ANA REFERENCE SERUM #12

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ARTHRITIS FOUNDATION - CENTERS FOR DISEASE CONTROL REFERENCE SERUM FOR HUMAN ANTIBODIES TO rRNP/ Ribosomal P

Intended Use

For in vitro immunodiagnostic use as a reference human serum having high levels of antibodies to rRNP/Ribosomal P to be used to confirm the specificity of local standards.

Description of Reference Serum

Defibrinated plasma from a single donor was donated by SLR Research in Carlsbad, CA. Volumes of 0.50 ml were dispensed into borosilicate vaccine vials, freeze-dried, and sealed with butyl rubber stoppers while still under reduced pressure. Vials are stored at the CDC at -20°C. The mean dry weight and SD of material in 10 randomly selected vials after freeze-drying was 18.5 ± 0.3 mg. The residual moisture content after freeze-drying was less than 1.0%.

The vial contents were sterile as determined by U.S.P. approved methods of sterility testing, negative for hepatitis-B surface antigen by radioimmunoassay, negative for antibody to HIV by Western blot, and free of rheumatoid factors as determined by latex agglutination.

Reconstitution and Storage

Store the freeze-dried material, at -20°C until use. To reconstitute, the contents should first be shaken to the bottom by tapping the upper end. Before the stopper is removed, the vacuum should be broken by insertion of a hypodermic needle through the rubber stopper. Precisely 0.50 ml of distilled water should then be added, and the vial restoppered. The freeze-dried powder should dissolve readily with gentle swirling (avoid foam). Allow to stand for at least 1 h before use and store at 4°C until use, not later than 24 h after reconstitution. Although not recommended, the reconstituted material will withstand at least 8 weekly freeze-thaws without loss of activity. If future use of reconstituted material is contemplated, portions of the undiluted material sufficient for a single use should be stored at -70 °C and discarded after use.

α-Ribosomal P Antibody Content

This serum contains antibodies to rRNP/Ribosomal P. This antibody reacts strongly with cytoplasmic ribosomal P proteins (P0, P1, P2) as well as the synthetic C22 peptide. Nine international reference laboratories confirmed the specificity of this serum. Methods used include EIA, Western Blot, Immunoprecipitation and Luminex Laser Bead Assays. Two laboratories found the serum negative in Immunodiffusion using rabbit and calf thymus extracts and three labs found no cytoplasmic or nuclear staining in the IFA ANA using HEP-2 cell substrate.
Suggested Procedure for Standardization of Quality Control Reagents

Since the amount of AF/CDC reference preparation is limited, it should be used to calibrate secondary standards which can be run each day along with other samples being analyzed. The validity of any secondary standard depends on its having the same specificity as the primary AF/CDC standard (anti-nDNA, anti-Sm, etc.). The specificity can be determined by EIA, Western blot, double immunodiffusion (anti-nRNP, anti-Sm, anti-SSB/La) or by analysis for anti-nDNA (Farr assay, millipore filter assay, Crithidia test).

To confirm the specificity of a secondary standard in double diffusion or counterimmuno-electrophoresis:

1. Reconstitute the ampule as described above.
2. Use both undiluted (neat) and 1:4 dilutions of the AF-CDC sera and the secondary standard.
3. Arrange antibody wells such that the secondary standards (neat and 1:4) are each next to an undiluted and 1:4 dilution of the AF-CDC serum.
4. Run the routine test procedure using nuclear extract antigen.
5. Lines of identity should be obtained without extra lines or ambiguous reactions.

To calibrate a secondary standard (double diffusion, counterimmunoelectrophoresis):

1. Reconstitute the ampule as described above.
2. Prepare undiluted and serial four-fold dilutions up to 1:256 of the secondary standard and AF-CDC serum.
3. Run the routine test on all dilutions.
4. Relative potency = \( \frac{\text{reciprocal of end point dilution of secondary standard}}{\text{reciprocal of end point dilution of AF-CDC serum}} \)

If the secondary standard is stable, the relative potency obtained should not change when periodically recalibrated. Any changes in day to day results with the secondary standard are more likely due to changes in test performance. Titer results may be particularly useful in assessing lot to lot changes in antigen preparations and for comparison of your results with the laboratories which initially evaluated these sera (see antibody content above).

**Caution**

This serum was found to be negative for HBsAg, HBeAb, HCV, RPR, HIV1&2, HIVAg, and HTLV1&2 by FDA approved methods. Since no test method can offer complete assurance that these or other infectious agents are absent, this serum should be handled at the Biosafety Level 2 as recommended for any potentially infectious human serum or blood specimen (Centers for Disease Control, National Institutes of Health. *Biosafety in Microbiological and Biomedical Laboratories, 1st Edition, 1984, 11-13.*)

**Supplementary Information**


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