Rabbit Antipeptide Antibodies against Restricted Domains of the Histocompatibility Complex
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Synthetic Peptides, Antipeptide Antibodies, Histocompatibility Antigens

The work reported here concerns the specificity of four antibodies elicited by synthetic peptides corresponding to three domains of the HLA-B7 glycoprotein, and one of the β2 microglobulin. One of the four peptides had been previously investigated, but the data obtained by its elicited antibodies were incomplete.

Of the three HLA-B7 peptides assayed, one was unable to raise an immune response. The other two produced antibodies in good titer that reacted with the antigen peptide, but were unable to recognize antigenic determinants when tested on membrane glycoproteins solubilized from human lymphoblastoid cells. In contrast, the antibody elicited by the only fragment from the β2microglobulin recognized the native β2microglobulin molecule as well as HLA/β2m complexes, and reacted with intact human cells, as determined by ELISA and by FACS analysis. This peptide, which contains one of the most hydrophobic fragment of the whole molecule, is likely an important antigenic site, since it is also recognized by traditional antisera raised against native β2microglobulin.

Introduction

HLA-A, -B, and -C antigens are membrane glycoproteins composed by two non-covalently associated polypeptide chains. The larger component (MW 45000 daltons) is a highly polymorphic glycoprotein encoded by the Mayor Histocompatibility Complex located on chromosome 6, and carries determinants responsible for allospecificity. The small component (MW 12000 daltons) is a serum protein encoded on chromosome 15, and which contains no allotypic determinants [1].

The biggest part of the heavy chain, including an oligosaccharide unit bound to Asn 86, lies on the external membrane of the cell: a small portion enters the lipid bilayer and the cytoplasm [2].

The β2microglobulin is entirely extracytoplasmic, non-covalently but tightly bound to the heavy chain, and contributes to the folding of this latter molecule. The precise role of the β2microglobulin in the immune system is unclear, although the binding of this molecule to the nascent heavy chain appears to be required for the complete cell expression of these membrane glycoproteins [3, 4]. Circular dichroism data [5] indicate that the heavy chain undergoes a conformational change after binding to the small molecule. However, the final 3-D structure of the complex, and the localization of its surface domains, is to day almost completely unknown.

In recent years a number of systems have been developed to predict protein conformations from the amino acid sequence [6, 7] as well as for locating antigenic determinants by analyzing the regions of greatest local hydrophilicity [8]. By this method, taking advantage of the availability of the amino acid sequences for the HLA heavy chains with different specificity [9, 10], as well as for β2microglobulins from different sources [11, 12], one could predict the localization of the most probable antigenic sites of the histocompatibility antigens. The direct proof of this theoretical approach has to be obtained by the determination of the binding capacity of the antibodies raised against every particular region, to the native, folded molecule. Preliminary experiments performed with this technique seem to indicate that fragment 39 to 50 of the HLA-B7 glycoprotein is buried or masked [13], while at least a portion of the fragment 61 to 83 must be exposed on the surface of the HLA/β2m complex, since the antipeptide antibody promptly recognizes the intact
complex in binding and immunoprecipitation assays [14].

In the present study, we selected new fragments from the HLA/β2microglobulin complex, and investigated the antibodies elicited by these peptides in their ability to bind to native membrane glycoproteins from a suitable cell line, with the aim of elucidating the antigenic structure of the histocompatibility antigens, and of selecting these antibodies that could be employed in future functional studies.

**Materials and Methods**

**Cell line and preparation of the glycoproteins**

The human homozygous lymphoblastoid cell line GM 3107 (HLA-A3,3: B7,7) was a generous gift of R. Tosi, Istituto di Biologica Cellulare, Rome.

A small amount of membrane glycoproteins were isolated from these cells by affinity chromatography on Lens culinaria lectin-Sepharose, as reported by others [15].

**Synthesis of peptides**

The sequence of the peptides used in the present investigation were chosen from the HLA-B7 glycoprotein [9] and from the human urinary β2microglobulin [11]. Three peptides were prepared by the solid-phase method [16] using a Beckman Synthesizer, Model 990 B, and one, HLA-B7(134-146) by manual synthesis. In all but this last peptide, a cysteine was added to the C-terminal amino acid of each fragment, to facilitate the coupling to the carrier protein and to the Agarose-DADPA resin used as immunoabsorbent.

**Coupling of the peptides of the carrier protein**

Three peptides were coupled to KLH with the aid of maleimidobenzoyl-N-hydroxysuccinimide ester (MBS) [17], through the cysteine at the COOH-terminal. Coupling efficiency, calculated by the determination of residual free -SH groups in solution after three hours reaction, [18], varied from 27% to 61%.

Peptide HLA-B7(134-146), which lacked a free -SH group, was coupled to KLH by the use of glutaraldehyde.

**Production of antisera**

Following an initial subcutaneous injection of 300 μg of peptide-carrier complex in complete Freund's adjuvant, each of four rabbits received bimonthly 150 μg of the complex in incomplete adjuvant. The animals were bled the first time after 30 days, and then regularly every second week.

Antiserum 4246 was a generous gift of Dr. William Church, and was obtained by immunization of a rabbit with bovine β2microglobulin purified from cow colostrum.

**Binding assays**

The binding of antisera or purified rabbit antibodies to the antigens was routinely assayed by the Enzyme Linked Immuno Sorbant Assay. The method has been described elsewhere [13].

**Preparation and use of the immunoadsorbents**

For the purification of antipeptide antibodies from the immune sera, specific immunoadsorbents were prepared coupling each peptide to agarose-diaminodipropylamino resins (AG-DADPA) (Pierce), through the C-terminal cysteine by the aid of MBS. Details of this method will be reported elsewhere (Chersi et al., in preparation). Different subclasses of antibodies from immune serum AC14, directed against different epitopes in the fragment β2m(32-50), were separated on an immunoabsorbent prepared from Agarose-diaminodipropylamine resins and membrane glycoproteins from GM 3107 cells. For that purpose, AG-DADPA resin (4 g) was reacted with 2 mg of glycoproteins and 5 ml of a 25% solution of glutaraldehyde in PBS, under stirring. After 90 min, the immunoabsorbent was washed by repeated centrifugations in PBS. The binding of the glycoproteins to the resin was only 27%, as determined by spectrophotometric measurements.

Partially purified antibodies from immune serum AC14 were separated on a 0.8 × 4 cm column of this resin, using an initial washing with the starting buffer (PBS), followed by 0.2 M glycine-HCl buffer pH 2.3.

**Preparation of the F(ab')2**

2 mg of purified antipeptide IgG were digested with pepsin at 37 °C for 18 hours, in 0.12 M acetate buffer pH 4.0 with an enzyme-to-substrate ration of
Samples were then centrifuged, dialyzed against PBS, and treated for 2 hours at 4 °C with a suspension of Protein A-Sepharose, to remove undigested IgG.

**Indirect immunofluorescence**

10⁶ GM3107 cells in 0.05 ml PBS containing 10⁻² M sodium azide were incubated with 100 μl of antibody (0.70 mg/ml) for 45 min at 2 °C, washed by centrifugation through FCS, treated with fluorescein-labeled goat anti-rabbit IgG (45 μg), centrifuged again through FCS, washed with PBS, and finally fixed in 10% formalin. Samples of this preparation were submitted to traditional fluorescence microscopy and to fluorescence-activated cell sorter analysis. Normal rabbit IgG was treated and tested in the same way as a control.

**Results**

Table I summarizes the sequences of the four peptides used as immunogens in the present investigation. All together they cover approximately 20% of the total amino acid sequence of the HLA-B7 heavy chain, and about the same of the β2 microglobulin. Their schematic localization in the primary sequences of both proteins is reported in Fig. 1.

The selection of two of the four fragments studied was based on the presence of regions with high local hydrophilicity, according to a recent method for locating the most likely antigenic determinants [8]. The peptide corresponding to pos. 186 to 200 of the heavy chain was selected because of its unusual amino acid composition, since it contained four histidine residues in a total of 15 amino acids, and it was assumed that it would elicit a rabbit antibody unable to show fortuitous crossreactivity with unrelated proteins. The small peptide HLA-B7(134–146) was chosen arbitrarily.

The four peptides were coupled to KLH by aid of MBS or glutaraldehyde, and injected into rabbits. The animals were bled the first time after three immunizations, and then regularly every second week. The amount of elicited antipeptide antibody in each immune serum was assayed by ELISA. Three out of four antisera promptly reacted with the inducing peptide, while antiserum AC 10, raised against the fragment HLA-B7(186–200) did not recognize its antigen, and therefore was not investigated further.

The three antisera AC07, AC11 and AC14 were then purified on the specific immunoadsorbent, and further characterized in their ability to bind to intact glycoproteins solubilized by nonionic detergents from human lymphoblastoid cell lines, using the ELISA. It was found that antibody AC07 did not react, antibody AC11 reacted poorly, while antibody AC14 against β2M(32–50) was able to recognize antigenic determinants located on the surface of the native complex. Moreover, it reacted with human urinary β2 microglobulin as well as with bovine β2 microglobulin purified from colostrum. On the other hand, a polyclonal antiserum (4246), raised in a rabbit against the bovine β2 microglobulin, was able to react with this fragment in ELISA, recognizing therefore an antigenic domain resembling the fragment used for immunization (Fig. 2).
Inhibition experiments confirmed that the binding of antibody AC14 to these antigens was selective and specific: increasing amounts of the inducing peptide \( \beta_2 \text{m}_{32-50} \) inhibited almost completely the reaction of antibody AC14 with membrane glycoproteins from GM3107 cells, and with human or bovine \( \beta_2 \text{m} \) (Fig. 3). The peptide, however, did not significantly inhibit the reaction of polyclonal antiserum 4246 with human or bovine \( \beta_2 \text{m} \) (data not shown).

However, it can be demonstrated that only a small subclass of the antibody population elicited by peptide \( \beta_2 \text{m}_{32-50} \) is effectively directed against antigenic sites of the intact HLA/\( \beta_2 \text{m} \) complex, the majority of the Ig molecules being directed against the synthetic peptide. A chromatography on a suitable immunoabsorbent (AG-DADPA-GP) allows to separate the antipeptide antibodies into two fractions, one reacting almost exclusively with the synthetic peptide, the other with the peptide as well as with intact membrane glycoproteins. This latter fraction accounts for about 20% of the total antibodies loaded, and this confirms preliminary studies in which the percentage of antipeptide antibodies reacting with native HLA-DR molecules was estimated by inhibition techniques (Chersi et al., unpublished results). A slight increase in the binding of antibody AC14 to intact glycoproteins can be obtained when the antigen is first subjected to full oxidation by performic acid, before being immobilized on microtiter plates; this procedure might cause a partial unfolding of the complex and exposure of buried or masked antigenic sites. The binding of this antibody to intact cells was evaluated by ELISA, using the F(\( \text{ab'} \))\(_2\) fragment of the
purified antibody, to minimize non-specific binding and background activity, and by Cell Sorter Analysis: Using the indirect immunofluorescence assay, from 33% to 39% of 3107 cells resulted stained by fluorescein-labeled goat antirabbit IgG, when previously incubated with antibody AC14, while only 7% of the same cells were labeled when normal rabbit IgG replaced antibody AC14 under the same conditions.

Discussion

Four antipeptide antibodies directed against different regions of the HLA/β2microglobulin complex were investigated in the present paper in their ability to bind to the antigen peptide and to native membrane glycoproteins and cells.

One of the four peptides used as immunogen, i.e. HLA-B7(186–200), was not able to elicit rabbit antibodies: this result may reflect an identity or a close similarity in sequence between this fragment, and one present in a protein of the animal. The immune response of the animal, in fact, was exclusively directed against the carrier protein, as evaluated by ELISA.

Antipeptide antibody AC07, characterized by a good titer in the binding to the antigen peptide, was not able to react with intact glycoproteins solubilized by non-ionic detergents from the cell line GM 3107. As for antipeptide antibodies described in previous papers [13, 14] the most likely explanation for this result is that the fragment selected for the immunization is buried in the interior of the native molecule, the HLA/β2microglobulin complex, where it is not available for interreaction with the antibody. An alternative possibility is that the coupling of this fragment to the carrier protein through the N-terminal amino and the C-terminal lysine ε-amino groups by glutaraldehyde prevented the peptide from assuming, for at least part of the time, a configuration that resembled that occurring in the intact protein.

Antibody AC11, raised against peptide HLA-B7(215–232)+, which is comparable to antibody 6153 partially investigated in a previous study [14], shows low but still measurable activity when assayed against native membrane glycoproteins. Although a partial exposure of this fragment could not be excluded, we think that its partial activity might be better explained in terms of crossreactivity of this antibody with a not-related antigenic site: one possibility is the region 32–50 of the β2microglobulin, since the comparison of the amino acid sequences of the HLA-B7(215–232) and β2m(32–50) peptides reveals fortuitous similarities, as a DGE fragment, a E–KVC terminal piece, and a D---D--E section. Crossreactions of antipeptide antibodies have been reported, as a result of identical or closely similar epitopes that may occasionally occur in otherwise unrelated proteins [19]. Partial crossreactivity can in fact be demonstrated between the two peptides and antibodies AC11 and AC14.

Antibody AC14, raised against β2m(32–50), reacts with human urinary β2microglobulin, and is the only one of the four antibodies tested able to bind in Enzyme Linked Immuno Sorbant Assay to intact membrane glycoproteins and to GM 3107 cells. The binding to these cells can be confirmed by conventional fluorescence microscopy and FACS analysis. The antibody reacts also with bovine β2microglobulin isolated from cow colostrum, as a result of the 75% homology in amino acid sequence between the human and the bovine proteins. On the other hand, a polyclonal antiserum raised in a rabbit against the intact bovine β2microglobulin recognizes the β2m(32–50) fragment in binding assays. This result is somehow unexpected, since antibodies to an intact protein very seldom react with its fragments, especially when presenting low molecular weights.

The binding data suggest the following considerations:

1) The domain 32–50 of the β2microglobulin is exposed (at least partly) on the surface of the native molecule.

2) This domain is also exposed on the HLA/β2m complex, is not involved in the binding to the heavy chain, is not sterically masked by neighbouring groups because of this interreaction.

3) This region likely contributes to the antigenicity of the whole molecule. However, the precise role of this domain in the intact complex has still to be elucidated.

The analysis of the binding data of the antipeptide antibodies to the HLA/β2m complex presented here, and in previous papers [13, 14] indicate that, as far as today, only two regions of the histocompatibility complex are effectively recognized by the antipeptide antibodies investigated: one is the hypervariable region of the heavy chain (res. 61 to
one is the fragment 32–50 of the β2-microglobulin. This scarce ability of the antipeptide antibodies in reacting with native structures seems to be contradicted by recent studies [20] that indicate that the majority of monoclonal antipeptide antibodies recognize native cognate proteins.

Unless one admits that the majority of our eight peptides investigated were unfortunately selected from internal regions of the HLA-β2m complex, and taking into account the special methodology by which monoclonal antibodies are selected, this result suggests that not all antipeptide antibodies might contain small, often undetectable populations of immunoglobulins able to react with native cognate structures: the scarce representation of these molecules could be due to the low probability by which a synthetic peptide may adopt in solution a conformation resembling the structure in the intact molecule. Further studies are in progress to evaluate this possibility.

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