RayBio[®] Rat Acute Kidney Injury Antibody Array 1 (G-Series)

Patent Pending Technology

User Manual (Revised June 6, 2014)

RayBio[®] Rat Acute Kidney Injury Antibody Array G-Series Cat# AAR-AKI-G1-4

RayBio[®] Rat Acute Kidney Injury Antibody Array G-Series Cat# AAR-AKI-G1-8

RayBio[®] Rat Cytokine Antibody Array G-Series Testing Service Cat# AAR-SERV-G

Please read manual carefully before starting experiment



We provide you with excellent Protein Array systems and services

Tel: (Toll Free) 1-888-494-8555 or +1-770-729-2992; Fax: +1-770-206-2393; Website: <u>www.raybiotech.com</u> Email: <u>info@raybiotech.com</u> RayBiotech, Inc., the Protein Array Pioneer Company, strives to research and develop new products to meet demands of the biomedical community. RayBiotech's patent-pending technology allows detection of up to 1,000 cytokines, chemokines and other proteins in a single experiment. Our format is simple, sensitive, reliable, reproducible and cost-effective.

Our product offerings include:

- 1. Protein (antigen) Arrays
- 2. RayBio[®] Cytokine Antibody Arrays
- C Series (Membrane, chemiluminescence detection)
- G-Series (Glass chip, fluorescence detection)
- 3. Pathway- and disease-focused antibody arrays
 - Angiogenesis Antibody Arrays
 - Apoptosis Antibody Arrays
 - Atherosclerosis Antibody Arrays
 - Chemokine Antibody Arrays
 - Growth Factor Antibody Arrays
 - Inflammation Antibody Arrays
 - MMP Antibody Arrays
 - Obesity Antibody Arrays
- 4. Quantibody® Multiplex ELISA Arrays
- 5. RayBio[®] L-Series Biotin Label-based Antibody Arrays
- 6. RayBio[®] E-Series Competition-based Antibody Arrays
- 7. RayBio[®] Phosphorylation Antibody Arrays
 o Receptor Tyrosine Kinases
 - EGFR and ErbB family (site-specific phosphorylation)
- 8. Over 1,300 different ELISA kits
- 9. EIA (Competitive ELISA) kits
- 10. Cell-based Phosphorylation Assay
- 11. Over 20,000 different antibodies
- 12. Recombinant proteins
- 13. Peptide
- 14. Recombinant antibodies



Protocol for RayBio[®] Rat Acute Kidney Injury Antibody Array G-Series 1

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RayBio[®] Cytokine Antibody Arrays are patent-pending technology. RayBio[®] is the trademark of RayBiotech, Inc.

I. Introduction

New techniques such as cDNA microarrays have enabled us to analyze global gene expression¹⁻³. However, almost all cell functions are executed by proteins, which cannot be studied simply through DNA and RNA techniques. Experimental analysis clearly shows disparity can exist between the relative expression levels of mRNA and their corresponding proteins⁴. Therefore, analysis of the proteomic profile is critical.

RayBiotech, The Protein Array Pioneer Company, introduced the first protein arrays to the market in 2001 and continues to lead in the development of innovative protein array technologies, such as the RayBio Human Acute Kidney Injury Antibody Array.

Acute kidney injury is a common complication among ambulatory and hospitalized patients. It is a rapidly progressive illness that independently predicts excess morbidity and mortality. It is critical to early detect acute kidney injury and distinguish it from prerenal azotemia and chronic kidney disease at the time of patient presentation to rapidly manage associated illness. However, serum creatinine, a standard marker of kidney function, does not distinguish acute kidney injury from prerenal azotemia⁵ or chronic kidney disease. In addition, the initial measurement of serum creatinine cannot reflect the extent of injury because its accumulation always lags behind the insult⁶.

The kidney is the primary organ responsible for the excretion of medications and their biotransformation products from the body. probing pharmacokineticlargely being used in Rat is pharmacodynamic (PK-PD) relationships for medications; in addition, it has been demonstrated to be a useful model for evaluating mechanisms of kidney toxicity. In recent years, numerous molecules have been described and investigated as candidate biomarkers of kidney injury. The United States Food and Drug Administration (FDA) has taken a active role in developing a process for qualification of biomarkers^{7, 8} that would potentially improve the drug development and regulatory review

process. In the gentamicin-induced rat model of acute kidney injury, based on histopathology, necrosis, or apoptosis scoring, kidney injury molecule-1 (KIM-1) was the best biomarker of overall renal injury⁹.

Traditionally, urine proteins or cytokines are detected by using ELISA. However, RayBio® Rat Acute Kidney Injury Antibody Array C Series can detect 7 protein biomarkers simultaneously with small amount of sample. It is a great tool in the acute kidney injury research areas and drug discovery area to hasten drug development.

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- 8. Goodsaid FM, Frueh FW, Mattes W. Strategic paths for biomarker qualification. Toxicology 2008; 245: 219-223.

 Rodney L. Rouse, Jun Zhang, Sharron R. Stewart, Barry A, Rosenzweig, Parvaneh Espandiari, and Nakissa K. Sadrieh3. Comparative profile of commercially available urinary biomarkers in preclinical drug-induced kidney injury and recovery in rats. Kidney International (2011) 79, 1186-1197

II. Product Information

A. Storage Recommendations:

For best results, we recommend storing the entire kit at -20°C or -80°C upon arrival and using the kit within 6 months of receipt. RayBiotech warranties this product for 6 months if stored in this manner.

Once thawed, store glass chips and 1X Blocking Buffer at -20°C or -80°C and all other component at 4°C. After thawing, the entire kit should be used within 3 months. RayBio[®] Antibody Array kits are robust and will retain full activity even if accidentally stored at room temperature (RT) for up to 24 hours.

B. Materials Provided

Item	Description	AAR-AKI- G1-4	AAR-AKI- G1-8
AAR-AKI-G1	RayBio [®] Rat Acute Kidney Injury Microarray Glass Chip*	1 chip with 4 Sub-arrays*	1 chip with 8 Sub-arrays*
0103002 -R- AKI1	Biotin-Conjugated Anti- Cytokines	1 ea	2 ea
0103004-H	1,500X HiLyte Plus™ 532 Streptavidin-Fluor†	1 ea	1 ea
0103004-B	1X Blocking Buffer	10 ml	20 ml
0103004-W‡	20X Wash Buffer I ‡	30 ml	30 ml
0103004-W‡	20X Wash Buffer II ‡	30 ml	30 ml
0103004-L	2X Cell Lysis Buffer (optional)	10 ml	10 ml

Other Kit Components:

Manual, Adhesive Plastic Strips, 30 ml Centrifuge Tube

* Kit contains 1 pre-assembled glass chip with either 4 or 8 printed sub-arrays per chip (in sealed plastic envelope)

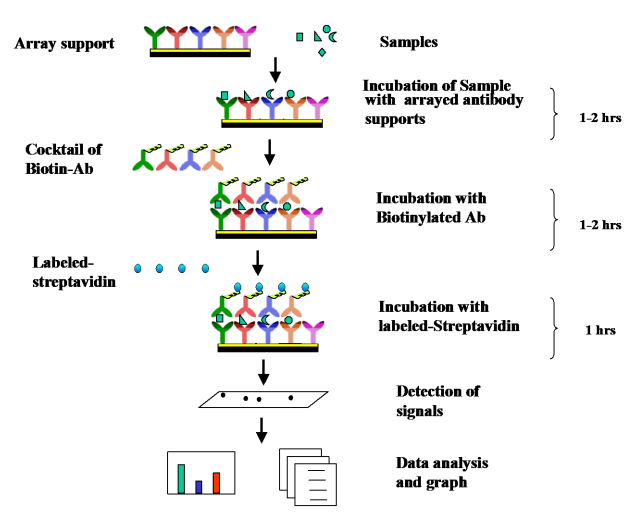
[NOTE: In some cases, 2 chips x 4 sub-arrays/chip may be substituted in kits containing 8 sub-arrays]

- † This fluor is patent-pending technology from Anaspec, Inc.
- ‡ Wash Buffers are sold as sets

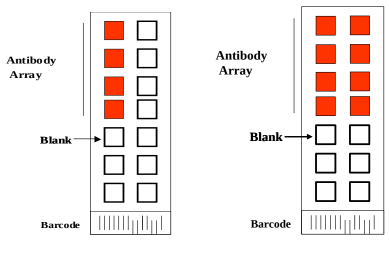
C. Additional Materials Required

- Small plastic boxes or containers
- Pipettors, pipette tips and other common lab consumables
- Orbital shaker or oscillating rocker
- Aluminum foil
- Gene microarray scanner or similar laser fluorescence scanner (see pages 9 & 15)

D. How It Works



E. <u>RayBio® G Series Glass Chip Layout</u>



4 arrays in one glass chip

8 arrays in one glass chip

III. Helpful Tips and General Considerations

A. Preparation and Storage of Samples

- 1. <u>General Considerations:</u>
 - Freeze samples as soon as possible after collection.
 - Avoid multiple freeze-thaw cycles. If possible, sub-aliquot your samples prior to initial storage.
 - Spin samples hard (5-10 minutes at 10K to 15K RPM) immediately prior to incubation of samples with array.
 - Optimal sample concentrations may need to be determined empirically based on the signal intensities of spots and background signals obtained.
 - Most samples will not need to be concentrated. If concentration is required, we recommend using a spin-column concentrator with a chilled centrifuge.
- 2. Recommended Sample Volumes and Dilution Factors

NOTE: All sample dilutions should be made using 1X Blocking Buffer. For all sample types, final sample volume = 50-100 μl per sub-array

- Cell Cultured Media: Neat (no dilution needed)
- Serum & Plasma: 5-fold to 10-fold dilution
- Most other Body Fluids: Neat or 2-fold to 5-fold dilution
- Cell and Tissue Lysates: Minimum 5-fold to 10 fold to equal concentrations of total protein in each lysate sample.
- You <u>must</u> determine the total protein concentration of each lysate/homogenate. We recommend using the BCA method (available from Pierce); it is insensitive to detergents commonly found in lysis buffers.
- Minimum Recommended Dilution of Lysates (prior to sample incubation): 5-fold to 10 fold with 1X Blocking Buffer. <u>Dilute all lysate samples to the same final concentration of</u> total lysate protein in 1X Blocking Buffer to 100 μl final volume.
- To start, we recommend using 10-100 μg of total protein in 100 μl of 1X Blocking Buffer (final volume) per sub-array.

- Optimal amounts of total lysate protein may range from 5-500 µg per sub-array. Based upon background and spots intensities, you may increase or decrease the amount of protein used in subsequent experiments.
- Other Liquid Sample Types: Most often Neat or 2-fold to 5fold. However, optimal dilutions should be determined empirically.

3. <u>Sample Preparation</u>

For tips on sample preparation, please visit our Website: <u>http://www.raybiotech.com/Tech-Support/SampleTips.pdf</u>

B. Handling Glass Chips

- Do not remove glass chip from assembly until Step 16.
- Hold the slides by edges only; do not touch the surface.
- Handle all buffers and slides with powder-free gloves.
- Dry glass chip completely before proceeding to Step 3.
- Handle and dry glass chip in clean environment.
- Avoid breaking glass chip when removing the chamber assembly.

C. Incubations and Washes

- Cover incubation chamber with adhesive film (included in kit) to prevent evaporation, particularly during incubation or wash steps >2 h or with liquid volumes <100 µl per well.
- Perform all incubation and wash steps under gentle rotation or rocking motion (~0.5 to 1 cycle/s).
- Wash steps in Wash Buffer II and all incubation steps may be performed overnight at 4°C.
 - Overnight sample incubations are the most effective at increasing sample spot intensities.
- Avoid cross-contamination of samples to neighboring wells
- To remove Wash Buffers and other reagents from chamber wells, you may invert the Glass Chip Assembly to decant, and aspirate the remaining liquid.

- In Wash Steps 6, 12 and 15, you may gently flush wells several times using a wash bottle filled with Wash Buffer I.
- D. Scanning and Data Extraction Tips:

For tips on scanning and data extraction, please visit our Website: http://www.raybiotech.com/Tech-Support/ScanningTips.pdf

For a list of recommended scanners, please visit our Website: <u>http://www.raybiotech.com/files/Tech-</u> Support/Laser Scanners for Glass Slide Arrays.pdf

See also page 18 of this manual.

IV. Protocol

A. Preparation and Storage of Reagents

NOTE: During this protocol, prepare reagents immediately prior to use and keep working dilutions of all reagents on ice at all times.

- 1. <u>Blocking Buffer</u> is supplied at 1X concentration. No dilution is required.
- 2. <u>Wash Buffers I and II</u> are supplied at 20X concentration.
 - a). For each glass chip (4 or 8 sub-arrays/chip), dilute 6 ml of 20X concentrate with deionized H₂0 to a final volume of 120 ml each of Wash Buffer I & Wash Buffer II.
 - b). Wash buffer reagents at working dilution (1X) can be stored at 4°C for up to 1 month. Stock solutions at 20X can be stored 4°C for up to 3 months.
- 3. <u>Biotin-conjugated Anti-Cytokines</u> are supplied at high concentration in a small liquid bead (typically ~2-5 μl).
 - a). Spin down the tube prior to reconstitution, as the concentrated liquid bead may have moved to the top of the tube during handling.

- b). Prepare stock reagent by adding 300 μl 1X Blocking Buffer to Biotin-Conjugated Anti-Cytokines. Mix well.
- c). 1X Biotin-Conjugated Anti-Cytokines may be stored for 2-3 days at 4°C.
- 4. <u>Streptavidin-Fluor</u> is supplied at 1500x concentration.
 - a). Mix the tube containing 1500X Streptavidin-Fluor well before use, as precipitants may form during storage.
 - b). Add 100 µl of 1X Blocking Buffer to tube containing 1500X Streptavidin-Fluor. Mix well.
 - c). Quantitatively transfer all of Streptavidin-Fluor reagent from the original tube to a larger one, and dilute with 1X Blocking Buffer to a final volume of 1500 μl (ie, 1.5 ml).
 - d). Wrap tube containing Streptavidin-Fluor with aluminum foil.
 - e). This working dilution can be stored for 3-5 days at 4°C.

B. Blocking and Incubations

NOTE: Please carefully read Section III of this manual before proceeding

NOTE: Prepare all reagents immediately prior to use as described above (Section IV.A) and before proceeding.

 Remove the package containing the glass chip assembly from the freezer. Place unopened package on the benchtop and allow the glass chip assembly to equilibrate to room temperature (RT), approx. 15 min. Open package, remove the glass chip assembly and place in laminar flow hood to dry for 1-2 hours.

NOTE: Be sure glass chip is completely dry before proceeding.

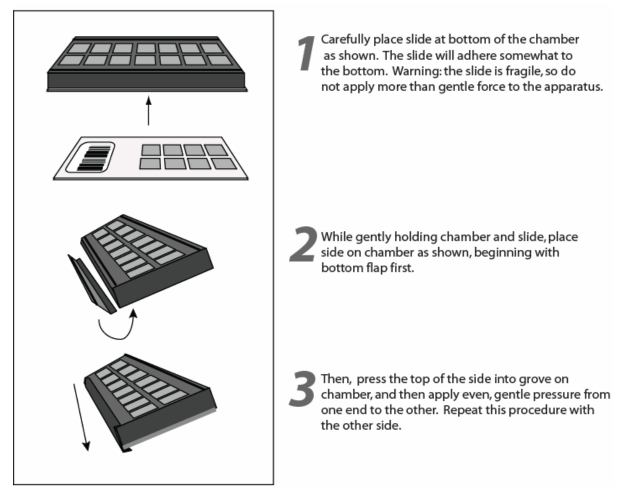
2) If necessary, assemble the glass chip into incubation chamber and frame as shown on page 11-12. (Note: if you slide is already assembled, you can proceed directly to Step 3). 3) Add 100 μ l 1 X Blocking Buffer into each well and incubate at RT for 30 min to block slides.

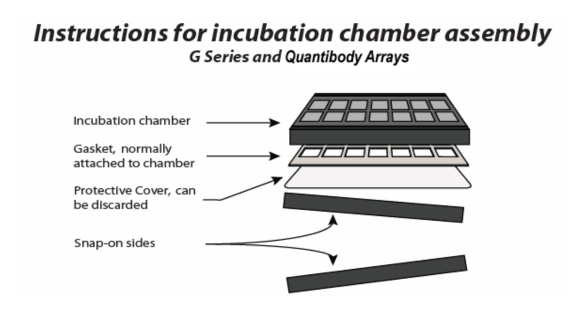
NOTE: Only add reagents or samples to wells printed with antibodies (see diagram on page 5)

4) Decant Blocking Buffer; then aspirate remaining liquid from each well.

NOTE: To aspirate liquid samples or reagents from wells, gently place the pipette tip only in the corners of the well. <u>Do</u> <u>not scrape the pipette tip across the surface of the chip.</u>

5) Add 50 to 100 µl of each sample to each sub-array. Cover the incubation chamber with Adhesive film (included in kit). Incubate arrays with sample at RT for 2 hours. Dilute sample using 1X Blocking Buffer if necessary.





6) Remove adhesive film, and carefully aspirate samples from sub-arrays, touching only the corners with your pipette tip.

NOTE: Try to prevent solution from flowing into neighboring wells.

- 7) Wash 3 x 2 min with 150 µl 1X Wash Buffer I at RT. Be sure to completely remove sample and Wash Buffer each time and use fresh buffer for each wash. Decant final wash solution before proceeding to next step.
- 8) Obtain a clean container (eg, pipette tip box or slide staining jar) and place glass chip assembly into the container. Add enough 1X Wash Buffer I to submerge the entire glass chip with frame intact (approx. 30-50 ml) and remove all bubbles in wells. Wash 10 min at RT with gentle rocking or shaking.
- Remove assembled glass chip from container and invert it to decant liquid. Decant buffer from container and replenish with 1X Wash Buffer I. Submerge the entire glass chip assembly and wash 10 min at RT with gentle rocking or shaking.
- 10) Remove assembled glass chip from container and invert it to decant liquid. Decant buffer from container and repeat Steps 8 & 9 with Wash Buffer II.

- 11) Remove assembled glass chip from container and invert it to decant liquid, then carefully aspirate wash buffer from wells, touching only the corners with your pipette tip.
- 12) Add 70 µl of 1X Biotin-conjugated Anti-Cytokines to each subarray. Cover incubation chamber with Adhesive film (included in kit). Incubate at RT for 2 hours with gentle rocking or shaking.
- 13) Carefully aspirate the entire Biotin-conjugated Anti-Cytokine reagent. Wash as described in Step 7 above, first with Wash Buffer I then with Wash Buffer II, making sure to completely remove buffer between washes and after final wash.
- 14) Add 70 μl of 1X Streptavidin-Fluor to each sub-array. Cover the incubation chamber with Adhesive film (included in kit), then cover entire assembly with aluminum foil to avoid exposure to light or incubate in dark room. Incubate at RT for 2 hours with gentle rocking or shaking.
- 15) Remove aluminum foil and adhesive film. Carefully aspirate Biotin-conjugated Anti-Cytokine reagent. Wash as described in Step 7 above, first with Wash Buffer I then with Wash Buffer II, making sure to completely remove buffer between washes and after final wash.
- 16) Remove the glass chip from the frame assembly. Place the whole chip in 30 ml centrifuge tube provided, or slide staining jar. Add enough Wash Buffer I to cover the whole slide (about 20 ml) and gently rock or shake at RT for 10 min.
- 17) Decant buffer and repeat wash as described in Step 16 (1 x 10 min with Wash Buffer I).
- 18) Decant buffer and repeat wash as described in Step 16, but this time using Wash Buffer II for only 2-3 minutes.

- 19) Decant buffer, remove the glass chip from the tube, then gently rinse the slide with de-ionized H₂O using a plastic wash bottle.
- 20) Remove water droplets by applying suction gently with a pipette tip.
- C. Obtaining Fluorescent Signal Intensities:
 - 21) Allow glass chip to dry in a laminar flow hood for 20 minutes or until slide is completely dry. Place chip under an aluminum foil tent to protect it from light. Make sure the slides are absolutely dry before scanning or storage.
 - 22) You may proceed immediately to scanning (Step 23), or you may scan at a later time. You may store the slides at RT indefinitely, provided they are protected from strong light.

Note: Unlike most Cy3 fluors, the HiLyte Plus[™] Fluor 555 used in this kit is very stable at RT and resistant to photobleaching on completed glass chips. However, please protect glass chips from strong light and temperatures above RT.

23) Scan the glass chip with a laser scanner (such as Innopsys' InnoScan®) using cy3 or "green" channel (excitation frequency = 532 nm). For tips on scanning, visit our Website: http://www.raybiotech.com/Tech-Support/ScanningTips.pdf

NOTE: If you do not have a laser scanner, for a nominal fee you can send your slide to us for scanning and data extraction using Innopsys' InnoScan, and we will return the results to you. Using using alternate protocols, RayBio® G-Series arrays are also compatible with Li-Cor's Odyssey and other microarray scanners.

V. Interpretation of Results:

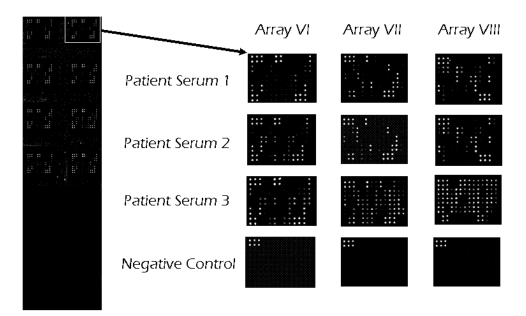
A. Explanation of Controls Spots

Positive Controls (POS1, POS2, POS3) are equal amounts of biotinylated IgGs printed directly onto the array. All other variables being equal, the Positive Control intensities will be the same for each sub-array This allows for normalization based upon the relative fluorescence signal responses to a known control, much as "housekeeping" genes or proteins are used to normalize results in PCR or Western blots, respectively.

<u>Negative Control (NEG)</u> spots are a protein-containing buffer (used to dilute antibodies printed on the array). Their signal intensities represent non-specific binding of Biotin-conjugated anti-Cytokines and/or Streptavidin-Fluor. Negative control signal intensities are usually very close to background signals in each sub-array.

B. <u>Typical results from RayBio[®] G-Series Antibody Arrays</u>

The following figure shows typical results obtained using RayBio® Antibody Array G-Series Arrays. The images were captured using a GenePix 4000B scanner.



In this example, sera from several patients were incubated with Human Cytokine Arrays 6, 7 & 8, (sold together as Human Cytokine Array G-Series 2000, AAH-CYT-G2000-4 or AAH-CTY-G2000-8) and processed using this standard protocol.

The 6 strong signals of the Positive Control spots in the upperleft corner are useful for proper orientation of the array image.

If scanned using optimal scan settings, 3 distinct Positive Control signal intensities will be seen: POS1>POS2>POS3. If all of these signals are of similar intensity, try increasing or decreasing laser power and/or signal gain settings.

Once you have obtained fluorescence intensity data, you should subtract the background and normalize to the Positive Control signals before proceeding to analysis.

C. <u>Background Subtraction:</u>

Most laser fluorescence scanner software have an option to automatically measure the local background around each spot.

As with spot signal intensities, we recommend using MEDIAN background signals. If your resulting fluorescence signal intensity reports do not include these values (eg, a column labeled as "MED532-B532"), you may need to subtract the background manually or change the default settings on your scanner's data report menu.

D. Normalization of Array Data:

To normalize signal intensity data, one sub-array is defined as "reference" to which the other arrays are normalized. This choice can be arbitrary. For example, in our Analysis Tool Software, the array represented by data entered in the left-most column each worksheet is the default "reference array."

You can calculate the normalized values as follows:

X(Ny) = X(y) * P1/P(y)

Where:

P1 = mean signal intensity of POS spots on reference array P(y) = mean signal intensity of POS spots on Array "y" X(y) = mean signal intensity for spot "X" on Array "y" X(Ny)= normalized signal intensity for spot "X" on Array "y"

The RayBio[®] Analysis Tool software is available for use with data obtained using RayBio[®] G-Series Arrays. You can copy and paste your signal intensity data (with and without background) into the Analysis Tool, and it will automatically normalize signal intensities to the Positive Controls.

To order the Analysis Tool, please contact us at +1-770-729-2992 or <u>info@raybiotech.com</u> for more information.

E. <u>Threshold of significant difference in expression</u>:

After subtracting background signals and normalization to Positive controls, comparison of signal intensities for antigenspecific antibody spots between and among array images can be used to determine relative differences in expression levels of each analyte (ie, protein detected) between samples or groups.

Any \geq 1.5-fold increase or \leq 0.65-fold decrease in signal intensity for a single analyte between samples or groups may be considered a measurable and significant difference in expression, provided that both sets of signals are well above background (Mean background + 2 standard deviations, accuracy \approx 95%).

NOTE: In the absence of an external standard curve for each analyte, there is no means of assessing absolute or relative concentrations of different analytes in the same sample using immunoassays. If you wish to obtain quantitative data (ie, concentrations of the various analytes in your samples), try using our Quantibody® Multiplex ELISA arrays instead.

Data Extraction Tips:

- Ignore any comet tails
- Define the area for signal capture for all spots as 110-120 micron diameter, using the same area for every spot.
- Use median signal value, not the total or the mean
- Use local background correction (also median value).
- Exclude obvious outlier data in its calculations.
- Scan all slides at same PMT

VI. <u>RayBio[®] Rat Acute Kidney Injury Antibody Array G Series Map:</u>

	А	В	С	D	E	F	G	Н
1	POS	POS	NEG	NEG	Cystatin C	FABP1	KIM-1	MCP-1
2	POS	POS	NEG	NEG	Cystatin C	FABP1	KIM-1	MCP-1
3	NGAL	TIMP-1	VEGF	NEG	NEG	NEG	NEG	POS
4	NGAL	TIMP-1	VEGF	NEG	NEG	NEG	NEG	POS

Detects 7 cytokines in one experiment

Abbreviations:

POS = Positive Control, NEG = Negative Control, L-FABP = Liver Fatty-Acid Binding Protein, KIM-1 = Kidney Injury Molecule-1, NGAL= Neutrophil Gelatinase-Associated Lipocalin (Lipocalin 2). All others use standard abbreviations.

VII. Troubleshooting guide

Problem	Cause	Recommendation		
No signal for any spots, including Positive Controls	Global detection failure	Adjust scanner settings or re- assemble chip into holder, wash slide 2 x 5 min with 150 μl Wash Buffer II and repeat Steps 12-21.		
Similar signal intensities for POS1/2/3	Improper laser power and/or PMT setting	Repeat scan using higher and/or lower laser power or PMT settings		
	Incomplete washes	Carefully follow wash protocols, and/or increase wash times		
High background signals	Sample concentration is too high	Repeat using lower sample concentration		
	Fluor and/or Anti- Cytokines are too concentrated	Review protocol for dilution of reagents		
	Bubbles present on chip during incubations	Be sure to completely remove all bubbles from chip surface		
Uneven	Evaporation during incubation steps	Cover chamber assembly during washes and incubations		
background and/or missing spots	Pooling/precipitation of sample or reagent; Incomplete washes.	Cover chamber assembly and use a rocker or shaker during washes and incubations; carefully follow wash protocols.		
	Sample is too concentrated	Repeat experiment using more dilute sample		
Randomly scattered high- intensity spots	Dust or other particulates	Dry slides in laminar flow hood and/or use clean containers and powder-free gloves.		

	Sample is too dilute	Repeat experiment using higher sample concentration
	Improper dilution of Anti-Cytokines or Streptavidin- Fluor	Re-assemble chip into holder, wash 2 x 5 min with 150 µl Wash Buffer II and repeat Steps 12-21. Spin down reagents before diluting and mix well.
Weak or no signals		Rescan at higher laser power or signal gain setting
antigen-specific pots + Low Background		Repeat using higher sample concentration and/or incubate wi sample O/N at 4°C
	Other Tips	Increase concentration of and/or length of incubation with Biotin-conjugated Anti-Cytokine (+ add'l large volume wash following Biotin-Ab incubation
		Review proper storage conditions for kit components

III. <u>Selected References Citing RayBio[®] Human G-Series Arrays</u>

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4-1BB	CNTF	GITR	IL-18 ΒΡα	ΜΙΡ-1δ	SAA
ACE-2	Cripto	GITR Ligand	IL-18 Rβ	MIP-3α	sgp130
Acrp30	CRP	GM-CSF	IL-1ra	ΜΙΡ-3β	Shh N
Activin A	CTACK	GRO	IL-2	MMP-1	Siglec-5
Adiposin	CXCL16	GROα	IL-2 Rβ	MMP-10	Siglec-9
Adipsin	DAN	GH	IL-2 Ry	MMP-13	Stree-9
AgRP	Decorin	HB-EGF	IL-2 Ra	MMP-13	sTNF RI
ALCAM	Deconn Dkk-1	HCC-4	IL-2 Ra	MMP-3	sTNF RII
α-Fetoprotein	Dkk-3	hCG (intact)	IL-21R IL-22	MMP-7	TACE
Amphiregulin	Dkk-4	HGF	IL-22 IL-28A	MMP-8	TARC
Angiogenin	DPPIV	HVEM	IL-20A	MMP-9	TECK
Angiopoietin-1	DR6	I-309	IL-3	MPIF-1	TGFα
Angiopoletin-2	Dtk	ICAM-1	IL-3	MSPα	TGFβ1
Angiostatin	E-Cadherin	ICAM-1	IL-31 IL-4	NAP-2	TGFβ2
ANGPTL4	EDA-A2	ICAM-3	IL-4 IL-5	NCAM-1	TGFβ2
ANGP 124 Axl	EGF	IFNy	IL-5 IL-5 Rα	NGF R	TPO
B7-1	EGFR	IGF-1 SR	IL-5 Ru IL-6		
BCAM	EG-VEGF	IGFBG-1	IL-6 sR	Nidogen-1 NrCAM	Thyroglobulin Tie-1
BCMA	EG-VEGF ENA-78	IGFBP-2	IL-0 SR IL-7	NRG1-β1	Tie-2
BDNF		IGFBP-2	IL-7 IL-8	NT-3	TIM-1
β2Μ	Endoglin Eotaxin	IGFBP-4	IL-8 IL-9	NT-3 NT-4	TIMP-1
β IG-H3	Eotaxin-2	IGFBP-6	Insulin	Oncostatin M	TIMP-2
bFGF	Eotaxin-2 Eotaxin-3	IGF-I	IP-10		TIMP-4
BLC	Ep CAM	IGF-I SR	I-TAC	Osteopontin OPG	TNFα
BMP-4	ErbB2	IGF-II	LAP	PAI-I	ΤΝΕά
BMP-5	ErbB2 ErbB3	IL-1α		PARC	TNFRSF21
BMP-6	EPO R	IL-1β	Leptin	PDGF Rα	TNFRSF6
BMP-7	E-Selectin	IL-1 R II	Leptin R LIF	PDGF Rβ	TRAIL R2
β-NGF	Fas	IL-1 R4/ST2		PDGF RP PDGF-AA	TRAIL R2
BTC	Fas Ligand	IL-1 RI	LIMPII	PDGF-AA PDGF-AB	TRAIL R3
CA125	For RIIB/C	IL-1 sRI	LIMPI L-Selectin	PDGF-AB PDGF-BB	
CA125 CA15-3	Ferritin	IL-10	LH	PECAM-1	Trappin-2 TREM-1
CA19-9	FGF-4	IL-10 Rα	Lymphotactin	PIGF	TSH
CA IS	FGF-6	IL-10 Rβ	LYVE-1	PF4	TSLP
Cardiotrophin-1	FGF-6	IL-10 Kp	Marapsin	Procalcitonin	Ubiquitin
Cathepsin S	FGF-7	IL-12	MCP-1	Prolactin	uPAR
CCL14a	FGF-9	IL-12 p40	MCP-2	PSA-free	VCAM-1
CCL21	Fit-3 Ligand	IL-12 p70	MCP-3	PSA-total	VE-Cadherin
CCL-28	FLRG	IL-13	MCP-4	RAGE	VEGF
CD14	Follistatin	IL-13 Rα-2	M-CSF	RANK	VEGF R2
CD23	Fractalkine	IL-13 RI	M-CSF R	RANTES	VEGF R3
CD30	FSH	IL-15	MDC	Resistin	VEGF-C
CD40	Furin	IL-16	MICA	S-100b	VEGF-D
CD40 Ligand	Galectin-7	IL-17	MICA	SAA	XEDAR
CD40 Ligand CD80	GCP-2	IL-17B	MIF	SCF	
CEA	G-CSF	IL-17C	MIG	SCF R	
CEACAM-1	GDF-15	IL-176	MIP-1α	SDF-1	
CK b 8-1	GDNF	IL-17R	MIP-1β	SDF-1β	
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