Reaction of 1-Fluoro-2,4-dinitrobenzene with the Free α Chains of Human Hemoglobin

EVALUATION OF THE pK OF THE TERMINAL AMINO GROUP*

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SUMMARY

The reaction of 1-fluoro-2,4-dinitrobenzene with isolated α chains in their mercaptide form with p-mercuribenzoate (PMB) was studied. From kinetic measurements with the change in absorbance at 353 nm upon substitution of the terminal valine, the pK of this group was calculated to be 7.30. The pH-independent reaction constant was estimated to be 0.282 m⁻¹ sec⁻¹. Comparing the proton-binding behavior of untreated α chains with that of the chains substituted in the terminal valine (αPMBDNP), the pK of this group was calculated to be 7.4. Histidines did not appear to be dinitrophenylated. However, from the rate of liberation of protons upon the reaction of the chains with 1-fluoro-2,4-dinitrobenzene we concluded that they do interact with the reagent without forming a stable derivative. A pK of 6.85 and a pH-independent reaction constant of 0.023 m⁻¹ sec⁻¹ per individual group was calculated for these residues. From oxygen equilibrium studies on a mixture in equal amount of αPMBDNP and β chains in their mercaptide form with p-mercuribenzoate we calculated a value of 1.35 for the Hill parameter n. The s10, w of this mixture was 4.95 while it is near 2.5 when the α chains are not dinitrophenylated (Antonini et al., J. Mol. Biol., 17, 29 (1966)), indicating that the dinitrophenylation strongly favors the formation of tetramers. This high sedimentation value is consistent with a partial specific volume $\bar{v} = 0.731$, as compared to $\bar{v} = 0.749$ commonly used for hemoglobin (Svedberg and Pedersen, The ultracentrifuge, Oxford University Press, New York, 1940).

A determination of this pK by the specific method as done by Hill and Davis (1) for human carboxyhemoglobin has not yet been carried out for the free α chains. In proton-binding studies of these chains, the pK of the terminal amino group was assumed to be near 8 (4, 5). However, Janssen, de Bruin, and van Os (5) pointed out that the large rotational freedom present for this group in the crystals of carboxyhemoglobin (6) might suggest that its pK is also low in the free α chains, where its rotational freedom should be at least as great.

A study of the pK of the NH₂-terminal group in the free chains is of particular importance because this group is considered to be one of the alkaline Bohr groups (1, 2, 6, 7). The pK shift upon oxygenation is thought to result from a change in interaction between this group and the carboxyl terminal of its partner α chain (6).

This paper presents a study of the reaction between the free chains and FDNB. The rate of reaction was measured by following the change in absorbance (8) at 353 nm or by measuring the release of protons with time upon reaction.

The proton-binding behavior of both unsubstituted and fully substituted α chains was consistent with the kinetic measurements, indicating that the pK of the amino group is indeed fairly low (near 7.4).

EXPERIMENTAL PROCEDURE

Materials

Adult hemoglobin was prepared by the toluene procedure of Drabkin (9). The isolated α chains were prepared by the method of Bucci and Fronticelli (10). Because of their higher stability, the chains were used without removal of the p-mercuribenzoate from the $\cdot$SH groups and will be designated as αPMB. The concentration of the αPMB solutions was measured with the molar extinction coefficient $\varepsilon_{450} = 14,000$ M⁻¹ cm⁻¹ at 540 nm for the CO derivative. FDNB was obtained from Sigma and was used without further purification. For all experiments a freshly prepared saturated FDNB solution (approximately 10

\* This work was supported in part by Grants HE 13178 and HE 13164 from the National Institutes of Health, United States Public Health Service.

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mm) in 0.1 m NaCl was used. The concentration was checked with the dinitrophenylbenzylamine method of Hill and Davis (1). Carbon dioxide-free solutions of NaOH were prepared from a saturated stock solution. All other chemicals were analytical grade.

The preparation of DNP-substituted αPMB chains was carried out in the following way. Solutions of αPMB (10-4 m in heme) were reacted with FDNB (3 × 10-4 m) in 0.2 m phosphate buffer at pH 7 for 1 hour at room temperature. The unreacted FDNB was removed by passage through Sephadex G-50. The reaction mixture was separated by chromatography on carboxymethylcellulose with a linear pH and ionic strength gradient made with two cylindrical bottles containing 1 liter of 0.05 m phosphate buffer at pH 6.2 and 0.1 m phosphate buffer at pH 7.0, respectively. The first bottle was connected to the column (2.5 × 20 cm), which had been equilibrated with 0.05 m phosphate buffer at pH 6.0. The protein solution was adjusted to pH 6.0, then dialyzed against the pH 6.0 buffer before absorption on the column. A typical elution pattern is shown in Fig. 1. The leading peak contained the DNP-substituted material. (see “Results”).

Methods

Measurement of Rate of Reaction Following Change in Optical Density at 353 nm—Of a deionized αPMB solution (2 × 10-4 m), 10 ml were mixed with 5 ml of 0.2 m phosphate buffer at the desired pH and 5 ml of 10-2 m FDNB in a jacketed vessel thermostated at 25°. At regular time intervals, 2-ml samples were pipetted into test tubes containing 6 ml of 0.2 m phosphate buffer at pH 5.4 at 0° kept in an ice bucket. In this way the reaction rate was lowered about 2000 times. The excess FDNB was removed by dialyzing the samples at 4° against three changes of 2 liters of 0.02 m phosphate buffer at pH 5.4. Both FDNB and 2,4-dinitrophenol dialyzed very rapidly out of the dialysis bags. Control FDNB solutions dialyzed in this way showed a residual absorbance near 0.002, at 353 nm and were used as blanks in the spectrophotometric measurements. A Cary 14 spectrophotometer was used to measure the absorbance of the solutions.

Measurement of Rate of Reaction Following pH Changes—For these measurements the same equipment was used as described for the hydrogen ion titrations. The procedure was as follows. Of a deionized αPMB solution (2 × 10-4 m), 3 ml were put into the titration vessel and adjusted with KCl to a μm of 0.1. The desired pH was obtained by adding known amounts of 0.1 m HCl. Of a 10-2 m FDNB solution roughly adjusted with CO2-free NaOH to about the same pH, 3 ml were put into a separate vessel at the same temperature (25°). This was done to avoid pH changes upon mixing caused by differences in temperature. At zero time the FDNB solution was transferred into the titration vessel through a plastic tube by nitrogen pressure.

Hydrogen Ion Titration—The hydrogen ion titration curves were measured with roughly the same titration equipment as described by de Bruin, Janssen, and van Os (3). Instead of a digital voltmeter a Sargent-Welch recorder model SRG was used; full scale deflection (10 inches) corresponded to 0.1 or 0.2 pH unit. All protein solutions were carefully deionized by passing them through a mixed bed ion exchange column. The solution was brought to an ionic strength of 0.1 with KCl and the titration was carried out at 25° under a stream of nitrogen.

Peptide Maps—These were obtained following the method of Baglioni (11). High voltage electrophoresis was carried out with a model D Gilson electrophorator.

Quantitative Amino Acid Analyses—Quantitative amino acid analyses were performed with a Beckman model 120C amino acid analyzer and the standard 4-hour run.

Oxygen Binding—Oxygen-binding curves were measured spectrophotometrically according to the method of Rossi-Fanelli and Antonini (12).

Sedimentation Velocity—Sedimentation velocity measurements were carried out with a Beckman model E ultracentrifuge with the schlieren optics.

Treatment of Kinetic Data

Spectrophotometric Kinetics—As given by Hill and Davis (1) the rate of reaction of FDNB with the unprotonated groups of Class i can be described by

\[
\frac{d[X]}{dt} = \sum_k \kappa_k [N] [P]
\]

in which [X] represents the product, the Ni represents the concentrations of the unprotonated groups of Class i, \( \kappa_k \) the corresponding reaction constant, and P the concentration of FDNB. As discussed later only the reaction of the terminal valine with FDNB is detected spectrophotometrically, so we are dealing with only one term in Equation 1. The concentration of the unprotonated α-NH2 group is related to the total concentration, P, of the unreacted αPMB chains by

\[
[N] = \alpha [P] = \frac{K}{K + (H^+)} [P]
\]

in which \( \alpha \) is the degree of ionization of the NH2-terminal amino group, K its ionization constant, and \( (H^+) \) the proton activity. Substituting Equation 2 into Equation 1...
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**RESULTS**

Characterization of DNP-substituted \( \alpha ^{PMB} \) Chains

Optical Density Measurements—In Fig. 1 also the ratio \( R = A_{434} : A_{400} \) of the various fractions is plotted. \( R \) is 1.75 in untreated \( \alpha ^{PMB} \) chains and has a maximum value \( R_{\text{max}} = 2.88 \) in the \( \text{NH}_{2} \)-terminal substituted chains as calculated from the molar extinction coefficient \( \varepsilon_{353} = 16,200 \) of the DNP-valylleucyl peptide (8). Fig. 1 shows that \( R \) reaches a plateau value of about 2.87 in the leading peak consistent with a complete substitution of the terminal valine, and drops to a value of 1.88 in the trailing peak, indicating the presence of unsubstituted chains. The value of 1.88 is somewhat higher than the expected value (1.75) probably because the leading peak tails into this peak. It is noteworthy that only a very small amount of material had a value for \( R \) of 3.3, which indicates a minor reaction of other groups with FDNB. The difference absorption spectrum of the CO derivatives of the \( \alpha ^{PMB} \text{DNP} \) and \( \alpha ^{PMB} \) chains were obtained by adjusting the path length of a variable path length cuvette (type X DC 14 Precision Cell) until the optical density of the solutions to be compared was the same at 540 nm. From the difference spectrum, which corresponded to the spectrum of the DNP-substituted terminal peptide with a maximum at 353 nm a molar extinction coefficient for this peptide was calculated equal to 16,000 \( \text{cm}^{-1} \text{M}^{-1} \).

Peptide Analysis—Peptide maps of the material eluted with the first peak in Fig. 1 showed the absence of Peptide \( \alpha T1 \) and the presence of only one additional yellow spot with a lower electrophoretic mobility and a larger \( K_{F} \) value in chromatography than the \( \alpha T1 \) peptide. The amino acid composition of the eluted yellow peptide was that expected for the \( \alpha T1 \) peptide. The amino acid analysis of the terminal peptide DNP-valylleucyl, extracted after partial hydrolysis as described by Hill and Davis (1), showed the presence of leucine with a recovery of about 80%.

**pH Measurements.** These showed that the isoinic pH of the material of the leading and second peak was in all preparations pH 7.38 and 7.55, respectively. The latter pH is equal to the isoinic point of untreated \( \alpha ^{PMB} \) chains.

Determination of \( pK \) of \( \alpha \)-Amino Group

Hydrogen Ion Titrations—In Fig. 2 the differential titration curves for unsubstituted and fully substituted \( \alpha ^{PMB} \) chains are shown. The analysis of such curves is described by de Bruin

\[
\frac{d[H^+]}{dt} = k_r [N] \quad (6)
\]
et al. (3) de Bruin and van Os (13). It represents the slope $\Delta pK/\Delta Z_H$ of the normal $pK$ against $Z_H$ curve, in which $Z_H$ is the mean proton charge. This slope is equal to the reciprocal of the buffer capacity. Without going into the details of the analysis of such curves it is enough to mention that the distance between the two peaks shown is ionic strength independent and equal to the number of groups titrated in the neutral region. In the curve for the $aPMB$ chains this distance is 8.1, which means that in $aPMB$ 7 histidines and the $\alpha$-amino group are titratable. This result is in full agreement with earlier reports (4, 5). The peak to peak distance in DNP-substituted $aPMB$ chains is 7.2 eq. So there is one group less titratable than in the $aPMB$ chains. Considering the results obtained in the previous section the group missing in the substituted $aPMB$ chains has to be the DNP-substituted NH$_2$ terminal. Substitution of lysines can be ruled out because this should have produced a shift of the right peak in the curve for $aPMB$DNP relative to the same peak in the curve for $aPMB$ in Fig. 2. It is also evident that the height of the right peak is higher in the substituted chains, meaning a diminished buffer capacity of the protein in this region of about 10%. This would indicate that the $pK$ value of the amino group is considerably lower than 8, because the disappearance of a group with a $pK$ near 8 would have changed the buffer capacity about 20% (3). In Fig. 3 the difference in protons bound by $aPMB$ DNP and $aPMB$ is shown. The curve shows clearly that the $pK$ of the amino terminal is near 7.4. The value for $w$, the electrostatic interaction factor obtained from a plot of $pH - \log a/(1-a)$ against $Z_H$ (14), where $a$ is the degree of ionization, was 0.060. This value is as usual (2) lower than that calculated on the basis of the radius of the macromolecule, $w = 0.078$ for the $aPMB$ chain at $\mu m = 0.1$.

Spectrophotometric Kinetic Measurements—Fig. 4 shows two examples of experiments done at pH 6.645 and pH 7.660, respectively. As is seen straight lines are obtained even to a degree of substitution of more than 70% indicating a homogeneous reaction. In Fig. 5 the kinetic data are plotted as $1/K_HpH$ against $(H^+)$ ($\Delta$). From the two intercepts we calculated $k^o = 0.282 \pm 0.01$ m$^{-1}$ sec$^{-1}$ and $pK$ 7.3. The $pK$ value found is slightly lower than the $pK$ calculated from Fig. 3, but the agreement is nevertheless satisfying. The $k^o$ value for this group in carboxyhemoglobin as measured by Hill and Davis (1) is somewhat lower, viz. $k^o = 0.239$ m$^{-1}$ sec$^{-1}$.

pH-Kinetic Measurements—Fig. 6 shows an example of the change in pH upon reaction of $aPMB$ chains with FDNB. Thirty seconds after addition of FDNB a straight line is obtained. The slope of this line was taken as the slope at zero time without any correction, as the spontaneous hydrolysis of FDNB was very slow in the pH region investigated. The initial change in pH was very probably caused by small differences in temperature and pH of the $aPMB$ and FDNB solution before mixing. $K_HpH$ values were calculated with the buffer capacity data available from the acid-base titrations. In spite of the fact that all data so far presented are consistent with the hypothesis that only the NH$_2$-terminal valine reacted with FDNB under our conditions, the $K_HpH$ values did not correspond to the...
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...substitution the oxygen affinity increases, for the tem. The value oxygen affinity was increased or at least equal in the substituted procedure. However, a few measurements performed on the substituted material completely denatured during the deoxygenation process. It is also higher than in hemoglobin for which we found $s_{40, w} = 4.45$ under identical conditions. This high sedimentation constant can be explained assuming that the PMB-DNP system is tetrameric and that the substitutions with PMB and DNP have changed the partial specific volume $\bar{\nu}$ of the hemoglobin molecule. Assuming a molecular weight of 67,000 for the DNP-PMB system and of 64,500 for normal hemoglobin and with $\bar{\nu}$ of 0.749 for hemoglobin, we calculated $\bar{\nu} = 0.731$ for the DNP-PMB system. This value is as expected somewhat lower than that reported by Edelstein et al. (18) for the PMB system, viz. $\bar{\nu} = 0.736$.

**DISCUSSION**

All the results concerning the degree of substitution of the material of the leading peak in Fig. 1 indicated that the NH$_2$ terminus of the α$^{PMB}$ chains was fully substituted. Since titration studies of the same material showed that only one group was no longer titratable, we conclude that this peak contained only α$^{PMB}$ chains homogeneously substituted with DNP in their terminal valine. The second peak in the same preparation appeared to be unreacted α$^{PMB}$ chains, as judged from the value of $R$ and of the isionic point. Also, in the chromatography it appears that the amount of material showing a value of $R$ higher than 3, (expected if some lysines were substituted) is very small. In other words the chromatographic analysis indicates that, at neutral pH, essentially only the terminal valine is substituted by FDNB. Substitution of histidines was in no way detectable, in agreement with the findings of Neer and Konigsberg (8) for carboxyhemoglobin. However the pH kinetics measurements showed that the release of protons by the α$^{PMB}$ chains upon the reaction with FDNB cannot be accounted for by the substitution of the NH$_2$ terminus alone. The rate of release of protons can only be explained assuming the presence of another "reactive" site with a pK 6.85 which does not produce a stable DNP derivative. The findings of Neer and Konigsberg (8) for carboxyhemoglobin.

The two curves measured at pH 7.65 and 6.5, respectively, coincided completely. Fig. 8 presents the sedimentation velocity data for the α$^{PMB}$DNP chains and the DNP-PMB system as a function of the concentration. The measurements were carried out on their carboxy derivatives in 0.05 M phosphate buffer at pH 6.9. The upper curve shows that the $s_{40, w}$ value for the α$^{PMB}$DNP chains at a concentration of 1 mg per ml is about 1.9. The $s_{40, w}$ value for the chain as reported by Antonini et al. (16) is 1.85 at a concentration of 2.5 mg per ml. The upper curve shows that the DNP-PMB system has an $s_{40, w}$ value near 4.95 at all concentrations. This value is higher than for the PMB system which shows an $s_{40, w}$ = 2.5 at pH 7.1 (16), when in the carboxy form. The plot in Fig. 5 of $K_B$ against $1/AK$ gave values of $K_B$ = 2.5 at pH 7.1 (16), when in the carboxy form. It is also higher than in hemoglobin for which we found $s_{40, w} = 4.45$ under identical conditions. This high sedimentation constant can be explained assuming that the PMB-DNP system is tetrameric and that the substitutions with PMB and DNP have changed the partial specific volume $\bar{\nu}$ of the hemoglobin molecule. Assuming a molecular weight of 67,000 for the DNP-PMB system and of 64,500 for normal hemoglobin and with $\bar{\nu}$ of 0.749 for hemoglobin, we calculated $\bar{\nu} = 0.731$ for the DNP-PMB system. This value is as expected somewhat lower than that reported by Edelstein et al. (18) for the PMB system, viz. $\bar{\nu} = 0.736$.

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...masked in hemoglobin in the α1S2 interface, as suggested by that H5 (122)a histidine, which is close to the NH2 terminus is...A...system indicates a strong tendency to form tetramers which...the terminal valine of the...disappearance of the heme-heme interaction in hemoglobin when...the value of...benzoate was overcome to the point that the PMB-DNP system...the partner dimer. The dissociating effect of the p-mercuri-phobic...in nature, of the DNP side chain with residues present in...subunits and in hemoglobin.

...With the model of Bjerrum described by Edsall and Wyman...contrasts with the easy dissociability of the PMB system. This...could result either from a change in the conformation of the...of 6 A between the two groups can be calculated from the difference...of 6.72 reported by Hill and Davis (1) for the histidines is somewhat...of 10 histidines can be protonated in hemoglobin...possible that while 10 histidines can be protonated in hemoglobin...chain.

...Aoid base titrations would assign a pK near 7 to those...their proton-binding characteristics.

...the value of...per histidyl residue. Also, it is...pK 6.4 chosen by Hill and Davis (1) for the histidines is somewhat...anomalies in the pH measurements. However, this seems improbable...the NH2-terminus of the PMB dimers or from a direct interaction, probably hydrophobic in nature, of the DNP side chain with residues present in...pK value of 7.35 for the terminal valine is remarkably higher than the value of 6.72 reported by Hill and Davis (1) for the same group in carboxyhemoglobin. This supports their hypothesis that this low pK is produced in hemoglobin by the proximity of the terminal arginine of the partner α chain. With the model of Bjerrum described by Edsall and Wyman (19) and assuming that the medium between the two groups has a dielectric constant equal to that of water an average distance of 6 A between the two groups can be calculated from the difference in pK of the terminal valine of the α chains in the isolated subunits and in hemoglobin.

Also, Hill and Davis (1) found that the pK of the NH3-terminal residue in carboxyhemoglobin was not pH dependent as if ω was very near zero, and they explained this by the absence of ionizable histidines near that group. In the free αPMB chains the value of ω estimated by us for the ionization of the terminal valine was 0.000. This finding could support the hypothesis that H5 (122)a histidine, which is close to the NH3 terminus is masked in hemoglobin in the αβ interface, as suggested by Perutz (20), but free to ionize in the isolated chains, so affecting the ionization of the NH3-terminal residue.

The oxygen-binding studies of the PMB-DNP system revealed a value of n = 1.3, only slightly lower than in the PMB system where n = near 1.5. This was unexpected in view of the total disappearance of the heme-heme interaction in hemoglobin when the terminal valine of the α chains was substituted with FDNB (8).

The high sedimentation coefficient found for the PMB-DNP system indicates a strong tendency to form tetramers which contrasts with the easy dissociability of the PMB system. This could result either from a change in the conformation of the αβ PMB dimers or from a direct interaction, probably hydrophobic in nature, of the DNP side chain with residues present in the partner dimer. The dissociating effect of the p-mercuri-benzoate was overcome to the point that the PMB-DNP system was no longer able to dissociate into α and β chains upon electrophoresis and chromatography.

Note Added in Proof—In a recent letter to one of us (S. H. B.) Dr. Perutz said that in collaboration with Dr. Ten Eyck he was able to locate, in human oxyhemoglobin, "the position where the guanidinium group of the Arginine 141α spends part of its time. It is in the internal cavity immersed in water without being bonded to any other group of the protein...Its distance from the α-amino group of the Valine 1 of the other α chain is near 6 A."

Acknowledgments—The authors wish to thank Dr. Clara Fron­ticelli for her advice in the fingerprinting and amino acid analy­sia and Mrs. Barbara Marrow for doing part of these experi­ments. The Health Science Computer Center of the University of Maryland (HSCC) in Baltimore generously granted free computer time to the investigators. The assistance of Mr. Robert Jones, Mr. Jerome Michener, and Mr. Javier Alharran in using the system at the HSCC is gratefully acknowledged.

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