INSTRUCTION MANUAL

OptiMax™ Evaluation Kit





Better Immunoassays through Innovative Microfluidics

INTENDED USE:

The OptiMax[™] Evaluation Kit (OPV-IL6) and the associated Instruction Manual are specifically designed for first time user to provide a comprehensive overview to the Optimiser[™] microplate system.

Section I serves as an Introduction to the Optimiser[™] microplate system and guides the user through correct pipetting techniques with the Optimiser[™]. The pipetting technique instructions are accompanied by a Tutorial that will allow users to evaluate their pipetting proficiency on the Optimiser[™].

Section I also includes detailed instructions for an illustrative IL-6 assay (all assay reagents and buffers included with OPV-IL6). Completing the associated Tutorial will allow users to learn the assay operation sequence for Optimiser^M microplate assays. Successful completion of this assay will also help users understand the **POWER OF MICROFLUIDICS^M** to deliver high sensitivity ELISA results with only 5 µL sample volume and a ~ 2 hour assay protocol.

Section II is a detailed method description to be used for migrating a validated assay from conventional 96-well plates to Optimiser[™] microplate. Section II is presented as a series of 3 experiments, where each experiment sequence is described in complete detail including reagent preparation steps, assay plate layouts, assay procedures, calculations and data analysis methods. Section II also describes an illustrative IL-6 assay and the results from the IL-6 assay experiments are used to illustrate the data analysis methods used in the assay transfer guide.

Instruction Manual

OptiMax™ Evaluation Kit

For evaluating the performance of Optimiser[™] Microplate System

Catalogue Numbers: OPV-IL6WDR

Manufactured by:

Siloam Biosciences, Inc. 413 Northland Blvd. Cincinnati, Ohio 45240 USA

FOR RESEARCH USE ONLY

Not for use in clinical diagnostic procedures.

Read the Instruction Manual in its entirety before using the OptiMax[™] Evaluation Kit

Optimiser[™] microplates are warranted to perform in conformance with published product specifications in effect at the time of sale as set forth in product documentation and/or package inserts. Products are supplied for Research Use Only. The use of this product for any clinical diagnostic applications is expressly prohibited. The warranty provided herein is valid only when used by properly trained individuals and is limited to six months from the date of shipment and does not extend to anyone other than the original purchaser. No other warranties express or implied, are granted, including without limitation, implied warranties of merchantability, fitness for any particular purpose, or non-infringement. Buyers' exclusive remedy for non-conforming product during the warranty period is limited to replacement of or refund for the non-conforming product.

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Symbol indicates mandatory step required to ensure proper operation

Symbol indicates helpful tips to achieve optimal performance

SECTION I:

Using the Optimiser™ Microplate System

INTRODUCTION:

Siloam Biosciences' OptimiserTM technology offers a rapid and sensitive chemifluorescent-based ELISA procedure that uses very small sample volumes. The speed, sensitivity, and small sample requirements are enabled by the unique microfluidic design of the OptimiserTM microplate. Standard immunoassay reactions such as analyte capture and detection occur within a ~ 5 μ L microfluidic reaction chamber. The unique microchannel geometry and small reaction volumes favor rapid reaction kinetics. The typical assay procedure utilizes a 5 μ L sample and each reaction step is completed in 10 - 20 minutes. With wash time, substrate incubation time, and read time accounted for, a typical assay can be completed within approximately 2 hours.

Please refer to the Optimiser[™] Technology page on Siloam's website for more details on the principles behind the Optimiser[™] microplate platform.



Figure 1. Optimiser[™] microplate:

The Optimiser[™] microplate is a revolutionary new microplate format. With an ANSI/SBS compliant 96-well layout, the Optimiser[™] integrates the **Power of Microfluidics** to allow for low volume, rapid, and sensitive immunoassay protocols. Figure 1 shows the Optimiser[™] microplate schematic with magnified view of one "cell" of the Optimiser™. Each cell of the Optimiser[™] has a loading well (only used to add reagents) and a microfluidic reaction chamber. Reagents/samples are added to the well and transported via capillary action to an absorbent pad (not shown). The unique design of the Optimiser[™] allows the well to be drained but each liquid is trapped in the channel by capillary forces. As the next liquid volume is added, the capillary barrier is broken and the liquid within the microchannel is drawn out by the absorbent pad and replaced by the new reagent. All assay reactions occur within the microfluidic reaction chamber.

Siloam Biosciences' OptiMAX[™] evaluation kit is designed to provide first-time users with a comprehensive introduction to the methods of use and the capabilities of the Optimiser[™] platform. Specifically:

- The Pipetting Instruction Section and Tutorial 1 are designed to guide users through the correct method for pipetting to the Optimiser[™] microplate. Although very similar to the conventional 96-well ELISA plate, pipetting to the Optimiser[™] requires careful attention to a few key details for reliable performance.
- Tutorial 2 is designed to allow users to complete a model IL-6 assay. Tutorial 2 illustrates that the workflow for Optimiser[™] based assays is similar but much simplified when compared to conventional 96-well ELISA plates by eliminating the traditional wash step. Tutorial 2 also shows the capabilities of the Optimiser[™] to deliver equivalent sensitivity to conventional 96-well ELISA plates while using only 5 µL sample volume.
- The Optimiser[™] Assay Transfer Guide contains detailed instructions for users to transfer a working assay from the conventional 96-well ELISA plate to the revolutionary Optimiser[™] platform. Following the step-by-step instructions, users can also successfully test their own assay on the Optimiser[™] microplate.

MATERIALS PROVIDED AND REQUIRED:

Materials Provided:

OptiMax[™] Evaluation Kit provides the critical materials and reagents necessary for the Tutorials described in this manual and for the user to develop and optimize an ELISA assay on the Optimiser[™] platform. Table 1 identifies the kit contents, their function, and their required storage temperature. It is recommended that the package be opened and various components stored separately (as listed in Table 1) to conserve refrigerator shelf space.

Material	Quantity	Function	Storage / Handling, (before and after opening)		
Optimiser™ Holder	1	Holds Optimiser™ Microplate and Optimiser™ Pad in proper alignment			
Optimiser™ Microplate	10	Contains microfluidic reaction chambers Six Optimiser™ microplates for protocols described in manual; Four spare Optimiser™ microplates	Room temperature		
Optimiser™ Pad	20	Absorbs used reagent volume, single use			
96-well polypropylene v-bottom plate	3	For dilutions and reagent reservoir			
OptiBind™, A-L	1 vial each (4 mL)	Coating buffer panel for screening to determine optimal coating buffer for capture antibody			
OptiBind™-H	1 vial (10 mL)	Coating buffer for model IL-6 assay			
OptiBlock™	1 vial (30 mL)	Blocking buffer and diluent for detection antibody and SAv-HRP			
OptiWash™	1 vial (60 mL)	Wash buffer	Refrigerated		
OptiGlow™ - A	1 vial (5 mL)		(2 – 8 °C)		
OptiGlow™ - B	1 vial (5 mL)	Components of chemifluorescent substrate			
OptiGlow™ - C	1 vial (1 mL)				
Red dye solution	1 vial (7 mL)	Dyed blocking buffer solution for pipetting exercise			
Green dye solution	1 vial (7 mL)	Dyed wash buffer solution for pipetting exercise			
IL-6 standard	1 vial	Lyophilized recombinant IL-6 protein for model assay standard curve	Refrigerated (2 – 8 °C)		
IL-6 Capture Antibody	1 vial	Captures IL-6 on solid-phase	After reconstitution,		
IL-6 Detection Antibody	1 vial	Binds captured IL-6, biotin conjugated	standard must be aliquoted and stored at ≤ -20 °C.		
SAv-HRP	1 vial	Binds detection antibody, interacts with substrate to yield chemifluorescence signal. 1:150 diluted with OptiBlock™ to make working solution	Avoid repeated freeze-thaw cycles for standard.		

Table 1. Materials Provided with the OptiMax[™] Evaluation Kit*

*Material Safety Data Sheets (MSDS) are available on the Siloam Biosciences' web site. (http://www.siloambio.com)

Materials Required for Testing but Not Supplied With OptiMax[™] Evaluation Kit:

- 1. Eppendorf or similar tubes for centrifugation and dilutions
- 2. Kimwipes[™] or other laboratory tissue paper
- 3. Reagent reservoirs (V-shape reservoir)
- 4. Pipet tips for delivering in the ranges of 1 -10, 10 -100, and 100 1000 μL

Equipment Required:

- 1. Pipette capable of accurately and precisely delivering liquids in the ranges of 1 -10, 10 -100, and 100 1000 μ L
- 2. Multichannel pipette capable of accurately and precisely delivering 5 μ L
- 3. Multichannel pipette capable of delivery of 30 μ L
- 4. Vortex mixer
- 5. Fluorescence plate reader and control software
- 6. Analytical software
- 7. Microcentrifuge
- 8. Timer

UNIQUE CONSIDERATIONS FOR OPTIMISER™ MICROPLATE

Optimiser™ Microplate and Assembly:



Figure 2. Optimiser™ microplate assembly

Position absorbent pad on holder, align the Optimiser™ microplate and press down gently to click-lock the plate in holder



The pad must be oriented correctly with the smooth surface (tape side) facing the holder and absorbent surface touching the microplate



THE USE OF PROPER PIPETTING TECHNIQUE IS CRITICAL TO AVOID AIR-BUBBLES.

Air bubbles will occlude the microfluidic channel and stop the flow of the Optimiser™.

Optimiser™ Microplate Pipetting Instruction:

Tutorial 1 provides hands-on training for first time users to practice pipetting with Optimiser[™]. Please read the entire Pipetting Instruction section before attempting Tutorial 1.

Avoiding Bubbles While Pipetting:

- 1. Bubbles will compromise the performance of assays on Optimiser[™] by interfering with the flow of liquid within the microchannels.
- 2. OptiBlock[™] reagent may form bubbles readily with standard pipetting techniques.
- 3. To avoid complications due to bubbles, Siloam Biosciences recommends the use of the "Reverse Pipetting" technique during all pipetting steps.
 - a. To aspirate liquid, press the operating button of the pipette to the second stop (refer to illustration below).
 - b. Immerse the pipette tip in the liquid to a depth of about 2 mm and steadily release the operating button completely.
 - c. Withdraw the tip from the liquid, touching it against the edge of the reservoir to remove excess liquid.
 - d. Dispense the liquid into the loading well of Optimiser[™] microplate by gently and steadily pressing the pipette's operating button to the first stop. Briefly hold the operating button in this position.
 - e. With the button in this position, move the tip from the loading well to the reagent reservoir, immerse the tip in the liquid and aspirate.

		Pipetti	ng step	
Ready position	1	2	3	4
First stop			Ļ	1
Second Stop	ł			-

Figure 3. Reverse Pipetting procedure

THE USE OF PROPER PIPETTING TECHNIQUE IS CRITICAL TO AVOID AIR-BUBBLES.

Accurate and Precise Delivery of 5 µL Volumes:

Assays on Optimiser^M require the accurate and precise delivery of 5 μ L volumes. The following guidance is offered to users.

- 1. Use pipette for which the upper limit of their operating range is $\leq 10 \mu L$.
- 2. Use pipette tips appropriate for 5 μ L pipetting.
- 3. To aspirate liquid, hold the pipette near vertical and immerse the pipette tip in the liquid to a depth of approximately 2 mm in the liquid. Withdraw the operating button steadily. Wait ~ 1 second. Withdraw the tip from the liquid.
- To dispense liquid, hold the pipette nearly vertical. With the pipette tips <u>touching the</u> <u>surface of the Optimiser™ well</u>, depress the operating button steadily until the liquid is dispensed.
- 5. <u>Note:</u> The pipette tip must make contact with the well surface for proper dispensing (see "RIGHT" frame below). Do not pipet directly into the hole at the bottom of the well (see "WRONG" frame).



Figure 4. Pipette tip positioning for dispensing in the Optimiser™

Additional Technical Considerations:

- 1. The Optimiser[™] system has been qualified with aqueous liquids only. Do not use solvent-containing samples.
- 2. The buffer reagents provided with the assay kit have been developed and validated for the Optimiser[™] microplate. **Do not substitute alternate buffers or reagents.**
- 3. The presence of particulates in liquids dispensed to Optimiser[™] wells may block liquid flow through the microchannels.
 - a. Centrifuge serum samples and serum-containing tissue culture supernates for 10 minutes at 13,000 rpm prior to testing.
- 4. Small flow rate variations (time to empty well) do not affect assay results.

Using Electronic Multi-channel Pipette:

An electronic multi-channel pipette is ideally suited for use with Optimiser[™] microplates since (a) it *eliminates* possibility of injecting bubbles and (b) can be used for convenient repetitive loads with single aspiration step for rapid reagent transfers.

General setup for using an electronic multi-channel pipette:

- Select pipette capable of delivery 5 μ L & 30 μ L (e.g., with volume range of 5-120 μ L).
- Choose "<u>Reverse Pipetting</u>" in function setting.
- Use "<u>Multiple Dispensing</u>" mode to transfer the solution into the Optimiser[™] microplate. For example, to transfer capture antibody solution in to a full Optimiser[™] microplate, set the program for 12 times dispensing, 5 μL per dispensing. Then the pipette will automatically aspirate 60 μL of solution and dispense 5 μL volumes 12 times. Users will not need to move pipette back and forth to transfer solution.



Multichannel pipette must be used for transferring solution into the Optimiser™ plate.



If the pipette tip is pushed inside the through-hole, the tip may cause the sealing tape at the base of the Optimiser™ to delaminate and lead to flow failure



If the pipette tip does not touch the surface of well, the solution may stick on the pipette tip end and not dispensed into the well OR may lead to air-bubbles.



Small variations in flow rates (time to empty well) do not affect assay performance. The incubation step smoothes out any flow variation differences.



An electronic multichannel pipette can allow for loading all reagents with a single aspiration step – Ideally suited for processing multiple Optimiser™ microplates in parallel

Almost all pipetting protocols specify users NOT to touch the well surface during pipetting. Why does the Optimiser[™] user guide suggest the exact opposite?

In conventional 96-well ELISA plates, if the pipette tip touches the bottom surface of the well, it may physically disrupt some of the bound bio-molecules. In the Optimiser[™] all the assay reactions occur *within the microchannel*. Hence, touching the pipette tip on the loading well of the Optimiser[™] has absolutely no effect on the assay performance.

For most dispensing steps in Optimiser[™] based assays, users are dispensing only 5 µl volumes. If the pipette tip does NOT touch the well surface, the dispensed well volume may "bead" and stick to the end of the tip. The well geometry of the Optimiser[™] is engineered to ensure smooth filling of well/microchannel provided the liquid is dispensed steadily and directly on the well surface.

See the Optimiser[™] Technology page on Siloam's website for instructional videos on pipetting techniques.

Why must all materials be transferred to the Optimiser[™] plate within one minute at each step in the assay procedure?

Optimiser™ incubation steps are from 10 to 20 minutes in length. Longer time to transfer material will cause time difference between each well in incubation, which may affect the assay accuracy.

I don't have a multichannel pipette - can I try the kit with a single channel pipette?

A multichannel pipette is essential to ensure that all dispense steps can be comfortably completed in 1 minute or less. With a single channel pipette it is very difficult to complete pipetting to even 3 columns in 1 minute.

How critical is the accuracy of 5 μl dispense volume?

The Optimiser^M is designed such that the 5 µl volume represents a slight excess compared to the microchannel internal volume. Provided that the dispense volume is greater than 4.5 µl, slight (even up to 10%) dispense volume variations will not affect assay results.

Why has the recommended operating volume been changed to 5 μ L? I remember seeing 10 μ L as recommended volume in earlier version of the FAQ.

- Minimizing the volume helps with improving the precision. When using the 10 μl protocol, there is higher variation in the "time to empty" for different wells on each plate. This is related to the flow rate of the microchannel and larger volume show more net effect on flow duration (and variation of the duration).
- 2) The new 5 μl protocol also reduces the incidences of "slow" or "stopped" flow. With proper pipetting technique and by use of the new protocol, our lab tests show that flow failure rate (well does not empty after 10 minutes) is now less than ~ 0.2%.
- 3) We have verified through extensive assay tests that change from 10 µl to 5 µl does not affect the assay sensitivity. This is partly owing to improvements made to the OptiMax[™] buffer formulations.

READER SETUP:

Optimiser[™] based assays are compatible with standard fluorescence plate readers and multi-mode plate readers with fluorescence reading capability. Below is the general guidance for setting up the readers. For further assistance, please contact Siloam's technical support.

Step 1: Selecting the wavelength for excitation and emission light:



Figure 5. Normalized absorption (left) and emission (right) spectra of OptiGlow[™] chemifluorescent substrate. Assays on Optimiser[™] uses OptiGlow[™] substrate, which can be detected using the appropriate excitation and emission settings (Figure 5). Quantitation does not require filters that precisely match the excitation/emission maxima. However, a non-overlapping filter set with a bandpass that includes the excitation/emission spectra is required. Wavelengths at 530-575 nm for excitation and 585-630 nm for emission can be used for detection. Below are examples for different types of readers:

- **Filter-based readers**: install 528/20 nm (or similar) filter for excitation, and 590/35 nm (or similar) filter for emission
- **Monochromator-based readers**: in wavelength setting, set excitation at 528/20 nm, and emission at 590/35 nm

• **Readers with pre-configured optical set:** select the wavelength setting for Rhodamine or Cy3.

Step 2: Selecting the plate type:

Optimiser[™] microplate fits 96-well SBS standard in all specifications. Please use "96-well standard" or similar in plate type setting.

Step 3: Selecting the probe direction:

Please use "top reading" for probe direction.

Step 4: Selecting the sensitivity/gain:

When defining reading parameters for fluorescence analysis, setting the PMT sensitivity (or "gain" in some types of fluorescence reader) is important for obtaining useful measurements. A manual sensitivity/gain setting is recommended for reading Optimiser™ microplates. The procedure is as described below:

- In a clean plastic tube, add 50 µL of OptiGlow[™] A, 50 µL of OptiGlow[™] B, 1 µL of OptiGlow[™] C, and 1µL of supplied SAv-HRP stock solution, mix well, and wait for 2 minutes. The substrate will be fully developed and stable for hours.
- Load 4 µL of mixture into well A1 of Optimiser[™] microplate (#1)¹ and wait until the well is empty (do not use pad/holder)
- 3) Read that well in reader with various gain setting.
- 4) Select the gain which gives the RFU reading closest to 11,000.
- 5) Use the same gain setting, read one blank well of Optimiser[™], the readout should be less than 50.
- 6) Save or record this gain setting.
- 7) This defines the max reading (RFU_{max}) that Optimiser[™] based assays can reach with this reader gain/sensitivity setting and the 50:50:1 substrate mix ratio. Note that in the trial IL-6 assay, RFU readings will be higher than 11,000 RFU owing to different substrate mix ratio.

The gain setting will be valid for all Optimiser[™] based assays. Repeat Step 4 if a) changing the reader or b) changing the optical unit such as light bulb, filters, etc.

The "Technical Support" section on Siloam's website offers detailed guidance on set up of several major brand instruments as illustrative examples.

¹: The protocols refer to the Optimiser[™] microplates with a numeric designation (#1 through #5) to identify plates that are re-used in certain experiments. Optimiser[™] #1 - #3 are required for completing the tutorials and the assay transfer protocol. Optimiser[™] #4 and #5 is provided as extra plates (if required for pipetting practice). **The 5 Optimiser[™] microplates are identical and are not labeled with numeric designator.**

TUTORIAL 1: PIPETTING TO THE OPTIMISER™ MICROPLATE:

Materials Required for Pipetting Tutorial and Supplied with Optimiser[™] Starter kit:

- 1. One Optimiser[™] holder
- 2. Same Optimiser[™] (microplate #1) used for reader setup, 3 (or more) unused columns will be used for pipetting
- 3. One new Optimiser[™] (microplate #2), <u>all 12 columns</u> will be used for pipetting
- 4. One Optimiser[™] pad, single use
- 5. OptiWash[™] buffer
- 6. Green dye solution
- 6. Red dye solution

Other Materials/equipment Required:

- 1. Kimwipes[™] or other laboratory tissue paper
- 2. Reagent reservoirs (V-shape reservoir)
- 3. Single channel pipette capable of delivering in the ranges of $100 1000 \,\mu$ L
- 4. Pipette tips for delivering in the ranges of $100 1000 \mu$ L
- 5. Multichannel pipette capable of accurately and precisely delivering 5 μ L
- 6. Multichannel pipette capable of delivery of $30 \ \mu L$

Procedure with Manual Multi-channel Pipette:

- Assemble the Optimiser[™] Microplate, Optimiser[™] Pad, and Optimiser[™] Microplate Holder as described on Page 4.
- 2. Transfer 1.5 mL of green dye solution into a V-shape reagent reservoir. Transfer 1.5 mL of red dye solution into another V-shape reagent reservoir. Transfer 3 mL of OptiWash[™] solution into a V-shape reagent reservoir.
- To aspirate liquid, hold the multi-channel pipette nearly vertical and immerse the pipette tip in the liquid to a depth of approximately 2 mm in the liquid. Withdraw the operating button steadily. Wait ~ 1 second. Withdraw the tip from the liquid.
- 4. Dispense into columns 2-4 (3-columns) for Optimiser[™] microplate #1 per the sequence illustrated below.
 - a. To dispense liquid, hold the multi-channel pipette nearly vertical. With ALL the pipette tips <u>touching the</u> <u>surface of the Optimiser™ wells</u>, depress the operating button steadily until the liquid is dispensed.
 DO NOT position pipette tips into the hole at the bottom of the surface.

5 µl Green dye;	OBSERVATIONS AND CONCLUSIONS
wait for 10 min. ↓ 5 μl OptiWash™; wait for 10 min.	 In each step, all wells should be empty within 10 minutes. If a well is not empty after 10 minutes, please inspect under low-power microscope and most likely a bubble will be evident near microchannel interface with well. This bubble was accidentally injected due to incorrect pipetting technique. Please refer to the pipetting guidelines and try again.
5 μl Red dye; wait for 10 min.	 Note that as the dye reagent is changed in the well; even a 5 μL volume will "clear" the previous reagent in the microchannel. This demonstrates the efficiency of the "flushing" action instead of the traditional wash step.
5 μl OptiWash™; wait for 10 min.	• Observe the wells as they drain out. Note the variation in time to empty each well. So far as each well drains out in 10 minutes, this variation has NO EFFECT ON ASSAY PERFORMANCE.
↓ 5 μl Red dye; wait for 20 min. ↓ 30 μl OptiWash™; wait for 10 min.	 Most assay protocols on Optimiser[™] recommend a 10 minute incubation interval (typically 20 minutes for sample/standard). The 20 minute incubation step with red dye shows that ALL incubations can be extended up to 20 minutes. This may be useful for processing multiple Optimiser[™] microplates in parallel. Incubation steps should be at least 5 minutes and no more than 30 minutes. Use at least 20 minutes incubation for sample/standard.

- 5. Assemble a new microplate (#2) and pad on the holder.
- 6. Repeat the dispensing protocol shown above again in <u>ALL 12 Columns of microplate#2</u>. Time the dispensing cycles and check that all dispensing steps are completed within 1 minute.
- 7. CHECK THAT ALL WELLS DRAIN WITHIN 10 MINUTES FOR EACH DISPENSING STEP.
- 8. If any wells take longer than 10 minutes the most likely cause is an error in pipetting causing a visually evident or micro-bubble. Please refer to the pipetting instructions and repeat steps 6 and 7. User MUST complete dispensing protocol on to entire Optimiser[™] plate with all wells on that microplate draining in 10 minutes (for all steps in protocol).



Optimiser[™] microplates are a powerful tool for ELISA's and require correct pipetting procedures to ensure repeatable results.

USERS MUST BE ABLE TO COMPLETE_THE DISPENSING PROTOCOLS FOR A <u>COMPLETE</u> <u>Optimiser™ MICROPLATE</u> WITH ALL WELLS DRAINING IN LESS THAN 10 MINUTES (FOR ALL STEPS OF THE PROTOCOL).

THIS SIMPLE STEP IS CRITICAL TO ENSURE USERS CAN ACHIEVE EXCELLENT ASSAY RESULTS ON OPTIMISER™.

Two additional Optimiser[™] microplates are included with the package to allow users to further practice and perfect pipetting to the Optimiser[™]. Excess volumes of the red/green dye and OptiWash[™] are also included.

(OPTIONAL) Procedure with Electronic Multi-channel Pipette (5-100 µL volume):

- 1. Repeat the protocol described for Manual multi-channel pipette with an Electronic multi-channel pipette.
- 2. Choose "<u>Reverse Pipetting</u>" in function settings for Electronic pipette.
- 3. Choose "<u>Multiple Dispensing</u>" mode, and program for 12 dispense cycles with 5 μl dispense volume per cycle (for dispensing to all 12 columns).

OBSERVATIONS:

- All observations and conclusions listed for previous protocol.
- Note the difference in time required to load 12 columns with an electronic multi-channel pipette.

If one well drains in (say) 1 minute and another in (say) 8 minutes, how is it possible that they provide comparable results?

Although it may seem that difference of minutes may have an impact on the assay precision, Siloam has demonstrated with multiple assays that <u>well optimized assays on Optimiser™ easily achieve CV < 6-10%</u>. The minimal effect of flow rate on precision is a combination of multiple factors:

- On the micro-scale reaction kinetics are vastly different compared to the macro-scale kinetics of conventional 96-well ELISA plate. In microfluidic channels, most surface binding reactions are saturated in ~ 5 minutes. Optimiser[™] characterization data shows that up to ~ 75% of peak adsorption is completed in only **10 seconds** and assay binding reactions saturate in ~ 5 minute. This is a result of two factors:
 - a. The diffusion distances in the microchannel are extremely small (the channel has a cross-section of only 200 μ m x 200 μ m) hence diffusion is no longer a limiting factor
 - b. The surface area of the microchannel is ~ 1.5 times the surface area at the base of a conventional 96-well ELISA plate. The volume contained in the microchannel is ~ 5 μ l leading to ~ 50x higher surface area to volume ratio which allows for extremely efficient binding reactions.
- Even for the well that drains in 8 minute, the initial section of the microchannel (towards the center) is filled up in ~ 2-3 minutes. Optimiser™ characterization data shows that the first few loops of the microchannel contribute ~ 95% of the optical signal hence even if the last 1-2 loops take significantly longer to fill, their contribution to the signal is almost negligible. Consequently, variations in signal from the last loops have little impact on overall (assay) signal variation.
- 3. For most reaction steps in the assay sequence (except for sample/standard loading step) the biomolecules are present in vast abundance and the binding reactions are completed extremely quickly. To ensure good precision, it <u>IS</u> recommended that the sample/standard incubation should be 20 min.
- 4. Finally, the incubation interval (when there is no liquid left in the well) "smooths out" the effect of flow rate variances.

How does the variance (CV) of Optimiser[™] microplates compare to conventional plates?

In most assays (conventional plates), raw signal variance for triplicates is <10% which is also true for Optimiser[™] microplates. Please see Siloam's website for a Technical Note detailing variance studies on the Optimiser[™] platform. For **first-time users**, it is common to see variances at ~ 15% and even up to 20%. In almost all cases, this is related to pipetting techniques and as any other platform "practice makes perfect" -- most users see noticeable improvement after running a few Optimiser[™] plates. The most common pipetting related issues that are resolved with careful attention to details and practice include:

- 1. Tips do not touch well surface leading to incomplete dispensing. This can lead to very high signal (e.g. if block buffer is not properly dispensed) or very low signals (e.g. if SAv-HRP is not properly dispensed).
- 2. Inconsistent load times as users learn pipetting procedures. With significant differences between loading intervals across the plate there is higher variance with practice users can establish a steady "rhythm" with improved precision.
- 3. Failure to change tips in reagent preparations and/or between dispensing steps for assay sequence.
- 4. Use of inappropriate pipette tips and/or pipettes

Most users after completing ~ 10-12 Optimiser[™] based assays can achieve (a) background signal <2-3% of peak signal (see Page 15) and (b) variance <10% for all points on standard curve. These can be used as metrics to determine user proficiency on the Optimiser[™] system.

TUTORIAL 2: IL-6 DEMONSTRATION ASSAY ON THE OPTIMISER™:

The Optimiser[™] starter kit also contains necessary reagents of an IL-6 sandwich ELISA Assay to demonstrate the capabilities of Optimiser[™] based assays under controlled conditions by a user. The representation of the expected data produced from **Optimiser[™] starter kit is not intended to be used as a routine use commercial assay kit**. Siloam offers an IL-6 assay kit under catalog# OMA-H-IL6; please contact customer support to order.

Materials Required for Demonstration Assay and Supplied with Optimiser™ Starter kit:

- 1. One Optimiser[™] holder
- 2. Optimiser[™] microplate (#3)
- 3. One new Optimiser[™] pad, single use
- 4. One 96-well v-bottom plate
- 5. OptiBind[™]-H buffer
- 6. OptiBlock[™] buffer
- 7. OptiWash[™] buffer
- 8. OptiGlow[™] substrate kit, contains component A, B and C
- 9. IL-6 capture antibody
- 10. Lyophilized IL-6 standard
- 11. IL-6 detection antibody, biotinylated
- 12. Streptavidin HRP

Materials Required for Assay But Not Supplied with Optimiser[™] Starter kit:

- 1. Eppendorf or similar tubes for centrifugation and dilutions
- 2. Kimwipes[™] or other laboratory tissue paper
- 3. Reagent reservoirs (V-shape reservoir)
- 4. Pipet tips for delivering in the ranges of 1 -10, 10 -100, and 100 1000 μL

Equipment Required:

- 1. Pipettes capable of accurately and precisely delivering liquids in the ranges of 1 -10, 10 100, and 100 1000 μL
- 2. Multichannel pipette capable of accurately and precisely delivering 5 μ L
- 3. Multichannel pipette capable of delivery of 30 μ L
- 4. Vortex mixer
- 5. Microplate fluorescence reader and control software
- 6. Analytical software

Assay Layout:

The plate layout of IL-6 standard concentration is shown in below. Each concentration will repeat 3 times. 3 columns of one Optimiser[™] microplate will be used.

 Table 2. Plate layout of IL-6 concentration (pg/mL) for demonstration assay

	1	2	3
Α	3000	3000	3000
В	1000	1000	1000
С	333.3	333.3	333.3
D	111.1	111.1	111.1
E	37.0	37.0	37.0
F	12.3	12.3	12.3
G	4.1	4.1	4.1
Н	0	0	0



Very small volumes of assay reagents are required and provided for Optimiser[™] based assays.

A quick-spin (minicentrifuge) is **CRUCIAL** to recover all material in Items 9-12.

Use a spin **each time** the assay reagents are to be used.

Reagent Preparation:

The incubation times for Optimiser[™] are only 10-20 minutes. Preparing all the reagents, samples, standards in advance will allow for proper timing (especially for first time users).

Always prepare extra volume of solution for easy transferring. Siloam suggest to prepare 30 µL extra volume each well in 96-well v-bottom plate. This volume can be reduced with careful pipetting if sample is very limited or precious.

Bring all reagents to room temperature before use and prepare all necessary dilutions before beginning the test procedure.

- 1. <u>OptiBind[™]</u>: OptiBind[™]-H is provided in a ready-to-use form. No further preparation is required. **Do not** substitute other coating buffers for OptiBind[™]-H.
- 2. <u>Capture Antibody</u>: The procedure requires 5 μL of capture antibody working solution for each assay well to be used.
 - a. Prepare a 1:62.5 dilution of the capture antibody stock in OptiBind[™] buffer in a clean plastic tube (Add 8 µl of capture antibody stock solution to 0.5 mL of OptiBind[™]).
 - b. Dispense 60 μ L of the working solution into each well of a single column in the polypropylene 96-well v-bottom plate.
- 3. <u>OptiBlock[™]</u>: OptiBlock[™] is provided in ready-to-use form and is used to block the surfaces of the Optimiser[™]'s microfluidic reaction chambers following their incubation with the capture antibody solution. OptiBlock[™] is also used as the diluent for the standard, detection antibody and SAv-HRP in this experiment.
- 4. <u>Recombinant IL-6 Standard:</u>
 - a. <u>Stock Solution:</u> The IL-6 standard is provided in lyophilized form.
 - i. Reconstitute the lyophilized standard by adding 420 µL OptiBlock™ blocking buffer.
 - ii. Mix by gentle swirling until all of the lyophilized material has dissolved.
 - iii. Vortex gently to ensure thorough mixing of the reconstituted standard.
 - iv. Use freshly prepared material on the day of reconstitution
 - b. <u>Working Solution</u>: The concentration of the reconstituted IL-6 standard is 4 ng/mL. Prepare a 3000 pg/mL standard (Standard 1) by mixing 120 μL IL-6 standard appropriately with 40 μL OptiBlock™ blocking buffer. Vortex the 3000 pg/mL standard briefly to mix.
 - c. <u>Standard Curve</u>: Prepare the remaining IL-6 standards by performing six serial three-fold dilutions in OptiBlock[™] beginning with the 3000 pg/mL standard as follows:
 - i. Dispense 120 μL of Standard 1 (3000 pg/mL) to well A1 of the 96-well polypropylene v-bottom plate.
 - ii. Dispense 80 µL OptiBlock[™] to each of the seven wells of the same column immediately below the 3000 pg/mL-containing well (wells B1 H1).
 - iii. Transfer 40 μ L of the 3000 pg/mL standard from well A1 to well B1 immediately below it. Mix the contents of well B1 gently. Then transfer 40 μ L from well B1 to well C1, **change tips**, and titrate.
 - iv. Continue serial dilutions **while changing tips after each 40 μL transfer** and before mixing until the 4.1 pg/mL standard has been created in the seventh well (well G1) of the column.
 - v. Do not transfer IL-6 solution to the eighth well (H1). It contains OptiBlock[™] only and will provide material for the blank wells.

	1			1
Α	120 µL Std 1		Α	3000
В	80 μL Blocking		В	1000
С	80 μL Blocking		∖ c	333.3
D	80 μL Blocking		D	111.1
E	80 μL Blocking		E	37.0
F	80 μL Blocking		F	12.3
G	80 μL Blocking	ECO	G	4.1
Н	80 μL Blocking		Н	0

- 5. <u>Detection Antibody</u>: The procedure requires 5 μL of the detection antibody working solution for each assay well to be used.
 - a. Prepare a 1:25 dilution of the detection antibody stock in OptiBlock[™] in a clean plastic tube (add 20 µL of detection antibody stock solution to 480 µL of OptiBlock[™]).
 - b. Dispense 60 μ L of the working solution into each well of a single column in the polypropylene 96-well v-bottom plate.
- 6. <u>SAv-HRP</u>: The procedure requires 5 μL of the SAv-HRP working solution for each assay well to be used.
 - a. Prepare a 1:150 dilution of the detection antibody stock in OptiBlock[™] in a clean plastic tube (add 4 µL of SAv-HRP stock solution to 0.6 mL of OptiBlock[™]).
 - b. Dispense 60 μ L of the working solution into each well of a single column in the polypropylene 96-well v-bottom plate.
- 7. <u>Substrate solution</u>: The procedure requires **10 μL** of the working substrate solution for each assay well to be used.
 - a. Prepare the working substrate solution no more than 30 minutes before the anticipated time for reading the completed assay.
 - b. To create the substrate working solution, combine OptiGlow[™]-A, OptiGlow[™]-B, and OptiGlow[™]-C in a ratio of **50:50:5** parts respectively in a clean plastic tube and vortex gently to mix (add 250 µl of OptiGlow[™]-A, 250 µl of OptiGlow[™]-B, and 25 µl of OptiGlow[™]-C).
 *OptiGlow[™]-C must be thoroughly thawed to function effectively. Warm the reagent in a 37° C incubator/oven/heater or by holding the vial gently in your hands.
 - c. Dispense 60 μ L of the working solution into each well of a single column in the polypropylene 96-well v-bottom plate.
- 8. <u>OptiWash™</u>: OptiWash™ is provided in ready-to-use form. No further preparation is required. The procedure requires 75 μL of OptiWash™ for each assay well to be used. Dispense 4 mL OptiWash™ buffer to a v-shaped reagent reservoir and use for all wash steps in the assay.

DO NOT SUBSTITUTE OTHER BUFFERS OR REAGENTS FOR THOSE PROVIDED WITH THE KIT.

OptiMax[™] buffers are specially formulated to work with the Optimiser[™] microplate and substitute buffers or reagents may lead to poor assay performance.

ENSURE THAT TIP CHANGES AS RECOMMENDED FOR STANDARD PREPARATION ARE FOLLOWED. Continued use of same tip may lead to errors in dilution and consequent assay signals.

Procedure:

- 1. Assemble the Optimiser[™] Microplate, Optimiser[™] Pad, and Optimiser[™] Microplate Holder as described on Page 4.
- 2. <u>Hint:</u> Optimiser[™] incubation steps are from 10 to 20 minutes in length. To achieve optimal assay performance, all materials must be transferred to the Optimiser[™] microplate within one minute at each step. To accomplish this, first place the materials to be transferred in the enclosed 96-well polypropylene v-bottom plate or v-shape reagent reservoir (*instructed in Reagent Preparation, page 12*). Then transfer the materials to the Optimiser[™] wells using a <u>multi-channel pipette</u>.
- 3. Dispense 5 µL capture antibody working solution to the required number of wells in the Optimiser[™] microplate. Incubate 10 minutes at room temperature (RT).
- 4. Dispense 5 µL OptiWash[™] to each well. Wait 10 minutes to proceed to the next step.
- 5. Dispense 5 μL OptiBlock[™] to the capture antibody-coated wells. Incubate 10 minutes at RT.
- 6. Dispense 5 μ L of the standard and blank to the required number of replicate wells of the plate. Incubate **20 minutes at RT**.
- 7. Dispense 5 µL OptiWash[™] to each well. Wait 10 minutes to proceed to the next step.
- 8. Dispense 5 μ L detection antibody working solution to each well. Incubate 10 minutes at RT.
- 9. Dispense 5 μ L OptiWashTM to each well. Wait 10 minutes to proceed to the next step.
- 10. Dispense 5 μL SAv-HRP to each well. Incubate 10 minutes at RT.
- 11. Dispense 30 μ L OptiWashTM to each well. Wait 10 minutes to proceed to the next step.
- 12. Again dispense 30 µL OptiWash[™] to each well. Wait 10 minutes to proceed to the next step.
- 13. Dispense **10 µL** OptiGlow[™] working solution to each well. Incubate for 15 minutes at RT.
 - a. <u>Caution</u>: Observe the wells <u>during the incubation</u>. When the substrate has completely drained from all wells, remove the plate and pad from the holder. Discard the pad. Wipe the bottom of the plate with a Kimwipe[™] to remove any liquid on the bottom surface of the plate. Step 13a will be completed within the 15 minute substrate incubation time.
- 14. Place the plate in the reading chamber of a fluorescence plate reader. Promptly at the conclusion of the 15 minute incubation, read the plate.

If all the assay reagent preparation steps and protocol are followed correctly, the "wells" (microfluidic channels) corresponding to top 2 (up to top 3) standards clearly appear pink (owing to developed substrate). If the pink color is not evident even for topmost standard, one or more reagent preparation steps or assay steps was not performed correctly.



It is common to see slight differences in the time required for different wells to empty. This difference has no impact on assay performance.



To facilitate work flow, incubations designated as 10 minutes may be extended to 20 minutes with no impact on method performance.



Optimiser™ "washes" are performed by simply <u>dispensing</u> OptiWash™ to the wells.



Wipe the plate bottom thoroughly. Any liquid residue on the bottom surface will cause false positive signal.



In rare cases (<0.2%), a well may not empty in 10 min. If so, blot the reagent from the well with a tissue. **Do not include data from this** well in calculations.

Calculations:

- 1. Calculate the mean background signal from the blank wells (wells containing OptiBlock[™] only at the sample incubation step).
- 2. Subtract the mean background signal from the signal of individual standard.
- 3. Create a standard curve by plotting the standard concentration (x-axis) vs the background-adjusted signal (y-axis). A five parameter logistic curve fit with appropriate software is recommended.

Typical Data:

The IL-6 standard curve ranges from **4.1 to 3000 pg/mL**. Concentration (x-axis) and signal (y-axis) are plotted on Log scales. A typical standard curve is presented below.



IL-6 (pg/ml)	Average	Blank- Subtracted
3000	36665	36506
1000	31722	31564
333.3	15764	15606
111.1	6254	6096
37.0	2572	2414
12.3	1041	883
4.1	514	356
0	158	

Figure 6. IL-6 Standard Curve with Tabulated Data

SIGNIFICANCE OF ASSAY BACKGROUND

The reader setup in this example sets a value of ~ 11,000 RFU as the high value for the 50:50:1 "saturated" substrate signal. *Regardless of the substrate ratio used*, the background RFU readings (blank signal) should not exceed ~ 350 RFU (~3% of max value in reader setup).

- Background signals higher than 3% of RFU_{max} (established during reader setup) indicate that one or more steps of the assay was performed incorrectly and users should repeat the assay.
- Background signals higher than ~ 6% of RFU_{max} (established during reader setup); corresponding to ~ 700 RFU in current example, indicate a failure and the assay <u>must</u> be repeated.

CAUSES:

- High backgrounds are most commonly a result of pipetting errors and can be resolved with careful attention to the procedure and additional practice.
- Another common cause for high background is use of alternate SAv-HRP or direct HRP labeled detection antibodies. Optimiser[™] based assays are exquisitely sensitive to HRP concentration and the SAv-HRP provided by Siloam has been carefully optimized to achieve best performance. Use of alternate buffers (especially blocking buffers) can also lead to high background signals.

Please consult with Siloam's Tech Support team before substituting any of the buffers provided with the starter, evaluation kits or as part of the OptiMax[™] assay buffer reagent sets.

TUTORIAL 2: TARGETED OUTCOME

First time users can run a complete assay on the Optimiser^M and confirm that they can generate similar data as listed in the User Manual. This Tutorial is intended to serve two purposes: (a) to familiarize users with the assay operation sequence on the Optimiser^M and ensure performance matches with Siloam's data and (b) to provide users an introduction to the capabilities of the Optimiser^M to deliver high-sensitivity assay data even when using only 5 µl sample volumes and a 2 hour assay protocol.

Assay Background

SECTION II:

Assay Transfer Guide

PROCEDURE FOR MIGRATING A VALIDATED ASSAY FROM CONVENTIONAL 96-WELL MICROPLATE TO OPTIMISER™

SANDWICH ELISA ASSAY TRANSFER GUIDE:

The OptiMax[™] ELISA procedure is a chemifluorescent immunoassay in which traditional ELISA reactions take place within the unique Optimiser[™] microplate architecture. Briefly, capture antibody is immobilized on the internal surfaces of the plate's microchannels. Following a wash step, any unreacted sites on the microchannel surface are blocked with a blocking solution. Standards and samples are dispensed to the Optimiser[™] wells. Antigen present in samples and standards will be specifically captured on the microchannel surface by the immobilized capture antibody. Following another wash, a biotin-labeled detection antibody is added to the wells. The biotin-labeled antibody will bind antigen that has been captured and immobilized on the microchannel surface thus "sandwiching" the antigen between the capture and detection antibodies. Following another wash, horseradish peroxidase-labeled streptavidin (SAv-HRP) is added to the Optimiser[™] wells. The streptavidin of SAv-HRP binds specifically to the biotin moiety of the biotin-labeled antibody if it is present in the [capture antibody+antigen+detection antibody] complexes formed and immobilized on the microchannel surface. Following two additional washes, a chemifluorescent substrate is added to the wells. If horseradish peroxidase has been captured on the microchannel surface during the sequence of reactions cited above, the enzyme will react with the substrate solution and will yield a fluorescent signal when excited at the appropriate wavelength. Within the linear portion of the curve, the light signal emitted will be directly proportional to the concentration of antigen in standards and samples, and will be quantifiable when the plate is read using a fluorescence plate reader.

In order to achieve best assay performance with Optimiser[™] platform, serial optimization tests for each type of assay need to be performed before measuring the real sample. A well-characterized and robust assay on 96-well platform is a mandatory pre-requisite for the Assay Transfer Process. The assay transfer is a 3-step process with step 1 being data collection for conventional assay. Second, run one experiment to screen 12 types of supplied OptiBind[™] coating buffers to determine the best coating buffer to use for this assay. Third, run a checkerboard titration experiment to determine the concentration of capture antibody and detection antibody which gives the best signal/noise ratio. Finally, run an assay with wide range of target protein concentration with the selected optimal coating buffer and antibodies concentrations to determine the measurable/dynamic range of the assay.



Figure 7. Schematic assay optimization procedure with Optimiser[™] microplate.

Requirements for Conventional Assay to be Transferred to Optimiser™

For transfer to Optimiser[™], a robust conventional (96-well plate) assay is expected with the following **minimum performance metrics:**

- Reasonable background (zero) reading: for an colorimetric assay, the OD450/630 absorbance reading of background shall be less than 0.15
- **Reasonable dose response** with various concentrations of standard: for an absorbance assay, the OD450/630 absorbance reading of highest concentration in detectable range shall be higher than 1.5

In addition to satisfying the minimum performance metrics, the following information is required for the assay transfer process:

- Known concentrations or dilution ratio for capture antibody and detection antibody working solution. The capture and detection antibody solutions must be in a format such that at least 4x concentration solutions (as compared to working concentration for conventional 96-well plate) can be prepared.
- HRP conjugate: the detection antibody must NOT be directly labeled with HRP; a biotin labeled detection antibody is most preferred (*see next page for details*)
- Known dynamic range for assay is used as a starting point for the assay transfer process.

IT IS STRONGLY RECEOMMENDED THAT ALL ASSAY MATERIALS USED SHOULD BE TESTED RIGHT BEFORE THE ASSAY MIGRATION PROCESS TO ENSURE MATERIAL QUALITY. DO NOT USE MANUFACTURER SPECS WITHOUT A CONFIRMATORY EXPERIMENT.

As an example, a working IL-6 colorimetric assay with conventional 96-well plate is shown below. The assay transfer guide uses this assay as an example to illustrate the transfer process.



IL-6 (pg/mL)	Average	Blank-Subtracted
250	1.586	1.561
125	0.823	0.798
63	0.441	0.416
31	0.247	0.222
16	0.145	0.120
8	0.089	0.064
4	0.059	0.034
0	0.025	

Figure 8. Standard curve of IL-6 assay run in conventional 96-well plate, using TMB substrate and colorimetric detection of absorbance at 450 nm and corrected with 630 nm. 2 μ g/mL each concentrations for Capture and Detection antibody.

Details for illustrative IL-6 assay:

- Assay metrics
 - Background OD = <u>0.025</u> (less than 0.15)
 - Max signal OD = 1.58 (more than 1.5)
- Known information:
 - Concentration of capture antibody = $2 \mu g/mL$
 - Concentration of detection antibody = $2 \mu g/mL$
 - Conjugate for detection antibody: biotin conjugated
 - Assay dynamic range = $\frac{4 \text{ pg/mL} 250 \text{ pg/mL}}{250 \text{ pg/mL}}$

HRP Concentrations in Assays on Optimiser[™]:

Optimiser[™] microplates are an exquisitely sensitive platform for high-sensitivity ELISA with minimal sample/reagent volume requirements. It is **CRITICALLY IMPORTANT** to follow the guidelines for HRP conjugate to ensure that the sensitive response from Optimiser[™] is not swamped by high HRP concentrations. The use of a biotinylated detection antibody is recommended with SAv-HRP provide by Siloam (Cat# OMR-HRP) to obtain the best response.

• For biotinylated detection antibody: a vial of appropriate formulized SAv-HRP is included in the kit. Please prepare the working solution at 1:150 dilution with OptiBlock[™].

It is strongly recommended that the SAv-HRP provided by Siloam (Cat# OMR-HRP) be used for all assays on Optimiser[™]. The concentration and activity have been characterized and optimized for use with the Optimiser[™] microplate system. Use of alternate SAv-HRP may lead to low signals or very high backgrounds.

- For using HRP conjugated secondary antibody (anti-mouse, anti-goat, etc.): use secondary antibody at 0.1-0.2 μg/mL concentration.
- For using HRP directly conjugated detection antibody: Characterization studies at Siloam with multiple direct labeled antibodies confirm a high background issue in most assays, which is mostly caused by high concentration of free HRP molecule in the solution. The use of direct HRP conjugated detection antibody in Optimiser[™] based assay except is not recommended (unless the conjugated antibody is purified thoroughly to remove all free HRP). Please contact Siloam technical support for further assistance.

Unit Conversions – Comparing Absorbance and Fluorescence Assay Readings

In order to compare the assay performance between Optimiser[™] and conventional plate, conversion between fluorescence readout (RFU) and absorbance readout (OD) is required. Please follow the guidance below:



Can I use Optimiser[™] plates for all the assay types that I run on a normal 96-well plate?

The current version of Optimiser[™] has been validated for ELISA applications, such as direct, indirect, sandwich and competitive immunoassays. For other applications, please discuss your application with Siloam tech support.

What detection modes can use with the Optimiser[™]?

- The current version of Optimiser[™] is well suited for fluorescence/chemifluorescence mode detection.
- We will have a version suitable for chemiluminescence mode available in the near future.
- Absorbance mode does NOT work with the Optimiser[™].

I am not sure if the 10 minute incubation in room temperature will work – I would prefer to incubate for at least 30 minutes or in 37° C. Is that OK?

Actually – NO. We strongly recommend that incubation times should not exceed 20 minutes. Incubating beyond 30 minutes or at 37 °C, will cause evaporative losses.

How critical is the 10 minute incubation window?

- Most binding reactions on Optimiser[™] microplates saturate in ~ 5 minutes. Users can actually use even 5 minute incubation steps (except for sample which should be at least 20 minutes). The Application note section (Technical Support Tab) of Siloam's website has an article that describes this in greater detail.
- We recommend that you start with at least 10 minute incubation cycles/step but you can certainly use longer (up to 20 minutes) incubation steps. This may be useful when you are processing multiple Optimiser[™] plates in parallel.
- All incubation steps must be at least 5 minutes (at least 20 minute for sample). Incubation steps should not exceed 30 minutes.

Can I use cell lysate supernatants or other biological fluids such as serum or urine?

• The flow does work in some circumstances even with particulates in the solution. However, we have seen that the flow is not very repeatable. For these fluids, we recommend using supernatant after centrifuging at 13,000 g for 10 minutes or pass through 0.2 µm filter.

How can I improve sensitivity of my assay using Optimiser™?

- In most cases, using the assay optimization protocol described in the "Assay Transfer Guide" you should be able to achieve slightly better sensitivity.
- A guaranteed method to significantly increase sensitivity is the use of repeat load process for sample/standard steps. Please see the Application Note (under the Technical Support Tab) on Siloam's website (authored by BioTek) that describes the use of a Precision automation station to increase assay sensitivity more than 100x! Please discuss your application with Siloam tech support and we can offer more accurate guidance.

Experiment 1 – Selecting the Best Antibody Coating Buffer:

In Optimiser[™] microplate, all assay reactions occur in the microfluidic microchannel. The high surface area to volume ratio and short diffusion distances of the microchannels allow rapid protein adsorption onto the surface. Unlike the assay in conventional plate, the capture antibody adsorption in Optimiser[™] is dominated by the reaction rate of protein adsorption, which is strongly affected by the coating buffer. The first step of assay development is to screen twelve types of OptiBind[™] coating buffer provided in the kit. It requires one assay experiment and uses one full Optimiser[™] microplate.

The assay sensitivity can vary as much as 10x depending on the coat buffer used for capture antibody coating. This additional assay optimization step is critical for Optimiser™ microplates to achieve best performance. OptiBind™ COAT BUFFER IS MANDATORY FOR OPTIMISER™ ASSAYS - DO NOT USE ANY OTHER COAT BUFFER.

Reagent Preparation:

The incubation times for Optimiser[™] are only 10-20 minutes. Preparing all the reagents, samples, standards in advance will allow for proper timing (especially for first time users).

Always prepare extra volume of solution for easy transferring. Prepare ~30 μ L <u>extra</u> volume in each well of 96-well v-bottom plate. The extra volume may be reduced with careful pipetting if sample is very limited or precious.

Bring all reagents to room temperature before use and prepare all necessary dilutions before beginning the test procedure.

- 1. <u>OptiBind[™]</u>: Twelve types of OptiBind[™] are provided in ready-to-use form. No further preparation is required. **Do not substitute other coating buffers for OptiBind[™]**.
- <u>Capture Antibody</u>: Use same concentration as conventional assay, prepare the capture antibody working solution by diluting the capture antibody stock in 12 types of OptiBind[™] to make 100 μL final working solution. Dispense the capture antibody working solutions in different OptiBind[™] coat buffers into a single row in the 96-well vbottom plate (one per well).

2 μg/mL capture antibody concentration is used in the example assay (Page 19). Hence, 100 μL (each) of 2 μg/mL capture antibody working solution in each of OptiBind™-A, OptiBind™-B, OptiBind™-C..... OptiBind™-L would be prepared for this step.

- 3. <u>OptiBlock[™]:</u> For blocking step, prepare **1 mL** of OptiBlock[™] in a v-shape reservoir.
- <u>High Concentration Protein Standard</u>: Prepare **1 mL** protein standard with concentration of 80% of top standard, diluted in OptiBlock[™] in a v-shape reservoir.
 250 pg/mL of IL-6 is the top standard in the example assay. Hence, 200 pg/mL IL-6 standard would be prepared for this step.
- Low Concentration Protein Standard: Prepare 1 mL protein standard with concentration of 20% of top standard, diluted in OptiBlock[™] in a v-shape reservoir.
 250 pg/mL of IL-6 is the top standard in the example assay. Hence, 50 pg/mL IL-6 standard would be prepared for this step.
- 6. <u>Blank (zero)</u>: Prepare **1 mL** of OptiBlock[™] in a v-shape reservoir.
- 7. <u>Detection Antibody</u>: Use same concentration as conventional assay, prepare the detection antibody working solution by diluting the detection antibody stock in OptiBlock[™] to make **1mL** final working solution in a v-shape reservoir.

2 μg/mL detection antibody concentration is used in the example assay. Hence, 1 mL of 2 μg/mL detection antibody working solution in OptiBlock™ would be prepared for this step.

- <u>SAv-HRP</u>: Use SAv-HRP stock solution provided in the kit. Prepare the SAv-HRP working solution by adding 8 μL of SAv-HRP stock solution to 1.2 mL of OptiBlock[™] (1:150 dilution) in a v-shape reservoir (mix well).
- Substrate solution: The procedure requires 10 µL of the working substrate solution for each assay well to be used. Prepare the substrate working solution in a v-shape reagent reservoir by mixing 0.9 mL of OptiGlow[™]-A, 0.9 mL of OptiGlow[™]-B, and 18 µL of OptiGlow[™]-C.
- 10. <u>OptiWash™</u>: OptiWash™ is provided in ready-to-use form. No further preparation is required. The procedure requires total of 75 μL of OptiWash[™] for each assay well to be used. Prepare **10 mL** of OptiWash[™] into a v-shaped reagent reservoir and use it for all wash steps in the assay.

Assay Layout:

OptiBind™ Type	Α	В	С	D	Ε	F	G	Н	I	J	К	L
	1	2	3	4	5	6	7	8	9	10	11	12
Α		High	concer	tratior	prote	in stand	dard. co	oncent	ration a	at 80%	oftop	
В		standard used in conventional assav										
С		010				onu u	,					
D		Low	concen	tration	protei	n stanc	lard. co	ncentr	ation a	t 20% (oftop	
E		stand	dard us	ed in c	onvent	ional a	sav					
F		Starr		cume	onvene		July					
G						Zero) blanl	<)				
Н							(,				

Procedure:

- 1. Assemble the Optimiser[™] Microplate, Optimiser[™] Pad, and Optimiser[™] Microplate Holder as described on Page 5.
- 2. <u>Hint:</u> Optimiser[™] incubation steps are from 10 to 20 minutes in length. To achieve optimal assay performance, all materials must be transferred to the Optimiser[™] microplate within one minute at each step. To accomplish this, first place the materials to be transferred in the enclosed 96-well polypropylene v-bottom plate or v-shape reagent reservoir (as *instructed in Reagent Preparation, page 22*). Then transfer the materials to the Optimiser[™] wells using a <u>10 µL multi-channel pipette</u>.
- 3. Dispense 5 μL capture antibody solution to the required number of wells in the Optimiser[™] microplate. Incubate 10 minutes at room temperature (RT).
- 4. Dispense 5 µL OptiWash[™] to each well. Wait 10 minutes to proceed to the next step.
- 5. Dispense 5 μL OptiBlock[™] to the capture antibody-coated wells. Incubate 10 minutes at RT.
- 6. Dispense 5 μ L of the standard and blank to the required number of replicate wells of the plate. Incubate 20 minutes at RT.
- 7. Dispense 5 µL OptiWash[™] to each well. Wait 10 minutes to proceed to the next step.
- 8. Dispense 5 μL detection antibody working solution to each well. Incubate 10 minutes at RT.
- 9. Dispense 5 µL OptiWash[™] to each well. Wait 10 minutes to proceed to the next step.
- 10. Dispense 5 μ L SAv-HRP to each well. Incubate 10 minutes at RT.
- 11. Dispense 30 µL OptiWash[™] to each well. Wait 10 minutes to proceed to the next step.
- 12. Again dispense 30 µL OptiWash[™] to each well. Wait 10 minutes to proceed to the next step.
- 13. Dispense **10 µL** OptiGlow[™] working solution to each well. Incubate for 15 minutes at RT.
 - a. <u>Caution</u>: Observe the wells <u>during the incubation</u>. When the substrate has completely drained from all wells, remove the plate and pad from the holder. Discard the pad. Wipe the bottom of the plate with a Kimwipe[™] to remove any liquid on the bottom surface of the plate. Step 13a will be completed within the 15 minute substrate incubation time.
- 14. Place the plate in the reading chamber of a fluorescence plate reader. Promptly at the conclusion of the 15 minute incubation, read the plate.



It is common to see slight differences in the time required for different wells to empty. This difference has no impact on assay performance.



To facilitate work flow, incubations designated as 10 minutes may be extended to 20 minutes with no impact on method performance.



Optimiser[™] "wash" is performed by simply **dispensing_**OptiWash[™] to the wells.



<u>Caution:</u> In rare cases, a well may not empty in 10 min. If so, blot the reagent from the well with a tissue. **Do not analyze signal from this well.**

Calculations:

- 1. Calculate the mean background signal from the blank wells for assay with each type of OptiBind[™] coating buffer (for example, average signal from well G1 and H1 for background signal of assay with OptiBind[™]-A coating buffer).
- 2. Calculate the mean sample signal from the high concentration protein standard wells for assay with each type of OptiBind[™] coating buffer (for example, average signal from well A1, B1, C1 for high signal of assay with OptiBind[™]-A coating buffer).
- 3. Calculate the mean sample signal from the low concentration protein standard wells for assay with each type of OptiBind[™] coating buffer (for example, average signal from well D1, E1, F1 for low signal of assay with OptiBind[™]-A coating buffer).
- 4. Create screening curves by plotting the OptiBind[™] coating buffer types (x-axis) vs the background-adjusted signal (y-axis).
- 5. Choose the type of OptiBind[™] coating buffer which gives highest signal (after subtracting background)

Selecting the Best Coating Buffer:

Select the OptiBind type which gives the maximum signal. This particular OptiBind[™] coat buffer should be used for all further experiments of this assay.

- Use the HIGH concentration curve ONLY if the peak RFU value is less than 90% of RFU_{max}.
- Use the LOW concentration curve ONLY if the peak RFU value is more than 10% of RFU_{max}.
- If both curves are valid per criteria listed above; usually the same OptiBind[™] formulation will show best results.
- If there is discrepancy in choice of OptiBind[™] formulation from LOW and HIGH concentration curves, use the HIGH concentration curve to make the selection.

Example Data:

Figure 9 shows screening test results for the illustrative IL-6 assay. Protein standard concentrations of 200 pg/mL (80% of max) and 50 pg/mL (20% of max) were used. 2 µg/mL of capture antibody and 2 µg/mL of detection antibody were used for this assay. Data read using Biotek Flx800 fluorescence plate reader with excitation filter at 528/20nm and emission filter at 590/35, sensitivity at 45 (based on RFU_{max} ~ 11,000; *see Unit conventions, page 20*). Data acquisition and analysis utilized Gen5TM software and Excel.



Figure 9. Results for coat buffer screening test for Illustrative IL-6 assay.

For the data shown in Figure 9, both curves for screening test result are valid and either curve can be used to select the best OptiBind[™] formulation. Both curves also show that OptiBind[™]-H is the best coat buffer for this assay.

Data logging and Calculation Worksheets Experiment 1 — Selecting the Correct Antibody Coating Buffer

Assay Target:		
Capture antibody information:	, and working concentration:	μg/mL
Detection antibody information:	, and working concentration:	μg/mL
Protein standard information:	, and top concentration:	
High protein standard, 80% of top standard concentration: _		
Low protein standard, 20% of top standard concentration:		
RFU _{max} :		

Test results

OptiBin	d™ type	А	В	C	D	E	F	G	Н	I	J	К	L
		1	2	3	4	5	6	7	8	9	10	11	12
	Α												
High	В												
standard	С												
	Mean												
	D												
Low	Ε												
standard	F												
	Mean												
	G												
Blank	Н												
	Mean												

OptiBind[™] type ______ gives highest signal, if that signal of high standard between 10% and 90% of RFU_{max}

With this type of OptiBind[™], protein standard at concentration ______ give signal between 10% and 50% of

 RFU_{max} . It will be used for experiment 2.

Tested By: _____ Date: _____

Experiment 2 – Selecting the Optimal Capture/detection Antibody Concentration:

The larger surface area and very high surface area to volume ratio in the microfluidic channel of Optimiser[™] microplate allows more capture antibody to be adsorbed onto the surface, which may improve assay sensitivity. This experiment uses a checkerboard titration pattern with 3 concentrations of capture antibody and 3 concentrations of detection antibody. Six rows (72 wells) in an Optimiser[™] microplate will be used for this experiment.

Based on the result from Experiment 1, concentration of protein standard which gives maximum signal between 10% to 50% of RFU_{max} will be used for this experiment.

Reagent Preparation:

Bring all reagents to room temperature before use and prepare all necessary dilutions before beginning the test procedure.

- 1. <u>OptiBind</u>[™]: Use the OptiBind[™] coat buffer selected from experiment 1.
- <u>Capture Antibody</u>: Three concentrations of capture antibody working solution will be tested: a) <u>same</u>, b) <u>2 times</u> and c) <u>4 times</u> as that used in conventional assay. Prepare the capture antibody working solution by diluting the capture antibody stock in *selected* OptiBind[™] to make **250 µL** final working solution. Use a single row in the polypropylene 96-well v-bottom plate, load **60 µL** of working solution to each well. Load wells 1-4 with 4x concentration capture antibody solution, wells 5-9 with 2x concentration, and wells 9-12 with "same as conventional" concentration.
- 3. <u>OptiBlock™</u>: OptiBlock™ is provided in ready-to-use form and is used to block the surfaces of the Optimiser™'s microfluidic reaction chambers following their incubation with the capture antibody solution. OptiBlock™ is also used as the diluent for the protein standard, detection antibody and SAv-HRP. For blocking step, prepare 1 mL of OptiBlock™ in a v-shape reservoir.
- Protein Standard: Prepare 1 mL of protein standard with OptiBlock™ in a v-shape reservoir. Select a concentration which gives the max signal between 10% to 50% of RFU_{max}.
 From results for Illustrative IL-6 assay; a concentration of 50 pg/mL would be used for this experiment.
- 5. <u>Blank (zero)</u>: Prepare **1 mL** of OptiBlock[™] in a v-shape reservoir.
- 6. <u>Detection Antibody</u>: Three concentrations of detection antibody working solution will be tested: a) <u>same</u>, b) <u>2 times</u> and c) <u>4 times</u> as that used in conventional assay. Prepare the detection antibody working solution by diluting the detection antibody stock in OptiBlock[™] to make **200 µL** final working solution. Use a single column in the polypropylene 96-well v-bottom plate, load **90 µL** of working solution in to each well. Load wells A and B with 4x concentration detection antibody solution, wells C and D with 2x concentration, wells E and F with "same as conventional" concentration.
- <u>SAv-HRP</u>: Use SAv-HRP stock solution provided in the kit. Prepare the SAv-HRP working solution by adding 8 μL of SAv-HRP stock solution to 1.2 mL of OptiBlock™ (1:150 dilution) in a v-shape reservoir.
- Substrate solution: The procedure requires 10 μL of the working substrate solution for each assay well to be used. Prepare the substrate working solution in a v-shape reagent reservoir by mixing 0.9 mL of OptiGlow[™]-A, 0.9 mL of OptiGlow[™]-B and 18 μL of OptiGlow[™]-C.
- <u>OptiWash™</u>: OptiWash™ is provided in ready-to-use form. No further preparation is required. The procedure requires total of 75 µL of OptiWash™ for each assay well to be used. Prepare **10 mL** of OptiWash™ into a v-shaped reagent reservoir and use it for all wash steps in the assay.

		Capture antibody concentration												
Detection antibo	dy	4 times as conventional assay				coi	2 tim nventio	ies as onal as	say	Same as conventional assay				
concentration	-	1	2	3	4	5	6	7	8	9	9 10 11 12			
4 times as	Α	Protein		n Blank		Pro	Protein		Dlank		Protein		Blank	
conventional assay	В	stan	dard	віапк		stan	standard		DIdIIK		standard		IIIK	
2 times as	С	Pro	tein	Bla	nk	Pro	tein	Blank		Protein		Bla	nk	
conventional assay	D	stan	dard	lard Blank		stan	standard		BIGUK		dard	BIGUK		
Same as	Е	Pro	tein	Blank		Pro	Protein		ank	Pro	tein	Bla	nk	
conventional assay	F	stan	dard			standard		DIdIIK		standard		Dia	BIANK	

Procedure:

- 1. Assemble the Optimiser[™] Microplate, Optimiser[™] Pad, and Optimiser[™] Microplate Holder as described on Page 5.
- <u>Hint:</u> Optimiser[™] incubation steps are from 10 to 20 minutes in length. To achieve optimal assay performance, all materials must be transferred to the Optimiser[™] microplate within one minute at each step. To accomplish this, first place the materials to be transferred in the enclosed 96-well polypropylene v-bottom plate or v-shape reagent reservoir (as *instructed in Reagent Preparation, page 26*). Then transfer the materials to the Optimiser[™] wells using a <u>10 µL multi-channel pipette</u>.
- 3. Dispense 5 μL capture antibody solution to the required number of wells in the Optimiser[™] microplate. Incubate 10 minutes at room temperature (RT).
- 4. Dispense 5 µL OptiWash[™] to each well. Wait 10 minutes to proceed to the next step.
- 5. Dispense 5 μL OptiBlock[™] to the capture antibody-coated wells. Incubate 10 minutes at RT.
- 6. Dispense 5 μ L of the standard and blank to the required number of replicate wells of the plate. Incubate 20 minutes at RT.
- 7. Dispense 5 µL OptiWash[™] to each well. Wait 10 minutes to proceed to the next step.
- 8. Dispense 5 μL detection antibody working solution to each well. Incubate 10 minutes at RT.
- 9. Dispense 5 µL OptiWash[™] to each well. Wait 10 minutes to proceed to the next step.
- 10. Dispense 5 μL SAv-HRP to each well. Incubate 10 minutes at RT.
- 11. Dispense 30 µL OptiWash[™] to each well. Wait 10 minutes to proceed to the next step.
- 12. Again dispense 30 µL OptiWash[™] to each well. Wait 10 minutes to proceed to the next step.
- 13. Dispense **10 µL** OptiGlow[™] working solution to each well. Incubate for 15 minutes at RT.
 - a. <u>Caution:</u> Observe the wells <u>during the incubation</u>. When the substrate has completely drained from all wells, remove the plate and pad from the holder. Discard the pad. Wipe the bottom of the plate with a Kimwipe[™] to remove any liquid on the bottom surface of the plate. Step 13a will be completed within the 15 minute substrate incubation time.
- 14. Place the plate in the reading chamber of a fluorescence plate reader. Promptly at the conclusion of the 15 minute incubation, read the plate.



It is common to see slight differences in the time required for different wells to empty. This difference has no impact on assay performance.



To facilitate work flow, incubations designated as 10 minutes may be extended to 20 minutes with no impact on method performance.



Optimiser[™] "wash" is performed by simply **dispensing_**OptiWash[™] to the wells.



Caution: In rare cases, a well may not empty in 10 min. If so, blot the reagent from the well with a tissue. **Do not** analyze signal from this well.

Selecting the Optimal Antibody Concentrations:

Use the following steps to determine the optimal combination of capture/detection antibody concentrations:

- 1) For most assays tested at Siloam to-date, using the **same** concentration of capture and detection antibody results in approximately the same dynamic (operating) range for the assay on Optimiser[™]
- 2) Select a higher capture (or detection) antibody ONLY when the data shows at least 1.5x higher signal than lower concentration combination. If the signal increase is less than 50% the net gain in assay sensitivity will not be significant. See example data below.
- 3) DO NOT select antibody concentrations which give background reading higher than 3% of RFU_{max}. *See example data below*.
- 4) Higher concentrations slightly increase the consumption (hence cost); although even with 4 times higher concentrations, Optimiser[™] assays still use 5 times less antibody than conventional plate. Select the higher concentrations only if your assay requires higher sensitivity.

Example Data:

As an example, an antibody optimization test has been performed for IL-6 assay. Protein standard concentration is 50 pg/mL. Three concentrations of capture antibody (2, 4, 8 μ g/mL) and three concentrations of detection antibody (2, 4, 8 μ g/mL) have been used for this assay. Use Biotek Flx800 fluorescence plate reader with excitation filter at 528/20nm and emission filter at 590/35, sensitivity at 45. Data acquisition and analysis utilized Gen5TM software and Excel. Results are shown as below:

Mean value of results

		Capture antibody concentration												
Detection antibody concentration			8 με	g/mL			4 μg	g/mL		2 μg/mL				
		1	2	3	4	5	6	7	8	9	10	11	12	
8	Α	27	2707		337		2339		348		1650		359	
δ μg/ mL	В	27												
4.ug/ml	С	24	0.476		207		2007		222		4720		200	
4 μg/ mL	D	2476		297		2097		233		1730		266		
2	Е	2244		220		2070		200		45.42		202		
2 μg/mL F		23	2344		238		2073		289		1543		292	

Converted to OD readings

		Capture antibody concentration													
Detection	ו		8 με	g/mL			4 μg	g/mL		2 μg/mL					
concentration		1	2	3	4	5	6	7	8	9	10	11	12		
9	Α	0.0	0.001		107	0.7		0.1	0.111		25	0.111			
δ μg/mL	В	0.801		0.107		0.744		0.111		0.525		0.114			
Aug/ml	С	0.7	0 700		0.005		0.667		0.074		0.550		0.095		
4 μg/ mL	D	0.788		0.095		0.007		0.074		0.550		0.085			
2 ug/ml	Е	0.746		0.076		0.6	0.000		0.000		0.401		02		
2 μg/mL	F					0.660		0.092		0.491		0.093			

• All background readings are less than 3% of RFU_{max} (RFU reading = 330 for RFU_{max} = 11,000).

 8 μg/mL capture; 2 μg/mL detection antibody concentrations selected (signal 50% higher than 2 μg/mL capture; 2 μg/mL detection).

Increasing capture antibody concentration has most significant effect on signal. Using 8 μg/mL capture; 8 μg/mL detection shows only ~15% signal increase. This will not have a significant effect on assay sensitivity.

Data logging and Calculation Worksheets Experiment 2 — Selecting the Optimal Capture/detection Antibody Concentrations

Assay Target:	_		
Capture antibody information:	,		
and working concentration: 1x=	µg/mL, 2x=	µg/mL, 4x=	_µg/mL
Detection antibody information:			
and working concentration: 1x=	µg/mL, 2x=	µg/mL, 4x=	_µg/mL
Protein standard information:		,	
and concentration will be used (from ex	periment 1):		

Test results

			Capture Antibody											
Detectio	n antibody	1	2	3	4	5	6	7	8	9	10	11	12	
1.	Α													
IX	В													
2x	C													
	D													
4x	E													
	F													

Calculated Mean

			Capture Antibody										
		4 times	times as conventional assay				s as conv	ventiona	al assay	Same as conventional assay			
Detection antibody		1	2	3	4	5	6	7	8	9	10	11	12
4 times as	Α												
conventional assay	В												
2 times as	С												
conventional assay	D												
Same as conventional assay	Ε												
	F												

Optimal capture antibody concentration: _____µg/mL

Optimal detection antibody concentration: _____µg/mL

With optimal antibody concentrations, the top concentration of protein standard for experiment 3 is: _____

Tested By: _____ Date: _____

Experiment 3 – Determine Assay Measurable Range:

The results from Experiment 1 and Experiment 2 are used for the final Experiment to to determine the dynamic range of the assay. This experiment will run a standard curve of the assay with wide range of concentrations which covers the expected dynamic range. Appropriate standard diluents must be used for this experiment. For example, use cell culture medium as standard diluent for measuring cell culture supernatant. 3 rows in one Optimiser[™] microplate will be used.

Reagent Preparation:

Bring all reagents to room temperature before use and prepare all necessary dilutions before beginning the test procedure.

- 1. <u>OptiBind[™]</u>: Use the optimal OptiBind[™] coat buffer selected from experiment 1.
- <u>Capture Antibody</u>: Use optimal concentration of capture antibody determined in experiment 2. Prepare the capture antibody working solution by diluting the capture antibody stock in *selected* OptiBind[™] to make **300 μL** final working solution. Use 3 wells in one column on the polypropylene 96-well v-bottom plate, load **90 μL** of working solution into each well.
- 3. <u>OptiBlock[™]</u>: OptiBlock[™] is provided in ready-to-use form and is used as blocking buffer. OptiBlock[™] is also used as a diluent for the detection antibody and SAv-HRP. For blocking step, prepare **1 mL** of OptiBlock[™] in a v-shape reservoir.
- 4. <u>Standard diluent:</u> Appropriate standard diluent shall be used for this experiment. <u>As an example</u>, use cell culture medium as standard diluent for measuring cell culture supernatant
- 5. <u>Protein Standards</u>: Prepare **100 μL** of protein standard at defined top concentration with standard diluent in a plastic tube. Using one row of 96-well v-bottom plate, load 45 μL of standard diluent in wells 2-12. Load 90 μL of top standard into the first well, transfer 45 μL of top standard to well 2, change tip and mix with the standard diluent to create a 1:2 dilution. Continue this process till well 11; **do not** transfer diluted standard to well 12; leave well 12 as blank.

Top concentration: Based on the converted OD reading from experiment 2 (and assuming the standard curve is linearly proportional to antigen concentration), calculate the concentration for which the signal will be higher than OD 3.5.

<u>As an example</u>, for the IL-6 assay, expected top concentration = (50 pg/mL x 3.5)/0.746 = 234 pg/mL. For easy calculation and dilution, 250 pg/mL would be used as top concentration for this experiment.

- 6. <u>Detection Antibody</u>: Use optimal concentration of detection antibody determined in experiment 2. Prepare the detection antibody working solution by diluting the detection antibody stock in OptiBlock™ to make **300 μL** final working solution. Use 3 wells in one column in the polypropylene 96-well v-bottom plate, load **90 μL** of working solution into each well.
- <u>SAv-HRP</u>: Use SAv-HRP stock solution provided in the kit. Prepare the SAv-HRP working solution by adding 8 μL of SAv-HRP stock solution to 1.2 mL of OptiBlock™ (1:150 dilution) in a v-shape reservoir.
- Substrate solution: The procedure requires 10 µL of the working substrate solution for each assay well to be used. Prepare the substrate working solution in a v-shape reagent reservoir by mixing 0.4 mL of OptiGlow[™]-A, 0.4 mL of OptiGlow[™]-B and 8 µL of OptiGlow[™]-C*.
- 9. <u>OptiWash[™]</u>: OptiWash[™] is provided in ready-to-use form. No further preparation is required. The procedure requires total of 75 µL of OptiWash[™] for each assay well to be used. Prepare **4 mL** of OptiWash[™] into a v-shaped reagent reservoir and use it for all wash steps in the assay.

*This represents a 50:50:1 mix ratio. Increase OptiGlow[™]-C in this mixing ratio (e.g. 50:50:5) will allow for wider dynamic range. The sensitivity is NOT improved but the range is extended on top of curve.

Assay Layout: Protein standards concentration

	1	2	3	4	5	6	7	8	9	10	11	12
Α	Ton											
В	TOP	1/2	1/4	1/8	1/16	1/32	1/64	1/128	1/256	1/512	1/1024	0
С	concentration											

Top concentration: Based on the RFU reading from experiment 2 (and assuming the standard curve is linearly proportional to antigen concentration), calculate the concentration for which the signal will be higher than RFU_{max}.

Procedure:

- 1. Assemble the Optimiser[™] Microplate, Optimiser[™] Pad, and Optimiser[™] Microplate Holder as described on Page 5.
- 2. <u>Hint:</u> Optimiser[™] incubation steps are from 10 to 20 minutes in length. To achieve optimal assay performance, all materials must be transferred to the Optimiser[™] microplate within one minute at each step. To accomplish this, first place the materials to be transferred in the enclosed 96-well polypropylene v-bottom plate or v-shape reagent reservoir (as *instructed in Reagent Preparation, page 30*). Then transfer the materials to the Optimiser[™] wells using a <u>10 µL multi-channel pipette</u>.
- 3. Dispense 5 μL capture antibody solution to the required number of wells in the Optimiser[™] microplate. Incubate 10 minutes at room temperature (RT).
- 4. Dispense 5 µL OptiWash[™] to each well. Wait 10 minutes to proceed to the next step.
- 5. Dispense 5 µL OptiBlock[™] to the capture antibody-coated wells. Incubate 10 minutes at RT.
- 6. Dispense 5 μ L of the standard and blank to the required number of replicate wells of the plate. Incubate 20 minutes at RT.
- 7. Dispense 5 µL OptiWash[™] to each well. Wait 10 minutes to proceed to the next step.
- 8. Dispense 5 μL detection antibody working solution to each well. Incubate 10 minutes at RT.
- 9. Dispense 5 µL OptiWash[™] to each well. Wait 10 minutes to proceed to the next step.
- 10. Dispense 5 μ L SAv-HRP to each well. Incubate 10 minutes at RT.
- 11. Dispense 30 µL OptiWash[™] to each well. Wait 10 minutes to proceed to the next step.
- 12. Again dispense 30 μL OptiWash[™] to each well. Wait 10 minutes to proceed to the next step.
- 13. Dispense **10 µL** OptiGlow[™] working solution to each well. Incubate for 15 minutes at RT.
 - a. <u>Caution</u>: Observe the wells <u>during the incubation</u>. When the substrate has completely drained from all wells, remove the plate and pad from the holder. Discard the pad. Wipe the bottom of the plate with a Kimwipe[™] to remove any liquid on the bottom surface of the plate. Step 13a will be completed within the 15 minute substrate incubation time.
- 14. Place the plate in the reading chamber of a fluorescence plate reader. Promptly at the conclusion of the 15 minute incubation, read the plate.



It is common to see slight differences in the time required for different wells to empty. This difference has no impact on assay performance.



To facilitate work flow, incubations designated as 10 minutes may be extended to 20 minutes with no impact on method performance.



Optimiser[™] "wash" is performed by simply **dispensing_**OptiWash[™] to the wells.



<u>Caution:</u> In rare cases, a well may not empty in 10 min. If so, blot the reagent from the well with a tissue. **Do not analyze signal from this well.**

Determine the Dynamic Range:

Use the following steps to determine dynamic (or operating) range of the assay:

- 1) The top measurable concentration should give signal between 60%-90% of RFU_{max} .
- For best linearity and easy operation, a 7-point, two-step dilution (with one zero) is most commonly used as standard curve. Using the selected top measurable concentration, determine the lowest concentration (6 columns to the right of the top measurable concentration).

Please check Appendix 1 to see a procedure for extending dynamic range of assays on Optimiser™.

Example Data:

As an example, a dynamic range test has been performed for IL-6 assay. Top concentration of IL-6 is 250 pg/mL. Use Biotek Flx800 fluorescence plate reader with excitation filter at 528/20nm and emission filter at 590/35, sensitivity at 45. Data acquisition and analysis utilized Gen5[™] software and Excel. Results are shown as below:

IL-6 (pg/mL)	250	125	62.5	31.3	15.6	7.8	3.9	2.0	1.0	0.5	0.2	0
Mean of RFU	9764	6396	3539	2116	1263	764	471	358	316	230	217	176
Converted OD	3.10	2.03	1.12	0.67	0.40	0.24	0.15	0.11	0.10	0.07	0.06	0.05

The (lowest) top measurable concentration of this IL-6 assay is 125 pg/mL. Hence, the lowest measurable concentration is at least 2 pg/mL.

Using 4x higher concentration of the capture antibody allows Optimiser^M based IL-6 assay to achieve higher sensitivity while still using less reagents than a conventional plate. The sample volume for the Optimiser^M assay is only 5 μ L (compared to 100 μ L on the conventional plate) and the total assay time for Optimiser^M based assay is only 2 hours (compared to 8 hours for conventional plate).

OTHER ASSAY FORMATS ON THE OPTIMISER™:

Indirect Immunoassay: Siloam has developed protocols for indirect ELISA on Optimiser[™]. These will be added to the Assay Transfer Guide in future revisions. Please contact Siloam's technical support for assistance.

Competitive Immunoassay: Siloam has also developed protocols for competitive ELISA on Optimiser[™]. These will be added to the Assay Transfer Guide in future revisions. Please contact Siloam's technical support for assistance.

TROUBLESHOOTING:

The Optimiser[™] technology and OptiMax[™] ELISA kits have been designed and manufactured to ensure problem-free sample analysis. However, Siloam Biosciences has prepared the following guidance for trouble shooting problems that might be encountered due to the unique features of the Optimiser[™] technology as well as problems that can be encountered with immunoassays in general.

Problem	Possible Cause	Solution						
	A bubble is in the well.	 Disrupt the bubble with a clean 26 gauge needle. Follow recommended pipetting guidelines. Prepare excess reagent to avoid aspirating air. Do not use detergents. 						
Liquid does not drain	Sample contains particulates.	 Centrifuge sample for 10 min at 13,000 RPM, or Filter the sample using a 0.2 μm filter. 						
from the Optimiser™ well or does not drain within 10 minutes.	Plate has lost contact with the absorbent pad or is positioned incorrectly.	 Ensure that the absorbent side (rough) of the pad is in contact with Optimiser[™] and the tape side (smooth) is facing down to touch holder. Ensure the topside of the pad is touching the bottom of Optimiser[™] plate by pushing down firmly on the 4 corners of the plate. Ensure the plate and pad are securely aligned in the holder. 						
	Standard has degraded.	 Use standard on the day of its reconstitution, or Thaw single use aliquots fresh on each test day. Avoid repeated freeze-thaws. 						
	Incorrect reader filters	Confirm filters meet requirements for substrate.						
No signal or unexpectedly low signal	Antibodies or SAv-HRP are degraded.	 Use within specified expiration period. Store according to recommended storage temperature. 						
	Substrate was prepared incorrectly.	 Thaw OptiGlow[™] - C thoroughly before preparing substrate working solution. 						
	Substrate working solution has degraded.	 Prepare substrate no more than 30 minutes before plate is read. 						
Unexpectedly high signal	Incorrect reader filters with overlapped wavelength bandwidth	 Confirm filters meet requirements for substrate. 						
	Reagent contamination	 Avoid cross contamination in reagents. Always change the pipet tips when handling different buffers/reagents. 						
Poor precision	Pipetting errors, use of alternate assay buffers or SAv-HRP	 Follow recommendations for pipetting small volumes (Page 5). Variance <10% and background < 3% of RFU_{max} (established during reader setup) are expected. Do not substitute provided assay buffers or Sav-HRP. 						
Signal of lower standard(s) are < 0 following	Degraded standard	 Use standard on the day of its reconstitution, or Thaw single use aliquots fresh on each test day. Avoid repeated freeze-thaws. 						
background subtraction.	Degraded capture antibody	 Use within specified expiration period. Store according to recommended storage temperature. 						

APPENDIX 1: ALTERNATIVE ASSAY PROCEDURES ON OPTIMISER™

Rapid Assay on Optimiser[™]:

The standard Optimiser[™] assay procedure, as described on page 15 of this Instruction Manual, requires approximately 2 hours (125 minutes) to complete. Most incubation steps are 10 minutes in length with the exceptions of sample incubation (20 minutes) and substrate incubation (15 minutes).

Siloam Biosciences has developed an alternative method that can be completed in 90 minutes. The sample incubation time (20 minutes), final two washes (10 minutes) and substrate incubation time (15 minutes) are unchanged. However, the remaining incubation times can be reduced from 10 minutes to 5 minutes. The plot in Figure 10 shows the adsorption kinetics on the OptimiserTM showing that in ~ 5 minutes, ~ 92% of peak adsorption (or binding) is completed. More importantly, from 5 – 30 min (next time point) the adsorption only changes from ~ 92% to ~ 96%.

In doing so, the total assay time is reduced from 125 minutes to 90 minutes with no change in <u>method performance</u>. Siloam strongly recommends that only users proficient in the use of the Optimiser[™] microplate system attempt the rapid test protocol. It is especially important to ensure that pipetting for each step is completed within ~ 30 seconds. It is also critically important to maintain consistency in pipetting and incubation intervals when using the accelerated protocol.

Contact Siloam Biosciences for additional details and specific guidance on running this alternate protocol.



Figure 10. Adsorption characteristics of capture antibody on the Optimiser[™] microchannel surface.



PLEASE CONTACT TECHNICAL SUPPORT FOR ASSISTANCE WITH THIS PROTOCOL. The description provided here should not be used a formal protocol.

Ultra-sensitive Assay on Optimiser™:

Because of the unique features of the Optimiser[™] plate and OptiMax[™] ELISA procedures, users can apply sample to individual microfluidic reaction chambers multiple times. The result is a significant improvement in assay sensitivity when ultralow sensitivity is required. The additional sample applications can be performed manually for a limited number of repeat sample loads but Siloam strongly recommends use of a laboratory sample processor for the ultra-high sensitive protocol.

The data in the figure below illustrates the sensitivity and dynamic range obtained using the standard OptiMax^M ELISA procedure (a single 5 μ L sample addition) and the improvement in sensitivity that is gained by performing 20 consecutive 5 μ L sample applications to individual reaction chambers using a laboratory sample processor.

Each additional sample incubation is 5 minutes in length. Thus with 95 additional minutes of assay time, the total assay time is approximately 3 hours with a corresponding increase in assay sensitivity of 20-fold.

The repeat sample loading methods is a reliable and simple method to "tune" the sensitivity of the assay to the desired range simply by adjusting the number of sample addition (and incubation steps).

Contact Siloam Biosciences for additional details and specific guidance on running this alternate protocol.







PLEASE CONTACT TECHNICAL SUPPORT FOR ASSISTANCE WITH THIS PROTOCOL. The description provided here should not be used a formal protocol. **Technical Assistance:** If you require assistance, please contact Siloam Biosciences, Inc. Technical Support at 513-429-2976 or <u>techsupport@siloambio.com</u>.

Additional technical assistance is available under the Technical Support tab on the Siloam Biosciences web site (<u>http://siloambio.com/</u>).

- Using Optimiser[™] Immunoassay Microplate Video
- Optimiser™ User's Guide
- Reader Settings
- Quick Reference Guide
- Frequently Asked Questions
- Application Notes

Two additional videos appear under the Technology tab of the web site.

- Optimiser[™] Principles of Operation
- Running an Assay with Optimiser™



Better Immunoassays Through Innovative Microfluidics

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