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Datasheet for ABIN625566 Human Cytokine Array C7

1 Image

73 Publications



Overview

Quantity:	2 samples								
Reactivity:	Human								
Method Type:	Sandwich ELISA								
Application:	Antibody Array (AA)								
Product Details									
Purpose:	C-Series Human Cytokine Antibody Array 7 Kit. Detects 60 Human Cytokines. Suitable for all liquid sample types.								
Brand:	RayBio®								
Sample Type:	Serum, Plasma, Cell Culture Supernatant, Cell Lysate, Tissue Lysate								
Analytical Method:	Semi-Quantitative								
Detection Method:	Chemiluminescent								
Specificity:	Adiponectin (ACRP30), AgRP, Angiopoietin-2, Amphiregulin, Axl, bFGF, beta-NGF, Betacellulin (BTC), CCL28 (MEC), CTACK (CCL27), Dtk, EGFR, ENA-78 (CXCL5), Fas (TNFRSF6/Apo-1), FGF- 4, FGF-9, GCSF, GITR Ligand (TNFSF18), GITR (TNFRSF18), GRO alpha/beta/gamma, GRO alpha (CXCL1), HCC-4 (CCL16), HGF, ICAM-1 (CD54), ICAM-3 (CD50), IGFBP-3, IGFBP-6, IGF-1 R, IL-1 R4 (ST2), IL-1 R1, IL-11, IL-12 p40, IL-12 p70, IL-17A, IL-2 R alpha, IL-6 R, IL-8 (CXCL8), I-TAC (CXCL11), Lymphotactin (XCL1), MIF, MIP-1 alpha (CCL3), MIP-1 beta (CCL4), MIP-3 beta (CCL19), MSP alpha/beta, NT-4, Osteoprotegerin (TNFRSF11B), Oncostatin M, PLGF, gp130, TNF RII (TNFRSF1B), TNF RI (TNFRSF1A), TECK (CCL25), TIMP-1, TIMP-2, Thrombopoietin (TPO), TRAIL R3 (TNFRSF10C), TRAIL R4 (TNFRSF10D), uPAR, VEGF-A, VEGF-D								

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Product Details

Characteristics:	 Easy to use No specialized equipment needed Compatible with nearly any liquid sample Proven technology (many publications) Highly sensitive (pg/mL) Sandwich ELISA specificity Higher density than ELISA, Western blot or bead-based multiplex
Components:	Antibody Array Membranes Biotinylated Detection Antibody Cocktail Blocking Buffer Wash Buffers 1 and 2 Cell & Tissue Lysis Buffer Detection Buffers C and D Plastic Incubation Tray Protease Inhibitor Cocktail (in select kits)
Material not included:	Pipettors, pipet tips and other common lab consumables Orbital shaker or oscillating rocker Tissue Paper, blotting paper or chromatography paper Adhesive tape or Saran Wrap Distilled or de-ionized water A chemiluminescent blot documentation system (such as UVP's ChemiDoc-It® or EpiChem II Benchtop Darkroom), X-ray Film and a suitable film processor, or another chemiluminescent detection system.

Target Details

Background:	Cytokines play an important role in innate immunity, apoptosis, angiogenesis, cell growth and
	differentiation. They are involved in interactions between different cell types, cellular responses
	to environmental conditions, and maintenance of homeostasis. In addition, cytokines are also
	involved in most disease processes, including cancer and cardiac diseases.

Application Details

Application Notes:	Perform ALL incubation and wash steps under gentle rotation or rocking motion (~0.5 to 1 $$
	cycle/sec) using an orbital shaker or oscillating rocker to ensure complete and even
	reagent/sample coverage. Rocking/rotating too vigorously may cause foaming or bubbles to
	appear on the membrane surface which, should be avoided. All washes and incubations should

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	be performed in the Incubation Tray (ITEM 10) provided in the kit. Cover the Incubation Tray									
	with the lid provided during all incubation steps to avoid evaporation and outside debris contamination. Ensure the membranes are completely covered with sufficient sample or									
	reagent volume during each incubation. Avoid forceful pipetting directly onto the membrane, instead, gently pipette samples and reagents into a corner of each well. Aspirate samples and									
	reagents completely after each step by suctioning off excess liquid with a pipette. Tilting the									
	tray so the liquid moves to a corner and then pipetting is an effective method. Optional									
	overnight incubations may be performed for the following step to increase overall spot signal									
	intensities:									
	- Sample Incubation									
	- Biotinylated Antibody Cocktail Incubation									
	- HRP-Streptavidin Incubation									
Comment:	The C-Series arrays feature chemiluminescent signal detection. The antibodies are spotted on									
	nitrocellulose membrane solid supports and are handled in a very similar manner to Western									
	blots.									
	All C-Series arrays work on the sandwich ELISA principle, utilizing a matched pair of antibodies:									
	an immobilized capture antibody and a corresponding biotinylated detection antibody.									
Sample Volume:	1 mL									
Plate:	Membrane									
Protocol:	1. Block membranes									
	2. Incubate with Sample									
	3. Incubate with Biotinylated Detection Antibody Cocktail									
	4. Incubate with HRP-Conjugated Streptavidin									
	5. Incubate with Detection Buffers									
	6. Image with chemiluminescent imaging system									
	7. Perform densitometry and analysis									
Sample Preparation:	Use serum-free conditioned media if possible. If serum-containing conditioned media is									
	required, it is highly recommended that complete medium be used as a control since many									
	types of sera contains cytokines. We recommend the following parameters for your samples:									
	50 to 100 μl of original or diluted serum, plasma, cell culture media, or other body fluid, or 50-									
	500 μ g/ml of protein for cell and tissue lysates. If you experience high background or if the									
	fluorescent signal intensities exceed the detection range, further dilution of your sample is									
	recommended.									
Assay Procedure:	1. Place each membrane into the provided eight-well tray (- means the antibody printed side). 2.									

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Do not let the membrane dry out during detection. The detection process must be completed within 40 minutes without stopping. 1. Proceed with the detection reaction. Add 250µl of 1X Detection Buffer C and 250µl of 1X Detection Buffer D for one membrane, mix both solutions. Drain off excess wash buffer by holding the membrane vertically with forceps. Place membrane protein side up (- mark is on the protein side top left corner) on a clean plastic sheet (provided in the kit). Pipette the mixed Detection Buffer onto the membrane and incubate at room temperature for 2 minutes. Ensure that the detection mixture is completely and evenly covering the membrane without any air bubbles. 2. Drain off any excess detection reagent by holding the membrane, protein side up, on a piece of plastic sheet (- mark is on the protein side top left corner). Cover with another piece of plastic sheet on the array. Gently smooth out any air bubbles. Avoid using pressure on the membrane. 3. Expose the array to x-ray film (we recommend to use Kodak x-omat AR film) and detect signal using film developer. Or the signal can be detected directly from the membrane using a chemiluminescence imaging system.

	Expose the membranes for 40 seconds and then re-expose the film according to the intensity of signals. If the signals are too strong (background too high), reduce exposure time (e.g. 5-30 seconds). If the signals are too weak, increase exposure time (e.g. 5-20 min or overnight). Or re incubate membranes overnight with 1x HRP-conjugated streptavidin, and redo detection in the second day. 4. Save membranes in -20° C to -80° C for future reference.							
Calculation of Results:	Visual comparison of array images may be sufficient to see differences in relative protein expression. However, most researchers will want to perform numerical comparisons of the signal intensities (or more precisely, signal densities), using 2-D densitometry. Gel/Blot documentation systems and other chemiluminescent or phosphorescent detection systems are usually sold as a package with compatible densitometry software. Any densitometry software should be sufficient to obtain spot signal densities from your scanned images. One such software program, ImageJ, is available for free from the NIH website along with an array plug-in.							
Assay Precision:	Inter-array Coefficient of Variation (CV) of spot signal intensities as low as 5% when run under optimal conditions.							
Restrictions:	For Research Use only							
Handling								
Handling Advice:	The antibody printed side of each membrane is marked by a dash (-) or number (#) in the upper left corner. Do not allow membranes to dry out during the experiment or they may become fragile and break OR high and/or uneven background may occur. Grasp membranes by the corners or edges only using forceps. DO NOT touch printed antibody spots.							
Storage:	-20 °C							
Storage Comment:	For best results, store the entire kit frozen at -20°C upon arrival. Stored frozen, the kit will be stable for at least 6 months which is the duration of the product warranty period. Once thawed, store array membranes and 1X Blocking Buffer at -20°C and all other reagents undiluted at 4°C for no more than 3 months.							
Expiry Date:	6 months							
Publications								
Product cited in:	Lee, Schierer, Blume, Dindorf, Wittki, Xiang, Ostalecki, Koliha, Wild, Schuler, Fackler, Saksela, Harrer, Baur: "HIV-Nef and ADAM17-Containing Plasma Extracellular Vesicles Induce and Correlate with Immune Pathogenesis in Chronic HIV Infection." in: EBioMedicine , Vol. 6, pp.							

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There are more publications referencing this product on: Product page

Images

Image 1.

		A	B	C	D	E	.F	G	н	1	J	K	L	M	N
2	1	POS	POS	NEG	NBG	BLANK	Adiponectia (ACRP30)	AgRP	ANGPT2	AR	Axi	bFGF	beta-NGF	BIC	CCL28 (MBC)
ertical	3 4	CTACK (CCL27)	Dtk	IGFR	ENA-78 (CXCL5)	Fas (Apo-1)	FGF-4	KGE-9	GCSF	GITR Ligand (TNESE18)	GITE (TNFRSF18)	GRO a/b/g	GPO alpha (CXCL1)	HCC-4 (CCL16)	HGF
duplicate v	5	KAM-1 (CD54)	ICAM-3 (CD50)	IGFBP-3	IGF87-6	KGF-1	IL-1 84 (ST2)	IL-1 81	llan.	IL-12 p40	IL-12 p70	IL-17A	IL-2 R alpha	IL o R	ILS (CICLS)
	7 8	FTAC (C)(L11)	Lympho- tactin	MIF	MIP-1 alpha (CCL3)	MIP-1 beta (CCL4)	MIF-3 beta (CCL19)	MSP alpha/beta	NT-4	OPG (TNFRSF11B)	NRO	PLGF	gp130	THE RII (THERSE1B)	TNF RI (TNFRSE1A)
	9 10	TECK (CCL25)	TIMP-1	TIMP-2	TPO	TRAIL R3 (TNERSETOC)	TRAIL R4 (TNFRSF10D)	sPAR.	VIGEA	VIGF-D	BLANK	BLANK	BLANK	BLANK	POS

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