

Procarta[®] Transcription Factor Plex Kit

User Manual

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Procarta Transcription Factor Assay Kit User Manual

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Citing Procarta in Publications

When describing a procedure for publication using this product, we would appreciate it if you would refer to it as the Procarta™ Transcription Factor Plex Kit.

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About the User Manual

Who Should Read this Manual Anyone that has purchased a Procarta Transcription Factor (TF) Plex Kit from Panomics to perform profiling of up to 44 different transcription factors per reaction in the following sample types:

- ◆ Nuclear extracts from cultured cells
- ◆ Cell lysates from cultured cells

What this Manual Covers This manual provides recommendations and step-by-step procedures for the following:

- ◆ Guidelines for assay design and data analysis
- ◆ Sample and assay preparation
- ◆ Set up and operation of the vacuum manifold system
- ◆ Assay procedure
- ◆ Troubleshooting

Safety Warnings and Precautions **CAUTION** All chemicals should be considered potentially hazardous. We recommend that this product and its components be handled by those trained in laboratory techniques and be used according to the principles of good laboratory practice.

CAUTION This kit contains small quantities of sodium azide. Sodium azide reacts with lead and copper plumbing to form explosive metal azides. When disposing, flush drains with a large volume of water to prevent azide accumulation. Observe all state and local regulations for disposal.

Note This product is intended for research use only. Not for diagnosis of disease in humans or animals.

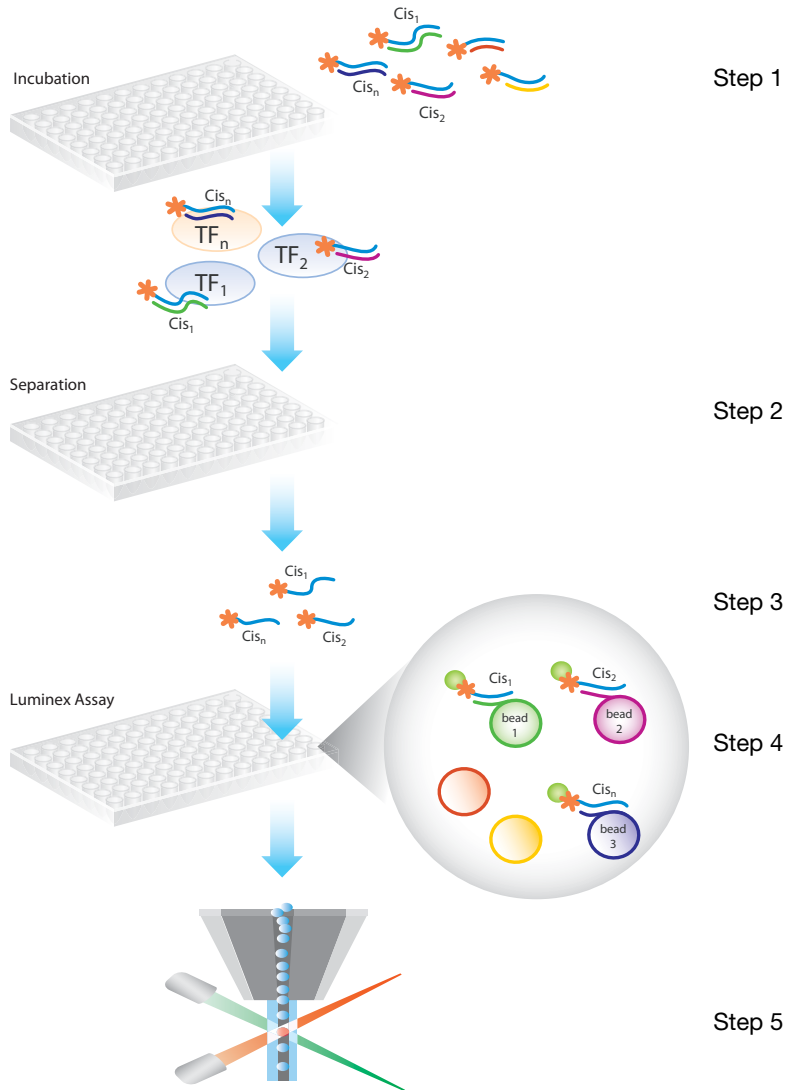
For More Information For information about the Procarta products mentioned in this manual, visit our website at www.panomics.com.

About the Procarta Transcription Factor Assay Kit

Fundamentals of Procarta Transcription Factor Assay

The Procarta TF assays combine two technologies: a patent-pending detection probe separation method and xMAP® (multi-analyte profiling beads). Together, these two technologies enable quantitative measurement of DNA binding activity of up to 44 transcription factors from as little as 500 ng of protein from a whole-cell lysate or 250 ng of protein from a nuclear extract. The xMAP technology, developed by Luminex® Corp., combines flow cytometry, fluorescent-dyed microspheres (beads), lasers, and digital signal processing to effectively allow multiplexing of up to 100 unique assays within a single sample.

Assay Overview



Assay overview:

Step	Action
1	Prepare nuclear extract or whole cell lysate and incubate with biotin-labeled DNA binding probes to form protein/DNA complexes. Bind protein/DNA complexes to the semi-porous filter using a Separation Plate.

Assay overview: (continued)

Step	Action
2	Wash unbound probes from the protein/DNA complexes utilizing the Separation Plate.
3	Chemically denature the protein/DNA complexes to release DNA from the filter in the Separation Plate and elute into PCR plate
4	Denature the double stranded eluted DNA using a thermal cycler and hybridize detection probes with TF-specific antisense conjugated beads.
5	Detect Streptavidin-conjugated R-phycoerythrin (SAPE) probe-bound beads with a Luminex instrument.

Available Kit Formats

Procarta TF Assay Kits are available in 1/4- and 1-plate 96-well formats for the standard 40-plex. Assay kits contain all the reagents required to detect transcription factors from prepared whole-cell lysates or nuclear extracts, including nuclear extract and whole-cell lysate control samples. Reagents for the preparation of nuclear extracts is included in this manual. The kit for the preparation of whole cell lysates is sold separately.

Additional Procarta TF Assay Kit are available as:

- ◆ User selected (2–43 plex) assay kits in 1-plate formats. Selected TFs are provided in premixed, ready to use format. All “By Request” assay kits must include NFκB in order to run the positive and negative controls provided in the kit and include reagents for the preparation of nuclear extracts.

Procarta TF Plex Kit Contents and Handling Conditions

Kit Contents and Storage

The Procarta TF Assay Kit contains the following components. Refer to the product insert for quantities and details of components supplied.

Procarta TF Plex: Box 1 shipped on dry ice, but store at -80 °C upon receipt

Component	Description
Positive Control Nuclear Extract	Untreated HeLa nuclear extract with addition of NFκB recombinant protein
Negative Control Nuclear Extract	Untreated HeLa nuclear extract
Positive Control Whole Cell Lysate	Untreated HeLa Whole Cell Lysate with addition of NFκB Recombinant Protein
Negative Control Whole Cell Lysate	Untreated HeLa Whole Cell Lysate

Procarta TF Plex: Box 2 shipped on dry ice, but store at -20 °C upon receipt

Component	Description
Buffer A	Aqueous buffered solution for the preparation of nuclear extracts.
Buffer B	Aqueous buffered solution for preparation of nuclear extract or dilution of nuclear extracts prior to running TF Plex Assay.
DTT, 100 mM	Aqueous solution for stabilizing protein

Component	Description
Protease Inhibitor	Aqueous solution for inhibiting protease activity
Phosphatase Inhibitor I	Aqueous solution for inhibiting phosphatase activity
Phosphatase Inhibitor II	Aqueous solution for inhibiting phosphatase activity
Detection Probes	Double stranded DNA oligos. Oligos contain consensus sequences for specific TFs. Sense strand is biotinylated.

Procarta TF Plex Kit : Box 3 Shipped on blue ice and store at 4 °C upon receipt

Binding Buffer	Aqueous buffered solution for protein/DNA binding
Separation Buffer	Aqueous buffered solution for separation of protein/DNA complex from free probes
Assay Buffer	Aqueous buffered solution for hybridization
Wash Buffer	Aqueous buffered solution for Capture Bead washing
Reading Buffer	Aqueous buffered solution for detection of Capture Beads
Capture Beads, premixed	Pre-mixed Luminex beads conjugated with anti-sense oligo corresponding to the Detection Probes
Streptavidin-PE	Streptavidin-conjugated R-Phycoerythrin for detection of bound biotin probes
Elution Buffer	Aqueous buffered solution for eluting the bound TF probes from protein
PCR Plate	96-well clear PCR plate for Protein/DNA Complex formation
Sample Collection Plate	96-well clear skirted-PCR plate
Utility Plate	96-well clear polystyrene plate for collection of waste during the separation of TF-Bound Detection Probes. Also used as a holder for the Filter Plate during the detection of eluted Detection Probes
Separation Plate	96-well white plate for the separation of Protein/DNA complexes from free probes
PCR Plate Seal	Plastic sealer for sample collection plate used to denature samples at 95 °C and other PCR plates
Plate Seal	Adhesive-backed foil for sealing filter plates during bead assay
Filter Plate	96-well sterile filter plate

- Kit Handling**
- ◆ Store all controls at -80°C and avoid multiple freeze/thaws.
 - ◆ If precipitates occur in the Elution Buffer, warm to 37 °C with gentle swirling.

Required Materials and Equipment Not Provided

Equipment

Item	Source
Vacuum filtration system	Millipore (P/N MAVM0960R and WP6111560)

Item	Source
PCR Instrument for controlled incubation of samples in PCR Plates	MJ Research Model PCT-200 or equivalent
Microplate centrifuge capable of working at 4 °C	Eppendorf 5415D or equivalent
Microplate shaker	Labline model 4625 or equivalent with 3 mm orbit
Shaking incubator with microplate adaptor	E&K Scientific (Vortemp 56 P/N S-2056) or equivalent
Luminex or Luminex-based instrument	MiraiBio, Bio-Rad or other Luminex instrument provider

Materials

Item	Source
Reagent Reservoirs, 25 mL and 100 mL capacities	Diversified Biotech (P/N RESE-3000, RESE-1000)
Procarta Transcription Factor Nuclear Extraction Kit or Whole Cell Lysis Kit	Panomics (P/N PC5101), (P/N PC5102)
Protein Determination Kit	Bio-Rad DC Protein Assay Kit (P/N 500-0112) or equivalent
DC Protein Assay	Bio-Rad
Phosphate Buffered Saline	Multiple Sources
Aluminum Foil	Multiple Sources

Guidelines for Assay Design and Data Analysis

Overview Here we provide information and guidelines on the following:

- ◆ Preparation of Nuclear Extracts
- ◆ Optimizing sample input
- ◆ Running assay controls
- ◆ Running replicate samples
- ◆ Analyzing data

Note An example experimental plate layout is provided in the Appendix.

Preparing Samples Protein concentration of sample inputs should be in the range of 0.4–2 µg/µL. As a starting point, we recommend that each 5 µL sample contain a total of 2 µg of protein. For example, dilute samples to 0.2 µg/µL, using Working Sample Dilution Buffer, and then load 5 µL.

Optimizing Sample Input We recommend running a 4-point, 2-fold serial dilution of the sample (diluting with Working Sample Dilution Buffer) to ensure you are operating in the linear range of the reader and assay.

For nuclear extracts, initially, we recommend preparing samples such that the addition of 5 µL of sample will contain a total of 2.0, 1.0, 0.5, and 0.25 µg of protein.

For whole cell lysates, initially, we recommend preparing samples such that the addition of 5 µL of sample will contain a total of 4.0, 2.0, 1.0, and 0.5 µg of protein.

Running Assay Controls

Assay Background Control

Assay background is the assay signal (median fluorescence intensity, MFI) generated by the assay components in the absence of sample.

We recommend you run an assay background control in every experiment and subtract the assay background MFI from each sample MFI when analyzing the data.

Whole Cell Lysate and Nuclear Extract Controls

Whole cell lysate and nuclear extract controls enable you to monitor assay performance. These controls are provided in the kit and are also available separately.

We recommend using at least one set of positive and negative controls in every experiment to monitor assay performance.

Running Replicates

Technical Replicates

These are replicates from a single extract or lysate sample.

Biological Replicates

These are replicates from different extracts or lysates but the extracts or lysates are biologically-equivalent.

Replicate Recommendations

For assays with 3–6 biological replicates, run 1 assay well/biological sample. For assays without biological replicates, run 3 technical replicate assay wells/biological sample.

Analyzing Data

Assay Precision

Run assays of the same sample in triplicate.

Calculate standard deviation and assay coefficient of variation (%CV = [std dev/mean] x 100%). Assay CVs are typical less than 15% for technical replicates.

Fold Change

Subtract average MFI background from all samples. Calculate fold change as the treated sample value/untreated sample value.

Assay Controls

For the positive control, calculate the ratio of the NFκB signal (MFI) divided by the assay background. For NFκB, this value should be >20.

For each set of positive and negative controls, subtract the NFκB assay background from the NFκB signals in the controls. Then, divide the background-corrected NFκB signal from the positive control by the background-corrected NFκB signal from the negative control. This ratio, or fold change calculation should be >5.

Set-Up and Operation of the Vacuum Manifold System

About Using the Vacuum Manifold

This topic describes how to set up and use the Millipore vacuum manifold. This includes how to calibrate the pressure and important guidelines that will help to ensure good assay reproducibility.

We recommend that you set up and calibrate the manifold before you start the assay to ensure the assay is performed without interruption.

Sealing Filter Plates

- ◆ Lay a Plate Seal over the Filter Plate and roll a 5 mL serological pipet (or equivalent) over the Plate Seal to seal the Filter Plate.

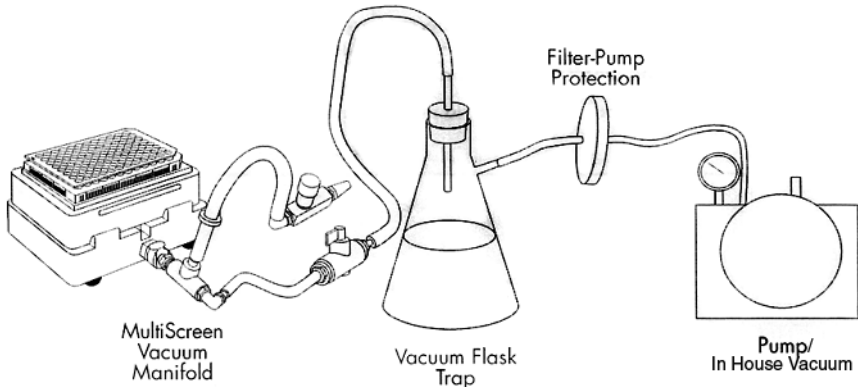
This ensures adequate plate sealing while avoiding any leakage due to capillary action.

IMPORTANT To avoid Filter Plate leakages, do not seal Filter Plates using a rubber roller (or equivalent) as they apply significant pressure resulting in leakage.

- ◆ Seal all unused wells with the provided Plate Seal to ensure proper vacuum pressure.

Setting Up and Calibrating the Manifold

To set up and calibrate the manifold:

Step	Action
1	<p>Set up the Filter Plate vacuum manifold as shown below. Follow the manufacturer's manual for details.</p>  <p>The diagram illustrates the vacuum manifold setup. On the left is the MultiScreen Vacuum Manifold. A tube connects it to a Vacuum Flask Trap (a flask with a side arm). This trap is connected to a Filter-Pump Protection device (a circular component). Finally, a tube connects the protection device to a Pump/In House Vacuum source, which includes a pressure gauge.</p>
2	<p>Calibrate the vacuum pressure using the Utility Plate:</p> <ol style="list-style-type: none"> Place the Utility Plate on top of the manifold. Turn on the vacuum. Press the corners of the Utility Plate to form a tight seal. Set the pressure to 2–3 mm of Hg. <p>IMPORTANT During filtration, maintain the vacuum between 2–3 mm Hg. Higher vacuum may result in the loss of Capture Beads.</p>

Operating the Manifold To operate the manifold:

Step	Action
1	Once the vacuum is set correctly, remove the Utility Plate. Check vacuum calibration periodically. As a general guideline, 200 μ L of solution should take approximately 2–5 seconds to clear the well of a Filter Plate.
2	For all filtration steps, turn the Filter Plate vacuum manifold on, transfer the Filter Plate to the vacuum manifold and then filter the solution. Avoid splashing and cross-contamination of wells during all wash steps. IMPORTANT During filtration, maintain the vacuum between 2–3 mm of Hg. Higher vacuum settings may result in loss of Capture Beads. IMPORTANT Do not allow the Filter Plates to air-dry following washes. Immediately add the next component following each filtration step.
3	Break the vacuum immediately after each solution has been completely filtered from all wells (approximately 2–5 seconds) by first turning off the vacuum, then removing the plate from the manifold. Note Wells typically filter at different rates.
4	Place the Filter Plate back on the Utility Plate.
5	Following the last wash in each series, blot the bottom of the Filter Plate thoroughly with a paper towel to remove traces of Wash Buffer. Avoid touching the bottom of the Filter Plate with your fingers or to the bench during manipulations.

Nuclear Extraction Procedure for Cultured Cells

Assay Guidelines **IMPORTANT** All components and PBS must be kept on ice at all times. Buffer A and B Working Reagents must be kept on ice and should be used within 2 hours of preparation.

Preparing Working Reagents To prepare working reagents:

Step	Action						
1	<p>Prepare 1 mL of Buffer A Working Reagent:</p> <ol style="list-style-type: none"> Combine: <ul style="list-style-type: none"> – 1 mL Buffer A – 10 μL DTT – 10 μL Protease Inhibitor – 10 μL Phosphatase Inhibitor I – 10 μL Phosphatase Inhibitor II Invert to mix. <p>Scale preparation of Buffer A Working Reagent based on experimental requirements. Use the table below as a guide.</p> <table border="1"> <thead> <tr> <th>Vessel</th> <th>Quantity of Buffer A Working Reagent</th> </tr> </thead> <tbody> <tr> <td>100 mm culture dish</td> <td>1 mL/dish</td> </tr> <tr> <td>6-well plate</td> <td>250 μL/well</td> </tr> </tbody> </table>	Vessel	Quantity of Buffer A Working Reagent	100 mm culture dish	1 mL/dish	6-well plate	250 μ L/well
Vessel	Quantity of Buffer A Working Reagent						
100 mm culture dish	1 mL/dish						
6-well plate	250 μ L/well						
2	<p>Prepare 1 mL of Buffer B Working Reagent:</p> <ol style="list-style-type: none"> Combine: <ul style="list-style-type: none"> – 1 mL Buffer B – 10 μL DTT – 10 μL Protease Inhibitor – 10 μL Phosphatase Inhibitor I – 10 μL Phosphatase Inhibitor II Invert to mix. <p>Scale preparation of Buffer B Working Reagent based on experimental requirements. Use the table below as a guide.</p> <table border="1"> <thead> <tr> <th>Vessel</th> <th>Quantity of Buffer B Working Reagent</th> </tr> </thead> <tbody> <tr> <td>100 mm culture dish</td> <td>150 μL/dish</td> </tr> <tr> <td>6-well plate</td> <td>150 μL/well</td> </tr> </tbody> </table>	Vessel	Quantity of Buffer B Working Reagent	100 mm culture dish	150 μ L/dish	6-well plate	150 μ L/well
Vessel	Quantity of Buffer B Working Reagent						
100 mm culture dish	150 μ L/dish						
6-well plate	150 μ L/well						

Preparing Nuclear Extracts From Adherent Cells

To prepare nuclear extracts:

Step	Action						
1	Remove the culture media from all wells and wash cells twice with an appropriate volume of cold 1X PBS.						
	<table border="1"> <thead> <tr> <th>Vessel</th> <th>Quantity of PBS</th> </tr> </thead> <tbody> <tr> <td>100 mm culture dish</td> <td>10 mL/dish</td> </tr> <tr> <td>6-well plate</td> <td>1 mL/well</td> </tr> </tbody> </table>	Vessel	Quantity of PBS	100 mm culture dish	10 mL/dish	6-well plate	1 mL/well
	Vessel	Quantity of PBS					
100 mm culture dish	10 mL/dish						
6-well plate	1 mL/well						
2	Following the second wash, make sure the PBS is completely removed.						
3	Add the appropriate volume of Buffer A Working Reagent to the wells.						
	<table border="1"> <thead> <tr> <th>Vessel</th> <th>Quantity of Buffer A Working Reagent</th> </tr> </thead> <tbody> <tr> <td>100 mm culture dish</td> <td>1 mL/dish</td> </tr> <tr> <td>6-well plate</td> <td>250 µL/well</td> </tr> </tbody> </table>	Vessel	Quantity of Buffer A Working Reagent	100 mm culture dish	1 mL/dish	6-well plate	250 µL/well
	Vessel	Quantity of Buffer A Working Reagent					
100 mm culture dish	1 mL/dish						
6-well plate	250 µL/well						
4	Transfer culture vessel(s) to an ice bucket and transfer ice bucket to a rocking platform at 200 rpm for 10 minutes.						
5	Release the cells from the bottom of the culture vessel: <ol style="list-style-type: none"> a. Using a sterile cell scraper, remove the cells b. Pipet up and down several times to disrupt the cell clumps. (Avoid creating bubbles). c. Rinse the bottom of the culture vessel with the cell suspension to maximize the cell yield. 						
6	Transfer each sample to a 1.5 mL microcentrifuge tube and centrifuge at 14,000 x g for 3 minutes at 4 °C.						
7	Remove and discard the supernatant(s) and keep the pellet(s) on ice.						
8	If you started with a 10cm plate, add 150 µL of Buffer B Working Reagent to each pellet and vortex at highest setting for 10 seconds. If you started with a 6 well plate, add 30 µL of Buffer B Working Reagent to each pellet. The pellet will detach from the microcentrifuge tube wall and may not disperse into a homogenous solution. This is normal. Do not attempt to disperse the pellet.						
9	Position the microcentrifuge tubes horizontally in an ice bucket and transfer ice bucket to a rocking platform at 200 rpm for 2 hours.						
10	Centrifuge samples at 14,000 x g for 5 minutes at 4 °C.						
11	Transfer supernatant(s) to a new microcentrifuge tube. This is your nuclear extract. It is good practice to prepare aliquots of your nuclear extract to avoid multiple freeze thaw events.						
12	Measure the protein concentration of each sample using a the DC Protein Assay from Bio-Rad. The BCA protein assay will not measure the concentration properly. Store samples at -80 °C or use immediately.						
	<table border="1"> <thead> <tr> <th>Vessel</th> <th>Typical Protein Yields</th> </tr> </thead> <tbody> <tr> <td>10 cm culture dish</td> <td>150–300 µg/dish</td> </tr> <tr> <td>6-well plate</td> <td>50–100 µg/well</td> </tr> </tbody> </table>	Vessel	Typical Protein Yields	10 cm culture dish	150–300 µg/dish	6-well plate	50–100 µg/well
	Vessel	Typical Protein Yields					
10 cm culture dish	150–300 µg/dish						
6-well plate	50–100 µg/well						

Preparing Nuclear Extracts From Suspension Cells

To prepare nuclear extract for suspension cells:

Step	Action						
1	Transfer cells to a 1.5 mL or 15 mL centrifuge tube as appropriate and centrifuge at 500 x g for 5 minutes.						
2	Remove the culture media and wash cells by resuspending in 1 mL of cold 1X PBS followed by centrifugation at 500 x g for 5 minutes. Repeat wash step. Following the second wash step, ensure that the 1X PBS solution is completely removed from the cells. Note Before the second centrifugation, if necessary, transfer contents from the 15 mL centrifuge tube to a 1.5 mL microcentrifuge tube.						
3	Immediately add the appropriate volume of Buffer A Working Reagent to cell pellets. Mix by pipetting up and down several times. <table border="1" data-bbox="592 667 1453 787"> <thead> <tr> <th>Vessel</th> <th>Buffer A Working Reagent</th> </tr> </thead> <tbody> <tr> <td>100 mm culture dish</td> <td>1 mL/dish</td> </tr> <tr> <td>6-well plate</td> <td>250 µL/well</td> </tr> </tbody> </table>	Vessel	Buffer A Working Reagent	100 mm culture dish	1 mL/dish	6-well plate	250 µL/well
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100 mm culture dish	1 mL/dish						
6-well plate	250 µL/well						
4	Transfer tube(s) to an ice bucket and transfer ice bucket to a rocking platform at 200 rpm for 10 minutes.						
5	Centrifuge samples at 14,000 x g for 3 minutes at 4 °C.						
6	Discard the supernatant(s) and keep the pellet(s) on ice.						
7	If you started with a 10 cm plate, add 150 µL of Buffer B Working Reagent to each pellet and vortex at highest setting for 10 seconds. If you started with a 6 well plate, add 30 µL of Buffer B Working Reagent to each pellet and vortex at highest setting for 10 seconds. The pellet will detach from the microcentrifuge tube wall and may not disperse into a homogenous solution. This is normal. Do not attempt to disperse the pellet.						
8	Position the microcentrifuge tubes horizontally in an ice bucket and transfer ice bucket to a rocking platform at 200 rpm for 2 hours.						
9	Centrifuge samples at 14,000 x g for 5 minutes at 4 °C.						
10	Transfer supernatant(s) to a new microcentrifuge tube. This is your nuclear extract. It is good practice to prepare aliquots of your nuclear extract to avoid multiple freeze thaw events.						
11	Measure the protein concentration of each sample using the DC Protein Assay quantitation assay from Bio-Rad (sold separately). The BCA protein assay will not measure the concentration properly. Store samples at -80 °C or use immediately. <table border="1" data-bbox="592 1476 1453 1596"> <thead> <tr> <th>Vessel</th> <th>Typical Protein Yields</th> </tr> </thead> <tbody> <tr> <td>100 mm culture dish</td> <td>150–300 µg/dish</td> </tr> <tr> <td>6-well plate</td> <td>50–100 µg/well</td> </tr> </tbody> </table>	Vessel	Typical Protein Yields	100 mm culture dish	150–300 µg/dish	6-well plate	50–100 µg/well
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6-well plate	50–100 µg/well						

TF Plex Assay Procedure

- Before You Start** ♦ When preparing the samples for the TF Plex assay, prepare an appropriate volume of Buffer B Working Reagent to appropriately dilute the sample. Based on

the calculations and the number of replicate samples needed, prepare Buffer B Working Reagent by combining 100 parts of Sample Dilution Buffer, 1 part DTT, 1 part Protease Inhibitor Cocktail, 1 part Phosphatase Inhibitor I and 1 part Phosphatase Inhibitor II. Store on ice.

- ◆ For 1-plate kits, there is sufficient reagents to process 1 entire plate using reagent reservoirs and multi-channel pipets. If you are running several partial plates, you may not have sufficient reagents for all steps. For 1/4-plate kits, there is not sufficient reagents to process the samples using reagent reservoirs and multi-channel pipets. For 1/4-plate kits, perform reagent additions using a single-channel pipet.

Forming Protein-DNA Complexes

To form Protein-DNA complexes:

Step	Action
1	Thaw Detection Probes on ice.
2	If you are not using a whole plate, trim a PCR Plate Seal to the appropriate size and seal any unused wells of the PCR Plate.
3	<p>Prepare the PCR Plate so that each of the control, sample and blank wells will contain 20 μL of either the following:</p> <p>Controls</p> <p>10 μL of Detection Probe to every experimental well of the PCR Plate</p> <p>10 μL of Nuclear Extract Controls and/or Whole-Cell Lysate Controls</p> <p>Total of 20 μL</p> <p>Samples</p> <p>10 μL of Detection Probe to every experimental well of the PCR Plate</p> <p>5 μL of prepared sample to each designated well + 5 μL of nuclease-free distilled water</p> <p>Total of 20 μL</p> <p>Blanks</p> <p>10 μL of Detection Probe to every experimental well of the PCR Plate</p> <p>5 μL of Working B Reagent to each designated well + 5 μL of nuclease-free distilled water</p> <p>Total of 20 μL</p> <p>Note As a starting point, we recommend that each 5 μL sample contain a total of 2 μg of protein. Dilute samples with Buffer B Working Reagent.</p>
4	<p>Mix samples by gently tapping the bottom of the PCR Plate and cover the plate with the aluminum seal.</p> <p>IMPORTANT Make sure all bubbles are removed from the wells. (Tapping the plate should accomplish this and if not, perform a quick spin in the centrifuge)</p>

To form Protein-DNA complexes: *(continued)*

Step	Action
5	<p>Incubate the plate at 15 °C for 30 minutes using a PCR machine</p> <p>IMPORTANT Make sure the temperature is maintained at 15 °C.</p> <p>IMPORTANT Samples can be kept at 4 °C for up to 1 hour before proceeding to the next step.</p>

Eluting TF-Bound Detection Probes

To elute TF-bound detection probes:

Step	Action
1	<p>Prepare Separation Plate:</p> <ol style="list-style-type: none"> Trim a foil Plate Seal to the appropriate size and seal any unused wells of the Separation Plate. Place the Utility Plate on the bottom of the Separation Plate.
2	<p>Pre-wet the Separation Plate:</p> <ol style="list-style-type: none"> Add 180 µL/well ice-cold Binding Buffer to each well of the Separation Plate. Centrifuge the Separation Plate/Utility Plate assembly at 563 x g for 2 minutes at 4 °C. Discard the flow through and dry the top surface of the Utility Plate with a clean paper towel to avoid contaminating the bottom of the Separation Plate. <p>IMPORTANT Do not place the Separation Plate directly on any surface other than the specified plates as this might result in cross contamination of wells. Place the Utility Plate on the bottom of the Separation Plate.</p>
3	<p>Transfer the samples to the Separation Plate:</p> <ol style="list-style-type: none"> Add 20 µL ice-cold Binding Buffer to each well of the protein-DNA complexes in the PCR Plate. Avoid pipetting up and down. Transfer 30 µL of the sample to the corresponding well of the Separation Plate. Make sure you apply the sample to the center of the filter.
4	<p>With the Utility Plate on the bottom of the Separation Plate, incubate on ice for 30 minutes.</p> <p>IMPORTANT Do not exceed the incubation time as this will result in high background.</p>
5	<p>Wash the Separation Plate:</p> <ol style="list-style-type: none"> Add 180 µL/well ice-cold Separation Buffer to each well of the Separation Plate. Incubate Separation Plate/Utility Plate assembly on ice for 5 minutes. Centrifuge the Separation Plate/Utility Plate assembly at 563 x g for 2 minutes at 4 °C. Discard the flow through and dry the top surface of the Utility Plate with a clean paper towel to avoid contaminating the bottom of the Separation Plate. Place the Separation Plate back on top of the Utility Plate.

To elute TF-bound detection probes: *(continued)*

Step	Action
6	Repeat step 5 (omitting step 5b) four more times for a total of 5 washes. IMPORTANT Do not reduce the number of washes as this will result in high background.
7	Centrifuge the Separation Plate/Utility Plate assembly at 563 x g for 3 minutes at 4 °C.
8	Elute TF-bound Detection Probes: a. Add 60 µL of Elution Buffer to the center of each experimental well in the Separation Plate. b. Seal unused wells of the Sample Collection Plate with a foil Plate Seal. c. Place the Separation Plate on top of the Sample Collection Plate and incubate at room temperature for 5 minutes. d. Centrifuge the Sample Collection/Separation Plate assembly at 563 x g for 3 minutes. You should have 60 µL/well in the Collection Plate. IMPORTANT Use a scale to balance the Sample Collection/Separation Plate assembly and ensure that the Sample Collection Plate is properly seated below the Separation Plate
9	Place the samples on ice and continue to the next step or, cover with a foil Plate Seal and store at –20 °C until you are ready to use. Thaw on ice before use.

Denaturing and Hybridizing the Detection Probes

IMPORTANT Before You Start, turn on the Luminex-based reader at least 30 minutes before you intend to read your plate.

To Denature and Hybridize the Detection Probes:

Step	Action
1	Seal the samples wells in the Sample Collection Plate with a PCR Plate Seal. Using another PCR Plate Seal, cover any unused wells. IMPORTANT Do not use a foil Plate Seal as it will stick to the plate at high temperatures.
2	Denature samples at 95 °C for 5 minutes using a PCR instrument or heat block. Place samples on ice for 5 minutes.
3	Place the Filter Plate on top of the Utility Plate. Cut a foil Plate Seal to size and cover any unused wells on the Filter Plate.
4	Pre-wet the Filter Plate by dispensing 150 µL of Wash Buffer into each well and incubating the Filter Plate at room temperature for 5 minutes.
5	Remove the Wash Buffer using the vacuum manifold. When all Wash Buffer has been filtered, remove the Filter Plate from the vacuum manifold and blot the bottom of the Filter Plate with paper towels and place the Filter Plate on top of the Utility Plate.

To Denature and Hybridize the Detection Probes: *(continued)*

Step	Action
6	<p>Add premixed Capture Beads to the Filter Plate:</p> <ol style="list-style-type: none"> Vortex both premixed Capture Bead solution at the highest setting for 30 seconds. Dispense 50 μL of premixed Capture Beads to each well of the Filter Plate. Remove the Capture Bead buffer using the vacuum manifold. Blot the bottom of the Filter Plate with paper towels and place the Filter Plate on top of the Utility Plate. <p>IMPORTANT Do not invert or tap the Filter Plate.</p>
7	<p>Wash the Capture Beads:</p> <ol style="list-style-type: none"> Dispense 150 μL of Wash Buffer to each well of the Filter Plate. Remove the Wash Buffer using the vacuum manifold. Blot the bottom of the Filter Plate with paper towels and place the Filter Plate on top of the Utility Plate.
8	Dispense 40 μ L of Assay Buffer to each well of the Filter Plate.
9	<p>Add denatured Detection Probes</p> <ol style="list-style-type: none"> Transfer 30 μL of each eluted sample to each well of the Filter Plate according to your assay plate map. <p>IMPORTANT Change the pipet tip after every transfer and avoid creating bubbles.</p> <ol style="list-style-type: none"> Seal the wells of the Filter Plate with a foil Plate Seal. <p>IMPORTANT Gently apply the Aluminum Plate Seal over the plate. Pressing too hard can cause the sample to leak through the filter plate. Place the Filter Plate on top of the Utility Plate.</p>
10	<p>Incubate the Filter Plate/Utility Plate assembly in a Vortemp shaking incubator set at 50 °C and 300–500 rpm for 30 minutes.</p> <p>Note If a shaking incubator is not available, wrap the Filter Plate/Utility Plate assembly with aluminum foil, shake the assembly at room temperature for 10 minutes, then transfer to a 50 °C incubator for 30 minutes without shaking.</p>
11	Proceed to Binding the SAPE .

**Binding the
Streptavidin-PE
(SAPE)**

To bind the Streptavidin-PE (SAPE):

Step	Action
1	<p>Prepare to wash the Capture Beads:</p> <ol style="list-style-type: none"> Remove the Plate Seal carefully to avoid splashing of samples. Remove the buffer using the vacuum manifold. Blot the bottom of the Filter Plate with paper towels and place the Filter Plate on top of the Utility Plate.
2	<p>Wash the Capture Beads:</p> <ol style="list-style-type: none"> Dispense 150 μL of Wash Buffer to each well of the Filter Plate. Remove the Wash Buffer using the vacuum manifold. Blot the bottom of the Filter Plate with paper towels and place the Filter Plate on top of the Utility Plate.

To bind the Streptavidin-PE (SAPE): *(continued)*

Step	Action
3	<p>Add the Streptavidin-PE:</p> <ol style="list-style-type: none"> Invert the Streptavidin-PE tube to mix. Dispense 100 µL of Streptavidin-PE to each well of the Filter Plate. Seal the Filter Plate with a foil Plate Seal. Wrap the Filter Plate/Utility Plate assembly with aluminum foil. Shake the assembly on a plate shaker at room temperature for 30 minutes at 300–500 rpm.

Detecting the Signal

To detect the signal:

Step	Action
1	<p>Prepare to wash the Capture Beads:</p> <ol style="list-style-type: none"> Remove the Plate Seal carefully to avoid splashing of samples. Remove the buffer using the vacuum manifold. Blot the bottom of the Filter Plate with paper towels and place the Filter Plate on top of the Utility Plate.
2	<p>Wash the Capture Beads:</p> <ol style="list-style-type: none"> Dispense 150 µL of Wash Buffer to each well of the Filter Plate. Remove the Wash Buffer using the vacuum manifold. Repeat step 2a-2b once more. Blot the bottom of the Filter Plate with paper towels and place the Filter Plate on top of the Utility Plate.
3	<p>Resuspend Capture Beads:</p> <ol style="list-style-type: none"> Dispense 120 µL of Reading Buffer to each well of the Filter Plate. Seal the wells with a Plate Seal. Wrap the Filter Plate/Utility Plate assembly with aluminum foil. Shake the Filter Plate at 300–500 rpm and room temperature for 5 minutes or until ready to read on the Luminex instrument. <p>Note The Filter Plate/Utility Plate assembly can be wrapped with aluminum foil and stored flat, in the dark at 4 °C, for up to 48 hours before proceeding. However, delay in reading the plate may result in decreased sensitivity for some analytes. Shake the plate at 300–500 rpm for 5 minutes before reading.</p>

To detect the signal: *(continued)*

Step	Action																								
4	<p>Analyze the plate following the respective operation manual for the Luminex or Luminex-based instrument.</p> <table border="1"> <thead> <tr> <th>Software</th> <th>Sample Size</th> <th>DD Gate</th> <th>Timeout</th> <th>Bead Events/Bead Region</th> <th>Statistic</th> </tr> </thead> <tbody> <tr> <td>Luminex</td> <td>50 μL</td> <td>8,000–15,000</td> <td>25 sec.</td> <td>50</td> <td>Median</td> </tr> <tr> <td>Bioplex</td> <td>50 μL</td> <td>4,300-10,000</td> <td>25 sec.</td> <td>50</td> <td>Median</td> </tr> <tr> <td>MiraiBio</td> <td>50 μL</td> <td>2,000-15,000</td> <td>25 sec.</td> <td>50</td> <td>Median</td> </tr> </tbody> </table> <p>When the Luminex beads are injected into the flow cell, a small percentage of the beads will have a tendency to clump and go through the flow cell as doublets. The DD Gate or Doublet Discriminator Gate will allow for discrimination of the doublets from the singlet beads. When setting the DD Gate, you can follow the appropriate settings from the table above. However, for the best results, you should adjust the DD Gate and center the gates around the largest peak which is the singlet beads. You can adjust the gates when processing the first sample.</p> <p>The BioPlex® Suspension Array System allows calibration using Low or High sensitivity settings. Perform the sensitivity selection during calibration, using predetermined values of CAL2 RP1 target, as provided by Bio-Rad. Using the RP1 Low target value will provide results comparable to those obtained from the Luminex 100. Using the RP1 High target value may increase detection sensitivity for low cytokine protein concentrations but sacrifices part of the linear dynamic range for high concentrations. We recommend RP1 Low target value for BioPlex.</p> <p>IMPORTANT Check to ensure the probe height in the Luminex instrument is adjusted appropriately for the Filter Plate.</p> <p>IMPORTANT We recommend that you calibrate the Luminex or Luminex-based instrument each day the assay is run.</p> <p>IMPORTANT If there is a malfunction of the machine or software during the run or there was a problem with the system, the plate can be re-processed on the Luminex machine. Remove the plate from the machine and vacuum filter the plate. Resuspend the beads in Reading Buffer and shake for 5 minutes then re-read. The plate may take longer to read since there will be less beads in the plate.</p>	Software	Sample Size	DD Gate	Timeout	Bead Events/Bead Region	Statistic	Luminex	50 μ L	8,000–15,000	25 sec.	50	Median	Bioplex	50 μ L	4,300-10,000	25 sec.	50	Median	MiraiBio	50 μ L	2,000-15,000	25 sec.	50	Median
Software	Sample Size	DD Gate	Timeout	Bead Events/Bead Region	Statistic																				
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Normalizing Data Using TFIID Normalizing Data Using TFIID

Step	Action																								
1	<p>As an example for analyzing the data using TFIID as a normalization tool, background subtracted fluorescent values of the untreated and treated sample data will be analyzed.</p> <p>To normalize Sample B, the ratio between the TFIID values of Sample A and Sample B is calculated and then multiplied by the fluorescent value of the TF measured in Sample B.</p> <p>To normalize sample C, the ratio between the TFIID values of Sample A and Sample C is calculated and then multiplied by the fluorescent value of the TF measured in Sample C.</p> <p>Below is some example Data from the Luminex machine that has been background subtracted:</p> <table border="1" data-bbox="548 722 1308 884"> <thead> <tr> <th>Description</th> <th>TF #1</th> <th>TFIID</th> </tr> </thead> <tbody> <tr> <td>Sample A Untreated</td> <td>300</td> <td>250</td> </tr> <tr> <td>Sample B Treated</td> <td>450</td> <td>180</td> </tr> <tr> <td>Sample C Treated</td> <td>200</td> <td>350</td> </tr> </tbody> </table> <p>To calculate the Normalized values for Sample B and Sample C</p> <p>Sample B_{NORM} = (TFIID_{sample A}/TFIID_{sample B}) x Sample B</p> <p>Normalized value for Sample B: 625 = (250/180) x 450</p> <p>Sample C_{NORM} = (TFIID_{sample A}/TFIID_{sample C}) x Sample C</p> <p>Normalized value for Sample C: 143 = (250/350) x 200</p> <p>Below is the update table with the normalized values for TF#1</p> <table border="1" data-bbox="529 1392 1276 1640"> <thead> <tr> <th>Description</th> <th>TF #1</th> <th>TFIID</th> </tr> </thead> <tbody> <tr> <td>Sample A</td> <td>300</td> <td>250</td> </tr> <tr> <td>Sample B normalized</td> <td>B = (250 / 180)x450</td> <td>B = (250 / 180)x180</td> </tr> <tr> <td>Sample C normalized</td> <td>C = (250/350)x200</td> <td>C = (250/350)x350</td> </tr> </tbody> </table>	Description	TF #1	TFIID	Sample A Untreated	300	250	Sample B Treated	450	180	Sample C Treated	200	350	Description	TF #1	TFIID	Sample A	300	250	Sample B normalized	B = (250 / 180)x450	B = (250 / 180)x180	Sample C normalized	C = (250/350)x200	C = (250/350)x350
Description	TF #1	TFIID																							
Sample A Untreated	300	250																							
Sample B Treated	450	180																							
Sample C Treated	200	350																							
Description	TF #1	TFIID																							
Sample A	300	250																							
Sample B normalized	B = (250 / 180)x450	B = (250 / 180)x180																							
Sample C normalized	C = (250/350)x200	C = (250/350)x350																							

Troubleshooting

Possible Problems and Recommended Solutions

Observation	Possible Cause	Recommended Action
Filter plate leakage	Vacuum pressure too high	Adjust the vacuum pressure to 2–3 mm Hg as recommended in “Denaturing and Hybridizing the Detection Probes” on page 18.
	Filter Plate is misaligned (at an angle) during incubation/processing	Set the Filter Plate/Utility Plate assembly on a flat, level surface during incubation/processing.
	Leakage from capillary action	After each vacuum step, blot the bottom of the Filter Plate using paper towels or absorbent paper.
	Plate Seal applied with too much force	Do not use a rubber roller to seal the plate.
High CV	Sample not prepared properly	Make sure the samples are eluted properly by using an accurate counterweight when centrifuging.
	Bottom of the Filter Plate is not dry	After each vacuum step, blot the bottom of the Filter Plate using paper towels or absorbent paper.
	Contamination from re-using the Plate Seal	Use a new Plate Seal for each incubation step.
	Contamination from Wash Buffer	Be careful not to splash Wash Buffer during wash steps into adjacent wells.
Low bead count	Volume of bead solution is incorrect	Make sure the volume of Capture Beads is correct.
	Beads are clumping	Vortex the bead solution well before using in the assay.
	Vacuum pressure too high	Use 2–3 mm Hg vacuum pressure.
	Filter ruptured due to excess vacuum time	Do not use the vacuum over 10 seconds in any of the steps.
	Dyes contained in the beads are photo-bleached from overexposure to light	Store bead solution and the assay plate in the dark
	Reader is clogged	Follow the instructions in the Luminex instrument user documentation.

Observation	Possible Cause	Recommended Action
Low signal or sensitivity	Low protein concentration	Increase the protein by at least 2-fold and retest.
	Samples were not kept on ice or Binding Buffer is not cold.	Keep samples and Binding Buffer on ice.
	Beads or reagents expired	Verify the expiration date of the kit.
	Beads stuck to the bottom of the plate	Make sure the plate is agitated at 300–500 rpm for the recommended time and for at least 5 minutes before reading.
High backgrounds	Incubation too long during elution of TF-bound Detection Probes	Do not exceed the 30 minute incubation time.
	Reduced number of wash steps during elution of TF-bound Detection Probes	Do not reduce the number of wash steps to less than 5.
Low or no fold induction observed	Protein concentration too low or too high	Optimize sample input.
	Signals from instrument are saturated	Reduce protein concentration of sample input.
	Target protein not activated (induced)	Review induction procedures. You may need to change cell lines, inducer, or induction conditions.

Contacting Panomics

Technical Help For technical questions, contact our technical support group by telephone at 1-888-362-2447 or by email at pqbhelp@affymetrix.com (US and Canada) techsupport_europe@affymetrix.com (Europe), or visit our website www.panomics.com for an updated list of FAQs and product support literature.

For Additional Services For information about Affymetrix products or for ordering information, contact your Regional Sales Manager, or visit our website at www.panomics.com.

Appendix I

Bead-Analyte Associations For Panel 1

The following tables provide the bead-analyte associations for setting your Luminex instrument for Panel 1 of the Procarta TF Plex Assay. Refer to your product insert for analytes included in your kit.

Bead	Analyte	Bead	Analyte	Bead	Analyte
7	ELK-1	33	AR	52	CREB
11	NFκB	34	ETS/PEA	53	ER
12	FAST-1	35	AP-1	54	GR/PR
17	Oct	36	E2F1	55	HIF-1
18	p53	37	MyoD	56	FKHR
19	PAX-3	41	STAT-3	61	GATA
20	NF-E2	42	PPAR	62	IRF
21	AP-2	43	SMAD	63	STAT-1
25	NF-E1(Y1)	44	RUNX/AML	64	HNF1
26	ATF-2	45	Brn3	65	STAT-4
27	NF-1	46	CEBP	66	STAT-5
28	ISRE	47	NF-Y	73	MEF-2
29	PAX-5	51	c-myb	76	NFAT
32	Nkx-2.5			39	TFIID

Bead Analyte Associations for Panel 2

The following tables provide the bead-analyte associations for setting your Luminex instrument for Panel 2 of the Procarta TF Plex Assay. Refer to your product insert for analytes included in your kit.

Bead	Analyte	Bead	Analyte	Bead	Analyte
7	HAS+HBS	26	LF-A1	46	AP-4
8	ALF-1/TAL-1	27	LvF	47	CDP
9	TREF-1/2	28	MTF	51	XBP-1
10	EGR	32	PUR-1	52	CEF-1
11	NFκB	33	RB	53	ANTIOXIDANT RE
12	c-MYC	34	TFE-3	54	COUP-TF
13	NeuroD1	35	AP-1	55	EVI-1
14	PDX-1	36	USF-1	61	GAG
15	TR	37	XRE	63	IKAROS
17	MRE	38	GFI-1	65	HBS/XBP
18	ELF-1	41	NPAS2	66	SIE
19	SRE	42	PIT-1	73	HSF
20	H4TF	43	SRY	76	TR (DR-4)
21	HiNF	44	E47	39	TFIID
25	KPF-1	45	CCAAT		

Appendix II

Sample and Blank Plate Layouts

	1	2	3	4	5	6	7	8	9	10	11	12
A	PC ^a NE ^b		Sample A, 2 µg (from NE), treated			Sample B, 4 µg (from WCL), treated						
B	NC ^c NE		Sample A, 1 µg (from NE), treated			Sample B, 2 µg (from WCL), treated						
C	PC WCL ^d		Sample A, 0.5 µg (from NE), treated			Sample B, 1 µg (from WCL), treated						
D	NC WCL		Sample A, 2 µg (from NE), untreated			Sample B, 4 µg (from WCL), untreated						
E	Assay Back-ground		Sample A, 1 µg (from NE), untreated			Sample B, 2 µg (from WCL), untreated						
F			Sample A, 0.5 µg (from NE), untreated			Sample B, 1 µg (from WCL), untreated						
G												
H												

- a. PC = Positive Control
 - b. NE = Nuclear Extract
 - c. NC = Negative Control
 - d. WCL = Whole Cell Lysate
-

	1	2	3	4	5	6	7	8	9	10	11	12
A												
B												
C												
D												
E												
F												
G												
H												

