Modification of Hemoglobin H Disease by Sickle Trait

KATHERINE K. MATTHAY, WILLIAM C. MENTZER, JR., ANDREE M. DOZY, YUET WAI KAN, and DOROTHY F. BAINTON, Departments of Pediatrics, Medicine, and Pathology, University of California, San Francisco, San Francisco, California 94143

ABSTRACT The rarity of hemoglobin (Hb) H disease in combination with sickle trait may be due in part to the absence of actual Hb H in individuals who, nonetheless, have inherited the deletion of three α-globin genes. We describe here a boy with persistent microcytic, hypochromic anemia despite adequate iron stores, who exhibited splenomegaly with a normal reticulocyte count and only rare inclusions in circulating erythrocytes. Starch gel electrophoresis and isoelectric focusing at age 5 yr showed 21% Hb S, persistent Hb Bart’s, but no Hb H. Recticulocyte α/non-α globin chain synthesis ratio was 0.58 at age 5. The mother (Asian) had laboratory evidence of α-thalassemia trait and the father (Black) had sickle trait. The nature of α-thalassemia in this patient was investigated both by liquid hybridization and by the Southern method of gene mapping, in which DNA is digested with restriction endonucleases and the DNA fragments that contained the α-globin structural gene identified by hybridization with complementary DNA. The patient had only one α-globin structural gene, located in a DNA fragment shorter than that found in normal or α-thalassemia trait individuals, but similar to that present in other patients with Hb H disease. Morphologic studies of bone marrow by light and electron microscopy revealed erythroid hyperplasia with inclusions in polymorphic and orthochromatic erythroblasts, suggesting early precipitation of an unstable hemoglobin. The lack of demonstrable Hb H may be the result of both diminished amounts of β^a available for Hb H formation (since one β-globin gene is β^s) and the greater affinity of α-chains for β^a than β^s-globin chains leading to the formation of relatively more Hb A than Hb S. The presence of a β^s gene may thus modify the usual clinical expression of Hb H disease.

INTRODUCTION

Simultaneous inheritance of α-thalassemia and structural abnormalities of the β-chain frequently modifies the usual clinical expression of the β-globin mutation (1–13). For example, observation of the nature of sickle cell anemia in several individuals who have also inherited α-thalassemia trait has suggested in some cases (4–6), but not all (7, 8), a favorable effect of the α-thalassemia gene on the clinical severity of sickle cell anemia. α-Thalassemia with sickle cell trait consistently results in a lower than usual percentage of hemoglobin (Hb)S (9–13). We have investigated a child with sickle trait and microcytic anemia whose Asian mother had α-thalassemia trait and whose Black father had sickle trait. The very low percentage of Hb S suggested that both α-thalassemia trait and sickle cell trait were present, but the presence of splenomegaly, extreme microcytosis, and bizarre erythrocyte morphology, features not seen in α-thalassemia trait, stimulated further investigation.

Analysis of DNA from this child by molecular hybridization and gene-mapping techniques revealed that three of the four α-globin genes normally found were deleted, thus unequivocally demonstrating the presence of the Hb H genotype. Paradoxically, Hb H was not identified in hemolsate from either peripheral blood or bone marrow, although inclusion bodies could be found in occasional erythrocytes in both places. The biochemical nature of these inclusions was not determined, but the appearance in early erythroid progenitors suggested a greater degree of instability than that associated with Hb H (14–16). This patient, who represents the unusual coincidence of Hb H disease with sickle trait, demonstrates how the inheritance of a β-globin structural mutation can modify the phenotype of Hb H disease sufficiently to alter the usual clinical
picture. In such patients, accurate genetic analysis is impossible unless the new techniques of molecular hybridization and gene mapping are used.

METHODS

Hematologic values. Hemoglobin levels and erythrocyte indices were obtained with a model S Coulter counter (Coulter Electronics Inc., Hialeah, Fla.). Bone marrow differential counts were performed by counting at least 500 cells. Cellulose acetate electrophoresis was performed in Tris-EDTA-borate buffer at pH 8.4 (17). Hb A2 was determined by microcrystallography (18). Fetal hemoglobin was determined by alkali denaturation (19). Hb H preparations were made by mixing one part of fresh whole blood with two parts of 1% brilliant cresyl blue (BCB)1 in 0.9% NaCl and incubating at 37°C for 1–3 h.

Hemoglobin and globin analyses. Globin chain synthesis was measured in vitro with fresh heparinized peripheral blood or bone marrow, as described by Kan et al. (20). L-[3H]leucine-labeled globin chains precipitated from cell-free hemolysate were separated on carboxymethyl cellulose columns in 8 M urea at pH 7.2 using a linear Na2HPO4 gradient. The relative synthesis of α-chain to β- and γ-chain was expressed as the ratio of the total radioactivity of the α- to non-α-globin peaks. Starch gel electrophoresis was carried out at 4°C for 3 h at 25 mA and 200 V according to the method of Smithies (21) with 0.006 M Na2HPO4 buffer for the gel and 0.04 M Na2HPO4 buffer for the electrodes, both at pH 7.0. The gel was then sliced horizontally and stained with benzidine.

Thin-layer isoelectric focusing in polyacrylamide gel was carried out with fresh hemolysate according to Basset et al. (22). Hemoglobin column chromatography was performed on carboxymethyl cellulose (CM-52, Whatman, Inc., Clifton, N. J.) according to Schroeder et al. (23). 1-ml of stroma-free hemolysate was prepared from fresh peripheral blood or bone marrow and then dialyzed for 18 h against 0.01 M Bis-tris [N,N-Bis-(2-hydroxymethyl)-iminotris-(hydroxymethyl)-methane], pH 6.1, plus 0.01% KCN. 50–60 mg of dialyzed hemoglobin was applied to a 0.9 × 20-cm column equilibrated with the dialysis buffer. The hemoglobin was eluted with 0.03 M Bis-tris and 0.01% KCN, pH 6.1, using a linear salt gradient (0–0.08 M NaCl, total volume, 1,200 ml) over 24 h. The position of each eluted hemoglobin type was determined by recording the optical density at 415 nm of each fraction as a function of salt concentration, measured from conductance. To identify each hemoglobin eluted, two or three fractions from each peak were then pooled and centrifuged for 18 h at 50,000 rpm in a Beckman ultracentrifuge (Beckman Instruments, Inc., Spinco Div., Palo Alto, Calif.) to concentrate the hemoglobin into a 0.5-ml vol. Globin was then prepared from each concentrated portion by acid-acetone precipitation, lyophilized, then resuspended and electrophoresed on cellulose acetate in Bis-tris-EDTA-borate-urea buffer at pH 6.4 according to the method of Salmon et al. (24). The strips were stained with Ponceau S and decolorized with 3% acetic acid.

Electron microscopy. Specimens of bone marrow and blood were immediately fixed in 1.5% buffered glutaraldehyde at 22°C, postfixed in OsO4, and processed for examination by transmission electron microscopy, as described (25). Other samples were prepared for scanning electron microscopy by critical-point drying and coated with gold-platinum.

1 Abbreviations used in this paper: BCB, brilliant cresyl blue; cDNA, complementary DNA; kb, kilobase pairs; Hb, hemoglobin.

**Liquid hybridization.** DNA from the patient, his mother, and controls with hydrops fetalis, Hb H disease, α-thalassemia trait, and normals was prepared from leukocytes, liver, or spleen by sodium dodecyl sulfate-pronase digestion and phenol extraction. The RNA was digested with ribonuclease and the DNA reduced to 2 × 106 daltons by limited depurination (26, 27). α-globin complementary DNA (cDNA) was prepared as described (26, 28). The α-cDNA (1,000 cpm) was incubated for 76 h at 78°C in duplicate 20-μl reaction mixtures containing 100 μg cellular DNA, 8 pg α-cDNA, 500 cpm of [3H]dCTP-labeled unique sequence HeLa DNA as internal control, 0.5 M NaCl, 0.002 M EDTA, and 0.04 M Tris-HCl (pH 7.4). The percentage of α-cDNA annealed was assayed by batchwise elution with hydroxylapatite (29).

**Restriction endonuclease analysis of DNA.** DNA was prepared from the leukocytes of the patient, his mother, α-thalassemia trait, silent carrier, Hb H disease, and normal controls, as described (30). 10 μg of human DNA and 1 μg of λ-DNA (as internal size marker) were digested for 4 h at 37°C with 1.25 U of Eco RI or Hpa I per microgram of DNA. The buffer for Eco RI digestion contained 100 mM Tris-HCl (pH 7.5), 50 mM NaCl, 6 mM MgCl2, and 6 mM 2-mercaptoethanol, and for Hpa I digestion, 6 mM Tris-HCl (pH 7.5), 26 mM NaCl, 6 mM MgCl2, and 6 mM 2-mercaptoethanol. The samples were precipitated in alcohol, dried and resuspended in 30 μl of 5 mM Tris (pH 7.5), 0.1 mM EDTA. The digested DNA's were electrophoresed in 0.8% agarose and then transferred to nitrocellulose filters (31). The filters were hybridized with λP-labeled α- and β-globin cDNA (2 × 106 cpm/μg) for 2 d and extensively washed and autoradiographed, as described (30).

**RESULTS**

**Case report.** S.H. is a 5/12-yr-old Black/Chinese male with marked microcytic anemia in the presence of adequate iron stores. Physical examination has been notable for persistent splenomegaly. Hematologic data obtained over a 5-yr period of observation are shown in Table I. Erythrocyte morphology has consistently resembled that of Hb H disease, with hypochromia, poikilocytosis, fragments, spherocytes, and target cells evident on the peripheral blood smear (Fig. 1A and B). BCB stains revealed no Hb H inclusions in the circulating erythrocytes at age 1 or 2 yr. An occasional inclusion-containing cell was evident at age 5 yr (Fig. 1C and D). Hemoglobin electrophoresis has been consistent with sickle trait, but with an unusually low percentage of Hb S; persistence of Hb Bart's and elevated Hb F (Table I). Bone marrow aspirate and biopsy showed erythroid hyperplasia (myeloid:erythroid ratio, 1:2), a finding in apparent contradiction to the repeatedly normal reticulocyte count. The percentage of early erythroid precursors (proerythroblasts, 5%; basophilic erythroblasts, 22%) was markedly increased, while polychromatophilic and orthochromatic erythroblasts were present in normal numbers. Methyl violet and BCB stains of the bone marrow showed the same frequency of hemoglobin inclusions as in the peripheral blood. Family studies confirmed that the Black
father had sickle trait, whereas the Asian mother had 
the hematologic features of a-thalassemia trait (Table I).

**Globin chain synthesis.** The reduced $\alpha/(\beta + \gamma)$
globin chain synthetic ratios in peripheral blood reticulocytes and bone marrow of the patient, shown in Table I, 
were consistent with either a-thalassemia trait or 
Hb H disease (32). $\beta^A:\beta^S$ activity ratios were 6:1, 4:1, 
and 3:1 at 10, 31, and 60 mo, respectively. The bone 
marrow $\beta^A:\beta^S$ activity (2:1) was more nearly normal. 
The mother’s peripheral blood $\alpha/\beta$-globin chain syn-
thetic ratio was 0.88, consistent with a-thalassemia 
trait.

**Hemoglobin analyses.** Starch gel electrophoresis of 
hemolysate obtained from peripheral blood, shown in 
Fig. 2, revealed a trace amount of Hb Bart’s. No Hb H 
or other abnormal fast hemoglobins were found. Thin-
layer isoelectric focusing also showed only hemoglo-
bins Bart’s, A, F, S, and $\alpha_2$. Column chromatography, 
(Fig. 3) also failed to detect measurable amounts of 
Hb H or other fast hemoglobins, except for Hb Bart’s. 
Globin electrophoresis was used to confirm the identity 
of each hemoglobin eluted. The first peak eluted con-
tained $\gamma$-chains; the second, $\alpha$- and $\gamma$-chains; the small 
shoulder and the third peak contained $\alpha$ and $\beta^A$; the 
fourth, $\alpha$ and $\delta$; and the last peak, $\alpha$ and $\beta^S$. $\beta^A$-chains 
were found only in the Hb S fraction, ruling out the 
possibility that $\beta^A$ tetramers were present elsewhere, 
but hidden because they eluted with another hemoglo-
bin. Only the Hb Bart’s fraction was composed of a 
single globin chain, which migrated to the $\gamma$ position, 
as expected. Hemoglobin analysis of a bone marrow 
lysate by starch gel and column chromatography 
gave virtually identical results to those found in peripheral 
blood. The bone marrow percentages were Hb Bart’s, 1.4%; 
Hb F, 6.1%; Hb A, 72.5%; Hb $\alpha_2$, 3.6%; 
and Hb S, 16.4%.

**Electron microscopy of bone marrow.** Numerous 
dense amorphous inclusions of varying sizes, compara-
tible to those seen in the bone marrow of patients with 
$\beta$-thalassemia major (33, 34) were seen in reticulocytes 
and orthochromatic (Fig. 4A) and polychromatic erythroblasts (Fig. 4B). Although these were generally 
not membrane-bound (Fig. 4A), in a few instances they 
were contained within autophagic vacuoles (Fig. 4B), 
as has also been described in $\beta$-thalassemia (33, 35). 
The frequency of inclusions corresponded to that seen 
on methyl violet stain. Many of the mature erythro-
ocytes appeared very thin and distorted, as can be seen 
in the scanning electron micrographs (Fig. 4C and D), 
because of poor hemoglobinization, a common finding

<table>
<thead>
<tr>
<th>TABLE I</th>
<th>Hematologic Data in Hb H Disease with Sickle Trait</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patient</td>
<td>10 mo</td>
</tr>
<tr>
<td>---------</td>
<td>-------</td>
</tr>
<tr>
<td>Hb, g/dl</td>
<td>9.4</td>
</tr>
<tr>
<td>PCV</td>
<td>0.29</td>
</tr>
<tr>
<td>MCH, pg</td>
<td>16.9</td>
</tr>
<tr>
<td>MCHC, g/dl</td>
<td>33.3</td>
</tr>
<tr>
<td>MCV, fl</td>
<td>50</td>
</tr>
<tr>
<td>Erythrocyte, $\times 10^{12}$/liter</td>
<td>5.67</td>
</tr>
<tr>
<td>Retic, %</td>
<td>0.6</td>
</tr>
<tr>
<td>Fe/TIBC, $\mu$m/liter</td>
<td>18.5/80.8</td>
</tr>
</tbody>
</table>

Abbreviations used in this table: PCV, packed cell volume or hematocrit; MCH, mean corpuscular hemoglobin; MCHC, MCH concentration; MCV, mean corpuscular volume; TIBC, total iron binding capacity.

* Relative hemoglobin percentages determined by densitometric scan of cellulose acetate strip, as well as separate quantitation of Hb $\alpha_2$ and Hb F.

† Peripheral blood.

§ Bone marrow.
in thalassemia (34). Occasionally small pits (Fig. 4C) could be detected. Similar pitting of erythrocytes has previously been noted with Nomarski optics in individuals with Hb H disease, even those with intact spleens (14).

\( \alpha \)-Globin–cDNA liquid hybridization. Because hematological, family, and globin chain synthesis studies were inconclusive, additional studies were done to analyze the \( \alpha \)-globin genes directly. Fig. 5 shows the percent annealing of \( ^{32} \)P-labeled \( \alpha \)-globin cDNA to DNA from the propositus, his mother, and controls. DNA from the patient showed the same percent annealing as six Asian Hb H disease controls, who lack three of four \( \alpha \)-globin genes. His mother showed the same percentage of \( \alpha \)-sequences as other \( \alpha \)-thalassemia trait individuals with a deletion of two of the four \( \alpha \)-globin genes.

Restriction endonuclease analysis of DNA. Figs. 6 and 7 show the diagrammatic and autoradiographic results of DNA digested with Eco RI and Hpa I and hybridized with cDNA. The \( \alpha \)-globin specific fragment in DNA from the patient was smaller than normal after Eco RI digestion and was identical to the 19-kilobase pairs (kb) \( \alpha \)-specific DNA of known Hb H disease controls. The patient was lacking the 4.0-kb fragment after Hpa I digestion, one of the two \( \alpha \)-specific fragments normally present, again corresponding to known Hb H disease controls. The autoradiogram of hybridized \( \alpha \)-globin DNA from the patient’s mother was identical to the pattern of controls with \( \alpha \)-thalassemia trait. The patient’s father was not available for testing.
A more precise definition of the nature of the thalassemia defect was obtained by molecular hybridization studies. Liquid hybridization with α-globin cDNA was clearly compatible with deletion of three α-globin genes, the typical genotype of Asian Hb H disease (27). Restriction endonuclease gene-mapping techniques also clearly delineated the presence of only a single α-globin gene. It is important to note that α-thalassemia trait, with deletion of the two α-globin genes in trans, also results in a restriction endonuclease digest of two identical, but smaller than normal, 19 kb α-globin DNA fragments, like the Eco RI pattern seen in the patient and in other individuals with Hb H disease (39). However, the patient could not have inherited α-thalassemia trait in trans, since the DNA digest from the mother showed only a normal size 23 kb α-globin DNA fragment, consistent with cis-α-thalassemia trait, where one chromosome has two intact α-globin genes and the other is devoid of α-globin genes. The patient, therefore, inherited his single intact α-globin gene from his Black father, who presumably was a silent carrier of α-thalassemia (genotype, α/αα). The silent carrier state is actually quite common in Blacks, with a frequency of 24% by restriction endonuclease gene mapping (40).

Hb H disease has never before been reported in a patient with sickle trait. Although the incidence of Hb H disease in Blacks is admittedly low (40–42), the apparent rarity of the combined defect may also be a result of the lack of a readily available means of detection, since Hb H is not always present in measurable amounts. Although other patients with a β-chain abnormality and a suspected Hb H genotype, but without β^2^ tetramers have been described (1, 5, 14, 43–46), direct demonstration of the α-globin gene defect has not previously been possible.

There are several possible reasons that Hb H disease, when inherited with the β-gene mutation of sickle trait, may produce a syndrome of hemolytic anemia without detectable β-globin tetramers. One is the decreased quantity of β^A^ globin chains available for Hb H formation, since only one β-globin gene is actually synthesizing such chains. Furthermore, α-globin chains have a greater affinity for β^A^ than for β^A^- chains. Thus, Hb A is formed more readily than Hb S, leaving excess free β^A^-chains whenever there is a relative deficit of α chains. Such a differential affinity of α-chains for β^A^-chains is suggested by the lower than expected percentage of Hb S seen in patients with α-thalassemia trait and with sickle trait (9–13). In vitro studies of the reassociation of α-chains with β^A^- and β^-chains show diminished formation of Hb S when decreased amounts of α-globin chains are present (47, 48).

Despite the lack of detectable Hb H, our patient had evidence of erythrocyte destruction with frag-
FIGURE 4 Transmission electron micrographs of developing erythroblasts from the bone marrow. (A) is an orthochromatic erythroblast and its cytoplasm contains abnormal amorphous inclusions (arrow). The nucleus and other organelles such as the Golgi complex (G), mitochondria (m), and polynribosomes (p) appear normal. ×15,000. (B) is a polychromatic erythroblast which also contains similar inclusions but within a membrane-bound organelle, probably an autophagic vacuole (AV). ×18,000. (C and D) Scanning electron micrographs of circulating blood erythrocytes. The cells are thinner than normal and there are many irregularities in shape. Note the knob-like projections in D. Occasional small pits (arrow) are evident. C, ×8,000; D, ×7,200.
The normal DNA has two α-specific fragments of 4.0 and 14.5 kb. The other fragments have been shown to be γ-, β-, and β-specific (30). The hydrops DNA lacks both α-specific fragments. The patient’s DNA lacks the smaller 4.0 kb α-specific fragment as does the Hb H control. (The single α-specific fragment in the patient’s DNA is larger than the usual 14.5 kb fragment, but this corresponds to a variant α-globin DNA fragment seen in some normal individuals without Hb H disease, probably because of polymorphism of the α-gene (Kan, Y. W., unpublished data). The very faint band appearing in the 14.5-kb position is not α-specific and was not present in a subsequent Hpa I digest of the patient’s DNA.) The patient’s DNA also contains a 13.0-kb β-specific fragment, which occurs in 80% of individuals with sickle hemoglobin (30). The mother’s DNA has both α-specific fragments, as seen in α-thalassemia trait.

tetramers that then precipitate rapidly, resulting in loss of erythroid precursors in the bone marrow and destruction of precipitate-containing peripheral blood cells by the spleen. Alternatively, the excess β⁰-chain might bind to the erythrocyte membrane, causing damage and subsequent hemolysis, since preferential binding of β⁰ to stroma has been demonstrated (49). The presence of some type of unstable hemoglobin was indicated by the occasional inclusion-containing cells seen in both peripheral blood and bone marrow by special stains and electron microscopy. The electrophoretic and chromatographic techniques employed are not sufficiently sensitive to unequivocally rule out the possibility that these occasional inclusions are composed of Hb H. However, the distribution of these inclusions was not typical of Hb H disease. β² tetramers precipitate in mature erythroblasts and in peripheral erythrocytes (15) causing membrane pitting and rigidity with gradual extramedullary hemolysis, rather than ineffect-

**Figure 5** Liquid hybridization with [α-³²P]globin cDNA.

**Figure 6** (A) Diagram representing the Eco RI restriction endonuclease cleavage sites of α-globin DNA in the normal individual and a person with Hb H disease. (B) Autoradiogram of Eco RI digestion pattern of DNA from patient, mother, and appropriate controls, using [α-³²P]globin cDNA as a probe. The normal DNA shows a single 23 kb α-specific fragment. The patient has a smaller 19 kb fragment, identical to the Hb H control. The DNA of the mother shows the 23-kb fragment seen in the α-thalassemia trait control, who has two α-globin genes in cis on one chromosome, the other chromosome being devoid of α-genes. The hydrops DNA has no α-specific fragment, while the silent carrier of α-thalassemia, with three out of four intact α-globin genes, has both a normal 23 kb and a small 19 kb fragment.

**Figure 7** (A) Diagram representing Hpa I restriction endonuclease cleavage sites of α-globin DNA in the normal and the individual with Hb H disease. (B) Autoradiogram of Hpa I digestion in DNA from the patient, his mother, and controls using both α- and β-cDNA as probes. The normal DNA has two α-specific fragments of 4.0 and 14.5 kb. The other fragments have been shown to be γ-, δ-, and β-specific (30). The hydrops DNA lacks both α-specific fragments. The patient’s DNA lacks the smaller 4.0 kb α-specific fragment as does the Hb H control. (The single α-specific fragment in the patient’s DNA is larger than the usual 14.5 kb fragment, but this corresponds to a variant α-globin DNA fragment seen in some normal individuals without Hb H disease, probably because of polymorphism of the α-gene (Kan, Y. W., unpublished data). The very faint band appearing in the 14.5-kb position is not α-specific and was not present in a subsequent Hpa I digest of the patient’s DNA.) The patient’s DNA also contains a 13.0-kb β-specific fragment, which occurs in 80% of individuals with sickle hemoglobin (30). The mother’s DNA has both α-specific fragments, as seen in α-thalassemia trait.
tive erythropoiesis. On the other hand, β-thalassemia is characterized by extensive ineffective erythropoiesis (14). Inclusions, presumably a result of unstable α-chains, can be seen in the bone marrow in immature erythroblasts with methyl violet stains (50) or electron microscopy (14), but do not appear in peripheral blood except after splenectomy. The distribution of inclusions in our patient to some degree resembles both β-thalassemia and Hb H disease. As in β-thalassemia, some inclusions are seen in the younger polychromatophilic erythroblasts, and may be responsible for rapid intramedullary hemolysis, as indicated by the large proportion of immature erythrocyte forms in the face of a low reticulocyte count. However, a few cells containing unstable hemoglobin do reach the peripheral blood (Fig. 1), where they are presumably destroyed in the spleen.

ACKNOWLEDGMENTS

We thank Dr. J. Rosa, Hopital Henri Mondor, Creteil, France, for kindly performing the thin-layer isoelectric focusing and Mr. Rick Trecartin for performing the globin chain synthesis studies and assisting with the column chromatography. We also gratefully acknowledge the secretarial assistance of Ms. Terry Tschopp.

This investigation was supported by grants from the National Institutes of Health (HL20985), the U. S. Public Health Service (5 K04 HL00086), and the National Foundation—March of Dimes (6-49).

REFERENCES


