Anemia      | Control, IDA, NIDA                                      | 50            | 17     | 33   | CHr, Ferritin                      |

ACD: Anemia of chronic disease, AI: Inflammation anemia, CHr: Reticulocyte hemoglobin content, CKD: Chronic kidney disease, CRF: Chronic renal failure ESRD: End-stage renal disease, ID: Iron deficiency, IDA: Iron deficiency anemia, MCV: Mean corpuscular volume, NIDA: Non-iron deficiency anemia, TM: β-Talasemia minor, TSAT: Transferrin Saturation
Publication Bias

Two reviewers evaluated independently the risk of bias in each study using the Diagnostic Precision Study Quality Assessment Tool (QUADAS-2) recommended (18). Publication bias is demonstrated by analysis with a funnel plot.

Results

Literature Search and Study Characteristics

Based on the search strategy, 113,536 studies were evaluated among the databases specified. A total of 13,116 studies were included in the study after the records, except for studies with open access and research articles were excluded. Thirteen thou-

Table II Meta-analysis of reticulocyte hemoglobin content (Chr), Mean Corpuscular Volume (MCV), ferritin, Transferrin Saturation (TSAT), sensitivity and specificity of Chr to diagnose iron deficiency anemia articles.

| Study                  | IDA (n) | IDA-Chr Main±SD  | Control (n) | Control-Chr Main±SD | Effect Size  | 95% CI Effect Size  | 95% CI Effect Size  | 95% CI Effect Size  | Sample size Proportion (%) | 95% CI Sample size Proportion (%) | 95% CI Sample size Proportion (%) |
|------------------------|---------|------------------|-------------|---------------------|--------------|---------------------|---------------------|---------------------|-----------------------------|---------------------------------|---------------------------------|
| Chaipokam et al. 2017  | 55      | 21.2±5.5         | 99          | 33.1±2.4            | -3.157       | -3.62 to -2.65      | -3.30 to -2.80       | -3.80 to -2.80       | 56                          | 85.0 to 91.9                   | 99 to 80.8 to 71.6 to 88.0      |
| Rehu et al. 2011       | 58      | 26.8±3.8         | 63          | 33.2±2.2            | -2.12        | -2.57 to -1.67      | -1.16 to -1.54       | -1.54 to -1.77       | 58                          | 82.7 to 91.4                   | 63 to 90.4 to 80.4 to 96.4      |
| Brugnara et al. 1999   | 24      | 24.2±2.7         | 186         | 26.8±1.8            | -1.349       | -1.79 to -1.08      | -1.08 to -1.51       | -1.51 to -0.64       | 24                          | 79.1 to 92.8                   | 186 to 74.7 to 67.8 to 80.8     |
| Bale et al. 2016       | 26      | 22.26±1          | 52          | 29.9±0.7            | -8.692       | -10.5 to -6.99      | -3.20 to -5.99       | -5.99 to -2.41       | 41                          | 85.3 to 94.4                   | 54 to 91.3 to 84.2 to 99.9      |
| Ceylan et al. 2007     | 41      | 21.8±5.3         | 54          | 28.2±1.7            | -2.349       | -2.94 to -1.75      | -1.85 to -2.40       | -2.40 to -1.50       | 41                          | 85.3 to 94.4                   | 54 to 91.3 to 84.2 to 99.9      |
| Cai et al. 2007        | 56      | 23.3±4           | 46          | 31.8±2.5            | -2.476       | -2.99 to -1.95      | -2.42 to -2.93       | -1.90 to -1.91       | 56                          | 87.5 to 94.8                   | 46 to 91.3 to 79.2 to 97.5      |
| Luo et al. 2007        | 30      | 23.5±3.1         | 71          | 52±1.1              | -4.036       | -5.15 to -3.65      | -3.609 to -4.27      | -2.94 to -2.93       | 59                          | 98.3 to 99.9                   | 145 to 97.9 to 94.0 to 99.5     |
| Ageel et al. 2013      | 100     | 22.9±2.9         | 60          | 50.9±1.3            | -5.72        | -3.76 to -2.79      | -4.095 to -4.68      | -5.55 to -3.55       | 100                         | 98.6 to 99.9                   | 145 to 97.9 to 94.0 to 99.5     |
| Butarello et al. 2016  | 58      | 24.4±4.8         | 164         | 35.1±2.2            | -5.23        | -3.66 to -2.80      | -2.141 to -2.50      | -1.78 to -1.51       | 58                          | 91.5 to 97.1                   | 164 to 94.5 to 88.9 to 97.4     |
| Uçar et al. 2019       | 52      | 21±4.1           | 54          | 36.6±7.2            | -2.69        | -3.22 to -2.15      | -2.996 to -3.55      | -2.43 to -1.10       | 52                          | 90.3 to 96.8                   | 54 to 100 to 95.2 to 100.0      |
| Toki et al. 2017       | 72      | 23.4±4.9         | 67          | 55.8±2.5            | -2.651       | -3.08 to -2.17      | -2.008 to -2.41      | -1.59 to -0.96       | 72                          | 90.9 to 99.9                   | 145 to 97.9 to 94.0 to 99.5     |
| Vázquez-L et al. 2019  | 13      | 24.5±5.0         | 1155        | 31.1±1.7            | -1.885       | -2.45 to -1.53      | -2.951 to -5.51      | -2.39 to -1.64       | 13                          | 91.5 to 97.1                   | 164 to 94.5 to 88.9 to 97.4     |
| Malczewska-L et al. 2017 | 55     | 29.3±1.8        | 87          | 31.4±1.0            | -1.567       | -2.02 to -1.12      | -0.457 to -0.86      | -0.05 to -2.44       | 55                          | 75.6 to 80.5                   | 155 to 96.7 to 92.6 to 98.9     |
| Chirudonowong et al. 2020 | 133  | 20.6±9          | 155         | 35.3±1.4            | -1.995       | -2.27 to -1.71      | -1.909 to -2.18      | -1.65 to -1.40       | 133                         | 75.6 to 80.5                   | 155 to 96.7 to 92.6 to 98.9     |
| Urechaga et al. 2011   | 126     | 22.5±3.7         | 90          | 35.7±1.4            | -3.827       | -4.28 to -3.57      | -5.255 to -5.64      | -2.82 to -2.44       | 126                         | 85.5 to 89.9                   | 980 to 91.8 to 85.6 to 96.4     |
| Rungngu et al. 2016    | 16      | 25.8±4.8         | 34          | 29.8±1.6            | -1.358       | -1.99 to -0.68      | -1.59 to -2.27       | -0.91 to -0.49       | 16                          | 45.7 to 70.1                   | 54 to 85.2 to 68.9 to 95.0      |
| Total (random effects) | 950     | 2540             |             |                     | -2.846       | -3.51 to -2.38      | -2.463 to -2.95      | -1.97 to -2.58       | 950                         | 76.1 to 89.9                   | 980 to 91.8 to 85.6 to 96.4     |

CI = Confidence interval, n = Sample size, SD = Standard Deviation
Figure 2A Forest plot of reticulocyte hemoglobin content (CHr), B Funnel plot of reticulocyte hemoglobin content, C Forest plot of mean corpuscular volume (MCV), D Funnel plot of mean corpuscular volume, E Forest plot of ferritin, F Funnel plot of ferritin, G Forest plot of transferrin saturation (TSAT), H Funnel plot of transferrin saturation.
sand twenty-two articles not suitable for the title and abstracts and published in languages other than English were excluded. From the remaining 94 pieces, 17 of them were included in the meta-analysis, excluding those with repeating records, animal studies, those that did not include a control group, and those related to chronic diseases other than iron deficiency. The flowchart and detailed literature search steps are shown in Figure 1.

Characteristics and Quality Studies

The articles on CHr, a biomarker used in patients with IDA, were included in our study, all in English until December 29, 2020. From the 17 articles, 2 of them were written in Thailand, 3 in Turkey, 2 in Spain, 1 in Finland, 1 in the United States, 2 in China, and one in Indonesia, Vietnam, Saudi Arabia, Italy, Poland, and Japan. The study contains 951 individuals with IDA and 5491 people, including 2540 control group members. The features of the included studies are shown in Figure 1.

Reticulocyte Hemoglobin Content

As a result of a detailed literature review, 17 articles including IDA data of CHr were included. In the meta-analysis performed with the low bias risk random-effects model in the included articles, the $I^2$ value of CHr was 93.52%, so intergroup heterogeneity was achieved ($P < 0.0001$), and random effect size value 2.84 (95% CI 2.36 to 3.31) is shown in Table II. Publication bias is evaluated with a forest plot in Figure 2A and funnel plot in Figure 2B.

Mean Corpuscular Volume

After the literature review, 16 articles with MCV data were included. In the meta-analysis performed with the low bias risk random-effects model in the included articles, the MCV $I^2$ value was 94.71%, so intergroup heterogeneity was achieved ($P < 0.0001$), and random effect size value 2.46 (95% CI 1.97 to 2.95) is shown in Table II. Publication bias is evaluated with a forest plot Figure 2C and funnel plot in Figure 2D.

Ferritin

12 articles with ferritin data were included. In the meta-analysis performed with the low bias risk random-effects model in the included articles, the ferritin $I^2$ value was 97.15%, so intergroup heterogeneity was achieved ($P < 0.0001$), and random effect size value 2.37 (95% CI 1.63 to 3.11) is shown in Table II. Publication bias is evaluated with a forest plot Figure 2E and funnel plot in Figure 2F.

Transferrin Saturation

5 articles with TSAT data were included. In the meta-analysis performed with the low bias risk model of random effects in the included articles, the TSAT $I^2$ value was 97.70%. Therefore, intergroup heterogeneity was achieved ($P < 0.001$), and random effect size value 3.76 (95% CI 2.14 to 5.38) is shown in Table II. Publication bias is evaluated with a forest plot Figure 2G and funnel plot in Figure 2H.

Discussion

Anemia is a global health problem that is quite common worldwide and affects 43% of children under the age of five, 38% of pregnant women, and 29% of non-pregnant women (19). IDA brings along many complications such as growth retardation, neurocognitive deficiencies, impaired immune system, increased risk of premature, and impaired learning ability (20–24). It is therefore important to ensure accurate and timely diagnosis of the disease by preventing such adverse effects (25–27). Although there is no single and best test for determining iron deficiency, bone marrow aspiration, which is accepted as the gold standard, and the method of staining bone marrow macrophages and erythroid precursors with Prussian blue is used. However, the method is not suitable for routine use because it is expensive, subjective, and invasive (26, 28).

There are many biochemical parameters used in the diagnosis of IDA. However, as these parameters are affected by certain conditions, it is not easy to evaluate them. Serum ferritin concentration, serum iron level, TSAT, and total iron-binding capacity (TIBC) are the most widely used biochemical tests. Although serum ferritin level reveals the iron concentration accumulated in the body, factors such as acute and chronic inflammation, malignancy, liver diseases, and excessive alcohol use increase independent of iron (29). Serum iron level decreases with infection, inflammation, and malignancy but increases with liver disease. Since the TSAT level is calculated on iron and TIBC, it is affected by changes in these values and does not always give an accurate result (29–32).

In recent years, CHr has become one of the parameters used to determine IDA (33, 34). Reticulocytes, as the first erythrocytes produced in the bone marrow, transform into mature red blood cells a day or two after entering the bloodstream. Reticulocyte parameters have become one of the parameters used to reflect the iron status in a short time due to their shorter lifespan compared to erythrocytes and ability to provide information about bone marrow erythrocyte production (12, 35). CHr has a higher specificity and a lower coefficient of variation since it is not affected by inflammation like some parameters used in the diagnosis of IDA (36). Reticulocyte hemoglobin content data can be obtained from a few millimeters.
Figure 3A Forest plot meta-analysis of the sensitivity of reticulocyte hemoglobin content to diagnose iron-deficiency anemia articles, B Funnel plot meta-analysis of the sensitivity of reticulocyte hemoglobin content to diagnose iron-deficiency anemia articles, C Forest plot meta-analysis of the specificity of reticulocyte hemoglobin content to diagnose iron-deficiency anemia articles, D Funnel plot meta-analysis of the specificity of reticulocyte hemoglobin content to diagnose iron-deficiency anemia articles, the reticulocyte hemoglobin content mean cut-off value.
of peripheral blood compared to bone marrow biopsy and is also advantageous because it is relatively inexpensive, convenient, and less invasive (37).

In this random-effect meta-analysis, studies on the efficiency of CHr in the diagnosis of IDA are summarized. For this meta-analysis, a total of 13,116 articles were examined, and 17 studies were included in the study according to the inclusion and exclusion criteria. The number of articles including parameters such as MCV, ferritin, and TSAT, which were used frequently in the past in the diagnosis of iron deficiency, varies (16 articles for MCV values, 12 articles for Ferritin values, and 5 articles for TSAT values were examined). The most effective examination is thought to be obtained by including each study that meets the criteria given in Figure 1 for meta-analysis. The literature review exhibited the fact that this study is the first meta-analysis of CHr to determine the diagnosis of IDA.

A total of 3491 individuals, of which 2540 people were in the control group, and 951 people were with IDA, were included in this study. The inclusion criteria for the current study were determined as CHr, which is one of the important parameters in the diagnosis of IDA and containing at least one of the parameters of MCV, ferritin, or TSAT.

Ferritin is an indicator of the total amount of iron stored in the body. When the serum ferritin level shows values less than 15 mg/L, it is highly suggestive for the diagnosis of IDA. At values below 30 mg/L, the sensitivity is 92%, and the specificity is 98% (38, 39). Again, a ferritin level below 45 mg/L and the sensitivity of 85% and specificity of 92% are particular to IDA (40, 41).

A low MCV value alone is not sufficient for a diagnosis of IDA. Unless the MCV volume is greater than 95 fL (95 fl), it should not be considered in IDA because this threshold has a sensitivity of 97.6% (40, 42).

TSAT is low in IDA, typically less than 10%, and in this case, the sensitivity is 48%, and the specificity is 88% (43, 44).

In the meta-analysis performed with the low bias risk random-effects model in the included articles, the sensitivity CHr I² value was 78.78% (95% CI 61.43 to 88.32), so intergroup heterogeneity was achieved (P<0.0001) (Figure 3A–B). The sensitivity of CHr to diagnose IDA was found as 83.5% (95% CI 76.1 to 89.8), and they are shown in Table II.

In the meta-analysis performed with the low bias risk random-effects model in the included articles, the specificity CHr I² value was 89.53% (95% CI 82.88 to 93.60), so intergroup heterogeneity was achieved (P<0.0001) (Figure 3C–D). The sensitivity of CHr to diagnose IDA was found as 91.8% (95% CI 85.5 to 96.4), and they are shown in Table II.

In conclusion, the meta-analysis study showed that CHr is a better marker than other more commonly used parameters in IDA. Many previous studies also support this conclusion. Also, this meta-analysis we conducted is important for being the first meta-analysis study regarding CHr and IDA.

There are important heterogeneity and bias problems in the studies. Patient selection and lack of reference methods are particularly important. There are important criteria differences for index tests. Also,
it is important to have significant group differences (pediatric patients, renal failure, etc.) in studies. It should be noted that there will be a significant change in methodologies over time (Figure 4).

All parameters of IDA, which are included in the research, have heterogeneity. CHr is a more effective marker in determining IDA compared to the routinely used MCV and ferritin levels. The effect size value of TSAT, one of the parameters examined, is above 0.80, and its selectivity is higher than CHr.

The study also has some limitations. These are as follows: like other meta-analysis studies in the literature, methodological differences are arising from combining studies conducted with different methods, and this may lead to bias. Since studies in which the diagnosis of IDA of CHr was evaluated in the meta-analysis were included, many parameters used routinely were excluded. IDA, erythropoiesis status, and chronic anemias were excluded from the study. In the study, no distinction was made according to gender and age.

**Conclusion**

This study is the first meta-analysis to evaluate the efficiency of CHr in the diagnosis of IDA. According to our results, CHr should be used additionally with the parameters used in the diagnosis of IDA. ‘The results of our study reveal the findings that CHr is a better biomarker than MCV and ferritin used in determining IDA, and its efficacy is lower than TSAT. It is very important to routinely use it for the pre-diagnosis of IDA, which is very important for public health.’ CHr alone provides important information about the current bioavailability of iron, but its use with other parameters removes uncertainty about the diagnosis and treatment of IDA. CHr is a very important parameter that can be used to evaluate a very common disease in the clinic, such as IDA. The heterogeneity index of the study results is quite high. Therefore, comprehensive studies with more homogeneous groups are needed to elucidate the relationship between IDA and CHr.

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**Conflict of interest statement**

All the authors declare that they have no conflict of interest in this work.

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