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Abstract. Thalassemia (Thal), sickle cell anemia, and iron deficiency anemia (IDA) are the most common blood disorders in many parts of the world, particularly in developing countries like India and Bangladesh. The well-established diagnostic procedure for them is the complete blood count (CBC); however, there is substantial confusion in discrimination between Thal and IDA blood samples based on such CBC. We propose a new spectral technique for reliable classification between the above two anemias. This is based on the identification and quantification of a certain set of fluorescent metabolites found in the blood samples of patients of Thal and IDA. © The Authors. Published by SPIE under a Creative Commons Attribution 3.0 Unported License. Distribution or reproduction of this work in whole or in part requires full attribution of the original publication, including its DOI. [DOI: 10.1117/1.JBO.19.2.027008]

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1 Introduction

Blood is the essential bodily fluid that delivers oxygen and nutrients to different organs of the body and carries away waste products from them. There are three types of blood cells: erythrocytes, leukocytes, and thrombocytes. By volume, the red blood cells constitute ~45% of whole blood, the plasma ~54.3%, and white cells ~0.7%. The normal hemoglobin (Hb) level varies for men from 13.5 to 17.5 g/dl and for women from 12.0 to 15.5 g/dl; a person with Hb level <10 g/dl is termed as anemic.

Anemia is the most common blood condition in the world and affects women more than men. Certain forms of anemia (e.g., thalassemia [Thal], sickle cell) are hereditary, and others such as iron deficiency anemia (IDA) mostly arise out of excessive blood loss, decreased red cell production, etc. Deficiency of iron or vitamin B12 in diet are some of the most common causes for IDA. These are temporary and could be easily rectified.

Beta Thalassemia (β Thal) is a genetic disorder caused by the reduced synthesis of the beta chains of hemoglobin. Individuals with β Thal major usually present with severe anemia, poor growth, and skeletal abnormalities within the first two years of life. This disease is most common in Mediterranean, west Asian, and north African countries as well as in India. The incidence of Thal disease is highest in Greece, but the number of Thal carriers is highest in the Maldives.

The most common methods of β Thal diagnosis are Hb electrophoresis, gel electrophoresis, and high-performance liquid chromatography (HPLC). These three techniques separate out different hemoglobin (e.g., HbA, HbS, HbA2, etc.) and quantify their concentration in term of percentage and classify as α Thal, β Thal, etc.1,2 Apart from the above methods, polymerase chain reaction is also used occasionally, which provides the gene profiles.3 Many studies had shown that IDA can cause misdiagnosis with heterozygote β-Thal. The hematologists then recommend rechecking of HbA2 level after overcoming IDA.4,5

Earlier, in a series of papers, we had shown the efficacy of spectral technique for diagnosis of different types of cancers.6 As a logical extension, we demonstrate here the potential of spectral differentiation between IDA and β Thal patients. This method could prove useful to identify and separate IDA samples from those of Thal.

2 Materials and Methods

Experiments were performed on blood samples from IDA and Thal patients and also age-adjusted healthy controls.

2.1 Controls

Exactly 5 ml of venous blood from each of 15 healthy volunteers (age range: 18 to 34 years) was collected in a violet sterile vial that contained the anticoagulant ethylenediamine tetra-acetic acid (EDTA). The vial was gently rocked five times to adequately mix the EDTA with the whole blood; then it was centrifuged at 3000 rotations per minute (rpm) for 15 min. A pale, greenish-yellow clear plasma supernatant was obtained by such centrifugation. A total of 1.5 ml of supernatant was removed from the top layer for spectrofluorometric analysis, leaving behind the buffy coat and the formed elements. The blood plasma sample thus obtained was subjected to synchronous spectral analyses without any other treatment.

Next, the buffy coat, which contained mostly the white blood cells, was removed and discarded, and exactly 1 ml of the thick formed elements from the bottom layer, which contained mostly red blood cells (RBCs), was removed and drawn into a sterile vial. It was then mixed with 2 ml of analytical-grade acetone.

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Proper care was taken to ensure that the formed elements did not develop lumps. After thorough mixing to enable the acetone to extract the fluorophores within and around the cells, the sample was centrifuged again (3000 rpm for 15 min). The resulting supernatant was subjected to fluorescence emission spectra analysis with an excitation wavelength of 400 nm.

2.2 Patients

The same protocol was used to process blood samples obtained from confirmed IDA and Thal patients. Samples were collected from all the Thal patients (β Thal) of Voluntary Health Service, Chennai, India, waiting for blood transfusion. Similarly, samples were collected from confirmed IDA patients of Kidwai Memorial Institute of Oncology, Bangalore, India. The patients were informed about the investigation, and proper consents were obtained. The IDA subjects (nine females and six males) had a median age of 26 years, and the Hb level for them varied from 6.2 to 9.5 with a median of 8.2 gm/dl. There were 15 Thal subjects (seven females and eight males) with a median age of 20 years, and the Hb level for them varied from 4.2 to 7.5 with a median of 6.2 gm/dl. The normal controls were students, researchers, and staff nurses (seven females and eight males) with an average age of 25, from whom informed consents were obtained. The Hb level for the control set varied from 11.2 to 15.5 with a median of 13.3 gm/dl.

The instrument used was a spectrofluorometer (Elico India LS142) capable of collecting excitation, emission, and synchronous spectra in the 200- to 800-nm range. An excitation and emission slit width of 10 nm and scan speeds of 1000 nm/min were used. Each sample was placed in quartz cuvettes and illuminated by a specified wavelength of light with a 10-nm spectral width and a spot size of 3 by 2 mm.

Three types of spectra are common in the field of fluorescence spectroscopy. In fluorescence emission spectra (FES), one particular wavelength is selected for the excitation of a molecule, and the fluorescence emission spectrum is obtained by rotating the emission grating over a predetermined range. The reverse is true of fluorescence excitation spectra (FXS), in which the peak emission band of a molecule is selected, and the excitation grating is rotated to scan the excitation spectra. Such FES and FXS are very useful for understanding the absorption and emission characterizing of a particular molecule. In contrast, for getting the information about the relative concentration about a set of molecules in a composite mixture like blood, synchronous excitation or emission spectra is more reliable. In synchronous excitation spectra (SXS), both gratings are synchronously rotated with an offset of 70 nm to obtain the fluorescence excitation band of every molecule in the predetermined range. The wavelength offset and scan range are not unique; they are determined empirically by trial and error for a given set of experimental protocols.

After analyzing other offsets, it was found in our experiment that the 70-nm offset provided excellent resolution and good contrast between the IDA and Thal samples because 70 nm was the Stokes shift of the most important biofluorophores [e.g., tryptophan, nicotinamide adenine dinucleotide (NADH), flavin adenine dinucleotide (FAD), etc.] found in blood plasma. When the offset was 10 nm, instead of 70 nm, the fluorescence emission bands of the same set of molecules present in the sample were obtained. This is known as synchronous emission spectra (SES).
Parameter is defined as \( R_2 = \frac{I_{465}}{I_{365}} \), i.e., the ratio of intensity at 465 nm to that at 365 nm. Such fluorescence intensity ratio parameter, instead of the actual intensity, reduces experimental and instrument variation. This ratio represents the relative concentration of fluorophores, such as FAD (with excitation/absorption peak at 465 nm) and NADH (with excitation/absorption at 365 nm). This ratio is \( \sim 0.98 \) for the normal control.

Figure 2(a) is to be compared with the SXS features of average spectra of Thal plasma, shown in Fig. 2(b). The two spectra are distinctly different with very low intensity at 365 nm for Thal. Once again \( R_3 \) is calculated and this is \( \sim 4.2 \), which means \( R_3 \) is about four times elevated for the Thal samples. Such changes are found due to decrease in intensity at 365 nm rather than the increase at 465 nm.

Figure 2(c) gives the average spectra of plasma sample of IDA patients. This set is markedly different from those of the normal control and also from those of the Thal subjects in terms of the ratio \( R_3 \), which is \( \sim 0.7 \). This ratio is 1.4 times smaller than that of the normal control and six times smaller than that of the Thal. Such distinct differences are mostly due to the band at 365 nm (due to NADH) and that at 465 nm (due to FAD). A scatter plot was obtained for the actual values of \( R_1 \) for all three sets, as shown in Fig. 3.

In an earlier paper, we had shown the spectral features to discriminate Thal samples from the normal control; hence, we focus more on the discrimination between Thal and IDA samples for the rest of the investigation.

Figure 4 gives the SES obtained with the offset of 10 nm between the emission and excitation grating. Here, Fig. 4(a) is the averaged intensity spectra for Thal and Fig. 4(b) for IDA samples. The two spectra are significantly different. For example, the first peak occurs at 325 nm and the most prominent peak occurs at 525 nm. In contrast, the first peak occurs at 360 nm for IDA; the next prominent peak at 500 nm is not much elevated. The ratio parameter \( R_3 = \frac{I_{325}}{I_{360}} \) gives the value 1.5 for Thal and 0.33 for the IDA. Another ratio \( R_4 = \frac{I_{585}}{I_{560}} \) is also taken for the Thal and IDA samples. The average value of \( R_4 \) is \( \sim 1.9 \) for the Thal and 0.62 for IDA.

As our main objective is discrimination between the Thal and IDA samples, receiver operating characteristic (ROC) curves were obtained for all the three ratio parameters \((R_2, R_3, R_4)\) obtained from the spectral features of plasma. For each of them, the area under the curve (AUC) is \( >0.8 \), indicating a significant level of separation. Note that for \( R_2 \), the curve runs parallel to both axes, showing a separation of 100%.

4 Discussion

When patients present with the symptoms of anemia, the clinician will send the blood samples of the patients for the complete blood count (CBC) analysis, as RBC indices <10 g/dl are crucial in anemia diagnosis as well as ferritin levels, and the next battery of tests would be hemoglobin electrophoresis and/or HPLC to identify blood disorders like Thal, IDA, sickle cell anemia, etc. These techniques are cumbersome and expensive.

Spectral diagnosis is a new technique wherein the detection and quantification of a certain set of fluorescent biomolecules have enabled diagnosis of liver cancer,10 cervical cancer,6 pancreatic cancer,11 etc., with sensitivity and specificity on the order of 90%. The same technique has been extended to the discrimination between two major inherited blood disorders: Thal and sickle cell anemia.12 This paper now pertains to obtaining spectral features of IDA and Thal blood samples to classify them as two distinct categories, as well as possible.

The Hb level for the IDA set under investigation had a range from 6.2 to 8.2 g/dl and for the Thal from 4.2 to 7.5 g/dl. Though Hb level was \( \sim 1.3 \) times greater for IDA subjects than for the Thal ones, there was considerable overlap between the Thal and IDA set from the point of view of hemoglobin level.

The fluorescence component in RBC was porphyrin, and its intensity was approximately proportional to the concentration of RBC in blood. Figure 1 shows a measure of porphyrin in three types of blood samples. It can be seen that the ratio parameter \( R_1 \) is \( \sim 1.4 \) for normal control, 0.5 for Thal, and 1.0 for IDA.

It is important to note that the spectral diagnosis establishes one-to-one correspondence to the conventional pathological findings. Figure 3 shows the scatter plot of ratio \( R_1 \) for all three sets. The spectral diagnosis as presented here also could not separate IDA from Thal completely as shown in Fig. 3.

However, the spectral features of blood plasma could accomplish this objective very well as shown in Figs. 2 and 4. Figure 2 is the SXS of blood plasma of the normal control, IDA, and Thal. It is easy to see the differences in the spectral features, the most important being the changes in the concentration of NADH and FAD. The ratio \( R_2 \) indicates the relative
concentration between FAD and NADH; this is $\sim 1.0$ for the normal control, 4 for Thal, and 0.7 for IDA. The ROC for this ratio alone runs parallel to both axes (see Fig. 5), indicating sensitivity and specificity each of 100%

The average lifetime of a normal RBC is $\sim 110$ days, and some of its final product is biliverdin and bilirubin. In this metabolic process, FAD is produced and NADH is consumed copiously. For normal controls, the ratio between these two is maintained in proper equilibrium as unity. This is not so for anemic patients. For Thal patients, not only the RBC concentration is about twice or thrice as low, but the average lifetime of RBCs is five times lower as well. This is reflected in the striking imbalance between NADH and FAD levels, with an abnormally low value for NADH and high value for FAD. In contrast, the RBC of IDA, though less in number, apparently does not decay as fast as that of the Thal patients. Due to some unidentified enzyme activity for the IDA, intensity of NADH is more elevated than that for Thal and normal controls. More work needs to be done to confirm this surprising feature, and also to identify the biochemical reason.

Figure 4 is the synchronous emission spectra (SES) of IDA and Thal plasma samples only. SXS and SFS are expected to be the two sides of the same coin, since the former one represents the excitation spectra and the latter one the emission spectra of the individual fluorophores in a composite system. This would be strictly true only when three pure biomolecules (tryptophan, NADH, FAD) are taken in saline water, but not as found in blood plasma. This is because blood plasma contains many biomolecules other than the above three and, hence, distortion is considerable.

In spite of these, there are 290-, 365-, and 465-nm bands in the SXS and 360-, 440-, and 500-nm bands in the SFS. These are the excitation and emission spectra of tryptophan, NADH, and FAD, respectively. In addition, there is a band at 325 nm due to tyrosine and 585 nm due to porphyrin in SFS (Fig. 4). These spectral band assignments agree well with the existing literature values. All these were confirmed by us too by adding small quantities of tyrosine, tryptophan, NADH, FAD, and porphyrin individually in saline water and taking the SXS and SFS (not shown here).

Ratio parameters $R_1 = I_{325}/I_{360}$ and $R_4 = I_{585}/I_{505}$ could also distinguish IDA from Thal reasonably well, with AUC for $R_1$ being 0.812 and that for $R_4$ being 0.924. Among the four parameters, it is $R_2$ which has the highest classification capability in terms of AUC.

The most important feature of the result reported here is that we could not separate IDA from Thal by CBC (based on RBC quantification), spectral features of plasma (which is generally discarded so far) could provide a reliable and easy classification strategy.

## 5 Conclusion

In this preliminary investigation, a fluorescence spectral analysis has been carried out with a limited number of blood samples drawn from patients with Thal and IDA. Our results show that the spectral analysis on plasma, better than on cellular component, would give more reliable results. Out of the two types of synchronous spectra reported here, SXS is better than SFS for classification of two sets with an accuracy of 100%. It is important to mention that a large number of such samples are being analyzed in a multicenter study to make this technique a clinically viable protocol.

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**Fig. 5** Receiver operating characteristic of ratio parameters of $R_2$, $R_3$, and $R_4$. This gives the classification efficiency of two data sets, IDA and thalassemia.
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