QA/QC for mIAA Assay

Establish of lab-cut-off

The cutoff index of mIAA was established as the 99th percentile in the control group (106 subjects, an index of 0.010). We constructed a receiver operating characteristic (ROC) curve among the newly diagnosed (within 2 weeks) patients with diabetes (n=104). The cutoff index of 0.010 corresponded to 66% sensitivity among diabetic patients and 99% specificity among normal controls.

*Intra-assay CV:* 12% (n=10)

*Inter-assay CV:* 16% (n=35)

**QA:**
1) All assays are run in duplicate, along with four standard samples (one high positive, one low positive, one ultra-low positive, and a negative control serum samples. Two-well duplicates for each sample will be aliquoted with two separated events (one-well aliquoting per sample for each process). Upon finishing sample aliquoting on the full plate, the plate should be hold against the light and checked from bottom for the wells missing samples. Every sample with index above 0.010 (above 99th percentile of upper limit of normal control) must be repeated in a separated assay. The 3rd will be run if 2nd disagree with 1st. The result will be reported as the 1st value of two agreement (+,+ or +,-,+).

2) The low positive, and one ultra-low positive must show positive, and negative control must be negative in each assay.

3) Shewart chart is plotted over time.

4) The lab should attend any national or international workshops or efficient evaluations if available.
Principle:
- Incubation of serum with labeled antigen with and without cold insulin overnight
- Precipitation of antibody-bound labeled antigens with protein-A/G Sepharose in a 96-well plate format, with each serum tested in duplicate
- Washing of the 96-well plates to remove unbound labeled antigens
- Counting of each well with a 96-well plate β counter.
- Results expressed as an index that adjusts the delta cpm of the test serum for the delta cpm of positive and negative control sera in a particular assay.

Plan for performing the assay:

Day 1:
- morning: retrieve and thaw sera to be tested
- mid-day: (1) set up incubation of sera in buffer 1 (Part III)
  (2) prepare protein-A/G Sepharose in buffer 1 (Part IV)
  (3) Coating the 96-well plates with buffer 1 (Part IV)

Day 2:
- morning: (1) add incubate to protein-A/G Sepharose in plates (Part V)
  (2) wash plates with buffer 2
  (3) dry plates
  (4) add scintillation liquid
- afternoon (4) count
  (5) analyse data (Part VI)
Part I: Reagents & Supplies

Reagents and Supplies

- Trizma Base   Fisher (BP152-5);
- NaCl    Fisher (BP358-212);
- Tween 20    Sigma (P-1379)
- Bovine Serum Albumin  Sigma (A-7906)
- Protein A-Sepharose  GE HealthCare (17528003)
- Protein G-Sepharose  GE HealthCare (17061803)
- 125I Insulin    Perkin Elmer (NEX420)
- NAP Column    GE HealthCare (17-0853-02 or 17-0854-02 if separating multiple labeling reactions)
- 5N HCl
- Parafilm    Sigma (P7793-1EA)
- 96-well round bottom plate  Fisher (08408220)
- 96-well filtration plates  Fisher (07200754)
- Bottle-Top 500 ml-Filter Units  Fisher (0974064A or B)
- TopSeal    Perkin-Elmer (6005185)
- Sealing Foil    USA Scientific (2923-0100)
- Microscint-20    Perkin-Elmer (6013621)
- Aluminum foil

Equipments

- TopCount β-counter (or similar) Perkin-Elmer
- 96-well Plate Shaker Wallac - Delfi
- Fume Hood
- Biological & radiation safety cabinets
- -20 °C freezers
- 4 °C refrigerator
- Pipette-Aid
- Water purification system
- Ice maker
- Radioactive contamination monitor
- Radiation sink
- pH meter
- Vortex mixer
- Steppe pipette
- Pipettes/tips
- Ice trays
- Vacuum-operated 96-well plate washer Millipore (MAVM0960R)

**Part II: Buffers**

(1) **buffer 1** (150 mM NaCl, 20 mM Tris-HCl, 1%BSA, 0.1%Sodium Azide pH 7.4)

- 30 ml 5M NaCl
- 10 ml 2M Tris-HCl pH 7.4
- 10 gm BSA
- 1 gm Sodium Azide (essential, to prevent bacterial contamination)
- 1.5 ml Tween-20
- up to 1000 ml

(2) **buffer 2** The same as buffer 1 except for 0.1%BSA

**Important Points:**
- Buffer should be filtered (0.45 micron filter) to prevent any particles blocking the membrane in bottom of the wells of the 96 well plate (which would decrease washing efficiency and increase the assay background)
- Store buffers at 4°C in a sterile bottle for up to 3 months
Part III: Incubation of Serum Samples with $^{125\text{I}}$-insulin

Each 96-well plate is sufficient for testing 24 samples in duplicate (24 duplicate with cold insulin and 24 duplicate without cold insulin. Usually, 4-8 plates can easily be run at one time (total of 96 to 192 samples). Six samples: High PC, nc, Low PC1, nc, Low PC2, nc, should be included in each assay.

1. **Spin down sera to remove fibrin clots when necessary** (otherwise these may partially block membrane in bottom of wells)

2. **Prepare the stock solution of $^{125\text{I}}$-insulin.**
   Use 1 ml of 5%BSA in PBS dissolve the powder of each 10 uCi of $^{125\text{I}}$-insulin.

3. **Calculate how much $^{125\text{I}}$-insulin and cold insulin are required.**

   6.4 ml of Washing Buffer for two plates:
   
   48 x 4.2 x 30 = 6 ml (48 samples, with 30 ul/well; in duplicate for both with and without cold insulin wells but multiply by 4.2 rather than 4 to allow for some extra)

   20,000 cpm is used for each well.

   \[
   \begin{align*}
   &3040 \text{ ul buffer 1} \\
   &160 \text{ ul } ^{125\text{I}}\text{-insulin} \\
   &3.2 \text{ ml}
   \end{align*}
   \]

   \[
   \begin{align*}
   &2784 \text{ ul buffer 1} \\
   &160 \text{ ul } ^{125\text{I}}\text{-insulin} \\
   &256 \text{ ul humulin (or Novolin)} \\
   &3.2 \text{ ml}
   \end{align*}
   \]

   **Keep the Buffer-labeled antigen mixture on ice.**

4. **Mix each serum sample with Buffer-antigen mixture in a PCR tube (or similar tube).**

   Serum: 6 ul
   
   Buffer: 30 ul

   Each sample for 2 wells.

5. **Vortex and incubate 2 hours at room temperature and overnight at 4°C.**
Part IV: Preparation of MultiScreen Filtration Plates and Protein A/G-Sepharose

(1) Coat the plate with BSA by adding 150 μl of Buffer 1 to each well.

    Incubate overnight at room temperature, after placing the plate on aluminium foil.

(3) Remove the washing buffer.

(4) The plates are now ready for running the assay, but can be stored at 4°C if necessary.

(5) Prepare Protein-A/G Sepharose:
   
   a: prepare Protein A-Sepharose
   - Use only plastic tubes because Protein-A sticks to glass
   - For each plate, take 5 ml Protein-A Sepharose in a 50 ml tube. Spin down and remove the fluid phase. Wash 2x times with buffer 1.
   - Finally add buffer 1 to give 62.5% concentration of Protein-A Sepharose by volume.

   b: prepare Protein G-Sepharose
   - Use only plastic tubes because Protein-G sticks to glass
   - For each plate, take 1 ml Protein-G Sepharose in buffer 1 in a 50 ml tube. Spin down and remove the fluid phase. Repeat once with buffer 1.
   - Finally add buffer 1 to give 40% concentration of Protein-A Sepharose by volume.

   c: mix Protein A/G Sepharose
   Mix Protein A/G Sepharose as 4:1 ratio (final concentration: 50% PA/8% PG).
Part V: Immunoprecipitation with Protein A-Sepharose

(1) Add 50 ul of Protein A/G-Sepharose mixture to each well. Use Eppendorf multi-step pipettor and re-suspend the Protein-A/G Sepharose after each row of the plate is done. (Will need 5 ml of Protein-A/G Sepharose per plate.)

(2) Add 30 ul of overnight incubate to each well (i.e., each serum will be tested in duplicate).

(3) Shake the plate on a Plate Shaker for 45 minutes at 4°C.

(4) Place the plate on Millipore plate washer device (with vacuum).

(5) Wash the plate three times in this way with 200 ul of Washing Buffer per well.

(6) Add 120 ul of Washing Buffer to each well. Shake for at least 5 minutes at 4°C.

(7) Wash the plate four times with 200 ul of washing buffer per well (change the plate direction after two times of washing at this stage).

(8) Place the plate in 37°C incubator for 15 minutes. Do not over-dry.

(9) Add 50 ul of scintillation cocktail (Microscint-20) to each well.

(10) Count on Top Count 96-well plate β counter.
Part VI: Data Analysis

(1) Delta cpm:
mean cpm of duplicate without cold insulin - mean cpm of duplicate with cold insulin

(1) CPM Index for each sample:

Sample delta cpm - NC delta cpm
-----------------------------
PC delta cpm - NC delta cpm

(2) Coefficient of Variation

For Duplicates:

(High CPM - Low CPM)/1.128
----------------------------- x 100
Mean CPM

For Triplicates:

(High CPM - Low CPM)/1.693
------------------------------- x 100
Mean CPM