Structural Analysis of Hepatitis B Surface Antigen by Monoclonal Antibodies

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Abstract

A method has been developed for the analysis of hepatitis B surface antigen (HBsAg) antigenic structure at the molecular level that creates “fingerprints” or “signatures” of various hepatitis B viral (HBV) strains. This technique employs high affinity IgM and IgG monoclonal antibodies (anti-HBs) directed against distinct and separate determinants on HBsAg. In performing this antigenic structural analysis, separate binding curves for different monoclonal anti-HBs are generated by measuring immunoreactivity in serial dilutions of HBsAg-positive serum by radioimmunoassay. Since the HBsAg concentration in serum is unknown, the binding profiles of groups of samples are aligned by an iterative least-squares procedure to generate the numerical signature characteristic of the viral strain. The numerical signatures are then displayed on a computer-graphic plot. The signature profiles of HBsAg subtypes are a true reflection of their antigenic structure, and in vertical and horizontal transmission studies the molecular characteristics of the viral epitopes are conserved. By signature analysis we found substantial antigenic heterogeneity among the ayw strain both in the U. S. and France, as well as in populations of the Far East and Africa. Populations in Ethiopia, Gambia, and the Philippines were infected with two antigenically distinct HBV strains. In some newly identified HBV strains, it was found that epitopes identified by some monoclonal antibodies were absent or substantially reduced, which suggested that a genetic mutation may have occurred. Thus this study suggests that there is far more antigenic heterogeneity in HBV than previously recognized. These variants are antigenically distinct from each other at the epitope level, and were heretofore unrecognized by polyclonal anti-HBsAg antibodies.

Methods

Monoclonal antibodies. The immunization protocols, cell fusion technique, and growth and cloning of hybridomas producing anti-HBsAg antibodies have been previously reported (8). The monoclonal anti-HBsAg antibodies have been characterized with respect to specificity for determinants on HBsAg, antibody class and subclass, and affinity for HBsAg-associated determinants (9-14). Monoclonal antibodies designated 5C3 (IgG2a), 5C11, 2C6, 1C7, 4E8 (IgG1), and 5D3 1F8, 2F11 (IgM), were selected from a library of monoclonal anti-HBsAg antibodies because of their following special properties: (a) several bind to all known subtypes of HBsAg (by definition a domain epitopes [antibodies 5D3, 5C3, 5C11, and 2C6] (15); (b) some demonstrate qualitative binding differences to ad and ay subtypes of HBsAg (antibodies 1F8, 2F11, 1C7, and 4E8) (8); (c) they recognize distinct and separate determinants on HBsAg by competitive binding studies (11); and (d) they possess very high affinity constants for HBsAg-associated epitopes (range, 4.8 × 10^9-4 × 10^11 liters/mol per molecule).

Radioimmunoassays (RIAs). We employed eight monoclonal “simultaneous sandwich” RIAs for analysis of HBsAg-associated binding activity in serum samples. In brief, polystyrene beads were coated with a monoclonal IgM anti-HBsAg designated 5D3, and the other eight antibodies, including 5D3, were radiolabeled to a specific activity of 10-12 μCi/μg of protein (1 Ci = 3.7 × 10^10 becquerels) with the Hunter-Bolton reagent (16). The 5D3 coated beads were incubated with serial 10-fold dilutions of serum samples (100 μl) and radiolabeled probe (100 μl) consisting of ~150,000 cpm of monoclonal anti-HBsAg for 4 h at 45°C. The beads were washed extensively with distilled water and the radioactivity bound to the bead was measured in a gamma counter.

Statistical data analysis. The antigenic structure of a HBV subtype is characterized by its binding activities to a panel of monoclonal antibodies over a range of viral concentrations. The statistical problem involved in establishing the antigenic structure of a HBV subtype is therefore one of estimating the multiple response functions \( f_i(\lambda) \) for \( i = 1, 2, \ldots, 8 \) between \( \log_{2} S/N \) (defined as mean cpm bound in the experimental samples/mean cpm of negative control serum) and \( \log_{10} \) dilution factor.

1. Abbreviations used in this paper: anti-HBs, antibody to hepatitis B surface antigen; CEP, counterelectrophoresis; HBsAg, hepatitis B surface antigen; HBV, hepatitis B viral.
(x > a reflection of viral concentration) for the eight selected antibodies, using serum samples from that subtype. It is hypothesized that each f^*(x) is a smooth growth curve which increases from zero to a unique local maximum with increasing viral concentration, and then decreases with further increase in concentration because of a "prozone" effect. We use the dilution scale in this paper instead of the opposite-running concentration scale because we know the relative dilution factors but not the viral concentrations.

Let y_i, A be the log S/N value obtained for antibody A (A = 1, 2, ..., 8) at log_10 dilution factor d (d = 0, 1, ..., 5) of the ith serum sample (i = 1, 2, ..., n) from a HBV subtype. Then the statistical model for the RIA's data can be expressed as follows:

\[ y_{i,A} = f^*(x_i + d) + e_{i,A}, \quad d = 0, 1, ..., 5; \]
\[ A = 1, 2, ..., 8; \quad i = 1, 2, ..., n, \]

(1)

where x_i is the unknown log_{10} concentration of the ith serum sample at d = 0, the e_{i,A} are the error terms with \sigma_i^2 = \sigma^2 and standard derivation \sqrt{\sigma_i^2} = \sigma, and the f^*(\cdot) are of the same functional form. The x_i values cannot be exactly determined because serum samples are complex protein mixtures. Since the x_i values are unknown, we cannot use standard curve-fitting methods to estimate the f^*(x) functions directly from the data. We have therefore developed an iterative least-squares procedure for aligning the concentrations of serum samples from the same subtype. The alignment is affected by first setting x_1 = 0, and then estimating the other x_i values (i = 2, 3, ..., n) using the statistical model given in Eq. 1. The aligned data are subsequently used to estimate the f^*(x) functions which collectively characterize the antigenic structure of a HBV subtype.

Iterative alignment-and-fitting procedure. The proposed procedure for aligning the unknown x_i values (i = 1, 2, ..., n) of serum samples from a HBV subtype, and subsequently estimating the multiple response functions f^*(\cdot) which characterize the subtype, consists of the following stages: (Stage 1) Initial alignment: select a "leader" whose undiluted log_{10} viral concentration x_1 is set to be zero. Using the eight piecewise linear functions defined by the log S/N values obtained for this leader at x = 0, 1, ..., 5, the other x_i values (i = 2, 3, ..., n) on this selected scale are then estimated using a least-squares method. (Stage 2) Curve fitting: estimate the response functions f^*(A = 1, 2, ..., 8) from the aligned data using multiple regression techniques. The overall sum of squared residuals is also computed. (Stage 3) Updating alignment: using the f^*(\cdot)'s obtained in step 2, the estimates of x_i values (i = 1, 2, ..., n) are updated. If the resulting decrease in the overall sum of squared residuals is less than a specified threshold, stop; otherwise, go to stage 2.

A detailed description of the proposed alignment method is given in the following, but it should be noted that this procedure only provides estimates of the (x_i - x_1) values, and not the x_i values themselves.

Initial alignment stage. (Step 1) The serum samples with the most 1 = i_{max} < n y_{i,A} values among the eight possible maxima is selected as the leader sample, whose viral concentration x_1 is set to be zero. (Step 2) For A = 1, 2, ..., 8, do the following: set y_{1,A} = 0 for d = 6, 7, 8, 11, because 10^6 or higher dilutions of practically all serum samples would contain effectively no viral proteins; then, for d = 0, 1, ..., 10, compute the intercept a_d^A and slope b_d^A of the line joining the two points, (d, y_{d,A}) and (d + 1, y_{d+1,A}). (Note that for A = 1, 2, ..., 8, the piecewise linear functions f^*_d(x) = a_d^A + b_d^A x, d \leq x < d + 1; d = 0, 1, ..., 11 defined on the log_{10} dilution scale is an approximation of the unknown f^*(\cdot)). (Step 3) For i = 2, 3, ..., n, we find the value of \hat{x}_i to minimize the sum of squared residuals: SSR_i = \sum_{d=0}^{5} (y_{i,A} - \hat{f}_d(x_i + d))^2. This is achieved by computing for j = 0, 1, 2, ..., 5, \hat{\hat{x}}_i = \sum_{d=0}^{5} \frac{b_d^A (y_{i,A} - a_d^A - d b_d^A)}{\sum_{d=0}^{5} b_d^A}, and its associated SSR_i values; \hat{x}_i is found by identifying the \hat{\hat{x}}_i with the smallest SSR_i value.

Curve-fitting stage. With the aligned data, a better approximation of the f^*(\cdot)'s than the \hat{f}_d^A's can be obtained by pooling the sample data from the same HBV subtype. Our objective is therefore to fit, for A = 1, 2, ..., 8, a function f^*_d to the aligned data (\hat{x}_i + d, y_{i,A}); d = 0, 1, ..., 5 and i = 1, 2, ..., n, with all the f^*_d's having the same functional form. For A = 1, 2, ..., 8, the fitting process is as follows: (Step 1) Resolve the log S/N values by using the transformation y_{i,A} = (y_{i,A} + 1) / 9; the resolved values range from 0 to 1 because the log S/N values range from -1 to +8. (Step 2) Compute the logistic transform of the resolved values:

\[ y^*(x_i + d) = \log \frac{1}{1 + (y_{i,A} + 1)^{-1}} = \log \frac{1}{8 - y_{i,A}}. \]

(Step 3) Do a multiple regression using y^*(x_i + d) as the target variable and \hat{x}_i, x_i, and \hat{d}_i, and \hat{d}_i as the set of predicting variables. The regression coefficients obtained are denoted, respectively, by \hat{\beta}_d^A, \hat{\beta}_d^A, \hat{\beta}_d^A, and \hat{\beta}_d^A. Therefore, we use the logistic curves of the form:

\[ f_d^*(x) = \frac{3}{9} \exp \left( \sum_{i=0}^{3} \beta_d^A x_{i} \right) \left( 1 + \exp \left( \sum_{i=0}^{3} \beta_d^A x_{i} \right) \right) - 1, \]

as approximations of the unknown f^*(\cdot)'s. The overall sum of squared residuals, \sum_{i=1}^{n} SSR_i = \sum_{i=1}^{n} \sum_{d=0}^{5} (y_{i,A} - f_d^*(\hat{x}_i + d))^2, is also computed for the sample from each subtype.

Updating stage. With the new \hat{f}_d^A's, the x_i values, i = 1, 2, ..., n are reestimated in turn by finding the value of \hat{x}_i that would minimize SSR_i = \sum_{i=1}^{n} \sum_{d=0}^{5} (y_{i,A} - f_d^*(\hat{x}_i + d))^2. Since the updated estimates will be in the neighborhood of the previous estimates, and SSR_i is a continuous function of \hat{x}_i, the new estimates are obtained using a simple numerical algorithm. It has been found that the solutions after the first few iterations are practically the same, and hence in order to avoid excessive computing, a threshold value T is specified so that when the difference in \sum_{i=1}^{n} SSR_i, is not greater than T (~2\sigma), the iterative procedure stops; otherwise, the iterations will continue by going back to stage 2.

Study design
Analysis of known HBsAg subtypes. 64 serum samples, derived from known chronic HBsAg carriers and representing HBV subtypes in the U.S. were selected for definition of viral antigenic structure by signature analysis. These specimens had been classified as ayw1, ayw2, ayw3, adw2, adw4, and adr HBsAg subtypes using conventional polyvalent anti-HBs antibodies (17-19). In addition, 39 serum samples from Centre National de Transfusion Sanguine, Paris, France (kindly provided by A. M. Courouce), were also analyzed by the eight monoclonal RIA's. These samples had been classified as ayw1, ayw2, ayw3, special ayw4, which is an intermediate between ayw2 and ayw4, adw2, adw4, and adr by counter electrophoresis (CEP) using absorbed polyvalent anti-HBs antiserum (20).

Analysis of the molecular characteristics of HBsAg-associated epitopes during vertical and horizontal transmission of HBV infection. (a) Two serum samples from each of the five individuals, drawn 9–12 mo apart, were analyzed in order to investigate the stability of viral epitopes. (b) The signature profile as exhibited by the eight monoclonal antibodies were compared in serum samples obtained from a mother and her 3-mo-old infant, who was presumably infected at birth or in utero. (c) Seven members of a family with a high incidence of HBV infection were investigated by signature analysis. It is noteworthy that the mother had acute hepatitis B, and her blood was tested during acute stage of the disease. It was found to be positive for HBsAg and HBsE, and contained IgM anti-HBC antibodies by conventional RIAs (Abbott Laboratories, No. Chicago, IL). Another serum sample tested 2 mo later was negative for HBsAg and anti-HBs antibodies were now present. It was therefore assumed that the HBV infection in this family had been transmitted horizontally. The distribution of HBV serologic markers in 21 family members living in the same household is depicted in Fig. 4.

Populations. HBsAg-positive serum samples from four different populations derived from various parts of the world were investigated. The composite signature profiles characteristic of HBV from each of these
populations, or subgroups, identified within a population, were compared with signatures of classic HBV strains from the U. S. and France. The following populations from various geographic regions of the world were studied:

(a) We studied 33 non-Jewish residents of Israel representing a stable permanent Middle-Eastern population. The HBsAg-positive sera were obtained from blood donors and other chronic carriers. These samples were identified by routine testing for HBsAg at the Rambam Medical Center, Haifa, Israel. Individuals were randomly selected from the population at large for signature analysis. In addition, 12 HBsAg-positive chronic carriers representing Ethiopian immigrants to Israel were investigated. Sera from these subjects were obtained between 1981 and 1983, a short time after their arrival to Israel from Ethiopia. The Ethiopians, therefore, were thought to represent a stable homogenous population. (b) 76 HBsAg-positive chronic carriers from Gambia with (21) and without (22) hepatocellular carcinoma were also investigated. Clinical details of this population will be reported elsewhere. (c) 55 patients from the Philippines were studied. HBsAg-positive individuals were randomly selected from the general population and were derived from patients with hepatocellular carcinoma (22), acute and chronic viral hepatitis (15), and asymptomatic chronic HBsAg carriers (10).

Results

Definition of a classic HBsAg subtype by signature analysis. As shown in a representative example depicted in Fig. 1, sera from 14 HBsAg-positive individuals of the adw2 viral strain from the U. S. have been studied with eight different monoclonal RIAs, each of which detects a separate and distinct epitope on HBsAg. There is considerable variability in HBsAg concentration in serum when comparing one individual with another, even though all individuals are of the same subtype as measured by polyvalent anti-HBsAg antibodies (17-19). This phenomenon is reflected by the scatter in the plots of loge S/N vs. log10 dilution factor (Fig. 1; top panel). However, when the data are aligned by the iterative least-squares procedure, a distinctive binding profile emerges for each of the monoclonal antibodies (Fig. 1; bottom panel). The aligned individual data points closely fit to the developed curve and a scatter of binding values is only observed at the lower part of the curve when loge S/N is <0.9 (S/N < 2.5). In the assay design, a sample is considered positive for

Figure 1. Computer- graphic plot of signature analysis. (Top panel) Unaligned binding values generated by eight monoclonal RIAs of HBsAg-positive serum samples from 14 individuals of the adw2 subtype. Note the scatter of binding values due to the differences in HBsAg serum concentrations among members of the group. (Bottom panel) Aligned binding values by the computer program (Methods) defines the signature of the adw2 HBV strain.
HBsAg when the S/N values of ≥2.5 (12) are observed. (S/N is defined as the mean cpm in the experimental sample divided by the mean cpm of the negative control.) Therefore, the scatter of binding values is observed only at dilutions negative for HBsAg binding activity. In subsequent figures we will include all the data points of the individual sample dilutions. However, binding values of log₁₀ S/N < 0.9 are considered negative. When the binding profiles of the eight different monoclonal antibodies are taken together, these curves form a numerical signature or fingerprint of the adw₂ subtype.

Structural stability of HBsAg-associated epitopes. The following studies were designed to establish the stability of HBsAg antigenic structure as measured by signature analysis and to evaluate the effect of host factors, such as glycosylation of HBsAg, on the binding profiles exhibited by the eight monoclonal RIAs. Two serum samples, drawn 9–12 mo apart from each of five individuals, were analyzed. The results of three representative comparisons are presented in Fig. 2. The binding profiles of the two separate samples obtained from each individual are identical, and align on the same binding curves for all eight monoclonal antibodies.

The antigenic properties of HBsAg derived from a mother and her 3-mo-old infant, who was presumably infected at the time of delivery, are compared in Fig. 3. There is a remarkable homogeneity of the two signatures, and one (infant) can be superimposed on the other (mother). Further analysis of the two signatures reveals that antibodies 5D3, 2C6, 5C11, and 5C3 demonstrate high binding to HBsAg-associated determinants, while viral epitopes recognized by antibody 4E8 appear to be present in low concentration. Viral epitopes recognized by antibodies 1C7, 1F11, and 1F8 are not present, or are in very low concentration on this HBV strain.

We studied a family with a high incidence of HBV infection; intrafamilial spread was presumed to be horizontal, since the mother was the last member of the family to acquire HBV infection (see Methods). The distribution of HBV serologic markers in 21 members of the family is depicted in Fig. 4 (top). Anti-HBs antibodies were present in 11 individuals. Eight others, including the mother, who had acute hepatitis B, were positive for HBsAg, and the remaining two were free of any serologic evidence of past or present HBV infection. The combined signatures of seven HBsAg-positive family members are presented in Fig. 4 (bottom). Indeed, all seven HBsAg-positive family members had the identical signature, and thus harbored the same viral strain.

Heterogeneity of viral antigenic structure within classic HBsAg subtypes. Fig. 5 depicts the substantial heterogeneity in the antigenic structure of the ayw₃ viral strain by signature analysis. Initially, the ayw₃ subtype had been identified by conventional polyclonal RIAs (17–19) or by CEP (20) as a single homogeneous viral strain. Two of the ayw₃ variants were from the U. S. and have been characterized as ayw₃ and subgroup ayw₃
by their signature profiles (7). The other two were derived from France and previously identified as a homogeneous ayw3 viral strain and ayw1-2-intermediate by polyclonal anti-HBsAg antisera (20). Substantial qualitative differences in the binding profiles were observed. The most striking quantitative and qualitative differences were observed between the classic ayw3 subtype found in France compared with the U. S. viral strains. HBsAg-associated epitopes measured by antibodies 1C7, 5C11, 4E8, and, to a lesser extent, 1F8, were found to be in higher density on ayw3 derived from France, as shown by higher binding values in the respective RIAs. The French strain classified as ayw1-intermediate gave a signature profile very similar to the U. S. ayw3 even though a qualitative difference was noted in the HBsAg-associated epitope recognized by 5C11, as expressed by a shift in the binding curve. Note that the greatest antigenic heterogeneity was demonstrated by the 5C11 epitope on the ayw3 viral strains.

**Structural diversity of HBsAg-associated epitopes**

To assess whether variations in the antigenic structure of HBsAg was detectable in HBV strains derived from different parts of the world, signature analysis was performed on HBsAg-positive individuals from four geographically separated populations.

**Non-Jewish population of Israel.** The binding profiles were generated by the eight monoclonal RIAs in 33 HBsAg-positive carriers of non-Jewish origin of Israel; such individuals represent a permanent Middle-Eastern population. The composite binding curves analyzed by the iterative least-squares method demonstrated striking homogeneity (Fig. 6, top); thus, all individuals appear to harbor the same HBV viral strain. The composite signature profile of this HBV strain most closely approximate

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**Figure 4.** Stability of the antigenic structure of HBsAg in intrafamilial horizontal transmission of HBV infection. (Top) Distribution of HBV serologic markers in 21 family members. (Bottom) A composite binding profiles generated by monoclonal RIAs from HBsAg-positive members of the family. Note that all are infected with the same HBV viral strain.

**Figure 5.** Comparison of signature profiles generated by four variants of the ayw3 strain, which was previously identified in the U. S. and France as a homogeneous HBV subtype by polyclonal anti-HBs antibodies. Note the qualitative and quantitative differences in the binding profiles exhibited by antibodies 5C11, 1C7, 4E8, and 1F8. (— — — 14 individuals with U. S. A. ayw3; (— — 8 individuals with U. S. A. subgroups ayw3 (see reference 7); (— — — — 5 individuals from France with ayw3; (— — — — — 5 individuals from France with ayw3 “intermediate” (see reference 20).

**Figure 6.** Signature analysis of 33 HBsAg-positive non-Jewish residents of Israel. (Top panel) Aligned binding profiles exhibited by eight monoclonal antibodies. Note the homogeneity of the binding profiles, indicating that all harbor the same HBV strain. (Bottom panel) Comparison of the HBV strain from Israel to the signature of 14 individuals from the U. S. A. who are infected with the classic ayw1 HBV strain. The residents of Israel are infected with a variant of classic ayw1. They differ from each other by the binding profile exhibited by antibodies 4E8 and to a lesser extent 2C6. Also note that the epitopes recognized by 2F11 and 1F8 are present in higher concentration on the Israeli strain. (—) ayw1 variant non-Jewish Israeli; (—) ayw1 from the U. S.
the signature of the ayw2 strain derived from the U. S. (Fig. 6, bottom). It is of interest that previous studies using conventional polyclonal antibodies classify such HBsAg-positive individuals as ayw2; this is the subtype most frequently found in the population (23) and in other regions of the Middle-East (20). However, viral heterogeneity was observed when comparing the Israeli ayw2 group to the classic U. S. ayw2 viral strain. The antigenic determinant recognized by monoclonal antibody 4E8 was present in higher density on the ayw2 viral strain in Israel, as shown by the higher binding values exhibited in the RIA. In addition, an increased density of epitopes detected by monoclonal antibodies 2F11 and 1F8 were also observed in the Israeli strain. Furthermore, qualitative differences in the epitope composition were demonstrated by shifts in the binding curves generated by antibodies 4E8 and, to a lesser extent, 2C6. The epitopes identified by antibodies 5D3, 1C7, 5C11, 5C3, and 1F8 were identical in antigenic composition in both groups. Finally, since the signature profiles had been previously generated for the classic HBV viral strains, which represent all of the nine Paris conference subtypes, it was possible to compare the non-Jewish population to each classic subtype. Thus, the composite profiles generated by the computer-graphic plots were compared with each other before the closest match was obtained. The non-Jewish group is therefore classified as a variant of ayw2 when compared with all known classic HBV subtypes previously determined by polyclonal anti-HBs antibodies.

**Ethiopian population of Israel.** 12 HBsAg-positive chronic carriers from Ethiopia, who immigrated to Israel, were analyzed by the eight monoclonal RIAs. Two distinct signature profiles representing different HBV strains were observed (Fig. 7, top, middle). Comparison between the two signatures as depicted in Fig. 7 (bottom) demonstrates considerable diversity in epitope antigenic structure between the two Ethiopian HBV viral strains. Thus, the geographical region from which the Ethiopians were derived appears to harbor two distinct HBV subtypes. The signature profiles of the HBsAg-positive Ethiopians most closely resembles the signature profile identified as HBsAg subtype adw2 and ayw3, respectively, of the U. S., when comparisons were made to other classic HBsAg subtypes (data not shown).

**The Gambian populations.** We studied 76 HBsAg-positive individuals from Gambia; two distinct signature profiles were observed. There were 16 individuals in the first group (Fig. 8, top panel), and 60 classified as a second distinct group (Fig. 8, middle panel). Fig. 8, (bottom panel) depicts a comparison of the signature profiles illustrating the antigenic difference between the two viral strains. For example, identical binding characteristics by antibodies 5D3, 5C11, 5C3, and 4E8 were observed. It was, however, possible to clearly distinguish between the two HBV strains by qualitative difference in epitope structure identified by antibody 2C6, and larger differences in epitope composition were observed by the binding profile generated with antibodies 1C7, 2F11, and 1F8. We observed that the signature of one Gambian strain most closely matched to the classic adw2 subtype found in the U. S. (Fig. 9, top panel). Identical binding profiles were generated in both groups by antibodies 5D3, 2C6, 5C3, 2F11, and 1F8, and only minor qualitative differences were noted with antibodies 5C11 and 4E8. The signature profile of the second group of Gambians was compared with other classic subtypes as defined by signature analysis. It was found that the ayw4 subtype previously identified by a polyclonal immunoassay in France was identical to the Gambian group as shown by nearly complete homology of the two binding profiles (Fig. 9, bottom panel). No ayw4 U. S. strain was available for comparison.

**Philippine population.** We studied 55 HBsAg-positive individuals from the Philippines in order to identify the prevalent HBV strains in a stable homogenous population of this country. 38 individuals were classified by signature analysis to harbor a single HBV strain, while 17 others clearly represent an infection with a different antigenically distinct virus (Fig. 10, top and middle panels). Comparison of the two signatures (Fig. 10, bottom panel) suggests very similar epitope composition as measured by antibodies 5D3 and 5C3; minor quantitative differences were observed with monoclonal antibodies 2C6 and 5C11. The most pronounced differences between the two HBV strains was dem-

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*Antigenic Variations of Hepatitis B Surface Antigen by Monoclonal Antibodies* 1343
Ben-Porath compared were strain found on strain ayw3, it was found that the signature profile was very similar to an ayw3 strain found in the U. S. Alternatively, the signature was virtually identical. Thus, the Gambians are infected with the classic ayw4 French strain.

Discussion

The hepatitis B surface antigen has three major antigenic domains: the group specific antigen α, and the subtype specific d or y and r or w. Thus, HBV was originally classified into four major subtypes, adr, adw, ayw, and ayr (24). However, additional subtype specificities have been described and the classification of HBsAg now include nine main subtypes, namely, ayw1, ayw2, ayw3, ayw4, ayw5, ayw6, and ayr, and the very rare ayr, found exclusively in the Far East; all of these groups belong to the y type, whereas adw2, adw4, adw5, adw6, and ayr belong to d type as measured by polyclonal anti-HBs antibodies (25). Therefore, most of the subtyping of HBV strains to date has been accomplished by polyclonal monospecific antiserum prepared by immunoadsorption of antiserum to one subtype with HBsAg representing another subtype (20). Monoclonal anti-HBsAg antibodies generated to the native antigen potentially represents a much more precise technique to investigate the diversity in antigenic structure of HBsAg or, alternatively, the interrelationship of various HBsAg subtypes (7, 26).

In the present and previous studies (7), we have described an approach, called signature analysis, for the study of HBsAg...
The most important features of this technique include: (1) the use of high affinity anti-HBsAg monoclonal antibodies; (2) the selection of monoclonal antibodies that recognize distinct and separate epitopes on HBsAg; (3) the linkage of an IgM antibody designated 5D3 to the solid-phase support; 5D3 recognizes all known subtypes of HBsAg; (4) the construction of M-RAs specific for different HBsAg-associated epitopes, and, most importantly, (5) a statistical technique that aligns the unknown HBsAg concentrations present in serum derived from different individuals who harbor the same HBV strain.

The main advantages of the signature analysis technique are the capability of distinguishing subtle and major antigenic differences between related viral strains heretofore undetectable by conventional polyclonal antibodies. In addition, the technique is easy to perform; once the physical characteristics of the monoclonal antibodies have been defined, analysis of a large number of samples is possible. Finally, and most importantly, the concentrations of viral protein in complex protein mixtures need not be known.

We are led to believe that this method will be quite useful in studying the antigenic characteristics of a number of RNA and DNA viruses either in cell culture, serum, or other biologic fluids. For example, by using this analytical method we have shown that the a domain, which is defined as an antigenic region common to all known HBV subtypes (27), is multideterminant. At least four antigenic sites are detected by four noncompeting antibodies on all known subtypes, although substantial quantitative differences in the magnitude of binding activity has been observed (7). The greatest antigenic diversity has been found when comparing the d and y domains. These findings are in accord with recent sequence data showing substantial changes in amino acid composition between HBsAg subtypes adw and ayw (28). The antigenic differences between w specificities were found to be very small, and signature analysis demonstrated remarkable antigenic homogeneity under these circumstances (7). This observation is not surprising, since the w variants are thought to be subdeterminants of the common a domain (29, 30), and have been subsequently assigned to w specificities for reason of simplicity (31). More importantly, evidence has been presented to support antigenic diversity within the classic HBsAg subtypes. Indeed, ayw3, heretofore thought to be a single homogenous viral strain, was found to be composed of at least three subgroups, as shown by signature analysis. More recently, subdivision of the ayw3 subtype has also been observed by other investigators using monoclonal antibodies to HBsAg (26).

Attempts were made to assess the stability of HBsAg-associated epitopes, at the molecular level, either in the same HBV-chronic carrier observed over time or in a naturally occurring setting of vertical or horizontal transmission. The results presented here demonstrated that in serially studied chronic HBV carriers, the signature, or fingerprint of the viral strain remained constant over 9–12 mo (Fig. 2), and suggests that these epitopes detected by the eight monoclonal antibodies are characteristic of HBV. Analysis of the antigenic characteristics of poliovirus vaccine strains by monoclonal antibodies revealed, after in vivo multiplication of the virus in infants fed the vaccine strain, a complete change in the composition and function of certain epitopes within a short period of time (22). Thus, in a highly mutable virus like poliovirus, there appears to be a rapid change in the viral antigenic characteristics as defined by monoclonal antibodies. In this situation, the use of monoclonal antibody analysis in studying the interrelationship between a poliovirus isolate and its parent strain would have to take these observations into account.

Nevertheless, evidence for the stability of the HBV antigenic structure by signature analysis was obtained when investigations were performed on HBV transmission in man. It was found that the signature profiles of a HBsAg-positive mother and her 3-mo-old infant were identical (Fig. 3). The infant was presumably infected at birth or in utero of HBV from the mother at the time of delivery, since HBsAg-positive mothers may often transmit
infection to their infants (32–34). Indeed, further evidence for the stability of HBsAg-associated epitopes at the molecular level was provided by the family with a very high incidence of HBV infection. Familial clustering of HBV infection has been previously reported (35–37), in which intrahousehold and person-to-person modes of transmission appear to have been responsible for dissemination of the disease. Chronic HBV carrier mothers may be the most important mode of transmission among children at an early age (32–34). However, other close contacts between HBV infected siblings (38), or chronic carrier fathers (21), or spouses and children exposed later in life (36), are also possible routes of transmission. In the family described here, HBV transmission at birth seemed unlikely, since the mother was the last individual to acquire infection. Therefore, an intrafamilial horizontal transmission seemed likely. The signature profiles of the seven HBsAg-positive family members were identical, suggesting that all were infected with the same viral strain (Fig. 4). Thus, we have demonstrated that in utero or horizontal intrafamilial transmission of HBV, the signature profile as a reflection of the viral antigenic structure “breed true.” In addition, since the characteristics of HBsAg-associated epitope did not change during viral replication in different individuals over time, it appears that the viral signature is independent of host factors such as glycosylation of HBsAg.

It has been previously shown that the different subtypes of HBV have a widespread geographic distribution (20, 39). This observation suggests that HBV may have undergone independent genetic evolution. It is well established that the HBsAg subtype reflects the country of origin rather than the present country of residence. Even offspring of immigrants carry the HBsAg subtype specific to their parents country of origin (20, 40, 41). The ayw4 subtype is detected almost exclusively in West Africa (20), and was identified in our study only in individuals from Gambia (Fig. 6, bottom panel). On the other hand, ayw3 is found in many regions of the world (20). It is possible that the ayw3 variants identified in the present study represents the most mutable of the HBV strains (7, 26).

We investigated homogenous and presumably genetically stable populations of the Middle East, Africa, and the Far East by signature analysis. In an attempt to study antigenic variations of HBV, we compared our findings with a similar analysis of classic subtypes. This technique proved useful in establishing the homogeneity or heterogeneity among HBV strains in a defined geographical area. In three of the populations, namely Ethiopians, Gambians, and Philippines, two distinct HBV strains were identified within groups of individuals living in close proximity of the same geographic region. The signature profiles characteristic of one group of individuals residing in these geographic regions was then compared with the antigenic structure of other known classic HBV subtypes. Differences in the monoclonal antibody binding profiles exhibited by certain monoclonal antibodies between the classic subtypes and the study population were observed. Quantitative variations were often observed as reflected by changes in the apparent density of viral epitopes, as illustrated by antibody 4E8; for example, compare the non-Jewish population of Israel and classic ayw2 (Fig. 6, bottom panel). These quantitative differences in monoclonal antibody binding activity are consistent with an absolute alteration in the number of epitopes detected by a given monoclonal antibody. However, most of the HBV antigenic differences observed between the population groups and the classic subtypes were qualitative, and manifested by shifts in the binding curves. An example of this phenomenon is demonstrated in Fig. 9 (top panel).

We believe that the shift in the binding curve when comparing different population groups may be due, in part, to conformational changes in epitope structure. Such conformational changes in antigenic structure, however, appear common to all individuals who harbor a particular HBV strain. It is unlikely that the differences in the monoclonal binding profiles are due to host factors, since serial studies have shown no change in the binding profile in the same individual over time. In addition, during vertical or horizontal transmission, HBV-associated epitopes “breed true.” That is, HBsAg antigenic structure appears unaltered by signature analysis when passed from mother to child and when transmitted horizontally among other individuals. One observation of note was the difference in the binding profile observed in the Philippine group with antibody 1C7 (Fig. 10, bottom panel). This HBV strain closely resembled the U. S. ayw4 subtype, except the epitope identified by 1C7 is missing. Such lack of antibody binding may be explained in part by a genetic mutation in which the epitope recognized by 1C7 is so drastically altered as to eliminate all binding activity.

Comment is required on the antigenic characteristics of HBV strains studied in various populations of the world. An ayw4-like virus was identified by signature analysis in the non-Jewish Israeli population; this observation is consistent with results available by polyclonal antibody conventional subtyping analysis (20, 23, 42). In the Ethiopian population, viral strains similar to adw2 and ayw4 were identified. HBV subtypes adw2 appears to be the most prevalent strain in East Africa (20), and the finding that approximately one-half the Ethiopians harbored an ayw4-like strain was a somewhat unexpected finding. 78% of the Gambians had an HBV strain with a characteristic signature identical to the ayw4 classic subtype. This observation is in accordance with findings of a high prevalence of ayw4 in other West African countries, and specifically in Senegal, which surrounds Gambia on three sides (20). However, the second prevailing HBV strain in the Gambians resembled a variant of the adw2 classic subtype by signature analysis, an apparently new observation since in Senegal and other West African countries, ayw2 was the second most common strain identified by conventional polyclonal assays (20). In the Philippines, 69% of the HBsAg-positive serum samples generated viral antigenic signature most closely resembling adw2, and the remainder were quite similar to ayw3. The adr strain which is common in Southeast Asia was not detected in the Philippines population.

At present, accurate antigenic analysis and characterization of HBV isolates may be accomplished by generating a signature profile with monoclonal antibodies. This signature is a basic property of a given HBV viral strain. Subtle quantitative and qualitative differences between closely related HBV strains may be determined, and we are led to believe these will assist in understanding the genetic evolution of HBV in various parts of the world. Indeed, new HBV viral strains have already been identified; it is likely there are many others. Thus, HBV represents an important virus with far more antigenic diversity than previously recognized.

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References

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