Identification of Pemphigus Vulgaris Antigen Extracted from Normal Human Epidermis and Comparison with Pemphigus Foliaceus Antigen

Russell W. Eyre and John R. Stanley
Dermatology Branch, National Cancer Institute, National Institutes of Health, Bethesda, Maryland 20892

Abstract

Immunoprecipitations of cultured keratinocyte extracts have shown that pemphigus vulgaris (PV) sera bind a polypeptide of 210,000 mol wt with disulfide-linked chains of 130,000 and 85,000 mol wt. To identify proteins in normal human skin recognized by PV antibodies, we performed immunoprecipitations of normal human epidermal extracts. All 22 PV sera tested immunoprecipitated a complex of polypeptides (PV complex) of 210,000, 130,000, and 85,000 mol wt, after reduction. One- and two-dimensional gel electrophoresis showed that the 130,000- and 85,000-mol-wt polypeptides of the PV antigen from both cultured keratinocytes and epidermis have identical charges and sizes. In addition to precipitating the PV complex, 14 of 22 PV sera also have antibodies to a calcium-sensitive epitope on a different complex of polypeptides (PF complex) which has previously been shown to be precipitated by all pemphigus foliaceus (PF) sera. The PF complex consists of polypeptides of 260,000, 160,000, 110,000, and 85,000 mol wt. Although the majority of PV sera also precipitate the PF complex, no PF sera precipitate the PV complex. Thus, PV and PF can be distinguished on a molecular level using the patients' autoantibodies. The PV and PF complexes, although distinct, have certain similarities. The 85,000-mol-wt polypeptide of each is identical. The 160,000-mol wt-peptide of the PF complex and the 130,000-mol-wt peptide of the PV complex have the same isoelectric point and both are capable of disulfide linkage to the 85,000-mol-wt polypeptide. The PV and PF complexes are closely related and may prove important in cell adhesion.

Introduction

Pemphigus, an autoimmune blistering disease, exists as two major types (1). In pemphigus vulgaris (PV) mucosal and skin blisters develop because epidermal cells come apart in a suprabasilar location through a process called acantholysis. In pemphigus foliaceus (PF), or superficial pemphigus, mucosal lesions are rare and skin blisters develop in a subcorneal location due to acantholysis of the superficial-most epidermal cells. Autoantibodies to the surface of keratinocytes throughout the epidermis are present in both types of pemphigus.

Recently, using an immunoprecipitation procedure with extracts of normal human epidermis, we have shown that all PV patients have antibodies to a complex of polypeptides with a calcium-sensitive epitope (2). This complex (PF complex) consists of polypeptides of 260,000, 160,000 (desmoglein I), 110,000, and 85,000 apparent molecular weight. The 160,000-mol-wt peptide of the PF complex was shown to be desmoglein I, a core desmosomal glycoprotein (2, 3).

There has been conflicting data regarding the identity of the PV antigen extracted from normal human epithelia. From esophageal mucosal extracts a 68,000-mol-wt protein was identified using an immunofluorescence absorption technique (4). A similar technique was used to identify a 20,000- and a 25,000-mol-wt complex from saliva (5). More recently a 33,000-mol-wt protein has been identified by immunoblotting of human skin extracts (6). A 140,000-mol-wt protein has been identified by immunoblotting of bovine tongue, but not bovine skin, extracts (7).

In our laboratory we have been unable to identify the PV antigen by immunoblotting of normal human epidermal extracts. However, we have characterized the PV antigen synthesized by cultured human keratinocytes (8, 9). Immunoprecipitation of metabolically labeled cultured keratinocytes has indicated that all PV sera bind a 210,000-mol-wt polypeptide with disulfide-linked chains of 130,000 and 85,000 mol wt (9). In addition, we have shown that the PV antigen from cultured keratinocytes no longer reacts with the antibody after the antigen has been denatured for immunoblotting experiments (9). With this finding in mind, we hypothesized that the PV antigen extracted from normal human skin might also, once denatured for immunoblotting, no longer bind (or barely very poorly) to the autoantibodies from PV patients. Thus, in this study, to identify PV antigen in human epidermis, we used a minimally denaturing extraction procedure and performed immunoprecipitation before denaturing the proteins for SDS-PAGE.

We present evidence here that all PV sera bind an antigen from epidermis that is very similar to that identified in cultured keratinocytes. We also show that most PV sera bind, in addition, a calcium-sensitive epitope on the PF complex and that the antigen complex identified by PV and PF sera are distinct but share certain biochemical similarities.

Methods

Sera. Sera from patients with PV, PF (including Brazilian PF), Darier’s disease (a nonimmunologically mediated acantholytic disease), bul-
lous pemphigoid, and normals were used. Brazilian PF sera were as previously reported (10). PV and PF sera were from patients with clinical and/or histological typical disease. All PV and PF sera demonstrated positive indirect immunofluorescence at titers $\geq 40$ with a cell surface pattern on normal human epidermis or monkey esophagus.

**Epidermal extraction.** Epidermal sheets were obtained by harvesting suction blister roofs raised on the volar forearms of normal volunteers (area $\sim 3 \text{ cm}^2$) (11). These sheets were immediately washed with ice-cold Tris-buffered saline (TBS) (0.15 M NaCl in 0.01 M Tris-HCl, pH 7.4) with the addition of 2 mM CaCl$_2$. The epidermal sheets were extracted on ice in 2 ml of 0.5% NP-40, 2 mM CaCl$_2$, and 1 mM phenylmethylsulfonylfluoride in TBS. The epidermal sheets were crushed with a blunt plastic rod, vortexed for 90 s, and ultrasonicated at setting 6 for 90 s (microultrasonic cell disrupter, Kontes Co., Vineland, NJ). The extract was centrifuged at 100,000 g for 1 h at 4°C and the resulting supernatant was preabsorbed with two pellets made from 1 ml each of protein A-bearing *Staphylococcus aureus* (Pansorbin, Calbiochem-Behring Corp., La Jolla, CA). In certain experiments 2 mM iodoacetamide was added to the TBS at all steps. The preabsorbed extracted epidermal proteins were then radiolabeled with $^{125}$I.

**Iodination.** Extracted epidermal proteins were iodinated using 1 mCi of Na$^{125}$I (Amersham Corp., Arlington Heights, IL) and iodo-beads (n-chloro-benzenesulfonyamide sodium salt, derivatized uniform, nonporous polystyrene beads) (Pierce Chemical Co., Rockford, IL) (2). 10 beads were washed 3 times with 5 ml TBS and then allowed to dry. Washed and dried beads were charged by adding the radiolabel in 0.5 ml of TBS for 5 min. The epidermal extract was added to the charged beads for 20 min. Unbound iodine was removed by dialysis at 4°C against 0.3% NP-40 in TBS.

**Cell culture.** Normal human epidermal cells from neonatal foreskins were cultured and radiolabeled with $^{14}$C-amino acids as previously described (8, 12).

**Immunoprecipitation.** Immunoprecipitations were performed as previously described (2, 8, 9, 12) except that in most experiments 2 mM CaCl$_2$ was added to all buffers and washing solutions. Approximately $2 \times 10^6$ cpmp of $^{125}$I-labeled epidermal extracts or $4 \times 10^6$ cpmp of $^{14}$C-labeled cultured keratinocyte extracts and 10–15 $\mu$l of sera were used per immunoprecipitation. As usual, immunoprecipitated proteins were only denatured after precipitation and washing by boiling the pellet for 2 min in 100 $\mu$l of sample buffer (2% SDS, 10% glycerol, and 0.01% bromphenol blue in 0.0625 M Tris-HCl, pH 6.8) and, if reduced, 0.1 M DTT. Precipitated polypeptides were resolved by SDS-PAGE or into two dimensions by isoelectric focusing followed by SDS-PAGE as previously described (3). Separated proteins were then visualized by autoradiography using enhancing screens (2).

**Diagonal nonreduced-reduced gels.** Immunoprecipitated proteins were eluted from protein A without DTT and then separated under nonreducing conditions by SDS-PAGE in 3.4-mm tube gels (13). Tubes were then incubated for 90 min at room temperature in sample buffer with 10 mM DTT. Each tube was sealed on a slab gel with agarose (1% agarose, 0.1% SDS, and 2.5 mM DTT in 0.0625 M Tris-HCl, pH 6.8) and separated in the second dimension under reducing conditions by SDS-PAGE. Proteins were then visualized using autoradiography with enhancing screens.

**Results**

**Immunoprecipitation of the PV antigen from extracts of normal human epidermis.** All PV sera tested precipitated a characteristic complex (PV complex) of polypeptides of $\approx 210,000, 130,000$, and 85,000 mol wt in epidermal extracts containing 2 mM calcium (Fig. 1). This complex is different from the complex of proteins (called here PF complex) precipitated by PF sera from epidermal extracts (Fig. 1 and reference 2) of 260,000, 160,000, 110,000, and 85,000 mol wt. All 22 PV sera tested to date have demonstrated this binding, and none of 17 PF sera tested have bound the PV complex. Neither the PV nor PF complex was recognized by 28 control sera. Controls included 2 patients with Darier’s disease (a nonimmunologically mediated acantholytic disease), 7 patients with bullous pemphigoid (a subepidermal immunologically mediated blistering skin disease), and 19 normal sera.

Although none of the sera from PF patients precipitated the PV complex, 14 of the 22 PV sera tested also precipitated the PF complex when extracts and immunoprecipitations were done in the presence of 2 mM calcium (Fig. 2). 7 of these 14 PV sera were tested on extracts that had calcium removed by chelation with EGTA and all then reacted only with the PV complex, although less intensely than if calcium were present (Fig. 3). These data indicate that some PV sera, in addition to having antibodies to the PV complex, have antibodies to a calcium-sensitive epitope on the PF complex. Similarly, we have previously shown that most PF sera also bind a calcium-sensitive epitope on the PF complex (2).

**Similarities between the PV and PF antigen complexes.** To more closely compare the PV complex with the PF complex,
we performed two-dimensional gel separation (isoelectric focusing followed by SDS-PAGE) of immunoprecipitated proteins using a PV sera that recognizes both complexes in the presence of calcium (Fig. 4). The 130,000-mol-wt polypeptide from the PV complex has the same isoelectric point as the 160,000-mol-wt polypeptide from the PF complex, as do the PV 210,000- and PF 260,000-mol-wt polypeptides. In addition, the 85,000-mol-wt polypeptide from both complexes co-migrate. The same results were obtained when the PF complex, precipitated by PF sera, and the PV complex, precipitated by PV sera, were compared on simultaneously run two-dimensional gels.

Further similarities between PV and PF complexes were revealed by nonreduced-reduced diagonal gels, as discussed below.

Diagonal gels indicate that peptides in both the PV and PF complexes are covalently (disulfide) and noncovalently linked. In two-dimensional diagonal gels, proteins are first separated by SDS-PAGE without reducing agents and then in the second dimension by SDS-PAGE after reduction of disulfide bonds. Proteins without disulfide bonds migrate equally in both dimensions and thus fall on the diagonal. Proteins with interchain disulfide bonds migrate below the diagonal in the second dimension at a point under the diagonal equal to their unreduced molecular weight. Diagonal gels of the PV and PF complexes look remarkably similar (Fig. 5). Some of the 210,000-mol-wt peptide of the PV complex is formed by disulfide linkage of the 130,000- and 85,000-mol-wt peptides. A similar linkage to form an ~ 260,000-mol-wt peptide also occurs between the 160,000- and the 85,000-mol-wt peptides of the PF complex. There are also some nondisulfide-linked peptides of the complexes present, migrating on the diagonal, which indicates that some peptides are also present in the complex as noncovalently linked peptides. After reduction, much of the 210,000- and 260,000-mol-wt peptides are reduced to their component chains; however, some of the 210,000-mol-wt polypeptide of the PV complex and some of the 260,000-mol-wt polypeptide of the PF complex persist on the diagonal. There are several possible explanations for this finding. First, there may be hidden nonreducible disulfide bonds that form between the smaller molecular weight polypeptides of each complex. Second, after disulfide bond formation, other non-disulfide covalent bonds may form. Finally, and less likely, the nonreducible higher molecular weight peptide of each complex may be unrelated to the disulfide-linked peptide, which just coincidently has the same molecular weight.

Some of the disulfide bonds that form are a consequence of the extraction and iodination procedure, which subjects proteins to a more oxidizing environment than in vivo. When iodoacetamide (which blocks free sulphydryl groups and prevents disulfide bonds from forming) is added to the epidermal extraction, solution more of the 130,000- and 85,000-mol-wt polypeptides of the PV complex migrated on the diagonal in nonreduced-reduced gels, indicating they are nondisulfide linked (data not shown). However, even in the presence of iodoacetamide, some of the peptides are still disulfide linked (and therefore migrate below the diagonal). Similar results with and without iodoacetamide were noted with the PF complex (data not shown).

These data indicate that the 130,000- and 85,000-mol-wt peptides of the PV complex and the 160,000- and 85,000-mol-wt polypeptides of the PF complex are present both as covalently and noncovalently linked peptides.

**PV antigen is similar in epidermal and cultured keratinocyte extracts.** The 130,000- and 85,000-mol-wt peptides immunoprecipitated by PV sera from either cultured keratinocyte or epidermal extracts co-migrate when separated by SDS-PAGE (Fig. 6) or into two dimensions by isoelectric focusing.
Figure 3. PV serum precipitates the PV and PF complexes in the presence of 2 mM CaCl₂, but only the PV complex if calcium is chelated with EGTA. Epidermal proteins were extracted with 0.5% NP-40 and 2 mM CaCl₂, then radiolabeled with ¹²⁵I. Immunoprecipitations were then performed either without (lanes labeled CA) or with 4 mM EGTA (lanes labeled EGTA) added to a portion of the extract. Both portions of the extract were precipitated by the same sera. PV 300 precipitated both the PV complex (marked by the Vs) and the PF complex (marked by the Fs) in the presence of 2 mM calcium, but when calcium was removed by chelation with EGTA then only the PV antigen was immunoprecipitated, although less intensely than when calcium was present. Normal human sera (N) do not precipitate either complex.

followed by SDS-PAGE (data not shown). As discussed above, the 210,000-mol-wt polypeptide seen only in the epidermal extract in these reduced gels may represent covalently linked (either nondisulfide linked or linked by hidden nonreducible disulfide bonds) 130,000- and 85,000-mol-wt peptides, or, less likely, a polypeptide unrelated to the smaller polypeptides of the complex. It has been shown that the 130,000- and 85,000-mol-wt polypeptides from cultured keratinocytes form disulfide bonds under nonreducing conditions to make a 210,000-mol-wt peptide (9), just as do the 130,000- and 85,000-mol-wt polypeptides extracted from normal human epidermis (Fig. 5). These data demonstrate the similarity of the PV antigen isolated from cultured keratinocytes to that isolated from normal epidermis.

Discussion

All PV sera tested have antibodies to a characteristic complex of polypeptides extracted from normal human epidermis, two peptides of 130,000 and 85,000 mol wt that are capable of forming disulfide and possibly other covalent bonds. About two-thirds of the PV sera tested also have antibodies to a calcium-sensitive epitope on the PF complex. The following observations suggest that the PV and PF complexes are separate and distinct, and that some PV patients have separate antibody systems against each: (a) all PF sera tested bind only the PF complex (reference 2 and Fig. 1); (b) some PV sera precipitate only the PV complex (Figs. 1 and 2); and (c) those PV sera that precipitate both complexes bind only the PV complex in the absence of calcium (Fig. 3).

The PV and PF antigen complexes, although distinct, have certain similarities. The 130,000-mol-wt peptide of the PV complex and the 160,000-mol-wt peptide of the PF complex have the same isoelectric point and both bind the same 85,000-mol-wt peptide through disulfide as well as noncovalent bonds.

No laboratory or clinical differences were noted among PV patients with sera that did or did not bind the PF complex. It appears that if antibodies are present to the PV complex, suprabasal acantholysis occurs whether or not antibodies are also present against the PF complex. Although both PF and PV antigens are usually present in a similar distribution by immunofluorescence (1) they are distinct complexes and may serve to mediate adhesion at different levels in the epidermis. In fact, in some cases, PF antibodies have been reported to be located in the superficial epidermis only (14, 15).

The PV antigen which we have previously characterized from extracts of keratinocyte cultures (8, 9) is similar to the
antigen we have described here extracted from human epidermis. We suspect that our inability to detect PV antigen by immunoblots of extracts of human epidermis (9) relates to the fact that antigen is fully denatured before being exposed to antibody in the immunoblot procedure. In contrast, in the immunoprecipitation procedure used here (and in the previous cell culture study) the antigen is not fully denatured until after it has been precipitated by the antibody. We have also noted this problem of antibody not binding denatured antigen on immunoblots with the majority of PF sera (3, 9, 10) and with some pemphigoid and epidermolysis bullosa acquista sera (16, 17). However, note that Peterson and Wuepper (6) were able to detect a PV antigen by immunoblotting if they concentrated the human PV sera that they used. This PV antigen had a molecular weight of 33,000 on SDS-PAGE. How this low molecular weight antigen relates to the PV complex described here is unclear and awaits further elucidation.

As suggested by Jones et al. (7), PV antigen may be a member of a large group of so-called cell adhesion molecules which includes molecules such as L-CAM, N-CAM, and uvo-morulin (18). PF antigen might also prove to be in this category. It is also possible that the PV and PF complexes may belong to a family of molecules called cytoadhesions, or integrins, which are important in cell–cell and cell–substrate adhesion and include molecules such as Mac-1, LFA-1, and platelet glycoproteins (GIIb/IIIa (19–22)). Integrins are comprised of noncovalently linked heterodimers with a small component of 95,000–130,000 mol wt and a large component with molecular weight between 130,000 and 210,000. The possible relationship of the PV and PF antigens to these molecular families awaits further studies.

Whatever the relationship, if any, of PV and PF antigens to these other molecular families, these antigens are clearly related to each other, and may prove to be important molecules in epidermal cell adhesion. Exactly how autoantibodies from these patients interact with these molecular complexes to cause loss of cell adhesion, whether directly by interfering with their function or indirectly through protease release by keratinocytes (23, 24), complement activation (25), or combinations of these processes, merits further study. In any case, the autoantibodies from these patients can now be used to absolutely distinguish PV and PF at a molecular level.

Figure 6. Comparison of PV antigen extracted from cultured keratinocytes and epidermis. Autoradiogram of SDS-PAGE of radiolabeled immunoprecipitated protein extracted from either cultured keratinocytes (C) or epidermis (E). The 130,000 (closed arrowhead) and the 85,000 (open arrowhead)-mol-wt peptides precipitated by PV sera from extracts of both epidermal and cultured keratinocytes co-migrate. The 210,000-mol-wt peptide (arrow) is present in this reduced gel only in the epidermal extract. The band present in the last lane (immunoprecipitation of the epidermal extract with normal human serum) migrates slightly faster than the 85,000-mol-wt peptide, and also does not co-migrate with the 85,000-mol-wt peptide on two-dimensional gels (data not shown).
Acknowledgments

We thank Dr. Stephen Katz for critical review of the manuscript, Dr. Richard Klausner for many excellent suggestions, and Ms. Vera Klaus-Kovtun for able technical assistance.

References


