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ProcartaPlex[™] Human Kidney Toxicity Panel 1 11-Plex USER GUIDE

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A.0	11 January 2023	New document for ProcartaPlex Human Kidney Toxicity Panel 1 11-Plex User Guide.

The information in this guide is subject to change without notice.

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Product Information



WARNING! Read the Safety Data Sheets (SDSs) and follow the handling instructions. Wear appropriate protective eyewear, clothing, and gloves. Safety Data Sheets (SDSs) are available from thermofisher.com/support.

Product Description

The ProcartaPlex™ Human Kidney Toxicity Panel 1 11-Plex has been optimized for detection of multiple analytes from urine, serum, plasma, and cell culture supernatants.

The panel is provided in a ready-to-use format with individual vials of 1X capture and detection reagents that require less pipetting and experimental setup. These reagents are not combinable with simplexes or other panels.

ProcartaPlex™ preconfigured panels are extensively tested for analyte combinability, interference and cross-reactivity to provide the highest level of validation and precision. All ProcartaPlex™ panels are supplied with the necessary reagents to perform the assay.

Table 1 Analytes

Calbindin	Clusterin (APO-J)	GSTA1 (Glutathione S-transferase A1)
IL-18	IP-10 (CXCL 10)	KIM-1 (TIM-1/HAVCR1)
MCP-1 (CCL2)	Osteoactivin (GPNMB)	RBP4 (Retinol-binding protein 4)
Renin	VEGF-A	

Contents and storage

Upon receipt, store the kit at 2°C to 8°C. When stored as indicated, all reagents are stable until the expiration date.

Contents	Amount
Standard Mix 15 (lyophilized)	2 each
Standard Mix 16 (lyophilized)	2 each
Biotinylated Detection Antibody Mix (1X)	1 x 3.5 mL
Capture Bead Mix (1X)	1 x 5 mL
Streptavidin-PE (SA-PE) (1X)	1 x 5 mL
Wash Buffer (10X)	1 x 25 mL
Reading Buffer (1X)	1 x 40 mL
Universal Assay Buffer, UAB (1X)	1 x 10 mL
8-Tube Strip	2 each
Flat Bottom 96-well Plate	1 each
Microplate Lid	1 each
Plate Seals	8 each

Retain the lot-specific Certificate of Analysis that contains the product expiration date. The Certificate of Analysis also contains important information such as bead number, analyte names, and highest standard concentration required for the assay setup on the xMAP™ instrument.

Required materials not supplied

Catalog numbers that appear as links open the web pages for those products.

- xMAP[™]-based instrument
- Hand-Held Magnetic Plate Washer (Cat. No. EPX-55555-000)
- Deionized water
- Vortex mixer (e.g., Cat. No. 88882010)
- Microcentrifuge
- Adjustable single and multichannel pipettes with disposable tips and low volume reservoirs (e.g., Cat. No. 95128093)
- Beakers, flasks, and cylinders necessary for preparation of reagents
- Orbital microplate shaker with at least 1.5 mm or 0.059 inch orbit diameter capable of maintaining a speed of 600 ± 50 rpm (e.g., Cat. No. 88882006)

Note: Use of rockers or large orbit shakers can cause adverse results.

Procedural guidelines

- Thoroughly read this user guide and the certificate of analysis before using the kit.
- All chemicals should be considered potentially hazardous.
- To avoid cross-contamination, do not invert the assay plate during the assay or allow contents from one well to mix with another well.
- Use a multichannel pipette and reagent reservoirs whenever possible to achieve optimal assay precision.
- This protocol was developed using the Hand-Held Magnetic Plate Washer (Cat. No. EPX-55555-000). Other washers should be validated by the end user.
- Ensure that the xMAP™ instrument has been properly calibrated and set up before preparing and running the assay.

Workflow

Assay protocol

Prepare antigen standard

Add capture beads

- 1. Vortex capture beads for 30 sec. Add 50 µL of the capture beads to each well.
- 2. Remove the liquid, then wash the plate.

Add samples and standards

1. Add the following according to sample type:

For serum, urine, and plasma samples: Add 25 µL of Universal Assay buffer, then add 25 µL of standards or samples. For background wells, add 50 µL of 1X UAB.

For cell culture supernatant samples: Add 50 $\,\mu$ L of standards or samples. For background wells, add 50 $\,\mu$ L of cell culture medium.

Note: Sample predilution of Clusterin (APO-J) in plasma/serum of 1:10,000 is recommended. Sample predilution of RBP4 in plasma/serum samples of 1 : 100 is recommended.

- 2. Seal the plate and incubate with shaking at room temp for 2 hr.
- 3. Wash plate two times.

Add detection antibody

- 1. Add 25 µL of Detection Antibody Mix (1X).
- 2. Seal the plate and incubate with shaking at room temp for 30 min.
- 3. Wash plate two times.

Add Streptavidin-PE

- 1. Add 50 µL of Streptavidin-PE.
- 2. Seal the plate and incubate with shaking at room temp for 30 min.
- 3. Wash plate two times.

Resuspend beads

- 1. Add 120 µL of Reading Buffer.
- 2. Seal the plate and shake at room temp for 5 min.

Acquire data on xMAP™ system

Methods



Prepare the samples

Dilution of serum/plasma samples

Clusterin (APO-J) and RBP4 typically have high plasma/serum concentrations. We recommend that you dilute samples in Universal Assay Buffer (1X) to ensure they fall within the range of the assay.

Dilution factor for Clusterin (APO-J) in serum/plasma samples is 1:10,000.

Dilution factor for RBP4 in serum/plasma samples is 1:100.

Dilute each sample according to the following scheme:

Tube	Sample volume	Universal Assay Buffer (1X) volume
Dilution 1	10 μL	90 μL
Dilution 2	10 μL of Dilution 1	90 μL
Dilution 3	10 μL of Dilution 2	90 μL
Dilution 4	10 μL of Dilution 3	90 μL

Prepare the reagents

Before starting with the assay protocol, define the plate map. Mark the standard, sample, and background wells to determine the number of wells used (see Appendix A, "Recommended plate layout").

Prepare 1X Wash Buffer

Bring the Wash Buffer Concentrate (10X) to room temperature and vortex for 15 seconds. Mix 20 mL of the Wash Buffer Concentrate (10X) with 180 mL ddH_2O . Mix gently to avoid foaming. Wash Buffer (1X) can be stored at 2–8°C for up to 6 months.

Note: Additional Wash Buffer Concentrate (200 mL, Cat. No. EPX-66666-001) can be purchased separately for automated plate washers.

Prepare Standard Mix

This kit is supplied with two lyophilized Standard Mixes for generation of standard curves. Two vials of each Standard Mix are provided to permit the user to run the assay twice if running a partial plate. For

serum, plasma, and urine use 1X UAB as diluent. For experiments measuring cell culture supernatant samples, use fresh cell culture medium as the diluent.

Note: Change pipette tips after each dilution step and avoid air bubbles.

- 1. Centrifuge the standard mix stock vial at 2,000 x g for 10 seconds.
- 2. Add 125 µL of diluent to each stock vial.
- 3. Vortex the vial at high speed for 30 seconds and centrifuge at 2,000 x g for 10 seconds to collect contents at the bottom of the vial.
- 4. Incubate on ice for 10 minutes to ensure complete reconstitution.
- 5. Combine the contents of each of the two Standard Mix vials into one vial.

Prepare 4-fold serial dilution

- 1. Label the tubes in the 8-Tube Strip: Std1, Std2, Std3, Std4, Std5, Std6 and Std7.
- 2. Add 200 μ L of the reconstituted standard mix into Std1 tube.
- 3. Add 150 µL of diluent into Std2-Std7 tubes.
- 4. Transfer 50 μL from Std1 tube into Std2 tube.
- 5. Mix by pipetting up and down 10 times.
- 6. Transfer 50 µL of the mixed standards from Std2 tube into Std3 tube using new pipette tip.
- 7. Mix by pipetting up and down 10 times.
- 8. Repeat steps 4–7 for tubes Std4–Std7, changing pipette tips between dilution steps.
- 9. Add 150 µL of diluent to the last tube of the 8-Tube Strip to serve as a background. See Figure 1

Chapter 2 Methods Assay protocol

10. Keep tubes on ice until ready to use.

Note: Use the reconstituted standard immediately. The reconstituted standard cannot be stored. Discard unopened standard vials if the entire plate was used in a single experiment.

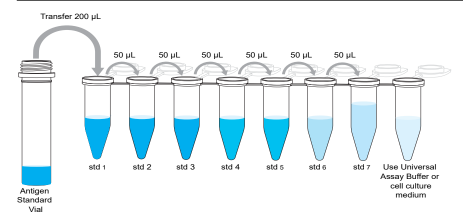


Figure 1 4-fold serial dilution

Assay protocol

- 1. Add Capture Bead Mix to the plate.
 - a. Vortex the 1X Capture Bead Mix vial for 30 seconds at high speed.
 - b. Using a multichannel pipette, add 50 µL of the Capture Bead Mix to each well of the plate.
- 2. Wash beads using a Hand-Held Magnetic Plate Washer.

Note: To avoid loss of beads, secure the plate using the clamps on both sides of the Hand-Held Magnetic Plate Washer during this procedure.

Note: This protocol was developed using the Hand-Held Magnetic Plate Washer (Cat. No. EPX-55555-000). Other washers should be validated by the end user.

- a. Place the plate on the Hand-Held Magnetic Plate Washer and wait 2 minutes to allow the beads to settle on the bottom of each well.
- **b.** Remove the liquid by quickly inverting the washer/plate assembly over a sink or waste container.
- **c.** Gently blot the inverted washer/plate assembly onto several layers of paper towels or absorbent surface to remove any residual liquid.
- d. Add 150 μL of 1X Wash Buffer into each well and wait 30 seconds.
- e. Remove the liquid by quickly inverting the washer/plate assembly over a sink or waste container.

- **f.** Gently blot the inverted washer/plate assembly onto several layers of paper towels or absorbent surface to remove any residual liquid.
- g. Remove the plate from the magnet and proceed to step 3.
- 3. Add samples and standards to the plate.
 - a. For serum, plasma, and urine samples: Add 25 μL of Universal Assay Buffer, then add 25 μL of standards or samples. For background wells, add 50 μL of 1X UAB. For background wells, add 50 μL of 1X UAB. For cell culture supernatant samples: Add 50 μL of standards or samples. For background wells, add 50 μL of cell culture medium.
 - **b.** Seal the plate using one of the provided Plate Seals and cover with the provided Microplate Lid. Shake at 600 rpm for 2 hours at room temperature.
- 4. Remove and discard the Plate Seal. Wash the plate following the steps below.
 - a. Place the plate on the Hand-Held Magnetic Plate Washer and wait 2 minutes to allow particles to settle on the bottom of each well.
 - **b.** Remove the liquid by quickly inverting the washer/plate assembly over a sink or waste container.
 - c. Gently blot the inverted washer/plate assembly onto several layers of paper towels or absorbent surface to remove any residual liquid.
 - d. Add 150 µL of 1X Wash Buffer into each well and wait 30 seconds.
 - e. Remove the liquid by quickly inverting the washer/plate assembly over a sink or waste container.
 - **f.** Gently blot the inverted washer/plate assembly onto several layers of paper towels or absorbent surface to remove any residual liquid.
 - g. Repeat steps 4d-4f once more for a total of two washes.
 - **h.** Remove the plate from the magnet and proceed to the next step.
- 5. Add Biotinylated detection Antibody Mix to the plate.
 - a. Using a multichannel pipette, add 25 μ L of the detection antibody solution to each well of the plate. Gently tap the plate to evenly distribute the solution in the wells.

Note: A narrow trough reservoir for multichannel pipetting is recommended to be used to prevent volume loss.

- **b.** Seal the plate using a new Plate Seal and cover with the provided Microplate Lid. Shake at 600 rpm for 30 minutes at room temperature.
- 6. Wash the plate following step 4.
- 7. Add Streptavidin-PE (SA-PE) to the plate.
 - a. Add 50 µL of SA-PE solution to each well.

Chapter 2 Methods Instrument settings

- **b.** Seal the plate using new Plate Seal and cover with the provided Microplate Lid. Shake at 600 rpm for 30 minutes at room temperature.
- 8. Wash the plate following step 4.
- 9. Prepare the plate for analysis on an xMAP™ instrument.
 - a. Add 120 µL of reading buffer into each well.
 - b. Seal the plate using new Plate Seal and cover with the provided Microplate Lid. Shake at 600 rpm for 5 minutes at room temperature.
- 10. Remove the Plate Seal and run the plate on an xMAP™ instrument.

Note: Alternatively, the sealed plate can be stored overnight at 2-8°C before running on an xMAP[™] instrument. Shake the plate at 600 rpm for 5 minutes at room temperature then remove the plate seal and run the plate.

Instrument settings

Follow the recommended guidelines and procedures for calibration and verification of the instrument. Laser-based systems require 30 minutes to warm up prior to use.

Instrument	Acquisition volume	Timeout (optional)	Bead type	DD gate	Reporter gain	Min. bead count
MAGPIX™	50 μL ^[1]	N/A	N/A	N/A	Standard PMT	50
INTELLIFLEX™	30 µL	40 sec	MagPlex™	7,000–17,000	Standard PMT	50
FLEXMAP 3D™	50 μL	60 sec	MagPlex™	7,500–25,000	Standard PMT	50
Luminex™ 100/200™						
Bio-Rad™ Bio-Plex™	50 μL	60 sec	MagPlex™	5,000–25,000	Standard PMT	50

^[1] MAGPIX volume can be changed during the run to optimize bead count.

Note: To assure a good bead count, the probe height must be adjusted to the plate provided in the kit. We recommend using two 5.08 mm spacer disks to adjust the sample probe height for Mylar-bottom plates.

Analyze results

The concentration of the samples can be calculated by plotting the expected concentration of the standards against the NET MFI generated by each standard. For Bio-Plex[™] Manager, plot standard

concentrations against FI-Bkgd. A 4PL or 5PL algorithm is recommended for the best curve fit. Analyze the assayed samples according to the operation manual for the Luminex[™] or Bio-Plex[™] instrument.

We offer a free and robust analysis software package for data analysis. To analyze the data, follow the instructions below or contact our technical support.

1. Export the run data in .csv format and navigate to the ProcartaPlex™ Analysis App on Thermo Fisher Connect: https://apps.thermofisher.com/apps/procartaplex

Note: Before exporting .csv raw data from Bio-Plex[™] Manager, please make sure to set 'Analytes Labels' under 'Document Export Properties' to 'Name (Region)'. The .csv raw data exported as Report Type 'xPONENT' from INTELLIFLEXTM instruments are supported.

2. Upload the .csv files to the ProcartaPlex™ Analysis App to analyze the run data. The intuitive software features 4PL/5PL curve fit optimization, group-wise statistical and heat map analysis. Users can export detailed reports including images for presentations and publications.

IMPORTANT! For ProcartaPlex[™] getting started guides, technical literature, protocol support tools, and common troubleshooting questions visit thermofisher.com/procartaplexsupport For more complete troubleshooting questions and answers, visit our FAQ database at thermofisher.com/procartaplexfaqs



Recommended plate layout

Standards			Samples								
1	1	1	1	9	9	17	17	25	25	33	33
2	2	2	2	10	10	18	18	26	26	34	34
3	3	3	3	11	11	19	19	27	27	35	35
4	4	4	4	12	12	20	20	28	28	36	36
5	5	5	5	13	13	21	21	29	29	37	37
6	6	6	6	14	14	22	22	30	30	38	38
7	7	7	7	15	15	23	23	31	31	39	39
Bkgd ^[1]	Bkgd	8	8	16	16	24	24	32	32	40	40

^[1] Background

	1	2	3	4	5	6	7	8	9	10	11	12
Α												
В												
С												
D												
Е												
F												
G												
Н												



Documentation and support

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 - User guides, manuals, and protocols
 - Certificates of Analysis
 - Safety Data Sheets (SDSs; also known as MSDSs)

Note: For SDSs for reagents and chemicals from other manufacturers, contact the manufacturer.

Limited product warranty

Life Technologies Corporation and/or its affiliate(s) warrant their products as set forth in the Life Technologies' General Terms and Conditions of Sale at www.thermofisher.com/us/en/home/global/terms-and-conditions.html. If you have any questions, please contact Life Technologies at www.thermofisher.com/support.

