
R J Desnick, …, P A Tishler, P Mustajoki

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To investigate the molecular pathology in acute intermittent porphyria (AIP), the nature of the defective porphobilinogen (PBG)-deaminase was determined in erythrocyte lysates from 165 AIP heterozygotes from 92 unrelated families representing 20 different ethnic or demographic groups. Immunologic and physiokinetic studies revealed the occurrence of four classes of PBG-deaminase mutations. In the majority of families studied, the amount of immunoreactive enzyme protein corresponded to the amount of enzymatic activity, indicating the absence of cross-reacting immunologic material (CRIM) produced by the mutant allele. In 78 of these CRIM-negative families (designated type 1), the affected heterozygotes had half-normal PBG-deaminase activity. In three families (designated CRIM-negative type 2), symptomatic patients had increased urinary excretion of delta-aminolevulinic acid and PBG, and normal levels of erythrocyte PBG-deaminase activity. In contrast, noncatalytic, immunoreactive protein was expressed in heterozygotes from 11 families, about one-eighth of those studied, consistent with mutations in the structural gene for PBG-deaminase. Two types of CRIM-positive mutations were identified: the type 1 mutation had a CRIM/activity ratio of approximately 1.7 and a crossed-immunoelectrophoretic profile in which all the enzyme intermediates were increased, with the B or monopyrrole-enzyme intermediate predominant (B greater than A much greater than C congruent to D greater than E). The mutation altered both the kinetic and stability properties of the noncatalytic immunoreactive enzyme protein. The second CRIM-positive mutation…

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Acute Intermittent Porphyria: Characterization of a Novel Mutation in the Structural Gene for Porphobilinogen Deaminase
Demonstration of Noncatalytic Enzyme Intermediates Stabilized by Bound Substrate

Robert J. Desnick, Ludmila T. Ostasiewicz, Peter A. Tishler, and Pertti Mustajoki
Division of Medical Genetics, Mount Sinai School of Medicine, New York 10029; Veterans Administration Medical Center, Brockton-West Roxbury, Massachusetts 02401 and Harvard Medical School, Boston, Massachusetts 02115; and Third Department of Medicine, University of Helsinki, 00290 Helsinki, Finland

Abstract

To investigate the molecular pathology in acute intermittent porphyria (AIP), the nature of the defective porphobilinogen (PBG)-deaminase was determined in erythrocyte lysates from 165 AIP heterozygotes from 92 unrelated families representing 20 different ethnic or demographic groups. Immunologic and physiokinetic studies revealed the occurrence of four classes of PBG-deaminase mutations. In the majority of families studied, the amount of immunoreactive enzyme protein corresponded to the amount of enzymatic activity, indicating the absence of cross-reacting immunologic material (CRIM) produced by the mutant allele. In 78 of these CRIM-negative families (designated type 1), the affected heterozygotes had half-normal PBG-deaminase activity. In three families (designated CRIM-negative type 2), symptomatic patients had increased urinary excretion of δ-aminolevulinic acid and PBG, and normal levels of erythrocyte PBG-deaminase activity. In contrast, noncatalytic, immunoreactive protein was expressed in heterozygotes from 11 families, about one-eighth of those studied, consistent with mutations in the structural gene for PBG-deaminase. Two types of CRIM-positive mutations were identified: the type 1 mutation had a CRIM/activity ratio of ~1.7 and a crossed-immunoelectrophoretic profile in which all the enzyme intermediates were increased, with the B or monopyrrole-enzyme intermediate predominant (B > A > C ≈ D > E). The mutation altered both the kinetic and stability properties of the noncatalytic immunoreactive enzyme protein. The second CRIM-positive mutation, type 2, had markedly increased levels of noncatalytic immunoreactive protein (CRIM/activity ratio ~ 5.7). Crossed-immunoelectrophoresis revealed markedly increased amounts of the substrate-bound intermediates, B, C, D, and E (B > C > D > E ≈ A). The accumulation of these noncatalytic enzyme intermediates presumably resulted from the enhanced binding and/or defective release of substrate molecules. The conformation of these enzyme-substrate intermediates apparently rendered the complexes more resistant to inaerythrocyte proteolysis. These findings provide evidence for the presence of different allelic mutations in the structural gene for PBG-deaminase and document molecular genetic heterogeneity in AIP.

Introduction

Acute intermittent porphyria (AIP) is a dominantly inherited inborn error of heme biosynthesis that results from the half-normal activity of porphobilinogen deaminase (PBG-deaminase; EC 4.3.1.8) (1–4). Clinical expression of the disease is highly variable, determined in part by environmental, metabolic, and hormonal factors that induce hepatic δ-aminolevulinic acid synthase activity and the subsequent increased production of heme precursors (5–7). Although the disease is clinically latent in many heterozygous individuals, those who are symptomatic usually have elevated levels of urinary PBG and its precursor, δ-aminolevulinic acid (ALA), particularly during acute attacks. Most affected individuals can be diagnosed enzymatically (i.e., decreased erythrocyte PBG-deaminase activity); however, kindreds have been described recently in which symptomatic patients have normal levels of the erythrocyte PBG-deaminase activity and markedly elevated urinary PBG and ALA levels (8, 9).

PBG-deaminase is encoded by a structural gene located on the distal portion of the long arm of chromosome 11 (11q23 → 11qter) (10). The erythrocyte enzyme has been purified to homogeneity and its physiokinetic properties have been characterized (11). The active enzyme protein is a monomer (Mr ~ 42,000) that catalyzes the sequential head to tail condensation of four molecules of PBG to form hydroxymethylbilane, a linear tetrahydroporphyrin that nonenzymatically cyclizes to uroporphyrinogen I. Hydroxymethylbilane is synthesized by a reaction mechanism involving the stepwise formation of stable enzyme-substrate intermediates (11–13). Five enzyme intermediates, A–E, have been identified by chromatographic and electrophoretic techniques, the A form being the free enzyme and the B, C, D, and E forms representing the mono-, di-, tri-, and tetrahydroxymethylbilane substrate–enzyme intermediates, respectively (11, 14). The overall Km for the reaction was 6 μM and kinetic studies indicated that the C intermediate, the dipyrrole–enzyme complex, was either the most stable intermediate or the rate-limiting step in the conversion of the monopyrrole to the tetrahydroporphyrin (11, 14).

In a previous communication, we reported the first immunologic characterization of the enzymatic defect in unrelated heterozygotes with AIP (14). Monospecific anti-human PBG-deaminase antibodies, which uniformly recognized each of the enzyme intermediates, were used to quantitate and characterize the amount of PBG-deaminase cross-reacting immunologic

Abbreviations used in this paper: AIP, acute intermittent porphyria; ALA, δ-aminolevulinic acid; CRIM, cross-reacting immunologic material; DTT, dithiothreitol; PBG, porphobilinogen.
material (CRIM) in erythrocyte lysates from 32 heterozygous individuals of 22 unrelated AIP families. Two classes of mutations were identified. In 21 of the AIP families, the amount of enzyme protein was directly proportional to the amount of enzymatic activity (i.e., the CRIM/activity ratio was ≈ 1.0); these mutations were classified CRIM-negative. In seven affected members from a family of Basque ancestry, the PBG-deaminase mutation was CRIM-positive; the CRIM/activity ratio of ≈ 1.65 indicated the presence of immunoreactive, noncatalytic enzyme produced by the mutant allele. Although only 1 of the 22 AIP families had a CRIM-positive mutation, this finding provided evidence for heterogeneity of the genetic defect in AIP.

In this communication, we report the immunologic and physiokinetic properties of erythrocyte PBG-deaminase in 165 AIP heterozygotes from 92 unrelated families representing 20 different ethnic or demographic backgrounds. Four classes of mutations were identified: two were CRIM-negative and two were CRIM-positive. The CRIM-negative mutations were divided into two types based on the level of erythrocyte PBG-deaminase activity. The majority of CRIM-negative families, designated type 1, had half-normal activity, whereas several CRIM-negative families, including a large Finnish kindred (8), had normal erythrocyte activity and were designated type 2. Notably, 11 AIP families, or about one-eighth of those studied, had CRIM-positive mutations. One type was similar to that previously described in the Basque kindred (designated CRIM-positive type 1). The second type, designated CRIM-positive type 2, was unusual in that it had markedly more CRIM than expected for a single mutant allele. Immunologic studies demonstrated that the increased amount of CRIM in the type 2 heterozygotes resulted from stabilization of the mutant protein by bound substrate.

Methods

Materials

PBG was synthesized as described previously (15). Uroporphyrin standards were purchased from Porphyrin Products, Logan, UT. Diaminobenzidine tetrahydrochloride, ALA (hydrochloride), DL-dithioerythritol, DL-dithiothreitol (DTT), bovine serum albumin (BSA), and human serum albumin were obtained from Sigma Chemical Co., St. Louis, MO. High-purity acetonitrile was purchased from Burdick & Jackson Laboratories, Muskegon, MI. Fluorescamine was obtained from Pierce Chemical Co., Rockford, IL. Goat anti-rabbit serum and peroxidase-conjugated sheep anti-rabbit IgG were obtained from Cappel Laboratories, Cochranville, PA. SeaKem agarose was purchased from Marine Colloids Div., FMC Corp., Rockland, ME. High-resolution Tris-barbital buffer was obtained from Gelman Instrument Company, Ann Arbor, MI. AgaroseEF, Pharmalyte ampholines (pH 5–8), and Gelbond film were purchased from Pharmacia Fine Chemicals, Piscataway, NJ. Bovine pancreas trypsin was from Worthington Biochemical Corp., Freehold, NJ, and pronase B grade (Lot 200191) was from Calbiochem-Behring Corp., San Diego, CA. Stractan was obtained from the St. Regis Paper Co., Tacoma, WA. All other materials and chemicals were of the highest quality available.

Human subjects and preparation of erythrocyte lysates

Subjects included heterozygotes with AIP from unrelated families representing a variety of different ethnic backgrounds or countries of origin. The clinical diagnosis of AIP in each subject was documented by demonstration of half-normal PBG-deaminase activity in erythrocyte lysates and/or increased urinary excretion of PBG and ALA.

Heparinized blood was collected from each AIP heterozygote and 35 age- and sex-matched normal individuals, with informed consent. After centrifugation at 2,500 g, the erythrocytes were removed, washed twice with 0.9% NaCl, and then used immediately or stored at −20°C. For specific experiments, human erythrocytes were separated into age fractions on discontinuous Stractan gradients according to the method of Corash et al. (16). Erythrocytes were lysed by the addition of 3 vol of 1.0 mM sodium phosphate buffer, pH 8.1, containing 1 mM DTT, 1 mM MgCl2, and 0.05% Triton X-100 (lysis buffer). The lysates were centrifuged at 30,000 g for 20 min and the supernatants were removed and used for assays of protein and PBG-deaminase activity. Samples were diluted and then reasayed to ensure that all lysates contained equal enzymatic activities prior to kinetic, heat denaturation, rocket immunoelectrophoretic and immunotitration studies. For protease digestion experiments, the protein concentrations of the lysates were equalized by dilution with lysis buffer.

Assays

PBG-deaminase assay. Enzymatic activity was determined by the quantitation of uroporphyrin fluorescence as previously described (15). One unit of enzymatic activity equaled that amount of enzyme that formed 1 nmol of uroporphyrin per h at 37°C.

Protein assay. Protein concentrations were determined by the fluorescamine procedure as previously described (17).

Kinetic, heat denaturation, and protease stability studies

For kinetic studies, 50 µl of erythrocyte lysate was incubated in the standard assay with the final substrate concentration ranging from 0 to 50 µM. The Km values were determined from Lineweaver–Burk plots.

To determine the effect of heat denaturation on erythrocyte PBG-deaminase activity, 500 µl aliquots of the lysates, which were equalized for activity, were placed in screw-capped vials and incubated at 65°C or 70°C for 2 h, and then cooled to 0°C. After centrifugation at 10,000 g for 10 min to remove precipitated proteins, 50 µl of the supernatant was removed and immediately assayed for PBG-deaminase activity.

To study the effect of trypsin and pronase on PBG-deaminase activity, erythrocyte lysates were equalized for protein concentration (to 80 mg/ml by dilution with lysis buffer). Aliquots (40 µl) of each lysate were placed at 0°C, then 10 µl of lysis buffer containing 125 µg/ml of pronase or 0.0625 µg/ml of trypsin was added to give a final protease concentration of 25.0 µg/ml or 0.0125 µg/ml, respectively. The reaction mixtures were preincubated for 0–60 min at 37°C, after which time 500 µl of 0.1 M Tris-HCl buffer, pH 8.1, containing 0.1 mM DTT, and 200 µl of 0.5 mM PBG were added immediately and the PBG-deaminase activity determined as described above.

Immunologic studies

Characterization of anti-PBG-deaminase. Anti-human PBG-deaminase was raised in New Zealand rabbits immunized with the homogeneous A intermediate and the IgG fraction was purified as previously described (14). This antibody preparation was shown to be specific for purified human PBG-deaminase (or the human enzyme in erythrocyte lysates or murine erythroleukemia–human fibroblast somatic cell hybrid) by Ouchterlony double-immunodiffusion and competitive immunoprecipitation studies (10, 14). Ouchterlony immunodiffusion gels showed a single arc of identity between the homogenous enzyme and that in human erythrocyte lysates or liver homogenate supernatants when stained for protein and activity using PBG as substrate (Desnick, R. J., et al., unpublished results).

Immunotitration of anti-human PBG-deaminase. Homogeneous PBG-deaminase intermediate A (40 ng of enzyme diluted with BSA to 50 µl; final protein concentration of 40 mg/ml) and 50 µl of rabbit anti-PBG-deaminase IgG (in serial dilutions) were mixed and incubated at 37°C. After 30 min, 50 µl of goat anti-rabbit IgG was added and the mixture was incubated at 37°C for 30 min. To assure quantitative precipitation of the anti-PBG-deaminase antibodies, the samples were left at 4°C overnight. The mixtures were then centrifuged at 4,000 g for 15 min, and the activity in the supernatant was determined by the
standard assay described above. For immunotitration of the immuno-
reactive PBG-deaminase in erythrocyte lysates from normal and AIP
heterozygotes, the lysate activities were equalized and then either 50-
μl or 25-μl aliquots were titrated against the antibody as described
above.

**Immunologic quantitation and characterization of PBG-
deaminase in erythrocyte lysates**

**Rocket immunoelectrophoresis.** Rocket immunoelectrophoresis of hu-
mans PBG-deaminase was performed as previously described (14). An
agarose solution (1%) was prepared in 0.06 M Tris-barbital buffer, pH
8.8, and bridges containing 4 ml of agarose solution were poured into
a 0.9% NaCl solution, and then 50 μl of antisera. a 0.5 M
H2SO4. The plate was
focusing. The gel was soaked in
a 0.5 cm-thick glass microscope slide
had been in the center. After the agarose had gelled, the slide
was removed and 4.0 ml of the agarose solution containing 15 μl of
rabbit anti-human PBG-deaminase IgG (11 μg protein) was poured
into the center area. Wells were cut at the cathodal end of the antibody-
containing gel and 5–10-μl samples were applied. Electrophoresis was
carried out in a tank (Chemtron model 200, Milan, Italy) containing
0.06 M Tris-barbital buffer, pH 8.8, at a constant current of 30 mA
for 4 h at room temperature. After electrophoresis, the gel was thoroughly
washed in 0.9% NaCl for 10–12 h with several changes and then the
antibody containing part of the gel was removed with 300 μl of
peroxidase-conjugated goat anti-rabbit IgG which had been diluted 1:2
with 0.9% NaCl. The plate was incubated at room temperature in a
moist chamber to prevent drying. After another overnight wash with
0.9% NaCl, the gel was stained for peroxidase with 50 ml of 0.1 M
Tris-HCl buffer, pH 7.6, containing 25 μg of diaminobenzidine
tetrahydrochloride and 0.15 ml of 3% hydrogen peroxide. After staining
for 10–20 min at room temperature, the rockets were visualized by
using an indirect light source, and the gel was photographed and stored
in 0.9% NaCl at 4°C.

**Isoelectric focusing and crossed-immunoelectrophoresis.** Isoelectric
focusing of the PBG-deaminase intermediates was performed on hori-
Zontal agarose slab gels using the Pharmacia system according to the
manufacturer's instructions. Gels were prepared by heating a mixture of
0.3 g of AgaroseIEF 3.6 g of sorbitol, and 27 ml of distilled water
in a boiling water bath. After cooling to 75°C, 1.9 ml of pH 5–8
Pharmalyte ampholines were added. The mixture was then poured
into a horizontal casting frame (11.4 × 22.5 cm; with Gelbond film
backing), which had been preheated to about 60°C using a portable
dryer. Gels were allowed to harden at least 1 h at 4°C or stored
overnight at 4°C in a moist chamber. Prior to isoelectric focusing,
erythrocyte lysates were heated at 60°C for 1 h and centrifuged at
10,000 g for 20 min, and then aliquots of the supernatants (50–100
μl) were pipetted on Whatman No. 17 paper strips (Whatman
Laboratory Products, Inc., Clifton, NJ) and placed 1 cm from the
cathode. The cathode contained 1.0 M NaOH and the anode 0.05 M
H2SO4. Focusing was carried out for 2.5 h at 10 W (constant power)
at 4°C. The filter paper strips were removed after the hemoglobin had
migrated about 1 cm from the origin. After focusing, the pH gradient
was determined by removing a 0.5-cm gel strip, cutting it into 1-cm
pieces which were soaked in distilled H2O, and then the pH values of
crossed-immunoelectrophoresis of the separated PBG-deaminase
intermediates was performed by placing an unstained lane from the
focusing gel at the cathodal end of a rocket immunoelectrophoresis
plate. The remainder of the plate was filled with 8 ml of 1% agarose
containing 40 μl of anti-PBG-deaminase IgG and rocket immunoelec-
trophoretic gels were stained for protein with Coomassie Brilliant Blue
R-250 as previously described (14).

**Results**

**Immunologic identification of PBG-deaminase mutant classes.** The
PBG-deaminase CRIM status was determined in 165 AIP
heterozygotes from 92 unrelated families representing at least
20 different ethnic or demographic backgrounds (Table I). For
these studies, equal amounts of erythrocyte PBG-deaminase
activity from each AIP heterozygote and normal individuals
were subjected to rocket immunoelectrophoresis. AIP hetero-
yzogotes whose rocket peak heights were essentially identical to

<table>
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<tr>
<th>Family ancestry</th>
<th>Number of families</th>
<th>Number of patients</th>
<th>PBG-deaminase mutant class</th>
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<td></td>
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| American Black‡ | 1                   | 2                  | ×      |        |        |        |
| Finnish‡        | 1                   | 5                  | ×      |        |        |        |
| Norwegian/Finnish‡ | 1                 | 2                  | ×      |        |        |        |
| Total           | 92                  | 165                | 5      | 6      | 78     | 3      |

* Both parents were from the same ethnic group or country unless otherwise indicated.
‡ AIP heterozygotes with normal erythrocyte activity.

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**Table I. Immunologic Characterization of the Defective PBG-
Deaminase in AIP Heterozygotes from Unrelated Families**
those of normal individuals were designated CRIM-negative (CRIM/activity ratio \(\leq 1.0\)). In contrast, AIP heterozygotes whose rocket peak heights were greater than those of normal individuals were designated CRIM-positive (CRIM/activity ratio > 1.0), i.e., the increased CRIM represented noncatalytic, immunoreactive PBG-deaminase expressed by the mutant allele. As shown in Fig. 1, four major PBG-deaminase mutant classes were readily identified: (a) CRIM-negative type 1 heterozygotes with half-normal PBG-deaminase activity, (b) CRIM-negative type 2 heterozygotes with normal erythrocyte activity, (c) CRIM-positive type 1 heterozygotes whose peak heights were slightly greater than those of normal individuals, and (d) CRIM-positive type 2 heterozygotes whose peak heights formed “railroad tracks,” indicating the presence of markedly more noncatalytic PBG-deaminase than that observed in the CRIM-positive type 1 heterozygotes. Review of the medical histories did not reveal notable differences in clinical disease expression in affected individuals among the four mutant classes.

**Quantitation of PBG-deaminase CRIM.** The amount of immunoreactive PBG-deaminase protein in each AIP heterozygote was determined by immunotitration. When equal amounts of erythrocyte enzyme activity from CRIM-negative type 1 and 2 heterozygotes and normal individuals were immunotitrated, the same antibody dilution (1:200) was required to precipitate 50% of the activity from each of these erythrocyte supernatants and their immunotitration profiles were essentially identical (Fig. 2). The identity of their immunotitration curves confirmed the rocket immunoelectrophoretic studies that indicated the absence of noncatalytic enzyme protein in either type of CRIM-negative heterozygote. In that the amount of CRIM detected by either technique corresponded to the amount of erythrocyte activity used, the CRIM/activity ratios for the CRIM-negative type 1 and 2 heterozygotes and normal individuals were 1.0.

In contrast, immunotitration of identical amounts of activity from the CRIM-positive type 1 and 2 heterozygotes required antibody dilutions of 1:116 and 1:35 to precipitate 50% of the supernatant activities, respectively (Fig. 2). The antibody dilutions required to precipitate 50% of the activity from 1:1 mixtures of normal and CRIM-positive type 1 or type 2 lysates were 1:146 and 1:60, respectively (data not shown). Based on these results, and assuming identical antibody avidities, the amounts of immunoreactive PBG-deaminase in these representative CRIM-positive type 1 and 2 heterozygotes were \(~1.7\)- and 5.7-fold that in normal lysates (i.e., CRIM/activity ratios of 1.7 and 5.7). Analogously, the CRIM/activity ratios determined by immunotitration for the 18 CRIM-positive type 1 and the 20 CRIM-positive type 2 heterozygotes ranged from 1.6 to 1.8 (mean=±1 SD = 1.7±0.07) and from 4.9 to 6.0 (mean=±1 SD = 5.7±0.3), respectively.

The amount of immunoreactive PBG-deaminase in the CRIM-positive type 2 heterozygotes also was determined by quantitative rocket immunoelectrophoresis. As shown in Fig. 3, the rocket peak heights for 7 and 10 mU of PBG-deaminase activity in the CRIM-positive type 2 erythrocytes were essentially equal to those obtained for 40 and 56 mU of normal lyase activity, respectively. Thus, the CRIM/activity ratio determined by rocket immunoelectrophoresis was \(~5.7\), consistent with the values for the CRIM-positive type 2 heterozygotes obtained by immunotitration.

**Characterization of the PBG-deaminase intermediates in the AIP mutant classes.** Crossed-immunoelectrophoresis of erythrocyte PBG-deaminase provided further information concerning the nature of the enzymatic defect in the four AIP mutant classes. For these studies, PBG-deaminase in erythrocyte lysates was first subjected to isoelectric focusing in agarose gels to separate the enzyme intermediates. Five major PBG-deaminase activity bands were observed in normal and CRIM-positive type 1 and 2 lysates (Fig. 4), as well as in CRIM-negative type 1 and 2 lysates (data not shown). There were no significant differences in the pi values for each activity band in lysates from the different mutant classes. After crossed-immunoelectrophoresis, a precipitin line of identity with peaks corresponding to each of the five intermediates was observed.

![Rocket immunoelectrophoresis of PBG-deaminase in erythrocyte lysates from unrelated AIP heterozygotes representing each of the four mutant classes. Equal enzymatic activities were applied to the gel, except in wells 9-11 in which the activity was diluted as indicated. Normal individuals, wells 1, 5, 7, 9, and 12; CRIM-negative type 1 heterozygote (with half-normal erythrocyte activity), well 2; CRIM-negative type 2 heterozygote (with normal erythrocyte activity), well 3; CRIM-positive type 1 heterozygote, well 4; CRIM-positive type 2 heterozygote, wells 6, 8, 10 and 11.](image1)

![Figure 2. Immunotitration of PBG-deaminase with rabbit anti-human PBG-deaminase IgG. Immunotitration curves are shown for 25-µl aliquots (equalized for enzymatic activity and protein concentration with BSA) of homogeneous enzyme (w) and erythrocyte lysates of a normal individual (n), a CRIM-negative type 1 heterozygote (j), a CRIM-negative type 1 heterozygote (n), and a CRIM-positive type 2 heterozygote (k). Note that the curves for 25 µl of the homogeneous enzyme, normal lyase, and the CRIM-negative type 1 lyase were essentially identical; the curve for 25 µl of the CRIM-negative type 2 lyase also was identical (data not shown). Antibody dilutions of \(~1\): 35, 1:116 and 1:200 were observed for 50% immunoprecipitation of enzyme from 25 µl of the CRIM-positive type 2 lyase, the CRIM-positive type 1 lyase and the CRIM-negative type 1 (or normal) lyase, respectively.](image2)
in lysates from normal individuals and AIP heterozygotes from each mutant class. The typical profiles obtained for erythrocyte PBG-deaminase from normal individuals and all CRIM-negative type 1 and 2 heterozygotes are shown in Fig. 4. In normal lysates, the A intermediate had the highest peak height, with B, C, D, and E intermediates having sequentially decreasing peak heights (upper gel). In contrast, lysates from CRIM-positive type 1 heterozygotes (middle gel) had increased amounts of all five intermediates, the immunoreactive B intermediate being most elevated (14). Lysates from the CRIM-positive type 2 heterozygotes (lower gel) had a distinctly different profile: the A peak appeared decreased whereas the B, C, D, and E peaks were markedly increased. For comparison, the amount of type 2 activity applied to this gel was one-fifth of that used for crossed-immunoelectrophoresis of the normal or CRIM-positive type 1 enzyme.

To further characterize the amount of immunoreactive PBG-deaminase intermediates in CRIM-positive type 2 heterozygotes, the erythrocyte enzyme intermediates from CRIM-positive type 2 and normal individuals were isolated by DEAE-cellulose chromatography (14), and equal activities of the A, B, C, and D forms were subjected to rocket immunoelectrophoresis. As shown in Fig. 5, each of the intermediates from the CRIM-positive type 2 lysate had an increased amount of immunoreactive enzyme protein when compared with that obtained with the respective normal intermediate. Notably, the CRIM-positive B, C, and D intermediates were markedly increased whereas the A form, the free enzyme, was only slightly increased.

Characterization of the immunoreactive PBG-deaminase in age-fractionated erythrocytes from the CRIM-positive type 2 heterozygotes. Erythrocytes from a CRIM-positive type 2 heterozygote and a normal individual were fractionated into age cohorts on discontinuous Stractan density gradients, according to the method of Corash et al. (16). As shown in Fig. 6, four age-cohort fractions were obtained from a CRIM-positive heterozygote and five fractions were obtained from normal erythrocytes. These fractions ranged in mean age from 30 to ~90 d. The lysates of the normal erythrocyte fractions (youngest to oldest) had PBG-deaminase activities of 0.159, 0.126, 0.102, 0.095, and 0.081 U/mg protein compared to 0.103 for the unfractionated lysate. The lysate activities of the fractionated CRIM-positive type 2 heterozygote (youngest to oldest) were
0.136, 0.077, 0.054, and 0.050 U/mg protein, whereas the activity in the unfractionated lysate was 0.068 U/mg. As shown in Fig. 7 (upper gel), rocket immunoelectrophoresis of equal amounts of activity from the unfractionated normal lysate and from the age-fractionated lysates (youngest to oldest) revealed no difference in rocket peak height, indicating that the amount of activity corresponded to the amount of immunoreactive enzyme protein in each age-fraction. In contrast, rocket immunoelectrophoresis of 0.2 activity units of the unfractionated lysate, and the four age-fractionated lysates from a type 2 heterozygote, demonstrated increasing amounts of cross-reactive enzyme protein with increasing erythrocyte age (Fig. 7, lower gel).

As shown in Fig. 8, crossed-immunoelectrophoresis of a lysate from the youngest type 2 age fraction revealed a profile in which the B intermediate had the highest peak height (middle gel), whereas crossed-immunoelectrophoresis of a lysate from the oldest type 2 age fraction (lower gel) revealed a profile (Fig. 7, upper gel) in which the C intermediate was the highest peak.

Presumably, the C or dipyrrrole-enzyme intermediate represents
the rate-limiting step in the sequential conversion of the monopyrrole to the linear tetrapyrrrole and/or the most stable enzyme-substrate complex (11, 14).

Kinetic, heat denaturation, and protease stability studies. The physicochemical properties of the PBG-deaminase in erythrocyte lysates from normal individuals and each of the AIP mutant classes are summarized in Table II. The $K_m$ values calculated from Lineweaver–Burk plots were essentially the same ($\sim 6 \mu M$) from all sources. However, heat denaturation and protease digestion studies revealed differences among the AIP mutant classes. When the lysates were heat denatured at 65°C and 70°C, the CRIM-positive type 2 enzyme was most stable and was slightly activated at 65°C. The CRIM-positive type 1 and CRIM-negative type 1 activities had mean heat inactivation values that were similar to those of the normal enzyme. Interestingly, the CRIM-negative type 2 activity was less stable than the normal enzyme after heat treatment at 65°C. When equal amounts of erythrocyte protein from each of the mutant classes were subjected to pronase or trypsin digestion, the activities of the CRIM-positive type 1, both CRIM-negative type 1 and 2 heterozygotes, and normal individuals were equally digested by either protease. In contrast, the CRIM-positive type 2 activity was more stable to pronase digestion and appeared slightly more stable to trypsin degradation. Rocket immunoelectrophoresis of equal activities of each mutant class did not reveal any differences in the peak height of the enzyme before or after pronase or trypsin digestion (data not shown), suggesting that the partially degraded enzyme proteins lost activity, but fully retained their antigenic properties.

Discussion

Four major mutant classes were identified by immunologic and physicochemical studies of PBG-deaminase in erythrocytes from unrelated AIP heterozygotes with this dominantly inherited, inborn error of metabolism. Two of these mutant classes had been defined previously by immunologic studies of heterozygotes from 22 unrelated AIP families (14). By using monospecific antibodies to quantitate immunoreactive enzyme protein, the heterozygotes from all but one of these families were found to be CRIM-negative (CRIM/activity ratio $\sim 1.0$),

Table II. Comparison of the PBG-Deaminase Mutant Classes

<table>
<thead>
<tr>
<th>Property</th>
<th>Acute intermittent porphyria</th>
<th>CRIM-negative</th>
<th>CRIM-positive</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Normal</td>
<td>Type 1</td>
<td>Type 2</td>
</tr>
<tr>
<td>Specific activity (U/mg protein)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean</td>
<td>0.063</td>
<td>0.030</td>
<td>0.060</td>
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<tr>
<td>±1 SD</td>
<td>0.008</td>
<td>0.004</td>
<td>0.008</td>
</tr>
<tr>
<td>(n)</td>
<td>(35)</td>
<td>(118)</td>
<td>(9)</td>
</tr>
<tr>
<td>CRIM/activity ratio</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean</td>
<td>1.0</td>
<td>1.0</td>
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<tr>
<td>±1 SD</td>
<td>0.06</td>
<td>0.05</td>
<td>0.06</td>
</tr>
<tr>
<td>(n)</td>
<td>(35)</td>
<td>(118)</td>
<td>(9)</td>
</tr>
<tr>
<td>Apparent $K_m$ (µM)</td>
<td>5.9</td>
<td>5.7</td>
<td>6.0</td>
</tr>
<tr>
<td>Heat stability* (% int. act.)</td>
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<td></td>
</tr>
<tr>
<td>65°C</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean (n = 6)</td>
<td>95</td>
<td>95</td>
<td>85</td>
</tr>
<tr>
<td>Range</td>
<td>(89–101)</td>
<td>(92–100)</td>
<td>(82–87)</td>
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<tr>
<td>70°C</td>
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<tr>
<td>Mean (n = 6)</td>
<td>62</td>
<td>60</td>
<td>58</td>
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<tr>
<td>Protease Stability (% int. act.)</td>
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<td></td>
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<tr>
<td>Pronase</td>
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<tr>
<td>Mean (n = 6)</td>
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<tr>
<td>Mean (n = 6)</td>
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<td>67</td>
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<td>Isoelectric focusing profile</td>
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<tr>
<td>Crossed-Immunoelectrophoresis</td>
<td>NI</td>
<td>NI</td>
<td>NI</td>
</tr>
</tbody>
</table>

* Stability after 120 min at 65°C and 70°C expressed as percent of initial activity (% int. act.). ‡ NI = A > B > C > D > E profile on isoelectric focusing and crossed-immunoelectrophoresis.
i.e., immunoreactive, noncatalytic enzyme protein encoded by the mutant allele was not detectable in erythrocyte lysates. In only one of 22 AIP families, the mutation was CRIM-positive (CRIM/activity ratio ~ 1.7), consistent with a structural gene mutation in an allele encoding PBG-deaminase. In this communication, two additional PBG-deaminase mutant classes, designated CRIM-negative type 2 and CRIM-positive type 2, were identified and characterized. Each of these mutations was novel and provided further insight into the nature of the defects underlying the PBG-deaminase deficiency in AIP.

Characterization of the CRIM-positive mutations was permitted by the recognition that PBG-deaminase catalyzed the formation of hydroxymethylbilane via five stable enzyme intermediates, designated A–E (11–13). Previously, a model was proposed (Fig. 9), which predicted the different types of mutations which might be revealed by immunologic characterization of the PBG-deaminase intermediates in CRIM-positive heterozygotes (14). This model assumes that the normal enzyme has two binding sites for PBG and one active site (18), with the normal reaction mechanism involving the stepwise formation of the mono-, di-, tri-, and tetrapyrroles by the enzyme–substrate intermediates, A–E. Mutations that render substrate-binding site 1 defective would not permit PBG binding, and would result in the accumulation of the free enzyme. A mutation in the second substrate-binding site would lead to the accumulation of both the B and A intermediates, presumably with more B than A. If a mutation resulted in altered binding and/or catalysis, a $K_m$ mutation would result and all of the enzyme–substrate forms would accumulate, perhaps the B or C form in greater amounts, depending on whether the substrate-binding or active site was more defective. If a mutation resulted in increased binding and/or defective release of the product, only the substrate-bound forms would accumulate and, presumably, the A form would not be increased.

Immunologic characterization of the CRIM-positive type 1 mutation revealed that each of the enzyme intermediates had elevated levels of cross-reactive material and the noncatalytic B enzyme–substrate intermediate was present in the greatest amount as previously described (14). These findings best fit the model for a $K_m$ mutation with a markedly altered, but functional, active site and an altered substrate-binding site 2 (Fig. 9). In addition, the immunoreactive gene product produced by the mutant allele was somewhat unstable, because the CRIM/activity ratio was 1.7, whereas a totally stable mutant protein would have had a CRIM/activity ratio of 2.0 in the heterozygous lysate.

In marked contrast, the CRIM-positive type 2 mutants had a CRIM/activity ratio of ~5.7 (range 4.9–6.0), an unexpected finding, and a crossed-immunoelectrophoretic profile in which the immunoreactive B, C, D, and E enzyme–substrate intermediates were markedly elevated, while the A intermediate, the free enzyme, was only slightly increased. These findings best fit the model, which predicted increased binding or defective release of the bound substrate molecules (Fig 9). It appears that the substrate-bound intermediates are particularly resistant to degradation in the erythrocyte, as evidenced by the increasing CRIM/activity ratio in the older age-fractionated erythrocytes (Figs. 7 and 8). Thus, the type 2 structural gene mutation represents a unique type of stability defect, one in which the nonfunctional mutant protein is more stable in vivo proteolysis than the normal, catalytically active erythrocyte enzyme. Of the numerous human protein mutations characterized to date, only two hyperstable variants have been identified, hemoglobins Hacettepe and Agenogi (19–21). In each of these mutant hemoglobins, a β-chain amino acid substitution resulted in the formation of new ionic bonds in the molecule which rendered the mutant proteins more stable to heat denaturation than the normal protein. By analogy, an amino acid substitution in the mutant PBG-deaminase mono-
mer could markedly alter substrate binding and/or product release rendering the mutant substrate–enzyme complexes more stable. Of particular interest was the observation that the presence of the immunoreactive, noncatalytic enzyme in these heterozygotes altered the heat and protease stabilities of the activity expressed by the normal allele. The mechanism by which the accumulated mutant enzyme stabilized the normal erythrocyte activity remains unclear. Because PBG-deaminase is an extremely hydrophobic protein, one possible explanation might be that the normal and mutant proteins aggregate, thereby permitting the hyperstable mutant protein to increase the stability of the normal enzyme.

The CRIM-negative type 2 mutation was differentiated from the CRIM-negative type 1 mutation by the fact that the level of erythrocyte PBG-deaminase activity in these individuals was within the normal range. The normal erythrocyte PBG-deaminase levels in these individuals did not result from elevated reticulocyte counts, nor were they due to the biologic range in the expression of a single normal allele (e.g., high heterozygote level), since the activity ratios of erythrocyte PBG-deaminase to δ-aminolevulinate dehydrase and uroporphyrinogen III cosynthase were normal (Desnick, R. J., unpublished observations). Several hypotheses have been advanced to account for the normal erythrocyte PBG-deaminase activities in these symptomatic patients. It has been suggested that these AIP heterozygotes have half-normal hepatic PBG-deaminase activities and normal erythrocyte levels (8). This hypothesis requires the presence of two structural genes encoding hematopoietic (e.g., erythrocytic) and nonhematopoietic (e.g., hepatic) isozymes or a regulatory mechanism which controls either the tissue specific expression of a single structural gene or the differential expression of hematopoietic and nonhematopoietic isozymes. Support for this concept will await determination of the hepatic activity in these patients. However, the fact that human PBG-deaminase has been assigned to a single, narrow, chromosomal location (10) and the finding that monospecific antibodies against the purified erythrocyte enzyme form a single line of identity with partially purified hepatic and erythrocytic PBG-deaminase activities, argue against this hypothesis (Ostasiewicz, L. T., and R. J. Desnick, unpublished results). Nevertheless, it is possible that two or more structural genes are present as a gene family in the same chromosomal region, analogous to the β-globin gene complex (22), and that the antityrocyte PBG-deaminase antibodies did not recognize unique determinants of a hepatic isozyme. Another explanation which also may depend on the occurrence of separate genes encoding hepatic and erythrocytic enzymes, is the suggestion that the defect in AIP results from a regulatory gene mutation (7, 23). Such a mutation could result in normal hematopoietic, but half-normal hepatic gene expression.

Finally, one could account for the rare individuals with normal erythrocyte activity by the segregation of a variant allele with high normal activity (i.e., a hypermorph). It is well known that there is a large range (about threefold) in normal erythrocytic PBG-deaminase levels and that this variation is genetically determined (24, 25). The occurrence of two co-dominant PBG-deaminase alleles was hypothesized recently to account for the distribution of the erythrocyte activities in over 200 normal individuals (26). For the two-allele model, the mean activities for the trimodal distributions were ~130, 170, and 220 pmol/30 min·mg protein, respectively. If an individual inherited such a hypermorphic allele and an AIP mutant allele, the level of erythrocyte PBG-deaminase activity may be within the normal range (but the nonhematopoietic activities could be half-normal). Symptomatic individuals in such a family have been described (27); normal individuals with the hypermorphhic allele had high normal activities of erythrocyte PBG-deaminase, whereas symptomatic heterozygous individuals, who had the hypermorphhic allele and a mutant AIP allele, had erythrocyte activities in the normal range. However, characterization of the erythrocyte enzyme in six asymptomatic, apparently normal members of the large Finnish CRIM-negative type 2 family did not reveal activity levels consistent with the presence of a hypermorphhic allele. Therefore, it is likely that insight into the precise nature of the defect in these CRIM-negative type 2 heterozygotes will be revealed by molecular studies of their PBG-deaminase genes.

In summary, immunologic and biochemical characterization of the enzymatic defect in a large series of unrelated AIP families identified four classes of PBG-deaminase mutations. At the molecular level, the two CRIM-positive mutations most likely result from single base substitutions in the exons of the structural gene. The CRIM-negative mutations may represent mutations which (a) markedly alter enzyme stability or anti-genicity, (b) result in early chain termination, or (c) alter mRNA transcription or processing. The classification of these mutations should facilitate the initial selection of appropriate candidates for future molecular genetic analyses, analogous to the characterization of the genetic defects in the human thalassemias (22).

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References


