

See discussions, stats, and author profiles for this publication at: <https://www.researchgate.net/publication/347516627>

# Mean corpuscular volume/mean corpuscular hemoglobin values are not reliable predictors of the $\beta$ -thalassemia carrier status among healthy diverse populations of Himachal Pradesh, In...

Article in *Asian Journal of Transfusion Science* · December 2020

DOI: 10.4103/ajts.AJTS\_109\_18

CITATION

1

READS

177

8 authors, including:



**Omesh Kumar Bharti**

Indira Gandhi Medical College

169 PUBLICATIONS 340 CITATIONS

SEE PROFILE



**Rajesh Kumar Sood**

Department of Health and Family Welfare, HP, India

35 PUBLICATIONS 110 CITATIONS

SEE PROFILE



**Rajinder Chauhan**

Bennett University

188 PUBLICATIONS 2,489 CITATIONS

SEE PROFILE



**Nitu Nigam**

King George's Medical University

78 PUBLICATIONS 175 CITATIONS

SEE PROFILE

Some of the authors of this publication are also working on these related projects:



UniDrug [View project](#)



108 emergency response services at GVKEMRI [View project](#)

|   |
|---|
| Access this article online  |
| Quick Response Code:  |
|  |
| Website:<br><a href="http://www.ajts.org">www.ajts.org</a>                        |
| DOI:<br>10.4103/ajts.AJTS_109_18  |

# Mean corpuscular volume/mean corpuscular hemoglobin values are not reliable predictors of the $\beta$ -thalassemia carrier status among healthy diverse populations of Himachal Pradesh, India

Omesh Kumar Bharti<sup>1</sup>, Rajesh Kumar Sood<sup>2</sup>, Hans Raj Sharma<sup>2</sup>, Harsharan Kaur<sup>1</sup>, Varinder Minhas<sup>1</sup>, Rajinder Chauhan<sup>3</sup>, Nitu Nigam<sup>4</sup>, Archana Phull<sup>5</sup>

## Abstract:

**BACKGROUND:** Himachal Pradesh is a hill state in North India in the Western Himalayas.  $\beta$ -thalassemia is a genetic disorder of hemoglobin inherited in an autosomal recessive manner that results in defective globin production leading to the early destruction of red blood cells.  $\beta$ -thalassemia has long been neglected in Himachal Pradesh due to popular belief that it runs along "Lahore-Gujarat-Punjab" belt in India. Therefore, there is no  $\beta$ -thalassemia testing facility currently in the state.

**METHODS:** To estimate the prevalence of  $\beta$ -thalassemia carriers, we calculated the sample size based on probability proportional to size self-weighting design. In each of 20 selected colleges, 111 students having an age of 18–25 were tested for high-performance liquid chromatography (HPLC) and complete blood count. Some were further tested for the mutations. We computed sensitivity, specificity, positive predictive value (PPV) and negative predictive value, and receiver operating characteristic curve for mean corpuscular volume (MCV) and mean corpuscular hemoglobin (MCH) red cell parameters.

**RESULTS:** Of the 2220 students, 57 were found to be  $\beta$ -thalassemia carrier by HPLC. The overall prevalence rate was 2.6% which translates to probable 180,000  $\beta$ -thalassemia carriers in Himachal Pradesh. Six districts bordering highly endemic Punjab had a higher prevalence. Hemoglobin D-Punjab, Heterozygous-Iran Trait, and raised fetal hemoglobin were found. Thalassemia major and sickle cell disease were not found. Anemic status or MCV/MCH parameters were not found to be reliable predictors of thalassemia carrier status among the healthy populations of HP. The predominant mutation found was IVS 1–5 G > C.

**CONCLUSION:** Popular ongoing strategy for screening with MCV and MCH has low-PPV and can miss upto 37% of true thalassemia carriers. HPLC is better strategy for screening carriers and reduces further spread of thalassemia.

## Keywords:

Complete blood count, hematological parameters, high-performance liquid chromatography, prevalence,  $\beta$ -thalassemia carriers

## Introduction

$\beta$ -thalassemia is a genetic disorder of hemoglobin dysfunction inherited in an

autosomal recessive manner that results in defective globin production leading to the early destruction of red blood cells (RBCs) and consequent microcytic hypochromic anemia. It is estimated that 1.5% of the

This is an open access journal, and articles are distributed under the terms of the Creative Commons Attribution-NonCommercial-ShareAlike 4.0 License, which allows others to remix, tweak, and build upon the work non-commercially, as long as appropriate credit is given and the new creations are licensed under the identical terms.

**How to cite this article:** Bharti OK, Sood RK, Sharma HR, Kaur H, Minhas V, Chauhan R, *et al.* Mean corpuscular volume/mean corpuscular hemoglobin values are not reliable predictors of  $\beta$ -thalassemia carrier status among healthy diverse populations of Himachal Pradesh, India. *Asian J Transfus Sci* 2020;14:172-8.

For reprints contact: [WKHLRPMedknow\\_reprints@wolterskluwer.com](mailto:WKHLRPMedknow_reprints@wolterskluwer.com)

<sup>1</sup>State Institute of Health and Family Welfare,  
<sup>2</sup>National Health Mission (NHM), Government of Himachal Pradesh, Shimla, Himachal Pradesh,  
<sup>3</sup>Prof. and Head Biotechnology, Bennett University, Greater Noida, Delhi,  
<sup>4</sup>Department of Centre for Advance Research (CFAR), King George's Medical University, Lucknow, Uttar Pradesh,  
<sup>5</sup>Independent Researcher and Master's from CSK Himachal Pradesh Agriculture University, Palampur, Himachal Pradesh, India

## Address for correspondence:

Dr. Omesh Kumar Bharti,  
State Epidemiologist,  
SIHFW and Principal Investigator Thalassemia Project NHM,  
Himachal Pradesh, India.  
E-mail: [bhartiomes@yaho.com](mailto:bhartiomes@yaho.com)

Submission: 06-09-2018  
Accepted: 02-12-2018  
Published: 19-12-2020

global population, i.e., 80–90 million people, are the carriers of  $\beta$ -thalassemia, and the South-East Asian region (India, Thailand, and Indonesia) account for approximately 50% of the world's carriers—approximately 40 million people and approximately half of the affected births.<sup>[1]</sup>  $\beta$ -globin gene (HBB) is located in the short arm of chromosome 11, and >200 HBB gene mutations are known to be associated with  $\beta$ -thalassemia globally and of these about 28 mutations have been documented in Indian patients.<sup>[2]</sup> It is estimated that there are 30 million  $\beta$ -thalassemia carriers in India only, but no population-based survey has been done in any state till date. It is also estimated that every year >7000 children are born  $\beta$ -thalassemic major in India and require frequent blood transfusions besides iron-chelating therapy. There are seven common  $\beta$ -thalassemia mutations found in India, and carrier rate may go up to 9.5%–15% in different regions in the general population in India.<sup>[3,4]</sup> Thalassemia has long been neglected in Himachal Pradesh due to popular belief that it runs along “Lahore-Gujarat-Punjab” belt in India. Therefore, no thalassemia testing facility is currently available in the state.

Himachal Pradesh is a hilly state of North India in Western Himalayan belt and has better health indicators than national averages, but there was no population-based data on genetic disorders in the state. The author while working in the State Blood Bank, Shimla, realized the threat of thalassemia from affected children who used to come for blood transfusion and submitted a project proposal of estimating the prevalence of  $\beta$ -thalassemia trait in the state of Himachal to the National Rural Health Mission. The project was approved with the objective of estimating the prevalence of thalassemia trait in Himachal Pradesh and to find suitable red cell based parameters for the establishment of a low-cost screening program in the state.

## Methods

Since, there is no prevalence study done in any state, a multicentric study<sup>[5]</sup> done in six cities of six states in India was taken as reference, and higher limit of prevalence of 2.94% was taken to calculate the sample size. Using public domain Epi Info software version 7.1.2.0 developed by the Centers for Disease Control and Prevention, Atlanta, USA with a cluster sample technique with a confidence limit of 1% and confidence interval of 95% with design effect of 2, with 20 clusters we got a sample size of 2220, i.e., 111 students in each cluster. To account for nonresponse, sample losses and deterioration during the transport we proposed to take additional 10% samples and total target sample size came out to be 2442. To maximize the benefits, we decided to take college-going young marriageable population of 18–25 years of age as the target population. The selection of colleges was done

by probability proportional to size self-weighting design. All the colleges including private ones were line listed, and a random number was selected using MS Excel, the sampling interval was added to this number to finalize 20 colleges comprising a population of 39,710 students. Before the screening, we sensitized the students of all 20 selected colleges through poster competitions and debates over the issue. A total of 45 Senior Secondary schools and one university were also involved in the awareness and stakeholder engagement. An awareness poster in the Hindi language, commonly used in this state, focusing on knowing the thalassemia carrier status before marriage was developed by the authors for sensitization, [Figure 1]. Interactive “Hello Doctor” radio talks were also organized. After sufficient sensitization over 2 years, blood sampling camps were conducted in consultation with director of higher education. The ethical clearance was taken from the Institutional Ethics Committee of J. P. University-wide “IEC/Project No-27-2015”, Dated 27-11-2015. Every student was asked to give informed consent in the form of written signed consent statement, whereas right to participate was voluntary. In each selected college, students were asked to pick slips for inclusion or exclusion and those included were tested for high-performance liquid chromatography (HPLC) and complete blood count (CBC) parameters. No one was denied the right for test, but only first 111 samples were included for analysis as per the sample size. Most of the colleges were Rural Government Colleges, and most of the participants were girls. The collected samples were analyzed locally same day or within 24 h for CBC by the Hematology Counter “Act Diff 5 CP Hematology Analyzer-Beckman Coulter Act Diff 5”, and the second samples were sent to Gurgaon for HPLC analysis by Bio-Rad variant II HPLC system by  $\beta$ -thalassemia short program. Seven type of Hb variants were tested with the help of HPLC, i.e., HbA, HbA2, fetal hemoglobin (HbF), hemoglobin (Hb D), HbS, hemoglobin D-Punjab (Hb D) Iran Trait, HbC, and unknown peaks with a disclaimer that this screening test does not rule out any Alpha-thalassemia/Hb variants that elute at similar retention times on HPLC. Receiver operating characteristic (ROC) curves were plotted for both mean corpuscular volume (MCV) and mean corpuscular hemoglobin (MCH) values using MedCalc easy to use the statistical software (Atlanta, Georgia). The methodology to prepare the ROC curve using the MedCalc software trial version was DeLong *et al.*<sup>[6]</sup> Confidence intervals and *P* values were calculated using Open Epi, Score (Wilson) software.<sup>[7]</sup>

Three common family counseling sessions were organized with the help of professional counselors to inform the thalassemia carrier students, the status they have and precautions they need before or after marriage. With some funds left from the project, we went ahead with the genetic analysis of some small samples that



**Table 2: Prevalence of thalassemia trait (n=57)**

|              | n  | N    | Prevalence (CI with P)         |
|--------------|----|------|--------------------------------|
| Sex (%)      |    |      |                                |
| Female       | 44 | 1596 | 2.8 (CI: 2-3.7, <0.0000001)    |
| Male         | 13 | 624  | 2.1 (CI: 1.2-3.5, <0.0000001)  |
| Religion (%) |    |      |                                |
| Hindu        | 55 | 2156 | 2.6 (CI: 2-3.3, <0.0000001)    |
| Muslim       | 1  | 27   | 3.7 (CI: 0.6-18.2, <0.0000001) |
| Sikh         | 1  | 26   | 3.8                            |
| Buddhist     | 0  | 10   | 0.0                            |
| Christian    | 0  | 1    | 0.0                            |
| Caste (%)    |    |      |                                |
| General      | 36 | 1381 | 2.6 (CI: 2-3.6, <0.0000001)    |
| OBC          | 9  | 327  | 2.8 (CI: 1.5-5.1, <0.0000001)  |
| SC           | 11 | 418  | 2.6 (CI: 1.5-4.7, <0.0000001)  |
| ST           | 1  | 94   | 1.1 (CI: 0.2-5.8, <0.0000001)  |
| Total        | 57 | 2220 | 2.6 (CI: 2-3.3, <0.0000001)    |

CI=Confidence interval

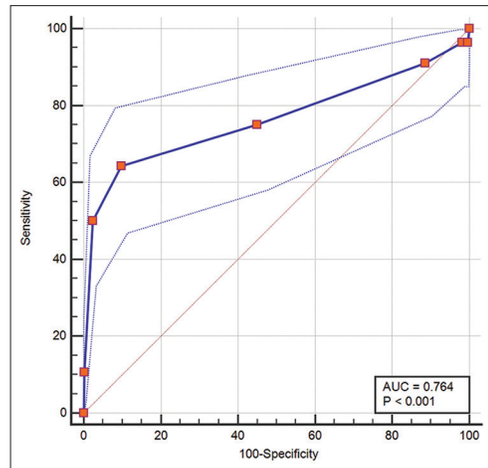
**Table 3: District bordering Punjab having a higher prevalence**

| District of birth | Carrier | Healthy | Total (%)  |
|-------------------|---------|---------|------------|
| Bilaspur          | 4       | 62      | 66 (6.1)   |
| Una               | 11      | 259     | 270 (4.1)  |
| Kangra            | 17      | 444     | 461 (3.7)  |
| Hamirpur          | 5       | 143     | 148 (3.4)  |
| Mandi             | 11      | 329     | 340 (3.2)  |
| Sirmour           | 3       | 116     | 119 (2.5)  |
| Solan             | 2       | 142     | 144 (1.4)  |
| Kullu             | 2       | 271     | 273 (0.7)  |
| Shimla            | 2       | 355     | 357 (0.6)  |
| Chamba            | 0       | 8       | 8 (0.0)    |
| Kinnaur           | 0       | 15      | 15 (0.0)   |
| L and S           | 0       | 10      | 10 (0.0)   |
| New Delhi         | 0       | 1       | 1 (0.0)    |
| Pathankot         | 0       | 3       | 3 (0.0)    |
| Punjab            | 0       | 1       | 1 (0.0)    |
| Roper             | 0       | 1       | 1 (0.0)    |
| Saharanpur        | 0       | 2       | 2 (0.0)    |
| Haryana           | 0       | 1       | 1 (0.0)    |
| Grand total       | 57      | 2163    | 2220 (2.6) |

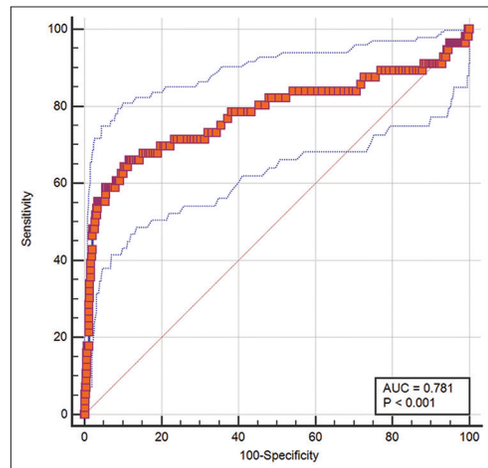
in normal range, therefore, further genetic analysis is required for the final diagnosis. Out of 11 siblings of the carriers who volunteered for testing, two brothers were found positive for thalassemia trait, i.e., KSG-118 B (HbA2 = 4.6) and HMR-084B (HbD = 37.0) [Table 9].

## Discussion

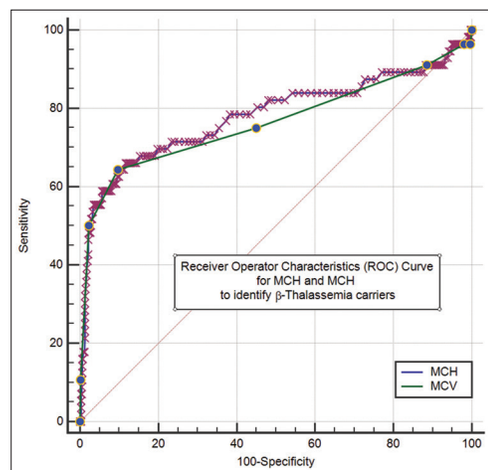
We found 57 samples positive for  $\beta$ -thalassemia trait, but there are borderline HbA2 between 3 and 3.9 that may be a carrier and may have lead us to underestimate the actual burden of the disease. Rosnah *et al.* found that 36/117 borderline samples (30%) were positive for thalassemia trait.<sup>[11]</sup> We had 610 such samples having borderline HbA2 between 3 and 3.9, and 23 were detected as a carrier by HPLC against expected 183 carriers as



**Figure 2:** Receiver operating characteristic curve for the ability of mean corpuscular volume to detect the  $\beta$ -Thalassemia carrier



**Figure 3:** Receiver operating characteristic curves for the ability of mean corpuscular hemoglobin to identify the  $\beta$ -thalassemia carrier



**Figure 4:** Receiver operating characteristic curve for mean corpuscular volume/mean corpuscular hemoglobin to identify the  $\beta$ -thalassemia carriers

per the study referenced above. Another study also flags the limitation of HPLC in borderline HbA2 values to predict  $\beta$ -thalassemia trait and suggest to combine

borderline HbA2 values with MCH <27 pg for better estimation.<sup>[12]</sup> Moreover, with this value, we found 23 possible carriers with HPLC than expected 31 out of 610 having borderline HbA2 values. However, sensitivity analysis we did for hematological markers in our study shows that although MCH <27 pg was found to be capturing more  $\beta$ -thalassemia carriers, but the positive predictive value (PPV) was low for both MCH <27 pg and MCV <80fl separately or collectively [Table 7]. Therefore, both MCH <27 pg and MCV <80fl are not reliable predictors of detecting  $\beta$ -thalassemia trait in mass screening camps as they can miss upto 37% true  $\beta$ -thalassemia carriers. The ROC curve provided an AUC value of 0.764 for MCV and 0.781 for MCH, suggesting only fair and not very good discrimination ability of these

parameters to detect  $\beta$ -thalassemia carrier status. However, we can say that an HPLC value of HbA2 >4 is a fairly good indicator for  $\beta$ -thalassemia carrier detection and borderline values (3–3.9) may need to be combined with the cell parameters (MCH <27) as detailed above. Another study observes that it appears difficult to differentiate the thalassemia major and intermedia by red cell morphology alone or with absolute values. Therefore, blood findings must be correlated with a clinical picture for an accurate diagnosis.<sup>[13]</sup> Most of the studies done to calculate the sensitivity of hematological markers as cut off for the detection of  $\beta$ -thalassemia trait have a small sample and are not population-based random sampling studies and, therefore, give higher sensitivity than observed by this study. Some of the other studies showing the sensitivity of cell indices, the reliability of cell indices was lost in averages that blurred the real picture or some of the studies were carried out were laboratory/hospital based.<sup>[14-16]</sup> A study<sup>[17]</sup> from Thailand advocated MCH <26.5 as cut off to get PPV of 40.4% but in this population-based study, this value could capture a total of 314 persons as possible carriers but only 37 actual carriers out of 57 carriers. Higher PPV values than we observed in this study for MCV, i.e., 30.85% were found in a study<sup>[18]</sup> on pregnant women in India, but cell indicators still are not reliable as they may still declare 70% potential carriers as noncarriers. In this study, on pregnant women, the PPV of the naked eye single tube red cell osmotic fragility test (NESTROFT) was found to be more (53.45%) than those of RBC count, however, we found making NESTROFT solution complicated and full of flaws and difficult to maintain uniformity/osmolality<sup>[19]</sup> in low-ionic strength solution over long durations. In another study,<sup>[20]</sup> the PPV of NESTROFT was calculated to be 35.3% while those of cell indices it was 32.5% for MCV <80 fl and 30.6% for MCV <75 fl. Out of 25.2% NESTROFT positive, actual positive captured was only 9.8%. In another study, Shewale *et al.* used readymade 0.36% buffered saline for NESTROFT and found that it missed 46 persons out of 1000 known  $\beta$ -thalassemia carriers and picked up 179 persons as  $\beta$ -thalassemia carriers out of 1000 healthy persons which has cost implications as well for getting HPLC or genetic evaluation for confirmation. False positivity in this study was attributed to the possible anemic status of the persons.<sup>[21]</sup>

In this study, anemia was no longer a factor in nondiagnosis of  $\beta$ -thalassemia trait, as 12% of anemic carriers were identified by the HPLC, this is in tune

**Table 4: Hemoglobin variants in the carriers as reported by the laboratory**

| Type Hb variant (n=57)   | Carrier, Noncarrier | n       |
|--|---------------------|---------|
| HB variant analysis shows BTT HBA2 window of $\geq 3.9$                                    |                     | 36      |
| HB variant analysis is suggestive of HBD Punjab heterozygous                               |                     | 8       |
| HB variant analysis shows borderline high normal HBA2 window of 3.3                        | 43                  | 5       |
| BTT with slightly raised HBF window  |                     | 3       |
| HBD Iran trait   |                     | 2       |
| HB variant analysis shows borderline high normal HBA2 window of 3.5                        |                     | 1       |
| HB variant analysis shows borderline high normal HBA2 window of 3.7                        |                     | 1       |
| HB variant analysis show borderline raised HBA2 window of 3.8%                             |                     | 1       |
| HB variant analysis appears to be normal with slightly raised HBF noted acquired elevation | 2                   |         |
| Normal   |                     | 2118    |
| Total  |                     | 57 2163 |

HB=Hemoglobin, HBF=Fetal hemoglobin, HBD=Hemoglobin D variant Punjab, BTT= $\beta$ -thalassemia trait, HBA2=Hemoglobin A<sub>2</sub>

**Table 5: Prevalence of thalassemia trait by anemia status**

|                      | Carrier | Healthy | Total | Prevalence (%), P             |
|----------------------|---------|---------|-------|-------------------------------|
| Anemia (<11 g/dl)    | 7       | 140     | 147   | 4.8 (CI: 2-9.5, <0.0000001)   |
| No anemia (>11 g/dl) | 50      | 2023    | 2073  | 2.4 (CI: 1.8-3.1, <0.0000001) |
| Total                | 57      | 2163    | 2220  | 2.6 (CI: 2-3.3, <0.0000001)   |

CI=Confidence interval

**Table 6: Distribution of level of hemoglobin A<sub>2</sub> among screened population**

| HBA2     | Carrier by HPLC | Healthy | Total population | Prevalence (%), P                |
|----------|-----------------|---------|------------------|----------------------------------|
| <3.0     | 11*             | 1565    | 1576             | 0.7 (CI: 0.39-1.28, <0.0000001)  |
| 3.0-3.4  | 2               | 591     | 593              | 0.33 (CI: 0.9-1.2, <0.0000001)   |
| 3.5-3.9  | 10              | 7       | 17               | 58.8 (CI: 36.1-78.39, 0.0000001) |
| $\geq 4$ | 34              | 0       | 34               | 100                              |

\*Mostly HbD >34.4 (as mutation seen in 34.4) and raised HBF window. HBF=Fetal hemoglobin, HBD=Hemoglobin D variant Punjab, HBA2=Hemoglobin A<sub>2</sub>, HPLC=High-performance liquid chromatography, CI=Confidence interval

**Table 7: Distribution of level of hemoglobin A<sub>2</sub> among screened population with respect to mean corpuscular volume/mean corpuscular hemoglobin values**

| HBA2           | The total screened population sample | Carrier by HPLC | Carrier by MCV $\leq$ 80 | Carrier by MCH $\leq$ 27 |
|----------------|--------------------------------------|-----------------|--------------------------|--------------------------|
| <3.0           | 1576                                 | 11              | 200                      | 332                      |
| 3.0-3.4        | 593                                  | 2               | 7                        | 25                       |
| 3.5-3.9        | 17                                   | 10              | 4                        | 6                        |
| $\geq$ 4       | 34                                   | 34              | 33 except HBD 38.1       | 34                       |
| <3.0- $\geq$ 4 | 2220                                 | 57              | 244                      | 397                      |

HBA2=Hemoglobin A<sub>2</sub>, HPLC=High-performance liquid chromatography, HBD=Hemoglobin D variant Punjab, MCV=Mean corpuscular volume, MCH=Mean corpuscular hemoglobin

**Table 8: Sensitivity and specificity of mean corpuscular volume/mean corpuscular hemoglobin to predict thalassemia carrier status**

|                     | Carrier | Healthy | Total | Sensitivity (%) | Specificity (%) | PPV (%) | NPV (%) |
|---------------------|---------|---------|-------|-----------------|-----------------|---------|---------|
| MCV <80             | 36      | 200     | 236   | 63.1            | 90.7            | 15.2    | 98.9    |
| MCV $\geq$ 80       | 21      | 1963    | 1984  |                 |                 |         |         |
| Total               | 57      | 2163    | 2220  |                 |                 |         |         |
| MCH <27             | 39      | 343     | 382   | 68.4            | 84.1            | 10.2    | 99.0    |
| MCH $\geq$ 27       | 18      | 1820    | 1838  |                 |                 |         |         |
| Total               | 57      | 2163    | 2220  |                 |                 |         |         |
| MCH $\leq$ 30       | 46      | 1020    | 1066  | 80.7            | 52.8            | 4.3     | 99.0    |
| MCH >30             | 11      | 1143    | 1154  |                 |                 |         |         |
| Total               | 57      | 2163    | 2220  |                 |                 |         |         |
| MCH <27 and MCV <80 | 36      | 195     | 231   | 63.1            |                 |         |         |

PPV=Positive predictive value, NPV=Negative predictive value, MCV=Mean corpuscular volume, MCH=Mean corpuscular hemoglobin

**Table 9: Detected mutations in some of the samples we could get voluntarily during counseling**

| Code    | HPLC                                    | Genetic mutation  |
|---------|---|---|
| CM-002  | 4.5 value for HBA2                      | IVS 1-5 G>C mutation ( $\beta$ +) )                                       |
| KUL-137 | 4.4 value for HBA2                      | IVS 1-5 G>C mutation ( $\beta$ +) )                                       |
| KSG-118 | 4.5 value for HBA2                      | IVS 1-5 G>C mutation ( $\beta$ +) )                                       |
| SL-016  | 3.7 value for HBA2                      | IVS 1-5 G>C mutation ( $\beta$ +) )                                       |
| KSG-120 | 4.8 value for HBA2                      | IVS 1-5 G>C mutation ( $\beta$ +) )                                       |
| HMR-84  | 34.4 value for HBD                      | IVS 1-5 G>C mutation ( $\beta$ +) )                                       |
| DHR-091 | 5.9 value for HBA2                      | 41/42 TCTT mutation ( $\beta$ +) )  |
| NPR-048 | 5.7 value for HBA2                      | Codon 8/9+G mutation ( $\beta$ +) )                                       |
| NPR-170 | 5.3 value for HBA2<br>3.6 value for HbF | Common five mutations not detected, need to be tested for other mutations |
| DHR-049 | 3.5 value for HBA2                      | Common five mutations not detected, need to be tested for other mutations |
| JHN-025 | 3.3 value for HBA2                      | Not detected  |
| SJP-047 | 3.3 value for HBA2                      | Not detected  |
| KSG-159 | 3.3 value for HBA2                      | Not detected  |
| NPR-029 | 3.2 value for HBA2                      | Not detected  |
| NPR-152 | 3.0 Value for HBA2                      | Not detected  |

HPLC=High-performance liquid chromatography, HBA2=Hemoglobin A<sub>2</sub>, HBF=Fetal hemoglobin, HBD=Hemoglobin D variant Punjab

with another study<sup>[22]</sup> done by P Sharma *et al.* that reported that iron deficiency is not a barrier for detecting  $\beta$ -thalassemia trait with HPLC.

Overall 6.3% of healthy students, mostly girls of marriageable age were found to be anemic that has implications for the maternal mortality rate and infant mortality rate. Severely, anemic girls were also counseled. Anemia was defined as per the National Family Health Survey-4 Manual.

National guidelines on for the prevention and control of hemoglobinopathies in India available on online<sup>[23]</sup> prescribe a screening protocol based on four steps, i.e., NESTROFT followed by MCV/MCH values <80/27 followed by HPLC followed by the DNA studies. We attempted to look for more sensitive red blood cell values for low-cost mass screening, but to our surprise, these cell indicators missed 37% of actual carriers despite nearly 99% of negative predictive value that the national policy refers to [Table 8] which is unacceptable. If we want to control thalassemia, we should start compulsory screening of college-going students and pregnant women with HPLC without thinking of cost implications to halt increasing carriers and birth of thalassemic children or children with other hemoglobinopathies. New genetic-based super kits can detect up to 20 types of genetic mutation and can be customized to suit local requirements. Such kits need to be developed in-house for better cost-effectiveness. A popular ongoing strategy for screening with MCV and MCH has low PPV and miss many true thalassemia carriers and has subsequent repercussions for the parents and the nation regarding costs<sup>[24]</sup> and to the thalassemic children, regarding clinical burdens.<sup>[25]</sup>

## Conclusion

Popular ongoing strategy for screening with MCV and MCH parameters has low-PPV and can miss upto 37% of true thalassemia carriers. HPLC is better and cost effective<sup>[26,27]</sup> strategy for screening carriers that would reduce further spread of thalassemia.

## Acknowledgments

We sincerely acknowledge the funding given by the National Health Mission along with voluntary office support given by Mr. Vinay Vashist, SIHFW, Parimahal, Shimla. We are grateful to SRL Laboratory for agreeing to do HPLC and CBC at government approved rates. We are thankful to all Directors of Health Services and Principals of SIHFW, Kasumpti for their support and continuous encouragement to complete the project.

## Financial support and sponsorship

National Health Mission, Government of Himachal.

## Conflicts of interest

There are no conflicts of interest.

## References

1. Weatherall D, Akinyanju O, Fucharoen S, *et al.* Inherited Disorders of Hemoglobin. In: Jamison DT, Breman JG, Measham AR, *et al.*, editors. Disease Control Priorities in Developing Countries. 2<sup>nd</sup> ed. Ch 34. Washington (DC): The International Bank for Reconstruction and Development/The World Bank; 2006.
2. Grow K, Vashist M, Abrol P, Sharma S, Yadav R. Beta thalassemia in India: Current status and the challenges ahead. *Int J Pharm Pharm Sci* 2014;6:28-33.
3. Colah R, Gorakshakar A, Nadkarni A. Global burden, distribution and prevention of  $\beta$ -thalassemias and hemoglobin E disorders. *Expert Rev Hematol* 2010;3:103-17.
4. Ambekar SS, Phadke MA, Mokashi GD, Bankar MP, Khedkar VA, Venkat V, *et al.* Pattern of hemoglobinopathies in Western Maharashtra. *Indian Pediatr* 2001;38:530-4.
5. Mohanty D, Colah RB, Gorakshakar AC, Patel RZ, Master DC, Mahanta J, *et al.* Prevalence of  $\beta$ -thalassemia and other haemoglobinopathies in six cities in India: A multicentre study. *J Community Genet* 2013;4:33-42.
6. DeLong ER, DeLong DM, Clarke-Pearson DL. Comparing the areas under two or more correlated receiver operating characteristic curves: A nonparametric approach. *Biometrics* 1988;44:837-45. Available from: <https://www.ncbi.nlm.nih.gov/pubmed/3203132>. [Last accessed on 2019 Jan 21].
7. Dean AG, Sullivan KM, Soe MM. OpenEpi: Open Source Epidemiologic Statistics for Public Health, Version. Available from: <http://www.OpenEpi.com>. [Last updated on 2013 Apr 06; Last accessed on 2018 Sep 05].
8. Mathew A, Sobti PC. The burden of thalassemia in Punjab: A roadmap forward. *Pediatr Hematol Oncol J* 2017;2:85-7. Available from: <https://doi.org/10.1016/j.phoj.2018.01.001>.
9. Arora S, Rana D, Raychaudhuri S, Dhupia JS. Coexistence of iron deficiency and thalassemia trait: A study in antenatal females. *Int J Res Med Sci* 2017;5:5362-6.
10. Mandrekar JN. Receiver operating characteristic curve in diagnostic test assessment. *J Thorac Oncol* 2010;5:1315-6.
11. Bahar R, Shahida NS, Nazri MH, Marini R, Noor Haslina MN, Shafini MY, *et al.* The diagnosis of beta thalassemia with borderline HbA2 level among Kelantan Population. *J Blood Disord Transfus* 2017;8:396. Available from: <https://www.omicsonline.org/open-access/the-diagnosis-of-beta-thalassemia-with-borderline-hba2-level-among-kelantan-population-2155-9864-1000396-97024.html#citation-btn>.
12. Al-Amodi AM, Ghanem NZ, Aldakeel SA, Ibrahim Al Asoom L, Rafique Ahmed N, Almandil NB, *et al.* Hemoglobin A2 (HbA2) has a measure of unreliability in diagnosing  $\beta$ -thalassemia trait ( $\beta$ -TT). *Curr Med Res Opin* 2018;34:945-51.
13. Khara R, Singh T, Khuana N, Gupta N, Dubey AP. HPLC in characterization of hemoglobin profile in thalassemia syndromes and hemoglobinopathies: A clinicohematological correlation. *Indian J Hematol Blood Transfus* 2015;31:110-5.
14. Rathod DA, Kaur A, Patel V, Patel K, Kabrawala R, Patel V, *et al.* Usefulness of cell counter-based parameters and formulas in detection of beta-thalassemia trait in areas of high prevalence. *Am J Clin Pathol* 2007;128:585-9.
15. Soliman AR, Kamal G, Walaa AE, Sallam Mohamed TH. Blood indices to differentiate between  $\beta$ -thalassemia trait and iron deficiency anemia in adult healthy Egyptian blood donors. *Egypt J Haematol* 2014;39:91-7.
16. Adlekha S, Chadha T, Jaiswal RM, Singla A. Screening of  $\beta$ -thalassemia trait by means of red cell indices and derived formulae. *Med J DY Patil Univ* 2013;6:71-4.
17. Pranpanus S, Sirichotiyakul S, Srisupundit K, Tongsong T. Sensitivity and specificity of mean corpuscular hemoglobin (MCH): For screening alpha-thalassemia-1 trait and beta-thalassemia trait. *J Med Assoc Thai* 2009;92:739-43.
18. Mendiratta SL, Bajaj S, Popli S, Singh S. Screening of women in the antenatal period for thalassemia carrier status: Comparison of NESTROFT, red cell indices, and HPLC analysis. *J Fetal Med* 2015;2:21.
19. Phillips PK, Bebbington C. The pH, conductivity and osmolality of low ionic strength solutions used within the U.K. for the antiglobulin test. *Transfus Med* 1991;1:155-8.
20. Manglani M, Lokeshwar MR, Vani VG, Bhatia N, Mhaskar V. 'NESTROFT' – An effective screening test for beta thalassemia trait. *Indian Pediatr* 1997;34:702-7.
21. Shewale SP, Meshram DP, Sameer MA, Deshpande SA, Sadhu D, Dager V. Study of effectiveness of NESTROFT and solubility test as a screening test for the detection of hemoglobin disorder at Nanded region of Maharashtra. *Int J Health Sci Res* 2014;4:9. Available from: [http://www.ijhsr.org/IJHSR\\_Vol.4\\_Issue.9\\_Sep2014/9.pdf](http://www.ijhsr.org/IJHSR_Vol.4_Issue.9_Sep2014/9.pdf).
22. Sharma P, Das R, Trehan A, Bansal D, Chhabra S, Kaur J, *et al.* Impact of iron deficiency on hemoglobin A2% in obligate  $\beta$ -thalassemia heterozygotes. *Int J Lab Hematol* 2015;37:105-11.
23. National Guidelines on for Prevention and Control of Hemoglobinopathies in India. Available from: [https://www.nhm.assam.gov.in/sites/default/files/swf\\_utility\\_folder/departments/nhm\\_lipl\\_in\\_oid\\_6/menu/schemes/Guidelines\\_on\\_Hemoglobinopathies\\_in%20India.pdf](https://www.nhm.assam.gov.in/sites/default/files/swf_utility_folder/departments/nhm_lipl_in_oid_6/menu/schemes/Guidelines_on_Hemoglobinopathies_in%20India.pdf). [Last accessed on 2018 Dec 23].
24. Mallik S, Chatterjee C, Mandal PK, Sardar JC, Ghosh P, Manna N, *et al.* Expenditure to treat thalassemia: An experience at a tertiary care hospital in India. *Iran J Public Health* 2010;39:78-84.
25. Gharaibeh H, Barqawi MA, Al-Awamreh K, Al Bashtawy M. Clinical burdens of  $\beta$ -thalassemia major in affected children. *J Pediatr Hematol Oncol* 2018;40:182-7.
26. Kantharaj A, Chandrashekar S. Coping with the burden of thalassemia: Aiming for a thalassemia free world. *Glob J Transfus Med [Serial online]* 2018;3:1-5. Available from: <http://www.gjtonline.com/text.asp?2018/3/1/1/229328>. [Last Accessed on 2019 Jan 24].
27. Koren A, Profeta L, Zalman L, Palmor H, Levin C, Zamir RB, *et al.* Prevention of  $\beta$  Thalassemia in Northern Israel - A Cost-Benefit Analysis; *Mediterr J Hematol Infect Dis* 2014;6. Available from: <http://mjhid.org/index.php/mjhid/article/view/2014.012/pdf>. [Last accessed on 2019 Jan 24].