Oxygen Equilibrium of Hemoglobin J Cape Town

SAMUEL CHARACHE and TREPOR JENKINS with the technical assistance of
MELODY F. WILDER

From the Department of Medicine, Johns Hopkins University School of
Medicine, Baltimore, Maryland 21205, and the Human Sero-Genetics Unit,
South African Institute for Medical Research, Johannesburg

ABSTRACT Polycythemia in carriers of hemoglobin J Cape Town or hemoglobin Chesapeake is thought to be produced by increased oxygen affinity of their blood. Both hemoglobins involve substitution of amino acid residue α FG-4. Measurements reported here, of the oxygen equilibrium of purified hemoglobin J Cape Town, permit direct comparison of the two hemoglobins. J Cape Town exhibits lower oxygen affinity, and greater heme-heme interaction, than Chesapeake; both exhibit normal Bohr effects. Substitution of one polar amino acid residue for another of opposite charge (arginine → glutamic acid) thus appears to create less disruption of the interface between α- and β-chains than substitution of a nonpolar residue (arginine → leucine).

INTRODUCTION

Some, but not all carriers of hemoglobin J Cape Town (Hb J), (a92(FG-4) Glu) are polycythemic (1, 2). Oxygen affinity of a hemolysate containing 35% Hb J was increased, while the Bohr effect was normal, suggesting that polycythemia was produced by decreased delivery of oxygen to tissues (3). Heme-heme interaction in Hb J was significantly decreased, with an n value of 1.8. These findings were of great interest, for hemoglobin Chesapeake (a92 Leu), which also is produced by a mutation at FG-4, exhibits increased oxygen affinity and decreased heme-heme interaction (n = 1.3) (4). We report here data on the oxygen equilibrium of purified Hb J, studied under the conditions used in previous studies of hemoglobin Chesapeake.

METHOD

Blood was collected in Johannesburg in ACD (acid citrate dextrose) solution under sterile conditions. It was shipped in ice to Baltimore and arrived in excellent condition. A portion of the blood was used for studies of ligand binding and mobility in the ultracentrifuge. Portions of the remainder were removed over the next 2 wk, maintaining sterility at all times. Red cells were washed three times, hemolysates were prepared with toluene and distilled water, and the hemoglobin components were separated by starch block electrophoresis at 4°C (5). After electrophoresis, the eluted hemoglobin solution was passed through a 4 x 40 cm column of Sephadex G-25, which had been equilibrated with 0.1M phosphate buffer. Samples were adjusted to a concentration of 0.1 g/100 ml, and oxygen affinity was measured at 10°C, using a modification of the techniques of Allen, Guthe, and Wyman (6) and Riggs (7). Methemoglobin formation was monitored by measurement of optical density at 630 nm; the only samples (pH 6.8) which showed more than a very slight increase in OD were discarded.

RESULTS

Oxygen affinity of Hb J was increased, and a normal Bohr effect was present (Fig. 1). Heme-heme interaction was decreased: the mean value of n from nine series of measurements, calculated by the method of least squares, was 2.22 (SD 0.20). Data for Hb J are compared with data for normal hemoglobin and Hb Chesapeake in Fig. 2: Hb Chesapeake has a lower p50 and a lower value of n than Hb J.

DISCUSSION

Heme-heme interaction of purified Hb J was not decreased to the degree found by Lines and McIntosh in a hemolysate of whole blood (3). The difference between their results and ours may be due to graphical summation of the properties of two hemoglobins with different affinities for oxygen: the true dissociation curve must be sinusoidal, since Hb J plays a disproportionate role in oxygen binding at low oxygen tensions. Effects of the two hemoglobins might be resolved using a continuously recording oxygen electrode and linear coordinates (8); the small departure from linearity would not be noted when the logarithmic form of Hill’s equation is utilized.
FIGURE 1 Bohr effect in hemoglobins A- and J-Cape Town. Each point represents the oxygen pressure required for half saturation of hemoglobin at 10°C in 0.1M PO2 buffer at a given pH. Points were derived from lines plotted according to Hill’s equation \((\log S/1-S = n \log P_0 - n \log p_{50})\), where \(S\) is per cent saturation, \(K\) is a constant, and \(n\) is related to the magnitude of heme-heme interaction. The oxygen affinity of Hb J is higher than that of Hb A, but the magnitude of the Bohr effect appears to be normal.

All abnormal hemoglobins with amino acid substitutions at the \(\alpha\beta\) interface between globin subunits have abnormal oxygen affinity, and Hb J is no exception [(9) and reference cited therein]. Analyses of X-ray diffraction data suggest that there are many differences between the structure of oxyhemoglobins A and Chesapeake (10). The oxygen equilibrium of Hb J is less abnormal than that of Hb Chesapeake, as are a number of physical properties: electron-spin resonance spectra (11), reactivity with ligands, and sedimentation in the ultracentrifuge. Substitution of a polar residue by another of opposite charge (arg \(\rightarrow\) glu) thus appears to create less disruption of the \(\alpha\beta\) interface than does substitution by a nonpolar residue (arg \(\rightarrow\) leu). Polycythemia produced by Hb J is less impressive than that produced by Hb Chesapeake, as might be expected from these findings (1, 2, 12).

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REFERENCES


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