RayPlex® Human Alzheimer's Disease Magnetic Array 1

Quantitative Measurement of 2 Human Proteins by Flow Cytometry

Catalog numbers:
FAHM-ALZ-1-48 (48 tests)
FAHM-ALZ -1-96 (96 tests)
FAHM-ALZ -1-192 (192 tests)
FAHM-ALZ -1-480 (480 tests)

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ISO 13485:2016

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Table of Contents

I.	Overview	2
II.	Introduction	3
III.	How it Works	4
IV.	Bead ID and Distribution	5
V.	Materials Provided	7
	Additional Materials Required	8
VI.	Assay Protocol	9
	A. Preparation of Samples	9
	B. Preparation of Reagents	10
	C. Preparation of Protein Standards	11
	D. Assay Procedure	12
	E. Flow Cytometer Set-up and Data Acquisition	14
	F. Bead Loss Prevention Check Procedure	16
	G. Data Analysis	17
VII.	Assay Information	18
VIII.	Troubleshooting Guide	21
IX.	Appendix	22
Χ.	Notes	27

I. Overview

Species	Human
Proteins Detected (2)	pTau217 and pTau181
Format	Bead-based
Detection Method	Flow Cytometry using a flow cytometer equipped with a blue (or green) laser (PE channel) and red laser (APC channel)
Minimal Sample Volume	25 μL per test after dilution. The optimal dilution mustbe determined empirically by the researcher
Number of Replicates	At least 2 replicates. Each replicate is considered a test
Reproducibility	See Section VII (page 20) for inter-CV and intra-CV percentages for each protein
Assay Duration	4 hours

II. Introduction

Alzheimer's Disease is a neurodegenerative disease affecting one's memory and cognitive function. The accumulation of beta-amyloid and phosphorylated Tau has been identified as having a strong correlation to the disease. This kit detects two forms of phosphorylated Tau: pTau217 and pTau181. Both forms are associated with the early stages of disease, but the role they play in Alzheimer's Disease progression is not fully understood. Research focused on these targets can aid in the study of the disease and patient prognosis.

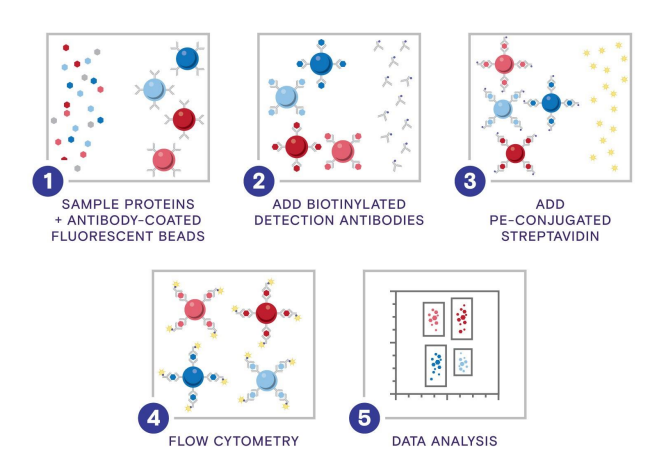
The traditional technique for detecting and quantifying proteins one at a time is the enzyme-linked immunosorbent assay (ELISA). However, using this method to detect multiple proteins – particularly for precious samples with small quantities – becomes challenging as the processing time and required sample volume increase.

RayPlex® is a multiplexed sandwich- and bead-based quantitative antibody array for the simultaneous detection of multiple proteins using small sample volumes. It couples the versatility of RayBiotech's vast antibody pair library with familiar, reliable flow cytometry methodology. Together this creates a multiplex bead-based array requiring no dedicated instrumentation, only common flow cytometers equipped with blue and red lasers.

This kit uses magnetic beads. If preferred, an automatic plate washing system with magnetic separation ability can be used for washing steps. For data in this manual, a BioTek 405 TS Washer and LifeSep™ 96F magnetic separator unit were used. Further details are included in Section VI. Assay Protocol.

III. How it Works

RayPlex® arrays are first prepared by immobilizing capture antibodies onto small beads of different sizes and different fluorochromes; there is only one target's capture antibody per bead size-fluorochrome combination. The capture antibodies bind to their specific protein targets during sample incubation, and unbound proteins are removed with washing. Biotinylated detection antibodies and PE-conjugated streptavidin molecules are added, thus enabling protein detection via the PE- streptavidin-biotin-antibody complex. Individual proteins are identified by their specific bead-fluorochrome combination, while the level of PE fluorescence reflects the amount of protein that has been captured onto the beads. The protein amount can be determined (i.e., quantified) by comparing the PE signal to a standard curve generated from purified protein standards at known concentrations.



A schematic showing how RayPlex® enables multiplex protein detection with flowcytometry.

IV. Bead ID and Distribution

1. General View of RayPlex® Multiplex Beads.

As many as two different sizes of beads are used with RayPlex® Magnetic arrays: small and large, (Figures 1 and 2). This kit uses **small** and **large** beads.

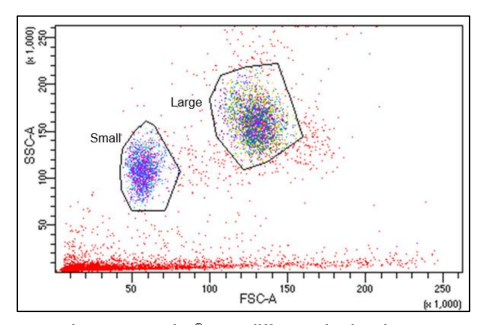


Figure 1. RayPlex® uses different size beads

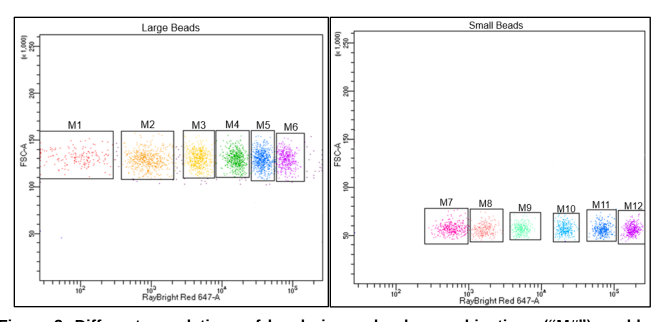
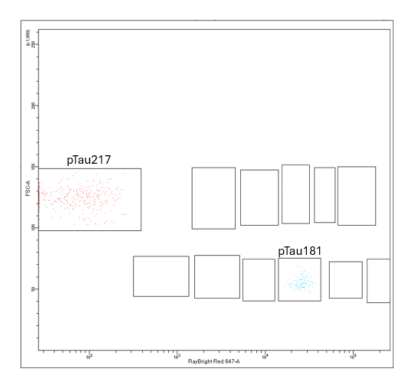


Figure 2. Different populations of bead size and color combinations ("M#") enable multiplex protein detectionwith RayPlex® arrays. Bead ID M1-6: Large Beads. Bead ID M7-12: Small Beads.

2. Target Proteins and associated Bead ID in this kit.

Bead ID	Target
M1	pTau217
M10	pTau181



V. Materials Provided

Upon receipt, the Lyophilized Protein Standard Mix (Item 5) should be stored at -20°C or below. Other components should be stored at 4°C. *Use reagents immediately after preparation*. Additional vials of the Lyophilized Protein Standard Mix (Item 5) can be purchased separately (cat # FAHM-ALZ-1-Item 5). If stored at the proper temperature, the kit will retain complete activity for at least 6 months.

Catalog #	Description	48 Tests	Per 96 Tests
FAHM-ALZ-1-Item1	RayPlex [®] Magnetic Multiplex Bead Cocktail	1.25 mL x 1 vial	1.25 mL x 2 vials
FAX-AD-1-Item2	1X Assay Diluent	25 mL x 1 vial	25 mL x 2 vials
FAX-WB-Item3	20X Wash Buffer	5 mL x 1 vial	5 mL x 2 vials
FAM-SD-1-Item4	RayBio [®] Serum Diluent	2.5 mL x 1 vial	2.5 mL x 2 vials
FAHM-ALZ-1-Item5	Lyophilized Protein Standard Mix	1	2
FAHM-ALZ-1-Item6	1X Biotinylated Detection AntibodyCocktail	1.25 mL x 1 vial	1.25 mL x 2 vials
FAX-PE-Item7	10X Streptavidin-PE	250 μL x 1 vial	250 μL x 2 vials
FAHM-ALZ-1-Item8	Flow Cytometer Setup Bead Cocktail**	200 µL x 1 vial	200 μL x 1 vial
FAX-FB-Item9	Flat-bottom 96-well Microplate	1	1
N/A	Manual	1	1

Notes:

- 1. ** Flow Cytometer Setup Bead Cocktail (Item 8) is an additional smaller aliquot of Item 1 provided to assist in the setup of the flow cytometer. It can also be used to test for bead loss during washing. A detailed procedure for bead loss prevention is outlined in Section VI.F.
- 2. RayBio® Serum Diluent (Item 4) is a pool of normal human serum from several donors with the proteins-of-interest depleted. Different lots may result in batch effects; thus, it is recommended that the researcher take this into consideration when planning their experiments. Contact info@raybiotech.com if a large lot of RayBio® Serum Diluent is required.
- 3. RayBiotech recommends using a Flat-Bottom 96-well Microplate (Item 9) in conjunction with an automated plate washing system which includes magnetic separation.

Additional Materials Required

- Orbital 96-well plate shaker (with ability to reach 1000 rpm)
- Flow cytometer with blue (or green) and red lasers capable of measuring phycoerythrin (PE) and allophycocyanin (APC) channels
- Aluminum foil or 96-well Plate Aluminum Sealers
- Deionized water
- 1.5 mL polypropylene microcentrifuge tubes or similar
- Microcentrifuge
- **Recommended**: Automated plate washing system with magnetic separation
- Optional: 96-well round bottom plate (to prepare samples)
- Optional: High Throughput Sampler (HTS) for 96-well plate reading
- Optional: Vacuum manifold compatible with a 96-well microplate if using a Filter 96-Well Microplate

VI. Assay Protocol

A. Preparation of Samples

- We recommend the following parameters for your samples: 25 µL of diluted serum, cell culture media, or other fluids. Actual sample dilutions may need to be empirically determined.
- It's recommended to use a 96-well round bottom plate to prepare samples before transferring the samples to the test plate or test tubes.
- We recommend analyzing samples at least in duplicate.

Serum Samples

- Serum samples: Dilute samples 2-fold with PBS (e.g., dilute 30 μL of serum with 30 μL of PBS), which is equal to 4-fold total dilution after mixed with beads.
- If further dilution is needed, RayBio[®] Serum Diluent (Item 4) should be used.

• Cell Culture Supernatant / Conditioned Media

- Samples: No extra dilution needed; total dilution will be equal to 2-fold after mixed with beads.
- The use of serum (e.g., fetal bovine serum; FBS) in cell culture can result in high background. To learn how to prepare samples, see our <u>Tips on Sample Preparation here:</u> https://www.raybiotech.com/tips-on-sample-preparation/.
- If cell culture supernatant from serum-containing conditioned media is tested, it is highly recommended that complete medium be used as a negative control since many types of sera contain proteins that may cross-react with the antibodies.

Note: Levels of target protein(s) may vary between different samples. Optimal dilution factors for each sample must be determined empirically by the investigator.

Note: Matrix effects are a common cause of non-linear dilution responses. This can occur when proteins or other components within the sample affect the immunoreactivity of the target molecule. These matrix components can also affect the ability of the antibody to recognize its target within the sample. Auto-antibodies, binding proteins, hemolysis, or certain disease states can contribute to this phenomenon. If matrix effects are suspected, centrifuge the sample(s), and dilute further such that matrix effects become negligible.

Note: If you experience high background or the readings exceed the detection range, further dilution of your sample is recommended.

Note: If you are using an automatic plate washing system that is not the same as defined in this manual, it is recommended to perform a bead loss prevention check as outlined in Section VI.F.

B. Preparation of Reagents

- Keep all reagents on ice.
- Protect fluorescent multiplex beads from light.
- 1. Dilute and prepare only what is needed to perform the tests for each experiment.
- 2. Vortex the RayPlex® Magnetic Multiplex Bead Cocktail (Item 1) for 30 seconds before use. Use 25 µL beads per test.
- 3. Dilute the 20X Wash Buffer (Item 3) 20-fold with de-ionized (DI) H₂O to prepare 1X Wash buffer (See Example note below).
- 4. Dilute the 10X Streptavidin-PE (Item 7) 10-fold in 1X Assay Diluent (Item 2) to prepare a 1X Streptavidin-PE working stock. Use 50 μL per test.

Example: A 20-fold dilution would be 1-part 20X Wash Buffer and 19 parts DI H₂O.

C. Preparation of Protein Standards

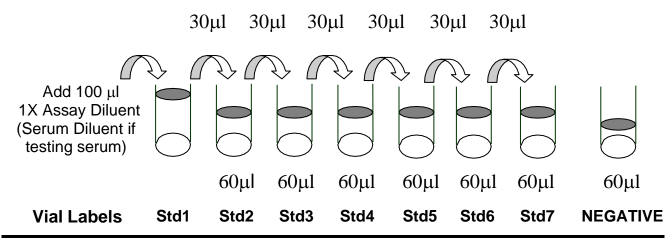
- 1. Centrifuge the Lyophilized Protein Standard Mix (Item 5) briefly (1-3 seconds) to collect contents to the bottle of the vial.
- 2. Reconstitute the Lyophilized Protein Standard Mix (Item 5):
 - For **conditioned media or non-serum fluids** add 100 μL 1X Assay Diluent (Item 2) to the tube. Dissolve the powder thoroughly and gently by pipetting up and down 5-10 times. Label this tube as Std1.
 - For **serum**, add 100 μL RayBio[®] Serum Diluent (Item 4) to the tube. Dissolve the powder thoroughly and gently by pipetting up and down 5-10 times. Label this tube as Std1.

Note: Use the Lyophilized Protein Standard Mix (Item 5) within 1 hour of preparation.

Note: When testing serum, RayBio® Serum Diluent (Item 4) should be used to reconstitute and serially dilute the standard mixture.

- 3. Obtain 6 clean 1.5 mL microcentrifuge tubes (or use a 96-well round-bottom plate). Label the tubes/wells as Std2 to Std7. Add 60 µL 1X Assay Diluent (Item 2), or RayBio[®] Serum Diluent (Item 4) as applicable to each tube.
- 4. Pipette 30 μ L Std1 into tube Std2 and mix gently. Perform 5 more serial dilutions by adding 30 μ L of Std2 to tube Std3, mix, and so on. (See also image below)

Prepare serial dilutions of protein standards



5. Add 60 µL 1X Assay Diluent (Item 2) or RayBio® Serum Diluent (Item 4) to another tube labeled as NEGATIVE. Do not add standard or samples to the NEGATIVE tube. This tube will be used as the negative control.

Note: The exact concentrations of each standard protein for the Std1 to Std7 serial dilutions in this kit can be found in Section VII.

D. Assay Procedure

1. Prepare a Flat-Bottom 96-well plate (Item 10), V-bottom plate or 1.2 μm filter-plate and mark positions for the standards and samples. Duplicate tests are recommended for all standards and samples.

Note: RayBiotech recommends using a Flat-Bottom 96-well plate (Item 10) in conjunction with a magnetic plate washing system. Data in this manual were obtained using a LifeSep™ magnetic separator unit and BioTek 405 TS Washer. Please note that if alternative systems are used, unforeseen bead loss may occur. If desired, investigators may follow the procedure outlined in Section VI.F to check for bead loss prior to beginning the assay.

Note: A Filter 96-well can be used for the washing steps if desired. The filter plate requires a vacuum manifold compatible with a standard 96-well microplate. Alternatively, a V-Bottom 96-well microplate can be used along with a centrifuge.

- 2. Add 25 µL of RayPlex® Multiplex Bead Cocktail (Item 1) to each well that willcontain the NEGATIVE, Standard, or Sample.
- 3. Add 25 μL of the NEGATIVE, Standard, or Sample to the appropriate well. The total volume in each well is now 50 μL (serum sample is diluted 4-fold totally with beads in 50 μL reaction system, cell culture supernatant / conditioned media is diluted 2-fold totally with beads). Place plate on an orbital plate shaker. Shake at 1000 rpm at room temperature for 2 hours.

Note: This step can also be performed overnight at 4°C, which may increase the signal-to-noise ratio, particularly for proteins with low concentrations. The optimal incubation length (i.e. room temperature for 2 hours or overnight at 4°C) for the experiment must be determined empirically by the investigator.

- 4. Wash the beads using 1X Wash Buffer (diluted Item 3; See Section B).
 - Automated Washing System: Place the Flat-Bottom 96-well Microplate (Item 9) on the magnetic separator unit for 5 minutes to allow the magnetic beads to migrate to the bottom. Place the Flat-Bottom 96-well Microplate and magnetic separator unit on the platform of an automated plate washer and perform one wash cycle.

One wash cycle should include two rounds of dispensing 200 μ L of 1X Wash Buffer and aspiration. It is recommended to adjust the settings such that the first aspiration leaves approximately 2 mm of Wash Buffer, and the final aspiration removes all liquid. It is pivotal that all liquid is removed at the end of washing. Remove the plate from the magnetic separator unit once washing is complete. Exact washing procedures may vary depending on the automated system used.

- V-bottom 96-well Microplate: Add 200 µL 1X Wash Buffer to each well. Spin the V-bottom microplate down at 1000 g for 5 minutes at room temperature and remove the supernatant using a multichannel pipet. Repeat once more.
- Filter 96-well Microplate: Add 200 µL 1X Wash Buffer to each well.
 Turn on the vacuum, open the valve, and place the filter plate on the
 vacuum. Do not allow pressure to exceed 10 inches Hg (254 mm Hg).
 Close the valve and remove the plate when the wells have drained.
 Repeat once more.
- 5. Add 25 µL of 1X Biotinylated Detection Antibody Cocktail (Item 6) to each well. Resuspend the beads by gently pipetting and incubate on an orbital shaker at 1000 rpm at room temperature for 1 hour.
- 6. Wash plate as outlined in Step 4.
- 7. Add 50 µL of 1X Streptavidin-PE (*diluted* Item 7; See Section B) to each well, incubate on an orbital shaker at 1000 rpm at room temperature for 30 minutes.
- 8. Wash plate as outlined in Step 4.
- 9. Resuspend in 150 μL of 1X Wash Buffer (diluted Item 3; See Section B).
- 10. Prepare samples for analysis on a flow cytometer:
 - Automated High Throughput Sampler (HTS): if using a flat-bottom plate, directly read samples from the plate. If using a filter-plate, transfer samples to a V-Bottom plate.
 - Manual reading: transfer samples to compatible tubes.
- 11. Analyze samples on a flow cytometer (see Section E).

E. Flow Cytometer Set-up and Data Acquisition

NOTE: The RayPlex[®] Human Alzheimer's Disease Magnetic Array 1 requires a flow cytometer that is equipped with blue (or green) and red lasers capable of detecting fluorescence in the PE and APC channels, respectively.

NOTE: Perform standard QC and optimization steps to prepare the flow cytometer, then set up the flow cytometer as you would for normal use. Manual adjustment of the PE and APC compensation may be necessary if the APC fluorescence is too intense, which sometimes causes a smiling effect (curved population grouping).

NOTE: These instructions are general for all RayPlex[®] assays. The target count, bead size, and populations may differ from the assay above. Please see your assay's specific manual Section IV and VII for the bead size and populations present in your array.

 Depending on the brand of the flow cytometer, you may need to start the acquisition software and run Quality Control beads before proceeding further.

Note: The Quality Controls beads are fluorescent particles that are used for flow cytometer setup, e.g., CS&T beads for a BD flow cytometer.

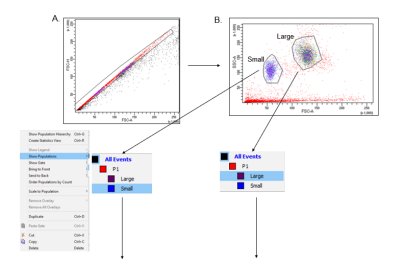
- 2. Start a new experiment with PE and APC channels.
- 3. Resuspend 100 μ L of the provided Flow Cytometer Setup Bead Cocktail (Item 8) in 200 μ L 1X Wash Buffer (*diluted* Item 3) and run them on the flow cytometer.
- 4. Create an FSC-H/FSC-A daughter population for "Single beads" selection to remove and limit doublets or higher complexes (Figure 3A).
- 5. Create a new dot plot from the "Single beads" parent gate, adjust voltage for FSC (forward scatter, linear mode) and SSC (side scatter, linear mode) so that the small, medium, and/or large bead populations are clearly defined. Then create a small and/or large beads gate (Figure 3B).
- 6. Create a new dot plot from the small and/or large beads' parentgate. Gate populations for all bead populations for the assay based on SSC(linear mode) and APC (log mode). Adjust APC channel PMT voltage so that all populations are evenly distributed and well separated throughout in a

prominent area (Figure 3C).

7. Create a dot plot of APC versus PE (using log scale for both) from the small, medium, and/or large beads' parent populations. Run a small amount of beads from the NEGATIVE control. Adjust the PE voltage so that the PE mean fluorescent intensity (MFI) is around 10^{2~3} for each bead population.

Note: To keep the testing consistent for each assay, we recommend use of Rainbow Particles, Rainbow Beads, or similar to standardize the assay if they are run prior to every analysis. The optimized MFI for both the APC and PE channels for the Rainbow Particles allow the experiment to be consistent from time to time and machine to machine.

- 8. Set the number of bead events to be acquired to at least 200 per target population for best results. More beads will improve data accuracy.
- 9. Following setup, run the standards and samples (Figure 3D).
- 10. Create a statistical view to show the PE MFI for each population (Figure 3E). The MFI of all analytes and samples can be exported in Excel format by "Batch Analysis" of a whole specimen in FACSDiva. Researchers can also export raw data as "FCS" files and analyze in FlowJo software.



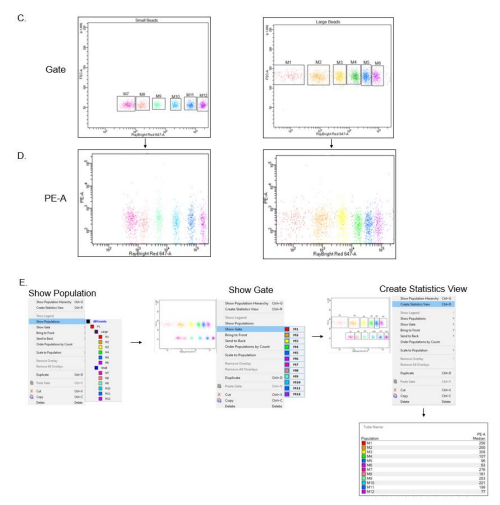


Figure 3. Representative Images of Gated Flow Cytometry Scatter Plots Using RayPlex® Arrays with small and large beads. (A) Single beads are (B) separated by size, then by (C) dye concentration. (D) Beads are read based on PE-A concentration. (E) All raw PE-A Median values based on Gate of (C).

Note: The above figures are for demonstration purposes only. This kit may not contain all bead populations shown.

F. Bead Loss Prevention Check

NOTE: Due to differences in automated plate washing systems, it is recommended to perform a bead loss prevention check prior to starting the assay to prevent loss of beads if an automated washing machine is being used.

- 1. Resuspend the contents of the Flow Cytometer Setup Bead Cocktail (Item 8) and add 25 µL to 2 wells of a 96-well plate
- 2. Add 25 µL of Wash Buffer
- 3. Perform one wash cycle

- 4. Repeat steps 2 and 3 two more times for a total of 3 wash cycles
- 5. Resuspend the beads in 150 μL of Wash Buffer
- 6. Run the samples either by transferring to tubes and running in "Tube Mode" or directly from the plate with HTS mode.

If the bead count is less than 100 bead events for any bead IDs, it is recommended to reconsider the washing procedure used for the assay. Bead events should ideally be more than 200 for each bead ID. Using a filter-plate for washing will prevent bead loss most effectively.

G. Data Analysis

- 1. The MFI of all analytes and samples can be exported in Excel format by "Batch Analysis" of a whole specimen in FACSDiva (See Appendix: RayPlex® Data Analysis Using BD FACS Diva Batch Analysis).
- 2. MFI data of each analyte for all samples can also be acquired by exporting the entire dataset as an FCS file. Analyze data in FlowJo or equivalent software and export the PE MFI into an Excel (or similar) format.

VII. Assay Information

1. 8-Point Standard Protein Concentrations (pg/mL)

The concentrations of target proteins corresponding to standard (Std) 1-7 serial dilutions are displayed below in pg/mL. These concentrations are used to produce a standard curve.

Target	STD1	STD2	STD3	STD4	STD5	STD6	STD7	CTRL
pTau217	50000.0	16666.7	5555.6	1851.9	617.3	205.8	68.6	0.0
ptau181	50000.0	16666.7	5555.6	1851.9	617.3	205.8	68.6	0.0

2. Representation of Standard Curve MFI Values

The values in this table are an example of MFI values that can be used to generate a standard curve. These values are for demonstration purposes only. A standard curve must be run with each assay since the MFI values will vary from experiment to experiment.

	pTau217	pTau181
Std 1 (pg/mL)	50000.0	50000.0
Standard 1 MFI	30126.7	15014.7
Standard 2 MFI	14184.3	7582.3
Standard 3 MFI	6869.0	3519.0
Standard 4 MFI	2319.3	1190.3
Standard 5 MFI	331.0	213.7
Standard 6 MFI	331.0	221.7
Standard 7 MFI	151.0	95.7
Negative MFI	65.3	65.0

3. Lower Limit of Detection (LOD) of Target Proteins

The LOD is the minimum detectable concentration (i.e., sensitivity) of the assay. It was determined by spiking each purified protein standard into RayBio[®] Serum Diluent (Item 4).

Target	LOD (pg/mL)
pTau217	14.67
pTau181	29.78

Note: The LOD may vary from run-to-run and batch-to-batch.

4. Cross Reactivity

No detectable or significant cross reactivity was detected between antibodies or targets.

5. Percent (%) Recovery of Spiked Standards

For spike recovery of standards, target proteins with concentrations corresponding to standard 2 (Std2) were spiked into RayBio® Serum Diluent (Item 4) across 9 replicates. The spiked proteins were assayed, and the measured concentrations were compared to the expected concentrations.

Ray Bío® Human Serum Diluent

Target	Percent Recovery
pTau217	99.5%
pTau181	101.8%

6. Linearity of Dilution

To test the linearity of dilution, samples (n = 12) were serially diluted in RayBio[®] Serum Diluent (Item 4) 1:3, 1:9, and 1:21. The measured concentrations of serially diluted samples were compared to that of the undiluted samples and averaged. See below for data.

RayBio® Human Serum Diluent

Target	Linearity of Dilution
pTau217	101.6%
pTau181	96.4%

7. Intra- and Inter-Assay Precision

1) Intra-Assay Precision

Spiked standard protein in RayBio[®] Serum Diluent (Item 4) was analyzed in triplicate. This was performed at two different concentrations per standard protein (i.e., Sample #2, Sample #3). The intra-assay precision, or coefficient of variation (CV), of the samples is provided below.

Target	Sample #	Mean (pg/mL)	Standard Deviation	CV %
pT21/217	2	16603.1	407.5	2.5%
pTau217	3	5611.2	623.9	11.1%
nTau101	2	16988.9	604.4	3.6%
pTau181	3	5071.7	774.4	15.3%

2) Inter-Assay Precision

Spiked standard protein in RayBio[®] Serum Diluent (Item 4) was analyzed in triplicate across three independent assays. This was performed at two different concentrations per standard protein (i.e., Sample #2, Sample #3). The inter-assay precision in (CV) of the samples is provided below.

Target	Sample #	Average Mean (pg/mL)	Standard Deviation	CV %
pT2u217	2	16477.2	178.0	1.1%
pTau217	3	5970.8	347.1	5.8%
nTou101	2	16309.5	960.8	5.9%
pTau181	3	6108.6	1103.0	18.1%

Note: All data in Section VII were obtained using a flat-bottom plate. Five-parameter logistic curve fitting was used for generation of standard curve and calculation of concentrations.

VIII. Troubleshooting Guide

Issue	Possible Causes	Recommendations
		Increase the length of the sample- bead incubation to overnight at 4°C (see Section D, step 3).
	Low protein content in sample	Decrease sample dilution.
	Low protein content in sample	Note that even if the incubation is increased or the sample dilution is decreased, it is not guaranteed that the protein will be detected.
	Detection antibody over diluted	Increase detection antibody concentration.
Wook Signal	diated	Optimize sample dilution.
Weak Signal	Sample matrix effect	For serum samples, prepare the standards in the provided "RayBio® Serum Diluent" (Item 4).
	Campio matrix oncot	Adjust the PE MFI using a small amount of beads from the negative control to around 10 ^{2~3} .
	Improper storage of kit	Store kit at suggested temperature and use within 6 months of receipt.
	Reagent evaporation	Cover the incubation plate with adhesive film during incubation.
	Cross-contamination from neighboring wells	Avoid overflowing wells during wash steps.
Poor standard curve	Lyophilized Protein Standard Mix (Item 5) not properly prepared	Serially dilute the Lyophilized Protein Standard Mix (Item 5) according to manual.
	Lyophilized Protein Standard Mix (Item 5) degraded or not property diluted	Reconstitute the Lyophilized Protein Standard Mix (Item 5) on ice <i>before</i> making serial dilutions.
	Improper flow cytometer setup and optimization	Run Setup Bead Cocktail (Item 8) before assay. Make sure top standard signal is not out of the linear range.
High background	PE voltage PMT voltage is not set up correctly	Set PE voltage using a small amount of beads from the negative control so that the PE MFI is around $10^{2\sim3}$.
	Insufficient washing	Increase wash time. Use more wash buffer.

IX. Appendix

Data Analysis using Batch Analysis

Batch analysis allows exporting PE MFI data of each bead population directly from FACS Diva. Before batch analysis, it's important to create statistics view from the dot plot of gated beads populations and choose "PE-A Median" to be shown (Figure 4). Batch analysis can be done in either manual tube mode or HTS mode. Both methods for batch analysis are described below (Figure 5 and 6).

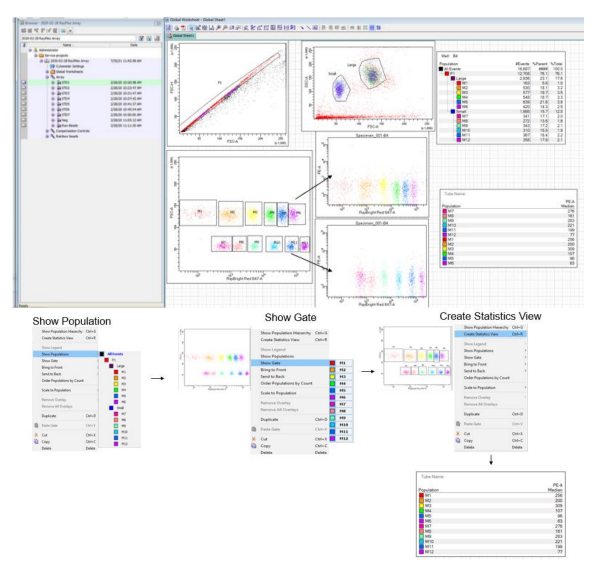


Figure 4. Screenshot of Creating Statistics View from the dot plot of gated beads populations. This is representative data only.

- 1. Create a dot plot by SSC-A (linear) and APC-A (log) to show all beads populations and make a gate for each population (see figure 4, "Gated Dot Plot"). Right click on the dot plot to "Create Statistics View", then edit the statistics view so that "PE-A Median" of gated beads populations are shown (figure 4, "Statistics View of Gated Plot").
- 2. Right click on the open folder containing your experiment data and select 'Batch Analysis.'
- 3. When the pop-up appears, select the desired destination folder, and then hit start.

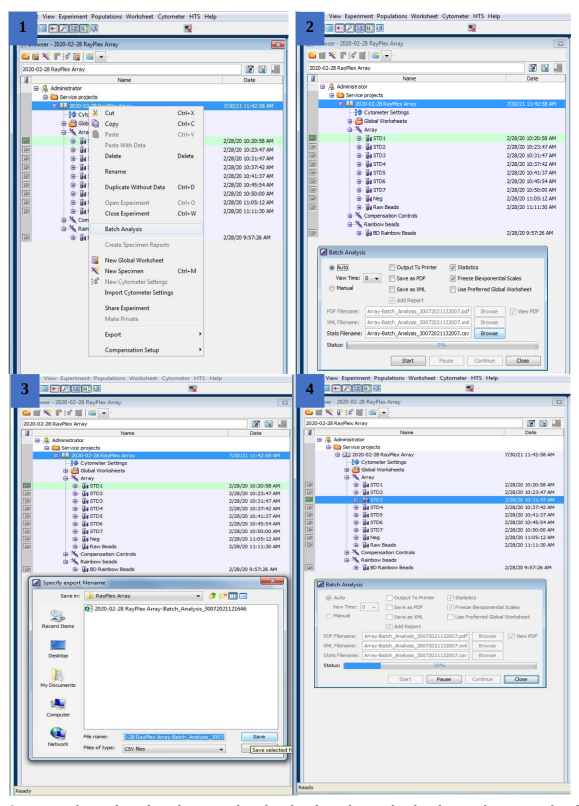


Figure 5. Screen shot showing how to begin the batch analysis through manual tube mode.

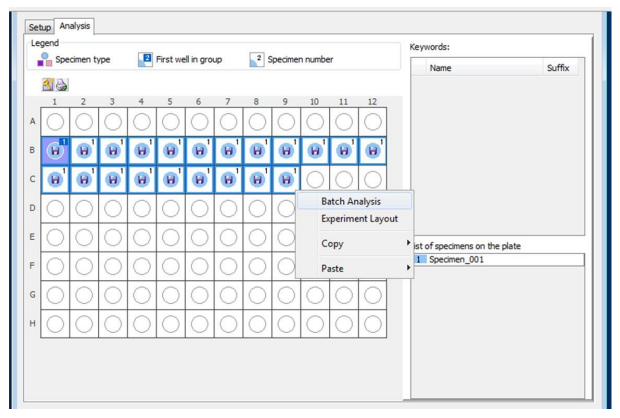


Figure 6. Screen shot showing how to begin HTS batch analysis.

- 4. When the HTS is done running samples, click on the 'Analysis' tab and highlight all of the samples. Right click and select 'Batch Analysis.' After selecting Batch Analysis through the HTS mode, the steps from Manual mode are the same.
- 5. Open the CSV file in the destination folder and select 'Save As' to save the data as an Excel workbook (Figure 7).

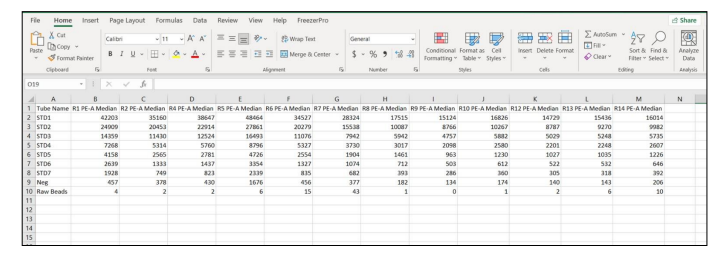


Figure 7. Representative raw data obtained from batch analysis. Rows = data from samples. Leftmost column = standard or sample name. (The names of beads are variable for different assays)

X. Notes

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