DISTINCT FUNCTIONS OF MONOCLONAL IgG ANTIBODY DEPEND ON ANTIGEN-SITE SPECIFICITIES

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Standard immune responses are the sum of many clonal responses, cross-reactive and specific. Any study of clonal expression as well as analysis of its products requires the separation of clones. There are several approaches to this goal (1–3). The one we used was to generate immune responses of restricted heterogeneity where a limited number of clonotypes determine the response (4, 5). The prerequisites for the induction of such responses are homogeneity of the immunogen and the genetic background of the animals immunized (1, 4–9). Bacterial cell walls carrying regularly spaced polysaccharide moieties on their surface meet the demand for homogeneity of the antigen (1). Certain breeds of rabbits and inbred mouse strains fulfill the genetic requirements (4–9). Considering both of these conditions antibody responses of predictable clonal patterns can be elicited (10).

Although most of the early work emphasized the restricted heterogeneity of anti-polysaccharide antibody responses (1, 4–9) we report here in detail that clonal IgG antibody patterns in response to polysaccharide determinants are rather complex. Despite this complexity it is possible to isolate, by a combination of methods, anti-streptococcal group A-variant antibodies in single-band purity (11). In using these antibodies, distinct functions of monoclonal IgG antibody could be ascribed to fine specificities for different determinants on a linear polysaccharide chain.

Material and Methods

Rabbit Hyperimmune Antisera and Antigen Preparations. The production of hyperimmune antisera in selectively bred rabbits to the streptococcal group A-variant polysaccharide (Av-CHO) and the identification and quantitation of predominant clonotypes has been described previously (5).

The Av-CHO was purified from K43 streptococcal cell walls with >98% purity (12, 13). Aliquots were labeled for radioimmunoassays with 125I or 131I (5). For coating sheep erythrocytes, (SRBC) the O-stearoyl-ester of Av-CHO was made (14).

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1 Abbreviations used in this paper: Aggl, agglutination; Av-CHO, streptococcal group A-variant polysaccharide; Av-CHO-SRBC, sheep erythrocytes coated with the O-stearoyl-ester of Av-CHO; 2,4-dinitrophenyl-hydrazone of Av-CHO; DTT, dithiothreitol; 125I-Av-CHO, 131I-Av-CHO, Av-CHO labeled in its tyraminated form with 125I or 131I; IEF, isoelectric focusing; KD, dissociation constant; PBS, 0.01 M sodium potassium phosphate; pIEF, preparative isoelectric focusing; Ppt, precipitation.
Preparation of 2,4-Dinitrophenylhydrazone of the Av-CHO. The method of Lloyd and Doherty was used for the preparation of this derivative (15). From 55 mg of starting material, 25 mg of 2,4-dinitrophenyl-hydrazone of Av-CHO (DNP-Av-CHO) were obtained.

Circular Dichroism Spectroscopy. Circular dichroism (CD) spectra of DNP-Av-CHO antibody complexes were measured in a Cary 60 instrument (Monrovia, Calif) rebuilt for CD measurements and equipped with a Jobin-Yvon modulator (Instruments SA, Longjumeau, France), using 1-cm quartz cells thermostated at 20°C. Otherwise, conditions were those described previously (16).

Affinity Chromatography. Av-CHO affinity columns were prepared by using epoxy-activated Sepharose 6B, Pharmacia Fine Chemicals, Uppsala, Sweden (17). Bound antibody was eluted by 0.15 N NaCl in 0.1 M glycine-HCl, pH 3.0.

Analytical Isoelectric Focusing. Analytical isoelectric focusing (IEF) was performed in 5% polyacrylamide gels, pH 5–10, (18). Patterns of antibodies were visualized both by staining with bromophenol blue and by autoradiography using the 125I-labeled Av-CHO (19).

Preparative Electrophoresis and Preparative IEF. Antibody was purified from whole immune sera first by preparative agarose block electrophoresis (4), and then by one or more runs of pIEF in horizontal layers (11).

Papain Digestion. Purified antibodies were digested by 2-β-mercaptoethanol activated papain (Sigma Chemical Co., St. Louis, Mo.), and Fab monomers were isolated (20).

Affinity Measurements. Binding of the Av-CHO by antibody from whole sera or purified fractions was determined by the quenching of antibody tryptophan fluorescence (21).

Equilibrium binding studies of 125I-labeled Av-CHO to antibody were performed with plastic microcels of 200 μl capacity (22). Glucose (5 mM in phosphate-buffered saline [PBS]) was added to eliminate weak unspecific binding of the Av-CHO to the dialysis tubing which had an average exclusion limit of 17,000 daltons. Under this condition equilibrium was complete after 32 h at 25°C. The specific radioactivity of hapten and antigen in these experiments varied from 11–45 cpm/nmol. Samples were counted in a Packard Gamma spectrometer (model 5130) (Packard Instrument Co., Inc., Downers Grove, Ill.). Dissociation constants (Kd) were obtained for the stoichiometry of ligand binding from replots of the data according to Scatchard (23).

Ultracentrifugation Analysis. Sedimentation experiments were performed in a Beckman model E analytical ultracentrifuge (Beckman Instruments, Inc., Fullerton, Calif.). The weight-average molecular weights were calculated employing a value of 0.73 cm³/g for the partial specific volume of antibody and antigen-antibody complexes (24).

Immunochemical Methods. Quantitative precipitin and inhibition analyses as well as binding of the 125I-Av-CHO by antibody was performed as described (6, 9). Inhibition studies of precipitation and binding analyses were conducted by addition of increasing amounts of L-rhamnose or a synthesized probe of L-rhamnosyl α-1 → 2-L-rhamnose (25). Double-diffusion analyses in gels were performed according to established methods (26).

Agglutination of O-searoyl-ester Av-CHO-coated SRBC (Av-CHO-SRBC) was performed in microtiter plates (14). Complement-dependent lysis (Cdl) was determined qualitatively by the spot assay (27) and quantitatively by reacting whole inactivated (56°C 30 min.) SRBC absorbed rabbit anti-Av-CHO antisera or purified antibody fractions in the presence of 0.1 ml 1:10 PBS diluted complement solution (guinea pig complement, Behring-Werke AG, Marburg/Lahn, West Germany). Av-CHO-SRBC and SRBC (OD, 540 nm = 0.4) were used in final test volumes of 1.1 ml at 37°C for 60 min. Lysis was expressed as a percent against water-lysed Av-CHO-SRBC and SRBC controls in supernates. These experiments were also performed in the presence of 10 mM dithio-threitol (DTT), Clelands reagent, Calbiochem for two low affinity antibody fractions of single-band purity (28).

Competition experiments between lytic and nonlytic antibodies for Av-CHO sites were performed at 10, 50, and 75% lysis in the Cdl assay using increasing amounts (up to 10-fold molar excess) of nonlytic Av-CHO specific antibody.

Binding of Av-CHO specific antibodies to Av-CHO-SRBC was determined qualitatively and quantitatively. Two preparations of Av-CHO specific antibody fractions of essentially single-band purity were used, the high affinity fraction K151-748II and the low affinity fraction K127-760II. Qualitatively, binding was determined by incubating 10 μg of antibody with 100
mu1 of 10% Av-CHO-SRBC and 100 mu1 of 10% SRBC as control, respectively, followed by three rapid washes in cold PBS, and subsequent treatment with a fluoresceine-conjugated goat anti-rabbit IgG immunoglobulin fraction.

Quantitatively, binding of antibody was determined as follows: using the chloramine T method (29), K151-748H was labeled with 125I (sp act 52 uCi/mg) and K127-760H with 131I (sp act 110 pCi/mg). To a constant number of isotope counts (corresponding to 128 and 113 ng of antibody per test, respectively), increasing amounts of Av-CHO-SRBC and SRBC were added, in a constant final volume of 50 u1 PBS containing 0.1% bovine serum albumin. Av-CHO-SRBC and SRBC treated in this fashion for 12 h at 4°C were either washed three times in PBS or layered on top of a 1-cm high fetal calf serum solution in conical polypropylene tubes. These were centrifuged quickly (5 min) to the bottom of the tube, then snap frozen in dry ice and the tips of the tubes cut off. The tips and supernates were counted (30).

Results

Clonal Restriction of Anti-Av-CHO Antibodies is a Quantitative Trait. High resolution analysis of 40 rabbit hyperimmune anti-Av-CHO antisera by IEF revealed simple spectrotype patterns (two to three band patterns) when stained with bromophenol blue (Fig. 1 A), confirming the rather restricted heterogeneity evidenced by microzone electrophoretic analysis (samples 7, 10–12, and 16). When, however, patterns were visualized by binding of 125I-Av-CHO (Fig. 1 B) according to Keck et al. (18) very complex spectrotype patterns emerged with up to 75 bands, corresponding to 20–30 clonotypes (31). The dominating clonotypes were no longer identified as such but
**Table I**

<table>
<thead>
<tr>
<th>Rabbit No.</th>
<th>Low affinity (pH 7.0) Ab</th>
<th>High affinity (pH 3.0) Ab</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. K6-137</td>
<td>28.1</td>
<td>93.4</td>
</tr>
<tr>
<td>2. K33-380</td>
<td>22.9</td>
<td>90.6</td>
</tr>
<tr>
<td>3. K34-413</td>
<td>13.2</td>
<td>94.3</td>
</tr>
<tr>
<td>4. K45-426</td>
<td>36.1</td>
<td>97.3</td>
</tr>
<tr>
<td>5. K49-501</td>
<td>30.2</td>
<td>97.1</td>
</tr>
<tr>
<td>6. K14-543</td>
<td>32.2</td>
<td>90.0</td>
</tr>
<tr>
<td>7. K14-545</td>
<td>24.8</td>
<td>89.5</td>
</tr>
<tr>
<td>8. K11-700</td>
<td>29.3</td>
<td>84.9</td>
</tr>
<tr>
<td>9. K13-730</td>
<td>37.1</td>
<td>97.6</td>
</tr>
<tr>
<td>10. K13-732</td>
<td>15.4</td>
<td>92.8</td>
</tr>
<tr>
<td>11. K15-745</td>
<td>31.8</td>
<td>97.2</td>
</tr>
<tr>
<td>12. K12-735</td>
<td>49.6</td>
<td>99.0</td>
</tr>
<tr>
<td>13. K12-760</td>
<td>51.7</td>
<td>99.2</td>
</tr>
<tr>
<td>14. K13-772</td>
<td>11.7</td>
<td>90.7</td>
</tr>
<tr>
<td>15. K15-776</td>
<td>13.0</td>
<td>89.0</td>
</tr>
</tbody>
</table>

*Concentrations of Av-CHO-specific antibody determined by the Farr assay (9) in pH 7.0 and pH 3.0 antibody fractions from Av-CHO-epoxy-Sepharose affinity columns. Ppt, precipitating activity with the hot formamide extracted Av-CHO at concentrations of 50 and 100 µg/ml, determined by the Lancefield precipitation test. Aggl., agglutination of Av-CHO-SRBC.*

The Av-CHO is a linear homopolymer of L-rhamnose with alternating α1 → 2 and α1 → 3 glycosidic bonds (32).
Table II

Dissociation Constants (KD), Precipitating (Ppt) and Lytic (CdL) Properties of Purified Anti-Av-CHO Antibody (Ab)

<table>
<thead>
<tr>
<th>Antiserum</th>
<th>Dominant Ab fraction</th>
<th>Minor Ab fraction</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. K81-543</td>
<td>2.0 × 10^-8</td>
<td>—</td>
</tr>
<tr>
<td>2. K116-700</td>
<td>1.71 × 10^-8</td>
<td>—</td>
</tr>
<tr>
<td>pIEF/17*</td>
<td>1.78 × 10^-8</td>
<td>—</td>
</tr>
<tr>
<td>pIEF/19-21</td>
<td>1.53 × 10^-8</td>
<td>—</td>
</tr>
<tr>
<td>pIEF/23-25</td>
<td>1.83 × 10^-8</td>
<td>—</td>
</tr>
<tr>
<td>pIEF/27-28</td>
<td>1.69 × 10^-8</td>
<td>—</td>
</tr>
<tr>
<td>3. K113-732</td>
<td>7.0 × 10^-3</td>
<td>—</td>
</tr>
<tr>
<td>4. K151-748</td>
<td>7.0 × 10^-3</td>
<td>—</td>
</tr>
<tr>
<td>pIEF*</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>pIEFII</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>pIEFIII</td>
<td>ND§</td>
<td>—</td>
</tr>
<tr>
<td>pIEFIV</td>
<td>ND</td>
<td>—</td>
</tr>
<tr>
<td>5. K128-755</td>
<td>3.9 × 10^-8</td>
<td>—</td>
</tr>
<tr>
<td>6. K127-760</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

* pIEF: Antibodies of single band purity by analytical isoelectric focusing and autoradiographic visualization by 125I-Av-CHO with the exception of pIEFIII; this fraction constituted a three-band pattern (see also Fig. 2).
‡ NS, no detectable signal of quenching of fluorescence upon addition of antigen.
§ ND, not determined.

The molecular weight of the Av-CHO was by ultracentrifugal analysis 5,000 daltons (data not shown).

The mode of antigen binding to low and high affinity antibodies was studied by ultracentrifugal analysis in the presence of increasing antigen concentrations (Table III). The binding of antigen to the low affinity antibody barely altered the sedimentation coefficient even under conditions near antigen saturation (KD = 7 × 10^-5 M). The slight decrease in the sedimentation coefficient may be a result of the smaller partial specific volume or the larger translational frictional coefficient of the antibody-antigen complex as compared to free antibody. Clearly contrasting to these findings were the sedimentation properties of high affinity antibody. The sedimentation coefficients increase in the presence of antigen confirming the presence of large, polydisperse complexes, some of which became insoluble (precipitation).

Weight-average molecular weights of high or low affinity antibody demonstrated that antigen binding to the former produced extremely large species, whereas antigen binding to the low affinity antibody resulted in a modest increase in molecular weight (19,000 ± 10,000), suggesting the formation of antigen-antibody complexes which contain only one molecule of antibody.

Quantitative measurements of the binding of antigen to low and to high affinity antibodies further clarified the nature of this binding. The results, now obtained by equilibrium dialysis on the binding of 125I-Av-CHO to a low affinity single-band antibody (K116-700 pIEF 17, Table II, Fig. 2A), yielded a value of 0.77 × 10^-6 M for the dissociation constant. This value was in good agreement with a KD = 1.78 × 10^-6 M by fluorescence titration.
Fig. 2. Scatchard plots of the fluorescence titration of low affinity antibodies of single-band purity with 5,000 daltons Av-CHO. \( Q/Qm \) is the fraction of total quenching, \( Qm \) being the maximal quenching and \( Q \) being the quenching of the free ligand concentration \( Lf \). The fractions pIEF17 (A), 21 (B), and 23-25(C) were isolated from antiserum K116-700 by preparative isoelectric focusing. The \( K_D \) of these preparations cluster within narrow limits. D. Scatchard plots of the monomeric Fab fragments of the high affinity anti-Av-CHO antibody fraction (K116-700) from the Av-CHO-epoxy-Sepharose column with 5,000 daltons Av-CHO (○) and with the DNP-Av-CHO (●). The data are expressed as number (n) of Fab molecules (mol wt 50,000) bound per 5,000 daltons Av-CHO molecule. The \( K_D \) calculated identify these Fab fragments as high affinity binders to the antigen. The DNP-Av-CHO is bound with comparable efficiency as the Av-CHO.

### Table III

<table>
<thead>
<tr>
<th>Molar ratio Ag:Ab</th>
<th>K113-732 (low affinity)</th>
<th>K116-700 (high affinity)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Sedimentation coefficient</td>
<td>Mol wt</td>
</tr>
<tr>
<td>0:1</td>
<td>6.6S</td>
<td>137,000</td>
</tr>
<tr>
<td>0.1:1</td>
<td>6.5S</td>
<td>ND</td>
</tr>
<tr>
<td>1:1</td>
<td>6.3S</td>
<td>ND</td>
</tr>
<tr>
<td>9:1</td>
<td>6.3S</td>
<td>ND</td>
</tr>
<tr>
<td>49:1</td>
<td>6.2S</td>
<td>ND</td>
</tr>
<tr>
<td>249:1</td>
<td>6.2S</td>
<td>156,000</td>
</tr>
</tbody>
</table>

* ND, not determined; ppt, indicates precipitation.

In addition, these measurements showed that two molecules of Av-CHO were bound per antibody molecule or one per combining site in the Fab. This demonstrated that the antigen is functionally monovalent for the low affinity antibody. Hence, low affinity antibodies have chain-terminal specificity, an assertion which was substanti-
Fig. 3. Complement-dependent lysis of Av-CHO-SRBC in the presence of lytic and nonlytic anti-Av-CHO rabbit antibodies. Lysis by increasing concentrations of pH 3.0 fraction K128-755 E.Ab and of essentially single band antibody K151-748II (Fig. 4A) was expressed as the percent of water-lysed Av-CHO-SRBC and SRBC as control at a total OD~om~ 0.4. No lysis was obtained with antibodies K151-748III and IV or with K116-700/27,28 and K127-760II up to concentrations indicated.

ated by the sedimentation behaviour of the low affinity antibody in the presence of increasing antigen concentrations (Table III).

In a complementary experiment, the stoichiometry of high affinity antibody binding to antigen was evaluated from fluorescence titration measurements (Fig. 2D). High affinity antibody K116-700 (Table II) bound with nearly identical affinity to the Av-CHO and to the DNP-Av-CHO \((K_D = 2.9 \times 10^{-8} \text{ M} \text{ and } K_D = 3.0 \times 10^{-8} \text{ M})\). Significantly, this evaluation yielded four antibody combining sites per antigen molecule. This value, when compared to the 1:1 stoichiometry of combining sites to antigen observed with the low affinity antibody, led us to conclude that the difference in the sedimentation behaviour of the low and the high affinity antibody is due to the polyvalency of the linear 5,000 daltons Av-CHO antigen for the high affinity antibody. This high valency naturally implies that high affinity antibodies recognize the internal determinants of the Av-CHO chain. Interaction of low affinity antibody with the DNP-Av-CHO induced neither a DNP cotton effect by CD spectroscopy nor enhancement of DNP fluorescence, so indirectly suggesting specificity for the nonreducing end of Av-CHO.

Functional Properties of Low and High Affinity Antibodies. It was possible to correlate the above fine specificities for the two different antibody populations with functional properties (Table I) part of which had been noted previously (6). Low affinity (pH 7.0 fraction) antibodies neither precipitated the free Av-CHO nor agglutinated or lysed Av-CHO-SRBC in CdL assays. Conversely, internally specific high affinity antibodies (pH 3.0 fraction) precipitated Av-CHO, agglutinated and lysed Av-CHO-SRBC in the presence of complement.

Quantitative CdL data were obtained for two high and four low affinity antibody fractions (Figs. 3, 4A,B). One of the high affinity antibodies was a pH 3.0 fraction eluted from an Av-CHO affinity column (K128-755 E.Ab.), and the other fraction was essentially of single band purity isolated by pIEF from a minor serum antibody (Table I, Fig. 4A). This latter, highly pure component yielded complete CdL at a concentration of 8 \(\mu\text{g/test}\) (Fig. 3).
Flu. 4. Autoradiograms (131-I-Av-CHO) of IEF analysis of antiserum K151-748(W.S.) and isolated fractions (A) and of antiserum K127-760(W.S.) and isolated fractions (B). Results of CdL analyses are indicated.

From antiserum K151-748 the dominant clonotype bands were also purified as three-band and single band patterns by pIEF (Fig. 4A). Neither of these fractions were active in CdL at a concentration comparable to that of the high affinity antibody (Fig. 3). When 125 and 135 μg of single-band Av-CHO-specific antibody of antiserum K116-700/27,28 (11) and K127-760II (Fig. 4B) were used, even upon the addition of 10 mM DTT (28), no CdL was observed, although binding of these monoclonal antibodies to the Av-CHO in solution was demonstrable by fluorescence titration and by radioimmunoassays.

In competition experiments, a 20-fold excess of the nonhemolytic antibody K127-760II failed to inhibit the lytic action of antibody K151-748II in the spot test (27, 33). Further, in the quantitative CdL assay (10, 50 and 75% lysis by antibody K151-748II), no inhibition of lysis was observed up to a 100-fold molar excess of the nonlytic antibody K127-760II, indicating that at these concentrations the low affinity antibody did not compete with the high affinity antibody.
### Table IV

**Binding of Single-Band Antibody Preparations to Av-CHO-SRBC**

<table>
<thead>
<tr>
<th>Av-CHO-SRBC</th>
<th>Molecules bound per Av-CHO-SRBC</th>
</tr>
</thead>
<tbody>
<tr>
<td>K127-760II (low affinity)</td>
<td>+</td>
</tr>
<tr>
<td>K151-748II (high affinity)</td>
<td>+</td>
</tr>
</tbody>
</table>

* For details see text. A sheep anti-rabbit IgG Ig fraction was used.

‡ Molecules bound per single Av-CHO-SRBC calculated after subtraction of binding to SRBC. This number was not significantly different upon the addition of 1,000-fold excess of cold antibody of the other specificity.

A subsequent set of experiments was performed to measure the ratio of binding of low versus high affinity antibodies to Av-CHO-SRBC. With the aid of a fluorescein-conjugated sheep anti-rabbit IgG immunoglobulin fraction antibodies K127-760II and K151-748II were shown to specifically bind to Av-CHO-SRBC but not to SRBC (Table IV). This specific binding was then quantitated (Table IV). High affinity antibody was bound by Av-CHO-SRBC with a 100-fold greater efficiency than low affinity antibody. Because neither binding of the labeled high affinity antibody by 1,000-fold excess of cold low affinity antibody nor binding of labeled low affinity antibody by 1,000-fold excess of cold high affinity antibody were inhibited we conclude that low and high affinity antibodies are directed to nonoverlapping determinants of the same antigen molecule. This experiment is, therefore, further support for the above described distinct terminal and internal antigenic sites recognized by these distinct antibody populations.

**Discussion**

In this paper the structural and functional correlates of two distinct antibody populations are described. These antibodies were identified in rabbit hyperimmune antisera to the streptococcal Av-CHO, initially recognized as precipitating and nonprecipitating Av-CHO specific antibodies (6). The data presented here aid in understanding the detailed immunochemistry of natural polysaccharides occurring as linear homopolymers on the surface of bacteria.

The Av-CHO has been identified as a linear homopolysaccharide of L-rhamnose consisting of alternating $\alpha 1 \rightarrow 2$, $\alpha 1 \rightarrow 3$ glycosidic bonds (32). The rather uniform molecular weight of 5,000 daltons suggests 30 rhamnoses/molecule of Av-CHO. In its extended form it would measure $\approx 200$ Å in length. Considering the Av-CHO content of streptococci the average surface density of Av-CHO/streptococcus calculates to $\approx 1.3 \times 10^6$ Av-CHO molecules at an average distance of 21 Å. In the pepsin-treated vaccines used for immunization, the Av-CHO is the outer most layer (1), and it is probably seen by the immune system as a thorn-apple like sphere. This recognition model is supported by 90% of the IgG antibody being of terminal specificity for Av-CHO.

Group polysaccharide-specific rabbit and mouse anti-streptococcal antisera are prototypes of antibody responses of restricted heterogeneity (1, 34). However, the resolution power of the modified technique of analytical IEF employed here, and described in detail elsewhere (19), indicates a substantial underestimation of the degree of heterogeneity of specific antibody raised by hyperimmunization reported.
previously (1, 5–9, 34). In analogy to the response by BASILEA rabbits (35) the number of phenotypically expressed clonotypes is similar in rabbits previously classified by the terms monoclonal, restricted, and heterogeneous. Therefore, these terms are operational ones, describing a quantitative feature documented by measurements of protein concentration at little resolution power, e.g. microzone electrophoresis, light chain banding patterns on polyacrylamide gel electrophoresis, or IEF using protein staining methods and the isotope overlay technique (1, 4–9, 34, 36, 37). Hence, a monoclonal or a restricted antibody response described phenotypic predominance of one or a few clonotypes accounting, on a weight basis, for up to 60–90% of the total group polysaccharide-specific antibody. The remaining 10–40% of group-specific antibodies are represented by many different clonotypes, each one of them at very different concentrations altogether making up a very heterogeneous sample.

This physico-chemically and, presumably, idiotypically very heterogeneous sample of anti-Av-CHO antibodies in rabbit hyperimmune sera can be divided functionally into two distinct antibody populations coexisting in all sera so far tested. It appears that these distinct functions relate to the nature of the antigenic sites recognized. However, the work presented here has not completely ruled out serologically or structurally that these two functions may be caused by different Cγ regions in the sense of subclass differences. This is, however, unlikely because IEF analysis (Figs. 4 A, B) precludes critical charge differences in many samples used for subclass determination (38–40); preliminary serological tests aiming at subclass differentiation between the two kinds of IgG antibodies have not disclosed differences (L. S. Rodkey, W. Schalch, and D. G. Braun); and genetically it is unlikely that different site specificities for the same antigen are linked to different heavy chain constant regions. The final proof for this reasoning must come from comparative structural analyses.

The data described in this paper require for hapten-carrier systems similar definitions of site specificities of antibody populations arising in immune responses, when claims for generation of antibody diversity by somatic mutation are raised which are based on changes in affinity or function (41, 42–46). Only the (DNP)2-gramicidin S system meets these requirements (47). In the context of this work, maturation of immune responses refers to the emergence of antibodies with different subsite specificities to one antigen presumably from a pre-existing pool of B-cell precursors. Accordingly, clonal hierarchy established as memory in vivo, which in antibody responses of restricted heterogeneity is measurable in mg ml⁻¹ of antiserum, persists in vitro under conditions where limiting numbers of lymphocytes were cultured, permitting an estimate of the frequency of responding units (27). These and additional results are evidence for the lack of maturation of anti-streptococcal group polysaccharide responses (48, 49).

Kabat and associates (50) have reported on binding properties of immunoglobulin combining sites binding either terminal or nonterminal antigenic determinants in dextran. In accordance with our findings their IgA mouse myeloma protein W3129 reacted with a synthetic a1 → 6 linked linear dextran without precipitating it. Protein W3129 was specific for the terminal nonreducing ends of this dextran, whereas a second IgA mouse myeloma protein QUPC52 precipitated the same linear synthetic dextran chain. This finding was consistent with the idea that QUPC52 recognized nonterminal antigen sites. The terminally specific protein W3129 had a high affinity constant (Kₘ = 1.0 × 10³ M⁻¹) whereas the internally specific protein QUPC52 had
a low association constant \( (K_A = 8.4 \times 10^9 \text{M}^{-1}) \). In marked contrast to this situation are the internally specific anti-Av-CHO antibodies. The remarkable high affinity for anti-polysaccharide antibodies of \( K_D < 10^{-8} \text{M} \) may be explained by hydrophobic interactions established via the C6 methyl groups. Chemically these methyl groups are the notable differences in comparing the Av-CHO and the dextran structures. The association constants of the two terminally specific antibodies W3129 and the low affinity anti-Av-CHO antibodies are very close, indicating similar kinds of interactions.

Immunization with cell surface-associated antigenic determinants (in this case of bacteria) may generally elicit a mixture of antibody molecules with very different, nonoverlapping site specificities for linear polysaccharide antigens. The Av-CHO system cogently suggests a critical role of simple mechanistic principles in antigen recognition (schematically shown in Fig. 5), where the most abundantly presented antigenic determinants are the nonreducing ends of single rhamnosyl chains. As a result, antibody of this specificity accounts entirely for dominant clonotypes. Conversely, the high affinity fraction of antibodies is not preferentially selected, as would be predicted by the maturation argument, and the mechanistic model of antigen presentation (Fig. 5) easily explains this failure. These data as well as those obtained with nonprecipitating equine antibodies of high affinity to the \( \beta \)-azophenyl-\( \beta \)-lactoside (Lac) haptenic group coupled to protein carrier molecules rule out univalence and low affinity as explanations for functional differences (51). Because mild reduction by 10 mM DTT did not convert the low affinity anti-Av-CHO antibodies into lytic ones, these antibodies do not correspond structurally to the incomplete anti-RH antibodies (28). Depending on the functional properties of anti-Av-CHO antibodies residing in the determinant specificity for this linear poly-rhamnosyl moiety, the associative model of complement activation and not the allosteric model appears to account for the phenomenon (52).
A final comment relates to the high degree of purity of the terminally and internally specific anti-Av-CHO antibodies used in this study. Although this is the highest degree of purity achieved for elicited antibodies functional criteria must additionally be considered. Single band patterns or triplets of bands upon IEF may still contain antibodies of two distinct functional properties. This caution is critical because analytical IEF is often considered to be the most stringent physico-chemical method to prove structural homogeneity of antibodies.

Summary

Intravenous hyperimmunization of selectively bred rabbits with streptococcal group A-variant vaccines elicits antibody responses of restricted heterogeneity at high antibody levels. All antisera contain two functionally distinct antibody populations, which can be isolated in single-band purity upon analytical isoelectric focusing. Typical examples of these two kinds of single-band antibodies were investigated in great detail for several parameters by a variety of methods. 85-99% of the streptococcal group A-variant polysaccharide (Av-CHO)-specific antibody in the antisera does not precipitate the isolated 5,000 daltons poly-L-rhamnose antigen, neither agglutinates nor lyses in the presence of complement Av-CHO-coated sheep erythrocytes (SRBC), binds the radio-labeled Av-CHO with an association constant in the range of $10^5$-$10^6$ M$^{-1}$, and is of terminal specificity (nonreducing end) for the linear Av-CHO. In contrast, the minor fraction of Av-CHO-specific antibody (1-15%) does precipitate the linear Av-CHO, both agglutinates and lyses Av-CHO-coated SRBC in the presence of complement, has an affinity range of $10^5$-$10^9$ M$^{-1}$, and is of internal specificity for the Av-CHO. The antigenic determinants of the Av-CHO for the antibodies are nonoverlapping, only one Fab of the low affinity antibody can be bound whereas four Fab of the high affinity antibody are accommodated. Hence, the determinant specificity explains the functional differences observed, for there is no indication of subclass differences. A mechanistic model of the A-variant carbohydrate presentation on the vaccine appears to account best for the unbalanced levels of low and high affinity antibody.

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