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Erythrocyte Membrane Proteins Reactive with Human (Warm-reacting) Anti-Red Cell Autoantibodies

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Abstract

Immunoglobulin G (IgG) autoantibodies of 20 patients with autoimmune hemolytic anemia (AHA) were used in immunoaffinity assays with surface-radioiodinated human red blood cells (RBCs), and detergent-solubilized products were analyzed by SDS-PAGE/autoradiography. Four membrane proteins were identified as candidate autoantigens: a nonglycosylated polypeptide with an apparent molecular mass of 34 kD (p34) that was expressed in all available RBC phenotypes except Rbnull but differed consistently in apparent molecular mass from the 32-kD Rh(D) polypeptide coisolated by IgG allo-anti-D; a heterogeneous 37–55-kD glycoprotein, also deficient in Rbnull RBCs, which disappeared after deglycosylation by N-glycanase, with the appearance of a sharp, new ~31-kD band distinct from p34 and from Rh(D) polypeptide; a ~100-kD major membrane glycoprotein identified by immunoblotting as the band 3 anion transporter; and glycophorin A (GPA), also confirmed by immunoblotting. GP37-55 was not seen in the absence of p34, and both proteins are likely to be members of the Rh family. Indeed, a 34-kD polypeptide band and 37–55-kD polypeptide “smear,” isolated concurrently from the same labeled RBCs by IgG allo-anti-e, were indistinguishable from their autoantibody-isolated counterparts and may well be the same protein identified at different epitopes by the auto- and allo-antibodies. Individual AHA patients’ autoantibodies isolated p34 and gp37-55, alone or in combination with band 3 (nine cases); strong band 3 alone (five cases); and combinations of band 3 with GPA (six cases). The autoantibodies of three additional patients whose AHA had been induced by a-methyldopa also isolated p34 and gp37-55. (J. Clin. Invest. 1993. 91:1672–1680.) Key words: erythrocyte autoantigens • autoimmune hemolytic anemia

Introduction

Autoimmune hemolytic anemia (AHA)1 was one of the first human diseases to be recognized as an authentic autoimmune disorder (2, 3). Although the effector mechanisms governing such antibody-mediated destruction of human red cells have been increasingly clarified over many years (4, 5), fundamental questions concerning the genesis of the erythrocyte-reactive autoantibodies have remained unanswered. With respect to the more common, “warm antibody” type of human AHA, knowledge of the red blood cell (RBC) membrane structures with which such autoantibodies react had long been based mainly on serological (indirect antiglobulin) testing with human RBCs of rare phenotypes (for review see reference 6). These autoantibodies characteristically react with essentially all human RBCs and with the RBCs of higher (old world) primates but not with RBCs of other species (2). In a significant proportion of AHA patients, specificity of their autoantibodies for one or more determinants of the rhesus (Rh) blood group had been deduced from serological observations including selectively weak or negative reactions with Rhnull RBCs, which lack any known expression of the Rh complex (6–10); or selective or preferential reactivity with RBCs expressing known Rh antigens such as “e,” “E,” or “c,” also present on the patient’s own RBCs (2, 6, 8, 9). The latter antibodies were usually found in combination with less well-defined autoantibodies. Cautious interpretation of these observations was warranted, however, by the knowledge that Rhnull RBCs have membrane anomalies other than deficiency of Rh antigens (11) and by the fact that careful serological analysis has suggested subtle differences between, e.g., auto-anti-e and allo-anti-e (12). Moreover, in many other AHA patients, the reactivity of their autoantibodies with Rhnull RBCs and RBCs of “normal” phenotype was equally strong, suggesting that in these patients the autoantibodies recognized RBC determinants distinct from known components of the Rh complex (6–10). Furthermore, through the use of human RBCs selectively deficient in other blood group antigens, evidence has accumulated in selected AHA patients for autoantibody specificity against serologically defined, non-Rh RBC antigens such as Wr6 (13), En4 (14), LW15, U16, or antigens of the Kell blood group (17). Efforts to elucidate the biochemical nature of the RBC autoantigens in AHA were initiated by one of us (J. P. Leddy) > 20 yr ago (18) but were set aside until more effective methods became available. More recent immunochemical studies have concluded that the major anion-transport protein, band 3 (19), or band 4.1 protein (20) serve as targets of these autoantibodies in AHA.

In this report we present evidence that four distinct membrane proteins may participate in the molecular composition of RBC autoantigens in AHA. Two appear to be minor RBC

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1. Abbreviations used in this paper: AHA, autoimmune hemolytic anemia; α-MD, alpha-methyldopa; GPA, glycophorin A; IAT, indirect antiglobulin test; NC, nitrocellulose; RBC, red blood cell; Rh, rhesus blood group; RT, room temperature.
proteins and members of the Rh family. The other two are major membrane glycoproteins, band 3 and glycoporphin A (GPA).

Methods

Anti-red cell antibodies. The autoantibodies used were in the form of concentrated eluates from the RBCs of patients with warm-antibody AHA (Table I). After informed consent, blood was drawn into 0.01 M EDTA and eluates were prepared either immediately or after no more than overnight storage at 4°C. The patients’ RBCs were washed six times (20 vol/vol packed RBCs) and membranes were prepared by lysis in digitonin, 0.25 mg/ml final concentration (Sigma Chemical Co., St. Louis, MO), as previously described (10). After five more washes of the membranes in cold saline, the antibodies were eluted in glycine-HCl, pH 3.0, and immediately dialyzed against cold PBS, pH 7.3. Up to five cycles of elution were required to dissociate all recoverable antibody. Indirect antiglobulin test (IAT)-positive eluates from the various cycles were pooled and concentrated by ultrafiltration. Patients were included in the study only on the basis of adequate strength of their eluates.

To prepare IgG anti-Rh(D) alloantibody eluates, group O RBCs, freshly obtained from healthy laboratory personnel whose probable Rh genotypes were Rr, Rr, (CDE/cDE) or Rr, Rr, (CDE/cDE), were washed and incubated (37°C, 1 h) with commercial hyperimmune human anti-D (Hyp-Rho; Cutter Biologicals, Elkhart, IN; or Rho GAM; Ortho Pharmaceutical, Raritan, NJ) in the ratio of 10 ml packed RBCs to 1 ml anti-D. This was slight antibody excess. The resulting anti-D-coated RBCs were then treated identically to autoantibody-coated RBCs. “Control eluates” were prepared by the same procedures from comparable numbers of unsensitized RBCs of the same healthy Coombs-negative donors. Eluates were stored at -20°C until used. Human serum containing IgG anti-e alloantibody from an individual donor was obtained from Sero-Tec Biologicals (North Brunswick, NJ).

Rabbit antisera to human erythrocyte band 3 (R334 and 1514B) were generously donated by Dr. Marguerite M. B. Kay (University of Arizona College of Medicine, Tucson, AZ). Murine IgG1 mAb to human glycoporphin A (10F17MNN) (21) was provided by Dr. R. J. Looney (University of Rochester) as culture supernatant from a hybridoma obtained from American Type Culture Collection (Rockville, MD).

Serological testing, by direct and indirect antiglobulin reactions, was performed by standard methods as described (10). Comparative reactivity of concentrated AHA eluates with normal and Rh null RBCs was determined by serial twofold titration using indirect antiglobulin reactions. The eluates for these assays were not always the fully concentrated eluates used in the radio-immunoprecipitation assays described below. It was our practice to set aside a small volume of partially concentrated eluate expressly for serological testing, to minimize freezing and thawing of the fully concentrated eluate.

Surface radioiodination of erythrocytes. Normal RBCs were freshly obtained (in citrate) from healthy group O donors of probable Rh genotypes Rr, Rr, Rr, Rr (CDE/cDE), or rr (cc/ce), immediately sedimented to allow aspiration of buffy coat, and washed twice in cold 0.15 M saline with further removal of the uppermost cellular layer. The RBCs were placed in Alsever’s solution, electronically counted (together with monitoring of leukocyte and platelet contamination), and used either the same day or, more commonly, after overnight storage at 4°C. Group O Rhnull red cells from two sibling donors (Sa and Na) were generously provided by Dr. Chantal Harrison, Blood Bank Director (University of Texas Health Sciences Center, San Antonio, TX). These cells were either freshly drawn, kept in their own plasma during shipping, and then placed in Alsever’s solution or reconstituted from frozen storage. These alternative methods of handling did not alter the results.

For radiolabeling, 10^8 previously washed RBCs in Alsever’s solution were washed twice in cold 0.15 M NaCl and the packed RBCs, after resuspension to 0.7 ml in PBS, were added to flat-bottomed glass vials precoated with 50 μg chloroglycoursil (Iodo-Gen; Pierce Chemical Co., Rockford, IL). 1 μCi carrier-free Na[125I] (Amersham Corp., Arlington Heights, IL) was added for 30 min at room temperature (RT) with gentle swirling every 10 min. The labeled RBCs were immediately transferred to a clean 15-ml conical tube containing 5 mM KI in PBS, washed twice more in 5 mM KI/PBS, transferred to a second clean tube, and washed once more in PBS. The packed cells were then brought to 0.4 ml with PBS. Alternatively, the same volume of RBCs was radioiodinated using Iodo-Gen beads (Pierce Chemical Co.) and 300 μCi of Na[125I]. By either procedure ~140 μCi[125I] was bound per milliliter packed RBCs.

Immunofluorescence isolation of RBC membrane proteins. Our general approach was adapted from earlier work on the Rh(D) protein (22, 23). In 0.65-ml microcentrifuge tubes (Click Seal; National Scientific Co., San Rafael, CA), 50 μl labeled RBCs (1.25 x 10^8 cells) plus 90 μl PBS were incubated with 40 μl AHA eluate (or with control eluate or anti-D eluate) for 90 min at 37°C on a rotator. The mixtures were transferred to 4-m1 polycarbonate tubes (Sorvall Instruments Div., part of DuPont Co., Newtown, CT) and washed three times with cold PBS. The washed RBCs were resuspended in 0.05 ml PBS and lysed by adding 1 ml cold 5 mM sodium phosphate, pH 8.0, containing 1 mM PMSF (Sigma Chemical Co.) and 10 mM EDTA, with Vortex mixing (30 s) and standing 30 min on ice. Lysates were centrifuged for 30 min at 35,000 g at 4°C (RC2-B, SS-34 rotor, Sorvall Instruments) to pellet RBC ghosts, which were then washed twice with cold hypotonic saline. Washed packed ghosts were solubilized in 125 μl freshly prepared peroxide-free 2% Triton X-100 (Boehringer-Mannheim Corp., Indianapolis, IN) in cold PBS containing 1 mM PMSF and 10 mM EDTA, with Vortex mixing for 10 min at RT. After centrifugation at 35,000 g for 30 min, the immune complex–containing supernatants were transferred to 1.7-ml microcentrifuge tubes (Click Seal) containing 25 μl packed protein G-Sepharose beads (Zymed Laboratories Inc., South San Francisco, CA), which had been prewashed eight times with 2% Triton X-100 in PBS. After overnight incubation of the solubilized RBC antigen–antibody complexes with the beads on a rotator at 4°C, the beads were washed twice with cold 2% Triton X-100 in PBS, twice with 1% Triton X-100 in PBS, and twice with 5 mM phosphate, pH 8.0. Substitution of goat anti–human IgG-Sepharose for protein G-Sepharose yielded comparable results.

The Sepharose pellets (25 μl) were suspended in 25 μl Laemmli sample buffer (24) containing 2% SDS and, in the experiments shown, 20 mM DTT, and incubated at 100°C for 2 min. After microcentrifugation, 20 μl of each supernatant was applied to a gradient polyacrylamide slab gel in 0.1% SDS using a Mini-Protein system (Bio-Rad Laboratories, Richmond, CA). Two types of gradient gel proved most effective: 5–15% with a 37:1 acrylamide/bis ratio; and 8–16% with a 42:1 acrylamide/bis ratio. In some experiments on deglycosylation by N-glycosidase (see below), a 10–20% gradient gel was chosen. The completed gels were dried and analyzed by autoradiography using XAR-2 film (Eastman Kodak Co., Rochester, NY) with Cronex intensifying screens (DuPont Co., Wilmington, DE). The presence or absence of DTT did not alter the positions of the proteins under study. Radiola beled (“rainbow”) molecular weight markers (Amersham Corp.) included myosin (200,000), phosphorylase b (92,500), BSA (69,000), ovalbumin (46,000), carbonic anhydrase (30,000), soybean trypsin inhibitor (21,500), and lysozyme (14,300). The apparent molecular mass of a given protein band was estimated graphically (25).

Immunoblotting. In selected experiments, radioiodinated RBC membrane proteins separated in SDS-PAGE gels were electrophoretically transferred to nitrocellulose (Transphor TE-50; Hoefer Scientific Instruments, San Francisco, CA). The nitrocellulose membrane was blocked with 3% BSA (2–3 h at RT) followed, in appropriate experiments, by sequential addition of 1:100 rabbit anti–band 3 (2 h) and 1:500 peroxidase-labeled goat anti–rabbit IgG (Kirkegaard & Perry Laboratories, Inc., Gaithersburg, MD), (2 h) or 1:10 mouse anti–GPA mAb and 1:250 peroxidase-labeled goat anti–mouse IgG (Kirkegaard & Perry Laboratories, Inc.). Development was with 4-chloro-1-naphthol/hydrogen peroxide substrate (Bio-Rad Laboratories). Autoradiographic

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Table I. Summary of Background Data and New Findings in Patients with Autoimmune Hemolytic Anemia

<table>
<thead>
<tr>
<th>Patient</th>
<th>Clinical diagnosis</th>
<th>Direct antihuman globulin test*</th>
<th>RBC membrane proteins identified by autoantibody affinity†</th>
<th>Serological reactions of autoantibodies with intact RBCs‡</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>p34</td>
<td>gp37-55</td>
</tr>
<tr>
<td>LM</td>
<td>1° AHA</td>
<td>IgG</td>
<td>Strong</td>
<td>Moderate</td>
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<tr>
<td>SH</td>
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<td>IgG</td>
<td>Strong</td>
<td>Moderate</td>
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<tr>
<td>JT</td>
<td>2° AHA; Hodgkins</td>
<td>IgG (C3nt)</td>
<td>Strong</td>
<td>Moderate</td>
</tr>
<tr>
<td>RE</td>
<td>1° AHA</td>
<td>IgG</td>
<td>Moderate</td>
<td>—</td>
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<tr>
<td>CR</td>
<td>1° AHA; SLE ?</td>
<td>IgG/C3</td>
<td>Moderate</td>
<td>Weak</td>
</tr>
<tr>
<td>RB</td>
<td>2° AHA; Hodgkins</td>
<td>IgG</td>
<td>Moderate</td>
<td>Weak</td>
</tr>
<tr>
<td>SM</td>
<td>1° AHA</td>
<td>IgG (C3nt)</td>
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<td>Weak</td>
</tr>
<tr>
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<td>1° AHA</td>
<td>IgG</td>
<td>Moderate</td>
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<tr>
<td>AL</td>
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<td>IgG</td>
<td>Weak</td>
<td>—</td>
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<tr>
<td>IB</td>
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<td>IgG</td>
<td>—</td>
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<tr>
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<td>IgG/C3</td>
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<td>BH</td>
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<td>IgG/C3</td>
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<tr>
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<td>IgG/C3</td>
<td>—</td>
<td>—</td>
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<tr>
<td>SS</td>
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<td>IgG/C3</td>
<td>—</td>
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</tr>
<tr>
<td>CV</td>
<td>2° AHA; SLE</td>
<td>IgG/C3</td>
<td>—</td>
<td>—</td>
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* DAT was typically performed with both rabbit anti-IgG and anti-C3, however, entries designated C3nt mean that test results with anti-C3 were not available in that case. A DAT result indicating IgG alone means that testing with anti-C3 was negative. † Listed here are autoradiographic bands that appeared to be clearly above background patterns obtained with control eluates (see text). ‡ Indirect antihuman globulin reactions were performed using twofold dilutions in isotonic saline with unmodified RBCs. Full titrations were not always performed because of the limited supply of RBC_mll cells. Eluate AL was only tested undiluted. Note that the eluates used for serological testing were not always the fully concentrated eluates used in the radioimmunoprecipitation assays (see Methods). The titers are the highest dilutions giving macroscopically detectable agglutination. Results listed as negative signify that there was no reaction over the full range of dilutions tested, including undiluted eluate. Weak, moderate, and strong refer to the strength of the agglutination at the highest autoantibody concentration tested. § Var., variably detected, depending on date of bleeding.
visualization of the same blot followed the procedures for direct autoradiography of the SDS-PAGE gels described above.

Deglycosylation. The effect of recombinant N-glycanase (Genzyme Corp., Cambridge, MA) could not be tested on RBC proteins that had been freshly eluted from protein G beads because of enzyme inactivation by 1% SDS. Therefore, a procedure was devised for in situ digestion of the RBC proteins while still bound to the protein G beads (before SDS elution). Our standard immunoaffinity isolation protocol (see above) was followed through the final washings in 2 and 1% Triton X-100, just before elution in SDS. At that point the beads were washed twice in cold “enzyme buffer” (20 mM phosphate, 100 mM NaCl plus PMSF and EDTA, pH 7.6). To each sample (25 μl) of pelleted beads was added 100 μl N-glycanase (0.25 U) in enzyme buffer. Control pellets received buffer alone. Except for kinetic assays (see below), the beads were incubated on a rotator for 4 h at 37°C followed by washing three times with cold buffer. The washed beads were then eluted in Laemmlli buffer and the eluted proteins subjected to SDS-PAGE/autoradiography.

Results

Rh-related membrane proteins. Two proteins in this category were isolated from surface-radioiodinated RBCs by AHA autoantibodies, one producing a dense autoradiographic band of 34 kD (p34) and the other a fainter, polydisperse “smear” (gp37–55) (Fig. 1). 9 of our 20 AHA patients’ autoantibodies mediated the isolation of p34 (Table I). (The mean measured M₁ in these nine cases was 33.5 kD.) In five of these patients (LM, SH, JT, RE, CR), p34 was the dominant band isolated (Table I). In the other four patients (RB, SM, LP, AL), the autoradiographs identified p34 plus at least one other relatively strong protein band of 95–100 kD or of ~ 41 kD, which, from evidence to be presented below, are band 3 and GPA, respectively. In dozens of experiments, p34 exhibited a small but very consistent difference in M₁ from the 32-kD Rh(D) polypeptide coisolated from the same labeled RBCs using allo-anti-D (see Figs. 1–3). A 34-kD protein band corresponding to p34 was difficult to detect in autoradiographs of unmodified whole RBC lysates (one example is shown below, see Fig. 3 b), suggesting that isolation of p34 by autoantibody-directed immuno-

Figure 1. Autoradiograph of SDS-PAGE separation of 125I-labeled membrane proteins immunoaffinity isolated from normal (O CDe/CDe) human RBCs by anti-Rh(D) alloantibody eluate (lane C); autoantibody eluate of patient LM (lane E); and control eluates (C) (lanes B and D). Molecular mass markers are shown in lane A.

Figure 2. SDS-PAGE autoradiograph of membrane proteins isolated from normal group O Rh(D)-positive (CDe/CDe) and Rh(D)-negative (ce/ce) RBCs by autoantibody eluate, patient LM (lanes A and B); anti-Rh(D) alloantibody eluate (lanes C and D); and autoantibody eluate, patient LP (lanes E and F). Both LM and LP eluates have isolated p34 and gp37–55.

affinity involves considerable concentration of a minor RBC protein. As illustrated in Fig. 2, by eluates LM and LP, p34 was isolated by such autoantibodies equally well from D-positive (O CDe/CDe) and D negative (O ce/ce) red cells, in contrast to the Rh(D) polypeptide that was isolated only from D-positive RBCs. Neither p34 nor the Rh(D) polypeptide, however, were detected in concurrent studies of radiolabeled Rhnull RBCs, as illustrated by eluate SH in Fig. 3 a. Each of six anti-p34 autoantibody samples tested failed to identify a 34-kD band in Rhnull RBCs. Allowances were made in such studies for possible unequal radioiodination of paired Rhnull and normal cells by loading gels, alternatively, with equal cpm or with equal volumes of the respective elutions from the protein G beads. Fig. 3 b demonstrates that the Rhnull RBCs were effectively radioiodinated. Moreover, other autoantibodies, e.g., eluate IB with pure anti-band 3 reactivity in Fig. 3 a, isolated strongly labeled ~ 100-kD bands from both Rhnull and normal RBCs.

Most of the p34-reactive autoantibody eluates (see Table I) produced an additional diffuse autoradiographic “smear” from ~ 37 to 55 kD (Figs. 1–3), suggesting variable glycosylation of the target protein. When labeled Rhnull RBCs were tested, this heterogeneous smear, like the p34 band, was not evident (Fig. 3 a). Precedent for Rh-related glycoproteins producing polydisperse autoradiographic images over a similar range of molecular mass is discussed below.

The effect of stripping N-linked carbohydrates by treatment of the immunoaffinity-isolated RBC membrane proteins with recombinant N-glycanase before SDS-PAGE was tested (see Methods). As expected (22, 23, 26), this treatment had no
effect on the Rh(D) polypeptide band obtained with allo-anti-D (not shown), nor was there a detectable effect on the mobility or density of the autoantibody-isolated p34 band (Fig. 4 a). However, the polydisperse 37-55-kD smear produced by these same autoantibody eluates disappeared after N-glycanase treatment, with the appearance of a sharp new ~31-kD band clearly below the usual position of p34 (Fig. 4 a) and, in experiments not shown, just below the Rh(D) polypeptide band. Resolution of this area to reveal this double-band pattern was aided by switching to a 10-20% gradient gel. The Rh phenotype of the RBC donor had no effect on the autoradiographic pattern observed after N-glycanase. The prominent new bands at ~69 and 90 kD appearing in lanes B and C have not been fully characterized but may result from the (4-h) action of N-glycanase on band 3 or its multimers. In separate experiments (not shown) bands of similar molecular mass became evident when whole lysates of unsensitized 125I-RBCs were treated with N-glycanase (4 h) before SDS-PAGE/autoradiography. Such bands were much less prominent after shorter incubations with N-glycanase (see below, Fig. 5).

Multiple experiments with N-glycanase were run both in 10-20% gels, in which the new ~31-kD band was consistently separable from what appeared to be typical p34 and in our standard 8-16% gels, wherein p34 and Rh(D) polypeptide maintain their typical relative positions but the “new” deglycosylated protein appeared as a downward broadening of the p34 band with incomplete resolution into two bands (not shown). We propose that the lower (~31-kD) band is the deglycosylated form of a variably glycosylated membrane glycoprotein (gp37-55) responsible for the polydisperse smear observed in untreated samples. This protein is apparently distinct from p34 since their deglycosylated polypeptides differ in Mr, under appropriate gel conditions (Fig. 4 a). The possibility that the 31-kD component in the double band (Fig. 4 a) represents a glycanase-induced shift of p34 itself whereas the 34-kD component arises by deglycosylation of gp37-55 is made most unlikely by kinetic studies with N-glycanase presented below.

Other investigators have demonstrated that alloantibodies to the E/e or C/c Rh proteins produce both sharp ~33-34-kD polypeptide bands (23, 27) plus polydisperse glycoprotein smears over a range of Mr similar to that produced by our autoantibodies (27) and that these E/e- or C/c-related patterns generally differ from the patterns observed with the Rh(D) polypeptide (23, 27, 28) or D-related glycoprotein (27). In our hands, allo-anti-e isolated both a 34-kD band indistinguishable from the autoantibody-isolated p34 polypeptide and a polydisperse smear that closely resembled autoantibody-produced gp37-55 (Fig. 4 b). Neither of these proteins was isolated by allo-anti-e from radiolabeled O cDe/cDe RBCs (not shown), whereas p34 and gp37-55 could be isolated by autoantibodies from RBCs of any common Rh phenotype. The effect of deglycosylation on these anti-e–isolated proteins was identical to that just demonstrated (Fig. 4 a) with autoantibody-isolated p34 and gp37-55. This is illustrated in Fig. 5 in which an autoantibody eluate (JT) recognizing p34 and gp37-55 was tested in parallel with allo-anti-e in a kinetic study with N-glycanase. The stepwise disappearance of the glycoprotein smears and emergence of the ~31-kD deglycosylated polypeptide displayed strikingly similar time courses (Fig. 5).

The three bands appearing in the 24-29-kD region of lane A in Fig. 4 b were not a consistent finding with anti-e (e.g., see Fig. 5). We have occasionally seen such 24-29-kD bands with other Rh allo- or autoantibodies but not in a reproducible fashion. Whether such lower mol wt bands are related in any way to the truncated Rh mRNA isoforms recently reported by Kim et al. (29) cannot be resolved without systematic study.

We were unable to obtain a sufficiently potent or high affinity anti-c or anti-C to permit us to compare the Rh C/c proteins in the same assays.

**Band 3.** A completely different pattern of membrane protein isolation was exhibited by the autoantibodies of 11 other AHA patients (IB through CV in Table 1). Their autoradiographs consistently displayed an intense ~100-kD band with no detectable isolation of p34 (eluate IB, Fig. 3 a). In contrast to p34, this 100-kD protein was isolated equally well from
\( \text{Rb}_{ab} \) and normal RBCs (Fig. 3a). This autoantibody-isolated 100-kD band appeared indistinguishable from the autoradiographic band isolated from the same labeled RBCs by rabbit anti-band 3 serum (not shown). Moreover, when this 100-kD protein was initially isolated from \( ^{125}\text{I}-\text{labeled} \) RBCs by appropriate human autoantibody eluates, subjected to SDS-PAGE, and then electroblotted to nitrocellulose (NC), development of the blot with specific rabbit antiband 3 (Fig. 6b) revealed a 100-kD band corresponding to the position of the autoantibody-isolated 100-kD band visualized by autoradiography (Fig. 6a). Such 100-kD bands, moreover, coincided electrophoretically with a radiiodinated protein of the same apparent molecular mass in whole RBC lysates (Fig. 6a), as would be expected for band 3. The 41-kD bands in lanes A and B (Fig. 6a) represent GPA that was not detectably isolated by ST eluate from this bleeding date.

It should be noted that essentially all of our autoradiographs contained a band of varying density in the band 3 area, including faint “background” bands observed with the following controls: substitution of normal control eluates (see Methods) for autoantibody eluates in the initial coating of the intact \( ^{125}\text{I}-\text{labeled} \) RBCs; use of D-negative RBCs in studies with anti-D eluates, a pairing that yields no detectable 32-kD Rh(D) band; and omission of all anti-RBC antibody sources and incubating unsensitized \( ^{125}\text{I}-\text{RBC} \) lysates with protein G beads. This experience agrees with that of other workers who have also commented on the ubiquity of background quantities of band 3 in autoradiographs (23). This does not mean, however, that the very dense autoradiographic bands obtained with many of the autoantibody eluates just discussed are nonspecific. These bands reproducibly stood out from the “background” bands observed with the controls. On the other hand, weak band 3 images obtained with other autoantibody eluates, which were not distinguishable from those produced by concurrently run control eluates, were omitted in compiling the summary data in Table I.

The very high mol wt bands seen in some autoradiographs presumably represent aggregates of band 3, which has a known propensity to aggregate under these conditions (30).

GPA. Another specificity was suggested by the reproducible appearance of sharp bands with a \( M_{r} \) of 41 kD in autoradiographs from certain AHA patients (Fig. 7a and Table I). The \( \sim 41\text{-kD} \) protein isolated by these autoantibodies coincided with the position of GPA when whole lysates of \( ^{125}\text{I}-\text{RBC} \) were analyzed in the same gels (Fig. 7a). The affinity-isolated proteins from two of these patients (SS and GF) were electrophoresed from the SDS-PAGE gel to NC and the 41-kD bands were confirmed as GPA monomer by reactivity with specific mAb (Fig. 7b). In addition, higher molecular mass bands of GPA dimer were now clearly recognizable in positions just below band 3 (Fig. 7b). N-glycanase treatment of autoantibody-bound GPA produced a shift to a lower molecular mass but had no effect on the intensity of the GPA bands in direct autoradiographs of SDS-PAGE gels (not shown).

None of the eluates immunoprecipitated GPA alone (Table I). For this reason we had some concern that its appearance in our autoradiographs might reflect nonspecific coprecipitation of this abundant RBC protein with specific immune complexes involving band 3 or p34 plus gp37–55. On the other hand, isolation of GPA was found only with certain AHA eluates and was reproducible. Moreover, GPA was not detectably coprecipitated by some of our most potent antiband 3 or anti-p34 eluates (Figs. 1–6 and Table I).

We will discuss below the possibility that eluates that isolate both GPA and band 3 might have specificity for the \( \text{Wr}^{b} \) antigen, which is thought to be formed by a GPA–band 3 interaction (31). The coisolation of these two proteins by a significant proportion of our patients (Table I) recalls their coisolation by mouse monoclonal anti-\( \text{Wr}^{b} \) (31).

\( \alpha \)-Methylidopa (\( \alpha \)-MD) associated autoantibodies. The autoantibody populations listed in Table I were from “spontaneously arising” AHA cases. Three additional autoantibody eluates were available, in frozen storage, from patients whose hemolytic anemia had been associated with \( \alpha \)-MD therapy. These eluates, which were less potent by IAT than the “sponta-

\[ \text{Figure 5. Kinetic study with N-glycanase comparing effects on membrane proteins isolated from O ce/ce RBCs by IgG allo-anti-e and autoantibody eluate JT. The arrowheads mark the position of p34 or e polypeptide and of the enzyme-induced product termed p31.} \]

\[ \text{Figure 6. Identification of band 3. (a) Autoradiograph of the nitrocellulose electroblot from SDS-PAGE separation of 1:5 and 1:10 dilutions of whole RBC lysate (lanes A and B) and of membrane proteins affinity isolated by autoantibody eluate ST (lane C) and by a control eluate (lane D). (b) ELISA development of the same NC blot by sequential addition of specific rabbit antiserum to human band 3, horseradish peroxidase-goat anti-rabbit IgG, and 4-chloro-1-naphthol substrate.} \]
neous" AHA eluates studied, each isolated weak but definite 34-kD bands that were electrophoretically indistinguishable from p34 coisolated from the same RBCs in the same gel by "spontaneous" AHA autoantibodies (not shown). Trace quantities of gp37-55 were also present; after N-glycanase treatment a sharp 31-kD band emerged in addition to p34.

Correlation with serological reactions against intact red cells. Table 1 (last two columns) summarizes data on IAT reactions of the autoantibodies with intact RBCs of normal and Rh<sub>Hall</sub> phenotypes. Eluates that strongly isolated band 3 protein alone or in combination with GPA in the immunoaffinity assays reacted strongly and equally with both intact normal and Rh<sub>Hall</sub> RBCs. Conversely, a majority of the autoantibody eluates that mediated a prominent isolation of p34 displayed either negative or clearly weaker IAT reactions with Rh<sub>Hall</sub> cells. There were two exceptions. The eluate of patient LP appeared to isolate both p34 and band 3 but did not react with Rh<sub>Hall</sub> RBCs by IAT in the dilutions tested. Conversely, patient CR displayed equal reactivity with Rh<sub>Hall</sub> and normal RBCs by IAT despite the fact that her immunoaffinity results with labeled RBCs had suggested both anti-p34 and anti-band 3 specificities. It is possible that anti-band 3 (in patient CR) and anti-p34 (in patient LP) are more dominant in the original eluate than was reflected in the autoradiographic patterns. Perhaps the autoradiographic image of band 3 produced by LP eluate is nonspecific despite our careful effort to distinguish background from specific bands. Alternatively, patient CR could have still another autoantibody population reactive with an RBC antigen that is not radioiodinated under our conditions. Not shown in Table 1 is the fact that in each of our three α-MD-associated cases the autoantibodies were selectively unreactive with Rh<sub>Hall</sub> RBCs, in keeping with their isolation of p34 and gp37-55.

Discussion

By means of an immunoaffinity procedure using purified autoantibodies (RBC eluates) and surface-radiolabeled human RBCs with analysis by SDS-PAGE/autoradiography, the IgG (warm-reacting) anti-RBC autoantibodies of 20 AHA patients have been used to identify erythrocyte membrane proteins potentially serving as target autoantigens. Our results indicate that at least four single-chain proteins may fulfill this role. Two appear to be members of the Rh family. One, which we have provisionally designated p34 on the basis of its apparent molecular mass, appears to be a minor RBC membrane protein; is lacking or severely deficient in Rh<sub>Hall</sub> RBCs; was consistently distinguishable by its slightly higher molecular mass from the well-studied 32-kD Rh(D) polypeptide; and, like the Rh(D) polypeptide, is judged to be a nonglycosylated polypeptide on the basis of the studies with N-glycanase (Figs. 4a and 5).

The second apparently Rh-related protein identified by these autoantibodies appears to be a variably glycosylated glycoprotein (termed gp37-55), which was also deficient in Rh<sub>Hall</sub> RBCs. Such a protein has a well-documented precedent in the Rh-related glycoproteins (27, 32, 33). The latter are reported to have differing NH<sub>2</sub>-terminal sequences from the nonglycosylated Rh polypeptides (32) and to produce polydisperse autodisperse radiographic patterns over a range of molecular mass encompassing the 37–55-kD smear isolated by our autoantibody eluates (Figs. 1–5). Such Rh-related glycoproteins have been isolated both by anti-D, anti-e, or anti-E alloantibodies (27) and by certain murine mAbs reactive with Rh proteins (32). For reasons not yet clear, such polydisperse radiographic images were not observed in our many studies with anti-Rh(D) eluates tested concurrently with our anti-p34 autoantibody eluates (Figs. 1, 2, 3a, and 4b).

In the present study, deglycosylation of gp37-55 yielded a polypeptide of Mr ~ 31 kD that was electrophoretically separable from N-glycanase-treated p34 and, therefore, gp37-55 is apparently a distinct protein rather than glycosylated variants of p34. GP37-55 was observed in our autoradiographs only when p34 was also evident. The converse, i.e., the appearance of the p34 band without the gp37-55 smear, occurred only with weaker anti-p34 eluates (Table 1) and could well be a quantitative phenomenon, reflecting the greater difficulty in visualizing a polydisperse pattern. Whether the autoantibodies reactive with p34 and gp37-55 recognize shared or differing epitopes on these two proteins is not yet clear. The possibility also exists that only one of these two proteins, p34 or gp37-55, carries the relevant autoepitope(s) and that the other protein is coprecipitated because of noncovalent bonding to the epitope-bearing protein. Alternatively, it is conceivable that the autoantigenic epitope results from an interaction of these two proteins (see discussion of Wr<sup>i</sup> below). Such points had been
raised earlier with respect to the glycoproteins isolated by Rh alloantibodies or Rh-related mAbs (11, 32, 33).

As noted earlier, Rhnull RBCs are deficient in proteins other than the Rh family. Thus, our inability to detect p34 and gp37–55 in Rhnull RBCs does not in itself provide unambiguous evidence that these proteins belong to the Rh family. However, there are other observations consistent with this interpretation. The Rh polypeptides are thought to be virtually unique among membrane proteins in being nonglycosylated (33) and p34 appears to have this property. On the other hand, gp37–55 is remarkably similar to the Rh-related glycoproteins, particularly those isolated by anti-c and anti-E (27) and by anti-e (Figs. 4 and 5). Moreover, a large body of serological data had strongly suggested that one or more proteins of the Rh complex serve as target antigens in many cases of AHA (6–10, 13) and in the great majority of α-MD–associated autoantibodies (34).

Extrapolating from our findings in the light of recent advances in our molecular understanding of the Rh system (33), we propose but have not proven that p34 and the 33–34-kD polypeptide carrying the Rh E/e alloantigens are the same molecule, but with the autoantibodies recognizing a public epitope distinct from the E/e alleloantigenic epitopes. The similarity of autoantibody-isolated gp37–55 to the diffuse glycoprotein isolated by anti-e alloantibody is striking (Fig. 5). We could not adequately test this hypothesis in relation to the Rh C/c proteins but, given the close structural similarity of C/c and E/e polypeptides (35) and their possible encoding by a single gene (36), we suspect that autoreactive epitope(s) may prove to be present on C/c as well as E/e. This concept would fit with observations that the C/c and E/e polypeptides share (with p34) a slightly higher apparent molecular mass than the D polypeptide (23, 27, 28) and with the accumulated serological experience that when RBC autoantibodies exhibit apparent specificity for individual Rh antigens (i.e., those defined by alloantibodies), it has frequently been for E/e or C/c but rarely for D (2, 6–8, 13). Alternatively, it cannot be excluded that p34 and gp37–55 could be distinct from any of the known alloantigen-expressing proteins of the Rh complex but still members of the Rh family. It is of much theoretical interest that, in each of three patients whose AHA was induced by α-MD therapy, autoantibodies to p34 and to gp37–55 were also found. Whether anti-p34 antibodies arising in the setting of α-MD therapy recognize precisely the same epitope(s) on these proteins as those occurring in "spontaneous" AHA remains to be seen.

Despite two exceptions noted above, autoantibody-mediated isolations of RBC membrane proteins correlated well with their serological reactivity against normal and Rhnull RBCs (Table 1). These observations may serve as a first step toward a molecular understanding of these long-recognized differences in serological behavior among AHA autoantibodies.

Two other RBC membrane proteins that are not members of the Rh family have also been identified as candidate autoantigens in our studies, the band 3 anion transporter and GPA. Band 3 appeared to be the sole or dominant specificity in seven cases and in many others was found in combination with anti-GPA or with anti-p34 or both (see Table 1). The issue of non-specific autoantibiotic images of band 3 was addressed in Results. Having used control eluates in nearly every experiment as "noise" monitors, we feel that the band 3 isolations illustrated in Figs. 3 and 6 and summarized in Table I are genuine. Victoria et al. (19) have also reported autoantibody reactivity with band 3 in 12 AHA patients. Their study did not implicate other RBC membrane proteins as potential autoantigens in AHA.

The final membrane protein identified by these patients’ autoantibodies was GPA. For the reasons given in Results, we do not regard the presence of GPA in our autoradiographs to be nonspecific. Isolation of this protein is consistent with evidence that GPA carries the En* antigen (37) and interacts with band 3 to form the Wr* antigen (31). Both had been identified serologically as non-Rh specificities in AHA (13, 14), with anti-Wr* being relatively common (13). Unfortunately, RBCs of the very rare Wr(a+b−) or En*(−) phenotypes were not available for concurrent study.

Murine monoclonal anti-Wr* isolates both GPA and band 3 from radiolabeled RBCs (31). On this basis the last six patients listed in Table I and, more weakly, patients JT and RE, all of whose autoantibodies isolated both GPA and band 3, might have anti-Wr* alone or in combination with other antibodies. In Issitt et al.’s (13) series, most auto-anti-Wr* antibodies were accompanied by other specificities. The presence of “pure” anti-Wr* in our own cases is perhaps made less likely by the finding that all five of the eluates that strongly isolated both GPA and band 3 were from patients who exhibited in vivo C3 binding to their RBCs (Table I) whereas Issitt et al. (13) concluded that none of their four examples of pure anti-Wr* autoantibodies was complement fixing.

In one of our patients (ST), from whom multiple eluates were available over a multiyear period, a change in reactivity with GPA or in the ratio of anti-GPA to antiband 3 appeared to occur without a major change in reactivity with band 3 or a correlating shift in total IgG content of eluates. This suggests that autoantibodies reactive with GPA and those reactive with band 3 may vary independently in some instances. Anti-Wr* (or another antibody reactive with GPA) might have been produced at certain times in the patient’s course but other antibodies apparently reactive with band 3 itself dominate throughout. Similarly, patients IB, AR, RL, BH, and FY (Table I), whose eluates consistently and strongly isolated band 3 alone, appear to have a distinct specificity whose serological counterpart, if there is one, remains to be clarified. The biochemical correlates of the entire group of non-Rh–related autoantibody specificities clearly require further study.

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Note added in proof. While this manuscript was under review, Barker et al. (Br. J. Haematol. 1992, 82:126–132) reported the isolation, by a nonradioisotopic method, of a 32-kD polypeptide plus a 38–51-kD diffuse protein zone by the autoantibodies of three AHA patients.

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patients. These proteins appear to be very similar to our p34 and gp37–55.


References


