Simultaneous Detection of Haemophilus influenzae Type b Polysaccharide-Specific Antibodies and Neisseria meningitidis Serogroup A, C, Y, and W-135 Polysaccharide-Specific Antibodies in a Fluorescent-Bead-Based Multiplex Immunoassay

Richard M. de Voer,1,2* Rutger M. Schepp,1 Florens G. A. Versteegh,3 Fiona R. M. van der Klis,3 and Guy A. M. Berbers1

Laboratory for Infectious Diseases and Screening, National Institute of Public Health and the Environment, Bilthoven, The Netherlands1; Department of Immunology, University Medical Centre Utrecht, Utrecht, The Netherlands2; and Department of Pediatrics, Groene Hart Ziekenhuis, Gouda, The Netherlands3

Received 3 October 2008/Returned for modification 4 December 2008/Accepted 26 December 2008

We expanded the meningococcal serogroup A, C, Y, and W-135 multiplex immunoassay (MIA) to simultaneously detect immunoglobulin type G antibodies directed toward Haemophilus influenzae type b polysaccharide (HibPS). The monoplex HibPS assay was compared to a HibPS-specific competitive enzyme-linked immunosorbent assay and showed a good correlation (R = 0.96). Furthermore, no cross-reactivity between HibPS and the four meningococcal serogroups was detected. This pentaplex meningococcal Hib MIA is a useful tool to investigate serological responses toward different childhood PS vaccines.

Childhood immunization programs are being expanded worldwide with various new vaccines: e.g., with Neisseria meningitidis serogroup C conjugate, pneumococcal conjugate, varicella-zoster virus, and hepatitis B virus vaccines. To reduce the number of injections, several vaccines have been developed that combine Haemophilus influenzae type b polysaccharide (HibPS), N. meningitidis serogroup C conjugate, diphtheria, tetanus, acellular pertussis, inactivated poliovirus, and/or hepatitis B virus vaccine, which in turn can be concordantly administered with other vaccines (6, 7, 9). The combination of these vaccines requires the evaluation or reevaluation of the immunogenicity of each vaccine component. Therefore, new methods have been developed, such as fluorescence multiplex immunoassays (MIAs) (1, 4, 10–13), in which the serological responses to the various vaccine antigens are determined simultaneously. MIA has the advantages of reduced laboratory time and the use of decreased amounts of specimen compared to those required by conventional methods such as enzyme-linked immunosorbent assay (ELISA).

Despite the fact that these new methods have many advantages, careful validation is needed before antibody responses toward different types of antigen (proteins or polysaccharides) can be measured in a single assay. Previously, an MIA was described in which meningococcal serogroup A, C, W-135, and Y polysaccharides were covalently attached to fluorescent beads via a poly-L-lysine (PLL) linker for the detection of antibodies against HibPS. Serum specimens used for evaluation of the HibPS MIA were from a study which evaluated the incidence of Bordetella pertussis infection during and shortly after pregnancy, executed during 2002 to 2006 (trial registration no. ISRCTN14204141). Blood samples were obtained from mothers directly postpartum and from the umbilical cord at the time of delivery. All participants had provided informed consent at the time of enrollment to use samples anonymously for future research. From this study, a subset of serum samples (n = 75) was selected which contain HibPS-specific antibodies over a wide concentration range, induced by natural exposure to Hib.

HibPS-specific antibodies were quantified in a competitive ELISA described in detail by Mariani et al. (5), with the modification that a different secondary conjugated antibody is used: alkaline phosphatase-conjugated goat anti-human immunoglobulin G (IgG) (Sigma, St. Louis, MO). This competitive ELISA reduces the overestimation of samples in the low concentration range compared to the more conventional noncompetitive method described by Phipps et al. (8). The free HibPS competition used in this ELISA allows elimination of day-to-day background variation typical in some sera; therefore, only values representing the real anti-HibPS response are determined (5).

We used two different approaches for the detection of HibPS-specific IgG antibodies in the MIA. Initially, HibPS conjugated to methylated human serum albumin (HbO-HA; Wyeth Lederle Vaccines, Pearl River, NY), similar to the antigen used in the ELISA (5, 8), was covalently attached to fluorescent carboxylated microspheres via a carbodiimide reaction as described by Pickering et al. (11). Secondly, HibPS (Chiron, Siena, Italy) was conjugated to PLL (Sigma-Aldrich, St. Louis, MO) and subsequently to fluorescent beads accord-

* Corresponding author. Mailing address: Laboratory for Infectious Diseases and Screening, National Institute of Public Health and the Environment, Antonie van Leeuwenhoeklaan 1, P.O. Box 1, 3720 BA, Bilthoven, The Netherlands. Phone: 31 30 2742496, Fax: 31 30 2748888. E-mail: Richarda.de.Voer@rivm.nl.

† Published ahead of print on 7 January 2009.
ing to the same procedure described for the meningococcal polysaccharides A, C, Y, and W-135 (3, 4). Subsequently, the MIA procedure for determination of total anti-HibPS IgG was performed as described for the meningococcal MIA (1, 4). Standardized reference serum lot 1983 (CBER/FDA) was used for quantitation of HibPS IgG, and standardized reference serum CDC 1992 (NIBSC, Potters Bar, United Kingdom) was used for quantitation of meningococcal serogroup A, C, Y, and W-135 IgG. For assay optimization, different types of serum diluents were used. One buffer contained 3% (wt/vol) bovine serum albumin (BSA; Sigma-Aldrich, St. Louis, MO) and 0.1% (vol/vol) Tween 20 (Merck, Darmstadt, Germany) in phosphate-buffered saline (PBS), pH 7.2 (13). A second buffer consisted of 50% (vol/vol) antibody-depleted human serum (ADHS; Valley Biomedical, Winchester, VA) in PBS (2, 4).

Intra- and interassay variation, the minimal level of detection, and the lower limit of quantitation were determined as described previously (1). Fluorescent beads conjugated with either HbO-HA or HibPS-PLL were subsequently tested using both serum diluents and compared to the results obtained by the HibPS competitive ELISA (Fig. 1). The results clearly indicate that when HbO-HA-conjugated beads with PBS containing BSA (Fig. 1A) were used, a reduction of nonspecific binding of antibodies with a concentration of <1 μg/ml was observed, compared to the use of the ADHS buffer (Fig. 1B) or compared to the HibPS-PLL beads with the BSA buffer (Fig. 1C). The best results, however, were obtained with HibPS-PLL-conjugated beads in combination with the ADHS buffer as a serum diluent (Fig. 1D). Using these beads, in combination with the ADHS buffer, the best correlation with the HibPS competitive ELISA was observed over a wide range of antibody concentrations.

We normally perform the meningococcal MIA with the ADHS buffer as a serum diluent (4). Therefore, to expand this assay for HibPS, we used the HibPS-PLL-conjugated beads. For implementation of HibPS in the meningococcal MIA, we tested 70 serum samples from the same panel (n = 75) in the meningococcal MIA (tetraplex), the HibPS MIA (monoplex), and these two assays combined (pentaplex), to examine if no interference between beads would occur. Results were compared by linear regression, and correlation coefficients were calculated ($R^2$ values). The individual $R^2$ values were 0.987, 0.970, 0.973, and 0.976 for meningococcal serogroups A, C, Y, and W-135, respectively, and 0.983 for Hib (Fig. 2). We observed no interference or cross-reactivity between the meningococcal and HibPS beads. For the determination of the sensitivity for HibPS, values obtained from blank wells were used and compared to the HibPS standard curve. The lower limit of quantitation was found to be 375 pg/ml anti-HibPS IgG. Intra- and interassay reproducibility was found to be high, with 5% correlation of variation and 19% correlation of variation, respectively.

Here we describe how the meningococcal tetraplex MIA can be expanded for the simultaneous measurement of HibPS-specific IgG. Due to the use of ADHS and conjugation of HibPS to PLL, we were able to measure IgG antibodies up to more than 100-fold more sensitively than in the HibPS competitive ELISA. The HbO-HA antigen used in conventional

**FIG. 1.** Comparison of anti-HibPS IgG levels measured by the competitive ELISA (cELISA) method with anti-HibPS IgG levels measured by the HibPS MIA. (A) ELISA comparison to HbO-HA-conjugated beads with 3% BSA serum diluent buffer. (B) ELISA comparison to HbO-HA-conjugated beads with 50% ADHS serum diluent buffer. (C) ELISA comparison to HibPS-PLL-conjugated beads with 3% BSA serum diluent buffer. (D) ELISA comparison to HibPS-PLL-conjugated beads with 50% ADHS serum diluent buffer.
ELISA methods can indeed also be used in MIA systems, for instance, in the assay described by van Gageldonk et al. (13). However, using HbO-HA for implementation in the meningococcal MIA, we observed a substantial amount of nonspecific binding in the low concentration range. By conjugating HibPS to PLL and using ADHS as a serum diluent, we bypass the fact that free HibPS competition is needed to overcome false-positive results. These data contribute to the growing awareness that MIAs are more sensitive and specific for measuring antibody responses than conventional ELISA methods.

We thank Pieter van Gageldonk for useful discussions and Mirte Scherpenisse for technical assistance (National Institute of Public Health and the Environment, Bilthoven, The Netherlands).

REFERENCES


