Characterisation of proposed reference materials for pertussis antiserum (human)
by an international collaborative study

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Abstract

Enzyme-linked immunosorbent assay (ELISA) has been widely used for evaluation of antibody responses to pertussis vaccination and infection. A common reference serum is essential for standardization of these assays. However, no internationally recognized reference serum is available. At the request of the Expert Committee on Biological Standardization (ECBS) of the World Health Organization (WHO), a set of four candidate international standards has been prepared.

These candidate materials have been assessed for suitability and compared with the widely used reference sera, US Human anti-pertussis reference sera lot 3, lot 4 and lot 5 by twenty-two laboratories from 15 countries in an international collaborative study. Laboratories measured IgG and IgA antibodies to pertussis toxin (PT), filamentous hemagglutinin (FHA), pertactin (PRN), and fimbriae (Fim2&3) using their established immunoassays. The results of this study showed each of the four candidates to be suitable to serve as an international standard. With the agreement of the participants, a recommendation has been made to the ECBS that the candidate material coded 06/140 be established as the First International Standard for pertussis antiserum (Human), with the following assigned International units (IU):

- IgG anti-PT: 335 IU/ampoule
- IgA anti-PT: 65 IU/ampoule
- IgG anti-FHA: 130 IU/ampoule
- IgA anti-FHA: 65 IU/ampoule
- IgG anti-PRN: 65 IU/ampoule
- IgA anti-PRN: 42 IU/ampoule
No formal unitage has been proposed for anti-Fim2&3 because most assays employed a mixture of fimbrial antigens. Additionally, the candidate material coded 06/142 has been proposed as a WHO working preparation for characterisation of assay systems.
Introduction

Serological analysis by Enzyme-linked immunosorbent assay (ELISA) has been widely used for evaluation of antibody responses to pertussis vaccination and infection. A quantitative measurement of concentration of serum antibodies in ELISA units (EU) per mL has been shown to be important in epidemiological studies (13, 19, 23), serodiagnosis of pertussis (1,2,5,7,17,24), and evaluation of responses to vaccines (3,21,22); however, the lack of internationally-recognized reference sera has hindered inter-laboratory comparisons and harmonization.

US Human anti-pertussis reference sera lots 3, 4, and 5 from the Center for Biologics Evaluation and Research (CBER), FDA, USA have been widely used and have played an important role in standardization of these assays (16, 18). However, only limited quantities of these sera remain. The WHO Working Group on the standardisation and control of pertussis vaccines recommended the preparation of a reference human antiserum to pertussis antigens with internationally-recognized status before the supply of the US preparations is exhausted (4, 25). As far as possible, continuity of unitage with that of the existing US reference preparations was recommended.

A set of freeze-dried candidate reference preparations has been prepared at the National Institute for Biological Standards and Control, UK (NIBSC), from sera obtained from German plasmapheresis donors. On behalf of WHO and in collaboration with members from CBER, FDA, USA and Institut für Infektiologie Krefeld GmbH, a collaborative study to compare these candidate international reference preparations with the US reference preparations was organized by NIBSC in 2007. The aims of the study were to characterize the candidate international reference preparations, to compare them to existing US and in-house reference preparations (IHR), and to define unitage for anti-PT, anti-FHA and anti-PRN for the candidates. In this manuscript we report the results of the collaborative study and additional studies evaluating the stability of the candidates.
Materials and methods

Participants. Laboratories actively performing serological assays measuring antibodies to pertussis toxin (PT), filamentous haemagglutinin (FHA), pertactin (PRN) and fimbriae 2 and/or 3 (Fims2&3) for the evaluation of human immunogenicity were invited to participate. A total of twenty two laboratories from fifteen countries, including vaccine manufacturers, diagnostic laboratories and research facilities participated in the study. Throughout this study, each laboratory has been identified by a randomly assigned code number from 1 through 22. All laboratories measured IgG anti-PT; but not all laboratories had assays for other antibodies.

Candidate preparations. (i)Collection and preparation of plasma samples.

Plasma samples were collected between June 2005 and November 2005 in Germany. All donors signed consent according to German law. A total of 2,500 donors of plasma or whole blood and 200 health care workers vaccinated with an acellular pertussis vaccine were screened for IgG-anti-PT with an ELISA in a single dilution protocol. Samples with values of $\geq 100$ EU/ml from a total of 72 donors and from 2 vaccinated adults were retested for the IgG anti-PT antibody concentration using previously published methods (24). According to their IgG-anti-PT antibody content two different groups of samples were defined: Samples with $\geq 200$ EU/ml IgG-anti-PT were classified as “high” and samples with a content $\geq 80$ EU/ml and $< 200$ EU/ml were classified as “low”. For the “high” IgG-anti-PT pool, we collected 28 plasma bags with ~250 ml and 4 plasma bags with ~750 ml. For the “low” IgG-anti-PT pool, we collected 46 plasma bags with ~250 ml and 9 plasma bags with ~750 ml.

Procedures for sample collection and preparation were carried out following the quality manual procedures in the blood bank of the HELIOS Klinikum Krefeld, where the plasma was obtained, processed and pooled according to the EU and German legal requirements for plasma intended for
human use. Plasmapheresis donations were frozen within a maximum time of 4h after plasmapheresis. Whole blood was processed within 24h after donation with an intermediate storage at +4°C and the separated plasma was frozen within 2h. All samples were then stored at -30°C.

Serum was prepared following a re-calcification procedure based on that used for the US reference sera. In brief, the plasma was thawed at 4°C. Then, 10 µl of sterile 2M CaCl₂ per mL of plasma was added and mixed in an isolator in a pharmaceutical class A laboratory. The re-calcified plasma in the bags was incubated in a stationary water bath for 30 min at +37°C. The bags were then gently shaken to dislodge the primary clots, and incubated in a shaking water bath at ~150 rpm for 120 min at +37°C. The bags were then incubated for 14 h at room temperature and subsequently centrifuged at 3,000 x g for 30 min at +4°C. The serum was transferred into a 450 ml bag and subsequently pooling was done by sterile connection of tubing (Terumo sterile tubing welder TSCD) using a manifold designed specifically for this study. The pooled serum samples were transferred into sterile plastic bags (3.5 l) and several 50 ml sterile bags (for pre-evaluation). All the frozen serum bags (approximately 9.5 L of high content and 19 L of low content) were transported frozen to the NIBSC by WorldCourier® in July 2006 and kept frozen (-20°C) until further processing.

(ii) Safety tests. Plasma samples were screened according to the German requirements. All samples were negative for Hbs-Ag, anti-HCV, anti-HIV 1-2 and TPPA. They also were negative for HCV-RNA and HIV-RNA by RT-PCR. The bulk materials were retested in NIBSC for HCV RNA (NAT test), Anti-HIV 1-2 and HBsAg. All samples were found to be negative.
(iii) **Lyophilization.** Four candidate reference preparations were prepared at NIBSC, UK from two pools of serum as detailed below.

On the day of filling, the bags were thawed at 37°C in a water bath. The material with no dilution or additions was filled in one ml aliquots into glass ampoules (meeting ISO 9187-1-2003) and then freeze dried according to the procedure described in WHO guidelines (26). Four batches of ampoules containing lyophilized serum were prepared on separate dates: the two batches prepared from the pool with the higher anti-IgG anti-PT content were coded 06/140 and 06/146 (4950 and 3300 ampoules respectively) and the two batches prepared from the pool with the lower anti-IgG anti-PT content were coded 06/142 and 06/144 (7800 and 7500 ampoules respectively).

**Baseline control serum.** Approximately 100 plasma donors were screened and one plasma sample from a single donor with antibody levels for IgG and IgA to PT and FHA, and antibody levels for IgG to PRN at/or below the assay detection limits was selected. This sample was dispensed into approximately 200 ampoules and freeze-dried for use as a baseline control (coded PM-06-047).

**US Reference Preparations.** Freeze dried ampoules of US Reference Pertussis Antiserum (Human) Lot 3 for IgG anti-PT and anti-FHA; IgA anti-PT and anti-FHA, lot 4 for IgG anti-PRN and lot 5 for IgA anti-PT, anti-FHA and anti-PRN were kindly donated by CBER, FDA, USA.

**Assay Method.** Enzyme-linked immunosorbent assay (ELISA) was used by all laboratories in the study (14, 15, 18), except for one, which carried out only CHO-cell neutralization assays. Two of the participants performed both ELISA and CHO-cell neutralization assays, and one participant performed complement fixation assays. Laboratories used their own methodology, reagents and
calculation methods. The majority of participants included in each assay at least one positive control serum from in house sources, typically as an in house reference preparation (IHR).

It was recommended that ampoules of lyophilised antisera be stored at -20°C. All materials were to be reconstituted in 1ml sterile distilled water and, if not used fresh, reconstituted antisera were to be stored in small aliquots at -20°C: if the reconstituted antisera were to be stored for more than 30 days, a temperature of -70°C is recommended. Repeated freeze-thaw cycles of reconstituted antisera were to be avoided. If feasible, a pilot study to determine suitable dilutions was suggested.

**Study Design.** Preparations included in the collaborative study are listed in Table 1. In this report, the study codes A and D have been used for ampoules coded 06/140 and 06/146 respectively which were prepared from the serum pool with the higher anti-PT IgG content. Study codes B and C have been used for ampoules coded 06/142 and 06/144 respectively, which were prepared from the serum pool with the lower anti-PT IgG content. Study code E has been used for the base line control (PM-06-047).

Each participating laboratory received three sets of ampoules comprising 5 samples of human serum coded by letter, together with the US lots lot3, lot4 and lot5 reference preparations (Table 1). Participants were also asked to include their in-house reference (IHR). Laboratories were asked to perform the three independent assays on three different days and to include all samples in each assay. For each assay, dilution curves for each reference and sample preparation were to have at least two replicates per assay and preferably at least five doses in the linear region.

**Thermally accelerated degradation study.** Ampoules of each candidate international reference preparation that had been stored at temperatures of 20°C, 37°C, 45°C and 56°C, were transferred at
1, 3, 6 or 12 months to storage at -20°C. All ampoules which had been stored at the different temperatures and times were reconstituted and compared in the same assay with freshly reconstituted aliquots from ampoules of the same candidate which had been stored continuously at -20°C (baseline) for IgG antibodies for each of PT, FHA, PRN and Fims2&3.

**In use stability study.** Ampoules of the candidate 06/140, study code A, and the candidate 06/142, study code B, were reconstituted and aliquots were stored at -20°C for up to 128 days and at 4°C for up to 14 days. At each of several time points, stored aliquots were compared with aliquots from freshly reconstituted ampoules of the same candidate for IgG antibodies for each of PT, FHA, PRN and Fims2&3.

**Statistical Methods.** All raw data, including plate layout, serum dilutions, and optical density (OD) values were returned to NIBSC for analysis, to ensure as far as possible consistent treatment. Each participating laboratory used their own design and assay format. Thus, the assays have been carried out using methods and layouts familiar to the participants, and as far as possible these data are representative of the way in which the reference preparations will be used in practice. ELISA was carried out as a quantitative assay, using a variety of different plate layouts. A number of assay layouts included consistent placement of samples on microtitre plates. Although this offers some practical advantages, it introduces the possibility of biases which may be caused by non-random positioning and order of assay, as well as other factors (8, 9,12). Where possible for the contributed data, assays have been assessed for the occurrence of any statistically significant positional or order effects.
All raw data were plotted and examined both graphically and statistically for any gross anomalies or outliers and to assess the overall consistency of the dose–response relations (10).

The dose–response curves were transformed to give approximately linear log dose – (transformed) response lines and to allow use of the methods of multiple parallel line bioassay analysis (6, 20). In the majority of assays, a four – parameter logistic curve provided a satisfactory fit to the dose – response curve and responses, expressed as proportion relative to upper and lower asymptotes, have been transformed to ‘logits’ and the bioassay analysis has been carried out using an in-house program (11). Where replicate titrations were performed, the classical analysis of variance has been used to provide an assessment of the linearity and parallelism of the log dose – (transformed) response lines. Where that was not the case, the linearity of the dose – response lines has been assessed graphically and the parallelism of the dose – response lines has been assessed by comparing the deviations from parallelism with the deviations from linearity. Additionally slopes from assays within a laboratory using the same antigen have been assessed using analysis of variance to determine any consistent differences among them.

In all cases, estimates of relative activity have been determined as the displacement of linear parallel log dose – response lines using the methods of multiple parallel line analysis. The log dose – response lines for some of the IHR and other control preparations showed apparent non-parallelism to the lines for the US and candidate reference preparations. To ensure that there is no effect of these preparations on the common fitted slope; data for the unique in house preparations have been omitted from the analysis. Estimates of the relative activities of the candidate reference sera are based on comparisons among the candidates and the relevant US reference preparations only.
Data have been combined to give a single set of estimates for each of the three sets of ampoules assayed for each antigen in each laboratory. All comparisons among estimates of relative activity have been made using analysis of variance of the logarithms of these estimates. The intra- and inter-laboratory variability for the various comparisons have been determined using these estimates. Estimates have been combined as geometric means (GM), and the variances have been summarized as geometric coefficients of variation (GCV).

A limited number of values (less than 0.1% of all data, and less than 0.2% of data in any individual laboratory) were identified as ‘outliers’ based on the occurrence of a gross discontinuity in the dose – response curve. Such values were omitted from all analysis. In some cases, a number of serum dilutions gave responses at or near the upper asymptote of the four – parameter logistic function, and in such instances, responses above a threshold, typically 95%, have been excluded from the final analysis.

Where positional effects could be assessed, these were typical of those frequently observed for this type of assay, confirming that such effects occur, and that the probability levels for the classical analysis of variance may not be accurate. Moreover, the possibility of bias in the estimation of slopes and relative activity cannot be excluded.

Three laboratories contributed CHO cell assays for anti-PT antibodies and one laboratory carried out assays by three additional methods for PT as detailed. Data from these assays were not quantitative and results have been summarized as mean titre and as relative activities for the CHO cell assays, and as positive or negative results for antibodies by the other methods.
Results

Dose – Response Relations

The logit transformed responses did not show significant ($p < 0.05$) deviations from linearity or parallelism for the majority of preparations in the majority of assays. Where apparently significant deviations from linearity were observed, these were not consistent across assays for the same antigen in the same laboratory and the apparent significance was considered to be the result of inaccuracies in the probability levels of the statistical test consequent on the failure of the assumptions which underlie the analysis of variance. Baseline control serum, study code E, gave assay response data at or near the detection limit in many assay systems. Thus, all assay systems showed a consistent lack of response to sample E. Data for this sample have not been included in detailed analysis.

Although the IHR and in-house control preparations, which are unique to the various laboratories, have been excluded from the final analysis, the data contributed to this study suggest that, depending on antibody and assay system, up to about 30% of these preparations show consistent slope differences when compared with the US reference and candidate reference preparations. A small number (less than 5%) of assays showed consistent significant deviations from parallelism among candidate reference preparations and CBER references. In these cases, the slopes for A and D (06/140 and 06/146, pool with higher anti-PT IgG content) tended to differ consistently from the slopes for B and C (06/142 and 06/144, pool with lower anti-PT IgG
content), with slopes for the IHR and/or for the CBER reference equally likely to be similar to those for A and D or to those for B and C. These differences were not concentrated within particular antibodies or laboratories. These differences also need to be considered in the context of positional effects for the assay, and the possibility of bias in estimates of slope.

**Comparison of the candidate reference preparations with US reference preparations for the various antibodies**

Comparisons of the candidate reference preparations with the US reference preparations, as appropriate for the antibody, generally showed good agreement between laboratories (Figures 1 – 3). The pooled results for the anti-PT, FHA, and PRN assays are provided in Tables 2 and 3.

The pooled intra-laboratory variability has been determined using the variabilities among the geometric mean estimates for the sets of ampoules within a laboratory for any of preparations A, B, C and D in terms of each other or in terms of the relevant US reference preparation. Based on these estimated variabilities, GCVs for individual estimates for an ampoule set range from 15% to 30%, with corresponding 95% limits 75-133% of estimated value to 57-176% of estimated value.

Preparations A (06/140) and D (06/146) were prepared from the same serum pool and should be similar. Similarly, preparations B (06/142) and C (06/144) are expected to be similar because they were prepared from the same serum pool. The data (Tables 2 - 4) support this, with no significant differences detected between dose – response lines for the paired preparations. For each antibody, the overall mean potency of the laboratory geometric mean estimates of the relative activity of D in
terms of A or C in terms of B do not differ significantly from 1 and analysis of variance showed no significant inter-laboratory differences relative to the pooled intra-laboratory variances.

The close similarity between A and D, observed for all antibodies, suggests that for estimation of potency, A and D might be treated as equivalent to one another, and similarly that B and C might be treated as equivalent to one another. Thus, where appropriate, estimates for A and D have been combined and estimates for B and C have been combined (Tables 2 and 3).

Based on the comparison of the IgG anti-PT and anti-FHA activity of preparations A and D with US3, the GM of the results were 335 EU anti-PT IgG and 130 EU anti-FHA IgG per ampoule of A or D. Based on the comparison of the IgG anti-PRN activity of preparations A and D with US4, the GM of the results were 65 EU IgG anti-PRN per ampoule of A or D (Table 2).

Comparison of the IgG anti-PT activity and IgG anti-FHA activity of preparations B and C with US3, and with A and D, gives an estimated IgG anti-PT content of 106 EU and IgG anti-FHA content of 122 EU per ampoule. Comparison of the IgG anti-PRN activity of preparations B and C with US4, and with A and D, gives an estimated IgG anti-PRN content of 39 EU per ampoule (Table 2).

This study shows that the IgA anti-PT, IgA anti-FHA and IgA anti-PRN dose – response lines for preparations A, D, B and C do not differ significantly from the respective antibody dose – response lines for US5. Estimates of the IgA anti-PT activity of preparations A and D in terms of US5 show significant inter-laboratory variability. However, if estimates from two of the eight laboratories contributing these assays are omitted there is no significant difference between laboratories. A similar finding was obtained for estimates of the IgA anti-FHA activity of these samples. Estimates
of the IgA anti-PRN activity of preparations A and D did not differ significantly among the five laboratories contributing these assays. It is therefore recommended that preparations A and D have an IgA anti-PT content of 65 EU, an IgA anti-FHA content of 65 EU and an IgA anti-PRN content of 42 EU per ampoule based on its comparison with US5 (Table 3).

Comparison of the IgA anti-PT activity of preparations B and C with US5, and with A and D, gives an estimated IgA anti-PT content of 18 EU, IgA anti-FHA content of 86 EU and IgA anti-PRN content of 38 EU per ampoule (Table 3).

**Comparison of the IgG anti-Fim activity of the candidate reference preparations A, D, B and C with one another:**

The US reference lots 3, 4 and 5 do not have an assigned anti-Fim activity, and thus no unitage for the candidate preparations can be determined for anti-Fim activity using the US reference lots. Although one laboratory carried out assays for Fim2 and Fim3 separately and one laboratory carried out assays for IgA anti-Fim2&3, all other participants described their assays as ‘IgG anti-Fim2&3’. The relative potency for IgG anti-Fim 2&3 in terms of US lot 3 are given in Table 4. Estimates of the IgG anti-FIM 2&3 activity of preparations A, B, C and D relative to each other (BC/AD) are consistent between laboratories with a geometric mean result indicating that 1.07 ampoules of A or D are equivalent to one ampoule of B or C. Potency relative to US lot 3 using the mixture Fim 2&3 was found to be 0.44 mL (GM) US lot 3 equivalent to 1 mL of preparation A (or D) (Table 4). However, these estimates are based on assays using a mixture of antigens and the results may differ if reagents or methods are modified.
Stability

Candidate preparations which had been stored at 56°C, especially for longer than 6 months, could not be readily reconstituted. Estimates showed an overall tendency for the preparations stored at the higher temperatures (e.g. ≥ 37°C) and for the longer times (e.g. 12 months) to have lower relative activities when compared to the samples stored at -20°C. Samples of the candidate 06/140, study code A, and of the candidate 06/142, study code B, which had been stored at temperatures of 20°C, 37°C and 45°C for 12 months were assessed in greater detail in additional assays. These assays showed no significant deviations from linearity or parallelism. Based on these data from one laboratory, predicted yearly losses of activity at -20°C for each antibody for each candidate were less than 0.02% in most cases. These data indicate that the candidate materials are sufficiently stable to serve as international reference preparations. Confirmation of the stability after 5 (and preferably within 10) years by comparison of ampoules stored at -20°C with ampoules stored at -70°C is suggested.

In use stability study showed that there was no evidence of loss of activity with time although variation in estimates of relative activity was observed in some cases (data not shown). These results suggest that aliquots of the reconstituted candidate reference preparations can be used if they have been suitably stored. However, since reconstitution and storage conditions may differ between laboratories, it is recommended that laboratories carry out validation under their own conditions.
Discussion

The value of accurate measurement of antibodies to *B. pertussis* antigens by ELISA has been demonstrated for serodiagnosis (1, 2, 5, 7, 17, 24), epidemiological investigation (13, 19, 23), and evaluation of vaccine responses (3, 21, 22). A common primary reference serum is an integral component of harmonization and inter-laboratory comparisons. This study reports the evaluation of four batches of ampoules, A (06/140), B (06/142), C(06/144), and D(06/146) that were prepared as candidate international reference preparations. The US reference preparations with their assigned ELISA units (EU) have been widely used. Hence the antibody contents of the candidate reference preparations have been estimated in terms of the EU assigned to the relevant US preparations.

Estimates of the activities of preparations A to D relative to that of US 3 (anti-PT, anti-FHA) or US 4 (anti-PRN) showed good agreement between laboratories. Estimates of the anti-FIM activity of preparations A to D relative to each other did not differ significantly between laboratories and potencies for A to D relative to US3 broadly agreed among laboratories. The majority of assays in this study have been carried out using designs with non-random assignment of preparations and dilutions to assay position and order. Differences between laboratories for estimates of IgG anti-PT, anti-FHA and anti-PRN are generally less than 20% and it is possible that to some extent, these differences may result from positional effects. Differences among laboratories are not consistent for the antibodies to the different antigens. Overall results are summarised in Tables 2 - 4.

In general, between laboratories variances appeared to be larger for IgA assays in comparison to the IgG assays, although the absolute range of estimates (on a log scale) from smallest to largest was not generally larger. The apparently larger variances may be in part an artefact resulting from the
smaller number of laboratories. Other factors which might lead to greater inter-laboratory variation for IgA assays are less experience in performing IgA assay in comparison to IgG assay in the participating laboratories, relatively lower IgA antibody contents in these preparations and differences among the specificities of the assay systems.

Suitability of a preparation to serve satisfactorily as a reference reagent requires similarity of dose response relationships for the preparations to be compared. In this study, the majority of log dose – response lines showed no significant deviations from parallelism for preparations A, B, C and D and the appropriate US reference preparations. However, up to 30% of the various IHRs or in-house control included in this study by the participants showed statistically significant non-parallelism when compared to the US and candidate reference preparations as indicated by differences in slopes of individual assays or consistent trends across assays (data not shown). The methods of production of the antibodies, including the specific antigens used for immunization, may produce dissimilarity among preparations. These data illustrate the importance of considering similarity when selecting reference preparations.

Seven laboratories performed assays on IgG anti-Fim using a mixture of Fim2&3 as coating antigen and additionally laboratory 21 used separate recombinant Fim 2 and Fim 3 as coating antigens. US reference lots 3, 4 and 5 do not have assigned anti-Fim activity since monovalent type 2 and type 3 fimbriae antigen preparations are not available. Thus no unitage for the candidate preparations can be determined for anti-Fim activity in terms of the US reference lots. However, comparison of IgG anti-Fim activity of the preparations A, B, C and D with one another showed good inter-laboratory agreement with no significant difference between laboratories. Potencies for A, B, C and D relative to US lot 3 for IgG anti-Fim 2&3 showed good agreement between laboratories. The
proportions of Fim2:Fim3 may differ both in the antigens used for immunization and in the antigens used for coating plates. It should be noted that some laboratories have assigned an arbitrary value for US lot 3 in their anti-Fim ELISAs (18); however such assignments are for in-house purposes only and have no official status. In this study, a mixture of Fim2&3 antigens was used by all except one participant and were apparently obtained from the same source. This study thus provides no information about whether the effect of different mixtures of antigens could be detected differentially by different ELISAs. When monovalent antigens become available for Fim2 and Fim3, these relationships will need to be re-examined and suitable unitages assigned.

Data for the CHO cell assays for IgG anti-PT from three laboratories give relative activities (data not shown) which are consistent with the results observed for the ELISAs, but this study provides data from only three laboratories and is not sufficient for reliable estimation of neutralizing titre. Other assays carried out comprised one assay using a test kit for IgA anti-PT and IgA anti-FHA, one assay using immunofluorescence for detection of IgA and one assay using complement fixation for anti-PT. These assays gave results which are broadly consistent with those obtained by ELISA and CHO cell assays.

In this study, there was generally good quantitative agreement among the laboratories in the estimates of unitage for the candidate preparations relative to the US reference sera. This agreement was observed even though there were differences in source, purity and characteristics of the coating antigen, the characteristics of the conjugate, the buffers used for blocking and washing, and other assay conditions that could affect consistency in estimation (8, 16). Based on information provided by the participants, at least 7 sources of PT and FHA, at least 4 sources of PRN and 2 sources of Fim antigens were used as coating antigens in the study. These data suggest that differences between the
antigen sources and other reagents do not appear to have significantly affected the estimates of relative activity in the present study. Importantly, however, common reference sera and calculation methods were used for all data, suggesting that these may merit particular emphasis in harmonization activities.

Data from this study indicate that preparation A is not distinguishable from preparation D, and that preparation B is not distinguishable from preparation C using these assay systems. This shows that the preparation of ampoules as separate batches on different days from a single pool of serum has not differentially affected activity in these assays. It is therefore recommended that for assignment of unitage in this study A and D be treated as equivalent and that B and C be treated as equivalent, and thus we have also calculated estimates based on combinations of data from the paired preparations.

This study shows that the parameters of the dose–response curves of preparations A, B, C and D for both the IgG anti-PT and the IgG anti-FHA do not differ significantly from that of US lot 3 and estimates of the activity of these samples in terms of US3 are consistent between laboratories. Similarly, the parameters of the IgG anti-PRN dose–response curves of preparations A, B, C and D do not differ significantly from that of US lot 4 and estimates of the activity of these samples in terms of US4 are consistent between laboratories. The candidate materials have also been shown to have satisfactory predicted stability.

A recommendation (with the agreement of the participants) was made to ECBS, 2008, that preparation A (ampoule code 06/140) be established as the First International Standard for pertussis antiserum (Human) with assigned international units (IU) per ampoule of

IgG anti-PT 335 IU/ampoule
IgA anti-PT    65 IU/ampoule
IgG anti-FHA    130 IU /ampoule
IgA anti-FHA    65 IU/ampoule
IgG anti-PRN    65 IU/ampoule
IgA anti-PRN    42 IU/ampoule

These proposed assigned IU are based on the estimates for A (06/140) determined in comparison with the relevant US reference preparation, so that as far as possible, continuity between studies can be maintained. It is additionally proposed that the candidate preparation coded 06/142 (sample B, lower anti-PT activity) be made available as a WHO working preparation for pertussis antiserum (Human) which might be suitable for characterisation of assay systems. Preparations D and C also appear suitable to serve as international standards and it is recommended that these materials be retained.

The recommendation was accepted by ECBS in October 2008, and, beginning in 2009, these reference preparations will be available from NIBSC, UK for distribution. They can be used to assist in the standardization of immunoassays used to measure human antibodies to *B. pertussis* in vaccine studies of products in current distribution, as well as those under development, for refinement of serological methods used for diagnosis or surveillance, and, potentially, for development of assays for measurement of antibodies to antigens other than the ones evaluated in this collaborative study.
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References


Table 1. Sample Information‡:

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<tr>
<th>Study Code</th>
<th>Sample/References Ampoule Code</th>
<th>Description</th>
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</thead>
<tbody>
<tr>
<td>A</td>
<td>06/140</td>
<td>Freeze dried preparation of ‘high’ anti PT IgG content human serum</td>
</tr>
<tr>
<td>B</td>
<td>06/142</td>
<td>Freeze dried preparation of ‘low’ anti PT IgG content human serum</td>
</tr>
<tr>
<td>C</td>
<td>06/144</td>
<td>Freeze dried preparation of ‘low’ anti PT IgG content human serum</td>
</tr>
<tr>
<td>D</td>
<td>06/146</td>
<td>Freeze dried preparation of ‘high’ anti PT IgG content human serum</td>
</tr>
<tr>
<td>E</td>
<td>PM-06-047</td>
<td>Baseline control serum; freeze dried preparation of human serum with ‘undetectable’ IgG anti-PT content.</td>
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</tbody>
</table>

US Lot 3
US Reference Pertussis Antiserum (Human ) Lot 3
Freeze dried reference preparation of serum with the following unitage
- IgG anti PT 200EU/ml
- IgA anti PT 15EU/ml
- IgG FHA 200EU/ml
- IgA FHA 100EU/ml

US Lot 4
US Reference Pertussis Antiserum (Human ) Lot 4
Freeze dried reference preparation of serum with the following unitage
- IgG anti-PRN (pertactin) 90EU/ml
- IgA anti – PRN (pertactin) 25EU/ml

US Lot 5
US Reference Pertussis Antiserum (Human ) Lot 5
A secondary reference with unitage assigned based on US Lots 3 & 4
- IgA anti-PT 140EU/ml
- IgA anti-FHA 280EU/ml
- IgA anti-PRN (pertactin) 90EU/ml

‡ For use in assay, all preparations were reconstituted with 1.0 ml of sterile water.
Table 2. Overall summary results of estimates, with 95% confidence limits, for samples A-D (IgG) in terms of US lot 3 (for anti-PT and anti-FHA) and US lot 4 (for anti-PRN). Number of estimates (one estimate per laboratory) combined is indicated in parenthesis.

<table>
<thead>
<tr>
<th>Antibody</th>
<th>Estimates as Units of US references</th>
<th>Estimates for B&amp;C as equivalent ml of A&amp;D</th>
<th>Estimates for B&amp;C in terms of A*</th>
<th>Proposed IU for A&amp;D</th>
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*Using proposed IU for sample A
Table 3. Overall summary results of estimates, with 95% confidence limits, for samples A-D (IgA) in terms of units of US lot 5. Number of estimates (one estimate per laboratory) combined is indicated in parenthesis.

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<td>06/142</td>
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*Using proposed IU for sample A
Table 4. Laboratory Geometric mean (GM) of IgG anti-Fim 2&3 estimates of relative activity in terms of equivalent ml of US3.

<table>
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<tr>
<th>Laboratory code</th>
<th>Samples (GM)</th>
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<td>0.36</td>
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<td>0.38</td>
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<td>0.40</td>
<td>0.39</td>
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<td>0.48</td>
<td>0.59</td>
<td>0.47</td>
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<td>12</td>
<td>0.73</td>
<td>0.57</td>
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<td>17</td>
<td>0.43</td>
<td>0.45</td>
<td>0.40</td>
<td>0.38</td>
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</table>

Geometric mean of laboratory GMs

|                  | 0.46‡        | 0.43‡      | 0.48     | 0.45     |
| GCV              | 33%          | 24%        | 25%      | 29%      |
| 95% limits       | 0.32-0.65    | 0.33-0.58  | 0.37-0.64 | 0.33-0.62 |

‡ GM = 0.44 mL
Figure legends:

Figure 1. Summary for laboratory mean estimates of IgG anti-PT activity in terms of US3.

Estimates of laboratory geometric mean activity, number=lab code followed by A or D (06/140 or 06/146), or B or C (06/142 or 06/144) to indicate ampoule code. a. Estimates for samples A and D in terms of US3; b. Estimates for samples B and C in terms of US3

Figure 2. Summary for laboratory mean estimates of IgG anti-FHA activity in terms of US3.

Estimates of laboratory geometric mean activity, number=lab code followed by A or D (06/140 or 06/146), or B or C (06/142 or 06/144) to indicate ampoule code. a. Estimates for samples A and D in terms of US3; b. Estimates for samples B and C in terms of US3

Figure 3. Summary for laboratory mean estimates of IgG anti-PRN activity in terms of US4.

Estimates of laboratory geometric mean activity, number=lab code followed by A or D (06/140 or 06/146), or B or C (06/142 or 06/144) to indicate ampoule code. a. Estimates for samples A and D in terms of US4; b. Estimates for samples B and C in terms of US4
Figure 1

a.

b.
Figure 2.

a.

b.
Figure 3.

a.

b.