PRE AND POST IRON THERAPY ABSOLUTE RETICULOCYTE COUNT- A RELIABLE TEST WITH EARLY RESPONSE TO TREATMENT IN IRON DEFICIENCY ANAEMIA

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ABSTRACT

BACKGROUND
Iron deficiency anaemia is usually treated using oral iron therapy as first line treatment. Several tests are available to monitor the response to oral iron treatment, but absolute reticulocyte count [ARC] is supposed to be the one showing one of the quickest responses, with an increase of ARC within 5 to 10 days of therapy initiation. Our aim is to verify that ARC is significantly increased following iron therapy and that the response is quick and cost effective.

METHODS

50 adults with recently diagnosed but untreated iron deficiency anaemia were included in this prospective cohort study. Haemoglobin [Hb], RBC indices, peripheral smear and ARC were done at baseline [T0], day 7 [T1] and day 30 [T2] after the start of oral iron therapy.

RESULTS

Mean (SD) Hb level at T0 was 8.93 (1.41) g/dl [Hb0]. It increased to 9.22 (1.33) g/dl [Hb1] and 10.59 (1.28) g/dl [Hb2] at T1 and T2, respectively. Mean (SD) ARC level at T0 was 19.6 x 10^9 (12.7 x 10^9) /l [ARC0]. It increased to 80.15 x 10^9 (28.1 x 10^9)/l [ARC1] at T1 and then reduced to 36.08 x 10^9 (17.9 x 10^9)/l [ARC2] at T2. Using Repeated measures ANOVA, there was a statistically significant (p<0.0001) increase in mean Hb2 compared to mean Hb0 (baseline), while there was no significant rise of mean Hb1 when compared with mean Hb0. 84% showed at least 1 g/dl increase of Hb by day 30. Mean ARC1 was statistically higher (p<0.001) than mean ARC0, while mean ARC2 was significantly lower (p value<0.001) than mean ARC1. Based on Spearman correlation test, there was a positive correlation (r = 0.282, n = 50, p<0.05 - significant) between relative rise of Hb [after 30 days] and relative rise of ARC [after 7 days].

CONCLUSIONS

Our study confirms that ARC is an ideal early predictor of response to iron therapy as it increases by 7th day compared with Hb which does not rise significantly till 4 weeks of therapy. Also, ARC can be done manually, requiring no special equipment and therefore cost effective especially in resource constrained settings.


Iron deficiency anaemia can be suspected by reduction in the following laboratory indicators: Haemoglobin [Hb], Haematocrit [PCV], RBC count, Mean Corpuscular Volume [MCV], Mean Corpuscular Haemoglobin [MCH], Mean Corpuscular Haemoglobin Concentration [MCHC] along with a raised Red cell Distribution Width [RDW]. Peripheral smear findings include microcytes, hypochromic RBC & anisopoliocytosis. Oral iron replacement is the most economical and preferred medication for the treatment of iron deficiency anaemia. Yet, failure of oral iron therapy may occur in case of i) Inadequate prescription ii) Continuing iron loss in excess of intake iii) Malabsorption of iron iv) Associated illness such as anaemia of inflammation in rheumatoid arthritis, Helicobacter pylori infection and Celiac disease. Failure of treatment may also occur in case of incorrect diagnosis of other conditions like Thalassemia, Sideroblastic anaemia, myelodysplastic syndromes as Iron deficiency anaemia. Since the routine laboratory tests are not sufficient to distinguish these conditions, they may go unnoticed until the ineffective response to treatment is found. So, we need other sensitive &more specific indicators in detecting IDA which include reduction in serum iron, serum ferritin, transferrin saturation [TS] and increase in Total Iron Binding Capacity [TIBC] & soluble transferrin receptor levels. But, these tests are relatively costly and lab facilities necessary for performing these tests are not available among...
all health care centres. Perl’s iron staining of bone marrow smears is a sensitive marker of body iron stores but not a first line of investigation since it is invasive.

Response to treatment can be identified by both clinical and laboratory investigations. Clinically, tongue and nails are the most responsive to treatment. In tongue, filiform papillae regenerate after 1 to 2 weeks and tongue returns to normal after 3 months. In nails, koilonychia takes 3 to 6 months to disappear.[10] Lab investigations useful in monitoring response include RBC count, Hb level, RDW, Red cell indices, Reticulocyte count, serum ferritin, TIBC and TS. Haematocrit reaches half way towards normal by 3 weeks and Haemoglobin reaches normal levels by 2 months after therapy is initiated.[10,12] Red cell indices may remain abnormal for some time even after the normal Haemoglobin level is restored. A study conducted among hospitalised patients revealed that MCV has low sensitivity whereas TIBC and TS have low specificity in detecting IDA.[13]

Of all, the earliest haematological evidence of response to treatment is an increase in the percentage of reticulocytes. Reticulocytes are juvenile red blood cells containing reticular network of rRNA. Once the bone marrow starts responding to treatment, an increase in reticulocyte count is expected. Reticulocyte count can be recorded either as a percentage [Normal count is about 0.5% to 2.5% in adults][9] or as an absolute reticulocyte count [normal range in adults: 50-100 x 10^9/L][9]. The reticulocytes attain a maximal value on 5th to 10th day after initiation of therapy.[10] The maximal value usually ranges from 5% to 10%. No special technology is required for this procedure which makes it more preferable.

To verify the reliability of this test, a prospective cohort study was planned to be conducted among the newly diagnosed cases of iron deficiency anaemia reporting in a tertiary health care centre. Previous study conducted in same regard among paediatric age group confirms the effectiveness of ARC in predicting the response to iron therapy.[14] This study was planned to assess whether pre and post iron therapy absolute reticulocyte count is a reliable & early method to monitor response to iron therapy in adults.

The primary objective of this study was to find whether ARC was a reliable laboratory indicator for predicting response to oral iron in iron deficiency anaemia patients. This was achieved by determining whether there was a significant increase in absolute reticulocyte count after initiation of iron therapy and whether the increase in absolute reticulocyte count was early and cost effective.

METHODS

After getting clearance from the Institutional Ethics Committee (IEC), Prospective cohort study was conducted among the newly diagnosed cases of iron deficiency anaemia (diagnosis was based on reduced Haemoglobin [Adult male: <13 g/dL; Adult female: <12 g/dL], raised RDW and peripheral smear findings of microcytic hypochromic RBCs) reporting in a Tertiary Health Care Centre in Perambalur district. 50 adults irrespective of gender were selected by simple random sampling method from the cohort of patients with proven iron deficiency anaemia. Sample size was taken random based on the convenience of the study. After obtaining informed written consent, all adult patients with laboratory evidence of iron deficiency anaemia who required oral iron therapy were included in the study. All patients in whom iron therapy had already been started, in whom treatment other than oral iron therapy was required, who were lost to follow up, paediatric patients (< 12 years age) and patients unwilling for the study were excluded.

Data collected included demographic details and baseline laboratory investigations comprising of Haemoglobin, Haematocrit, RBC count, MCV, MCH, MCHC, RDW [all tests done using Horiba 5-part haematology analyser], peripheral blood smear findings and ARC. One week and one month after start of iron therapy, blood sample was collected and Haemoglobin, Haematocrit, RBC count, MCV, MCH, MCHC (CBC) and ARC investigations were carried out. The same pathologist (to avoid individual variation) reported all the reticulocyte counts and he was blinded of the study. Reticulocyte count [both % and ARC] was done using standard procedure routinely done in the tertiary care centre in which two to three drops of commercially available reticulocyte staining fluid (New methylene blue) was taken in a 75x10 mm plastic tube by means of a plastic Pasteur pipette. To this 2-4 volumes of the patient’s EDTA anticoagulated blood was added and incubated at 37°C for 20 minutes. The mixture was gently mixed, and films were made on glass slides and examined under the microscope. Reticulocytes were identified as red cells with blue precipitate, granules or filaments in the cytoplasm. Reticulocyte count is given by the formula:

\[ \text{Reticulocyte count [\%]} = \frac{\text{Number of reticulocytes in "n" fields} \times 100}{\text{L}} \]

For severe anaemia the reticulocyte count was corrected {corrected ARC = Reticulocyte count [\%] \times \frac{\text{(Patient Haematocrit/45)}}{\text{ARC}}}.[10] Samples [from normal healthy volunteers without anaemia] were used as quality control for ARC.

Statistical Methods

Data were entered in Excel and statistical analysis done using SPSS 20.0 software. A p value of < 0.05 was considered as statistically significant. The ARC before iron therapy, 1 week and 1 month after iron therapy were compared using repeated measures ANOVA test and its significance determined. Similarly, the Haemoglobin level before iron therapy, 1 week and 1 month after iron therapy were also compared by the same statistical test.

The correlation between ARC and Haemoglobin levels was done by Spearman correlation test. The response to iron treatment was evaluated by improvement in haematology parameters [> 1 g/dL increase of Haemoglobin,[15] Haematocrit, RBC count, MCV, MCH, MCHC] one month after start of therapy.

RESULTS

Fifty patients of proven iron deficiency anaemia, predominantly female patients, had a mean (SD [Standard Deviation]) age of 36.18 yrs. (13.87), and ranging from 20 to 80 years. The three blood samples from study participants were labelled as T0 [before initiation of iron therapy], T1 [1 week after iron therapy] and T2 [1 month after iron therapy] respectively, the corresponding Haemoglobin values were designated as Hb0, Hb1 and Hb2, and ARC as ARC1, ARC2 and ARC3.
Hb and ARC for all the three samples are presented in figures 1 and 2 respectively which show Hb0 values ranging from 5.30 g/dL to 10.90 g/dL, Hb1 values ranging from 6.1 g/dL to 11.5 g/dL, Hb2 values ranging from 6.6 g/dL to 12.7 g/dL. ARC0 values ranged from 3.6 x 10^9/L to 63 x 10^9/L, ARC1 values ranging from 23.8 x 10^9/L to 168 x 10^9/L, and ARC2 values ranging from 7.2 x 10^9/L to 94.3 x 10^9/L. Mean (SD) Hb0 was 8.93 (1.41) g/dL, Hb1 was 9.22 (1.33) g/dL, and Hb2 was 10.59 (1.28) g/dL (Fig. 3). Since both Haemoglobin and ARC follow normal distribution, Repeated measures ANOVA was applied to compare the means. Data analysis using repeated measures ANOVA comparing the different RBC parameters and their significance are given in Table 1.

### DISCUSSION

Hb2 was significantly higher compared to Hb0 and Hb1 (p < 0.001) whereas the rise of Hb from Hb0 to Hb1 was not that significant (p<0.01) whereas there was a statistically significant (p < 0.001) increase in ARC from ARC0 to ARC1 and a decrease in ARC values by 30th day (ARC2). 82% [41 cases] showed an increase of 1 g/dL Haemoglobin by 30th day.
after iron therapy i.e., 82% responded to treatment with oral iron (15). Relative rise of Hb (Hb2-Hb0/Hb0) after a month and relative rise of ARC (ARC1-ARC0/ARC0) after a week were compared using Spearman correlation test to identify the correlation between relative rise of Hb and ARC. The analysis showed that there was a positive correlation between relative rise of Hb after a month and relative rise of ARC after a week ($r = 0.282, n = 50, p=0.05$ - significant).

Okam et al(16) in their analysis of 738 cases had a mean Hb of 9.3 g/dL while in our study the baseline Hb was 8.93 g/dL which was almost similar. 72.8% cases had an increase of Hb of >1 g/dL after 14 days of iron therapy, whereas in our study 82% of cases had an increase of Hb of >1 g/dL after 30 days of treatment. Okam et al found that the rise of ARC had an 81.9% positive predictive value in predicting a rise of > 2 g/dL Hb at day 42-56 interval. In our study, in 82% of cases the rise in ARC1 (Day 7 ARC) accurately predicted a Hb2 (day 30 Hb) increase of at least 1 g/dL, while in 18% cases the ARC increase was not associated with a concomitant increase of Hb. This requires further evaluation.

Parodi et al(14) did a study on 34 cases of children with IDA measuring both Hb and ARC at baseline, 7 days & 30 days after oral iron therapy. In their study, mean (SD) Hb level before treatment (T0) was 6.84 (2.42) g/dL which increased to 7.36 g/L (1.31) and 10.56 g/L (1.62) at T1 and T2 respectively, while the Hb parameters in our study were 8.93(1.41) g/dL, 9.22(1.33) g/dL and 10.59(1.28) g/dL at T0, T1 and T2 respectively. Mean (SD) ARC in the Parodi et al study was 72.8 x 10^6/L (35.155) before treatment (T0). It increased to 168.6 x 10^6/L (101.2) at T1 and then decreased again to 76.3 x 10^6/L (38.9) at T2, respectively. In our study Mean (SD) ARC level at T0 was 19.6 x 10^6/L (12.7 x 10^6) which increased to 80.15 x 10^6/L (28.1 x 10^6) at T1 and then reduced to 36.08 x 10^6/L (17.9 x 10^6) at T2. On comparing both studies, we found that both had a significant increase of mean ARC by 7th day after iron therapy. We deduce from our study as well as the studies by Parodi et al and Okam et al that ARC rises very early around 7th day of treatment. Okam et al found that the rise of ARC on day 7 on all those cases. Similarly, the rise of Hb was < 1 g/dL after 30 days and all of them also had a rise of ARC at 7th day. This finding requires further evaluation of the reason for non-rise of Hb even though ARC increased.

A population study done to monitor response to iron therapy like serum ferritin, transferrin saturation, total iron binding capacity and bone marrow evaluation. In our study, manual routine used lab procedure was used. This further raises a question about the difference in ARC values in different experimental setup which needs further evaluation by larger studies. This study also found that other indices were not much significant in predicting response to iron replacement therapy, but ARC was positive well in advance of the traditional increase of at least 1 g/dL of Haemoglobin after 1 month of iron treatment.

**CONCLUSIONS**

Firstly, our study confirms the fact that oral iron improves haemoglobin level as we noted that 82% of our cases improved by at least 1 g/dL in 30 days. Secondly, within one week of therapy there is no significant increase of Hb compared to baseline values whereas absolute reticulocyte count increased significantly in the same period. Hence this conclusively proves that ARC is an early indicator of response to oral iron therapy. Thirdly, as is already known, the cost of ARC is very less compared to other tests for monitoring the response to iron therapy like serum ferritin, transferrin saturation, total iron binding capacity and bone marrow evaluation. Therefore, ARC is a cost-effective alternative to monitor iron therapy response.

Since our study was of a short duration, we limited our study to 50 cases, and we could not identify cases in which there was no increase of ARC. More studies on larger cohorts is required to identify cases which do not have an increase of ARC so that we could confirm if lack of ARC increase corresponds to failure of response to iron treatment. Also, some cases (16%) in which there was no Hb response however showed an ARC increase. This requires further study on maturation of reticulocytes to RBCs to understand the reasons behind this finding. Further studies are required to analyse the differences between manual and automated methods of reticulocyte counts.
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REFERENCES