



Q-Plex™ Human Pneumococcal IgG

Version 2.0
For Research Use Only
Not For Use in Diagnostic Procedures

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Symbol	Explanation
	Catalog Number
	Lot Number
	Use By YYYY-MM-DD
	Temperature Limitation
	Manufacturer
	Keep Away from Sunlight

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NAME AND INTENDED USE

Q-Plex™ Human Pneumococcal IgG

Quansys Biosciences Catalog Numbers

Q-Plex Human Pneumococcal IgG (9-Plex) **482149HU**

Q-Plex Human Pneumococcal IgG (14-Plex) **482249HU**

Kits containing extra diluent:

Q-Plex Human Pneumococcal IgG (9-Plex) **482149HU-30**

Q-Plex Human Pneumococcal IgG (14-Plex) **482249HU-30**

The Q-Plex Human Pneumococcal IgG is a chemiluminescent Enzyme-Linked Immunosorbent Assay (ELISA) intended for quantitative concurrent measurement of IgGs against Pneumococcal serotypes 1, 2, 3, 4, 5, 6B, 7F, 8, 9N, 9V, 10A, 11A, 12F, 14, 15B, 17F, 18C, 19A, 19F, 20, 22F, 23F and 33F in serum and plasma samples. The kit accompanying this manual will contain a subset of the above analytes which are identified on the In-Kit Certificate of Analysis.

PRINCIPLE OF THE ASSAY

This multiplex assay is based on the microplate indirect immunoassay technique for the measurement of human IgG antibodies reactive to pneumococcal polysaccharide antigens.

In addition to pneumococcal polysaccharides, the microplate is arrayed with 3 control spots.

The first control spot is for cell wall polysaccharide (CWPS) negative control. Anti-CWPS antibodies can bind to the specific pneumococcal antigens that the assay aims to detect creating non-specific false positive results. CWPS is added to the sample diluent as an adsorbent to prevent these interactions. Thus, if signal over the assay-designated limit is detected the user should consider the data from that well as potentially affected by false positives. Quansys recommends re-running affected samples at a higher dilution factor.

The second control spot is a human serum loading positive control. This control spot produces signal if sample sourced from a human patient is loaded and thus acts a positive control for proper performance of the protocol. If Q-View Software reports this control is out of range, it is possible that human sample was not loaded or is extremely diluted.

The third control spot is the reference spot positive control. The reference spot is used by Q-View Software to properly align and adjust the spot matrix overlay. This spot will produce signal in each well in which both the detection and substrate mix were added correctly.

Samples or calibrators are pipetted into wells of a 96-well plate arrayed with immobilized polysaccharides that capture human IgGs specific for the respective serotypes. After washing away any unbound antibodies, an HRP-conjugated anti-Human IgG antibody is added. Following an additional wash, the amount of HRP-conjugated antibody bound to each location of the array is proportional to the amount of serotype-specific antibody initially captured.

The amount of conjugated enzyme on each location of the array is measured with the addition of a chemiluminescent substrate.

SAMPLE COLLECTION AND STORAGE

The sample collection and storage conditions are intended as general guidelines. The Clinical and Laboratory Standards Institute (CLSI GP44-A4) recommends the following storage conditions for samples: Samples should be stored at room temperature no longer than 8 hours. If the assay will not be completed within 8 hours, the samples should be refrigerated at +2°C to +8°C. If the assay will not be completed within 48 hours, or if the samples will be stored beyond 48 hours, samples should be frozen at -20°C or lower. Samples should not be repeatedly frozen and thawed. Frozen samples must be mixed well after thawing and prior to testing. Do not use bacterially contaminated samples. It is the responsibility of the individual laboratory to use all available references and/or its own studies to determine its own specific stability criteria.

Serum – Use a serum separator tube (SST) and allow samples to clot for 30 minutes before centrifugation for 15 minutes at 1000 x g. Remove serum and assay immediately or aliquot and store samples at $\leq -20^{\circ}\text{C}$. Avoid repeated freeze-thaw cycles.

Plasma – Collect plasma using EDTA, Citrate, or Heparin as an anticoagulant. Centrifuge for 15 minutes at 1000 x g within 30 minutes of collection. Assay immediately or aliquot and store samples at $\leq -20^{\circ}\text{C}$. Avoid repeated freeze-thaw cycles.

WARNINGS AND PRECAUTIONS

1. Read all instructions before beginning test.
2. For research use only. Not for use in diagnostic procedures.
3. The kit should not be used beyond the expiration date on the kit label.
4. **This kit is sensitive to saliva. Wear a mask during preparation and running of the kit.**
5. If running multiple kits, a calibration curve must be included for each 96-well plate. The curve from one plate cannot be used to calculate sample values from other plates.
6. Do not mix or substitute reagents with those from other kits or lots.
7. Pre-wet pipette tips 3 times by drawing up the liquid into the pipette and then dispensing back into the original vessel prior to the addition of samples, calibrators, or external controls to the microplate.
8. Load all calibrators, controls, and samples into the microplate within 4 minutes of each other.
9. Be exact with incubation times
10. Be exact when mixing Substrate A and B+ and mix thoroughly.
11. Do not allow the substrate or HRP-conjugated detection antibody to be exposed to UV light, as this may degrade these reagents.
12. Do not allow the plate to dry out between steps.
13. All products are carefully validated; however due to the variability encountered in biological sample matrices, the possibility of interference or sample matrix effects cannot be excluded.

Warning: The calibrator contains components of human origin. These components have been tested at the donor level and found to be negative for HBsAg, HIV-1 and HIV-2 antibodies, and HCV.

KIT CONTENTS & STORAGE

Unopened Kit -  Store at 2-8°C.  Do not use past kit expiration date.
Do not re-use.

Part/Description	Storage of opened/reconstituted material
Q-Plex™ Array Microplate  482252HU (14-Plex) 482152HU (9-Plex) 1 arrayed and blocked 96-well polystyrene microtiter plate	 2-8°C until kit expiration
Wash Buffer Concentrate (20X)  101158GR Liquid, 50 mL/vial of a concentrated solution of buffered surfactant	
Sample Diluent  481157HU Liquid, 2 vials with 10 mL/vial each of a buffered protein solution with CWPS	
Detection Mix  482156HU Liquid, 6 mL/vial of an HRP-labeled anti-human IgG antibody in a buffered protein solution with preservatives	
Calibrator  481155HU Lyophilized, human IgG antibodies specific for Streptococcus pneumoniae capsular polysaccharides in a buffered protein base	Discard unused reconstituted calibrator.
Substrate A  101159GR Liquid, 3.5 mL/vial	 Do not expose to UV light.  Store mixed substrate solution at room temperature (20-25°C) for up to 1 week.  Store unmixed solutions at 2-8°C until kit expiration.
Substrate B+  101120GR Liquid, 3.5 mL/vial	 Do not expose to UV light.  Store mixed substrate solution at room temperature (20-25°C) for up to 1 week.  Store unmixed solutions at 2-8°C until kit expiration.
Plate Seals (3)  S1064 Adhesive strips	Non-perishable

MATERIALS REQUIRED BUT NOT SUPPLIED

In addition to the kit contents listed, the following materials are required to run this assay:

1. Multichannel pipette (20-200 μ L) and/or single channel pipettes (20-1000 μ L) with appropriate tips
2. 10 mL serological pipette
3. Polypropylene tubes or polypropylene 96-well plate(s) for sample and calibrator preparation
 - a. Example: Nunc® MicroWell™ 96-Well Plates, Polypropylene, 249944; Eppendorf® LoBind Protein or Genomic Microcentrifuge Tubes, 022431102
4. Q-View™ Imager Pro or Q-View™ Imager LS and Q-View™ Software
5. Deionized water
6. Microplate washer (An optional multichannel pipette protocol is provided in Appendix A)
7. 1 liter graduated cylinder for the preparation of wash buffer.

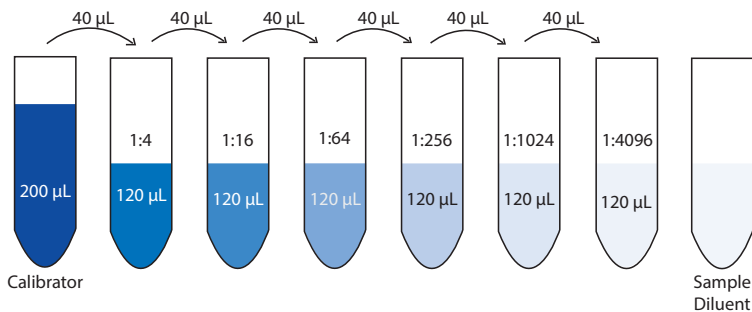
ASSAY PREPARATION

1. Install Q-View Software on any computers that will be used for analysis or operating a Q-View Imager.
2. Set up the imager. For imager-specific instructions, see www.quansysbio.com/manuals.
3. Allow all kit components to come to room temperature (20-25°C) prior to use.
4. Prepare Wash Buffer: Place 50 mL of Wash Buffer Concentrate (20X) into 950 mL deionized water and mix thoroughly.
5. Prepare Calibrator: Reconstitute lyophilized Calibrator using Sample Diluent with the volume specified on the In-Kit Certificate of Analysis accompanying the kit. Allow to sit for 5 minutes. Mix thoroughly. Use prepared Calibrator within 30 minutes.
6. Prepare chemiluminescent substrate: Combine 3 mL of Substrate A with 3 mL of Substrate B+ and mix gently. **Do not expose to UV light. Store at room temperature (20-25°C) after mixing.**

ASSAY PROCEDURE

Allow all reagents to equilibrate to room temperature (20-25°C) before use and prepare as directed in the previous sections. It is recommended that all calibrators and samples be assayed in duplicate.

- Using the previously prepared Calibrator (see “Assay Preparation” step 6), prepare an 8-point calibration curve (7 points plus 1 blank) in either polypropylene tubes or a polypropylene 96-well plate (see “Materials Required but Not Supplied” for example tubes and plates).
 - Pipette 200 μL of prepared Calibrator into the first tube or well.
 - Pipette 120 μL of Sample Diluent into the other 7 tubes or wells.
 - Transfer 40 μL of undiluted prepared Calibrator from the first tube or well into the second, mix thoroughly, and repeat the transfer from tube to tube or well to well for 5 more points, leaving the last tube or well without any prepared Calibrator. This process is diagrammed below. The undiluted prepared Calibrator serves as the high point of the calibration curve. The Sample Diluent serves as the blank.



- Prepare samples by diluting 1:50 (1 part sample to 49 parts Sample Diluent) in either polypropylene tubes or a polypropylene 96-well plate in a volume sufficient to provide 50 μL of diluted sample per well.

Note: If you anticipate that your analyte concentrations will be greater than the high point of the calibration curves as reported on the In-Kit Certificate of Analysis, use the prepared sample diluent to dilute your samples further.

3. Add 50 μL per well of the prepared calibration curve and samples to the Q-Plex™ Array Microplate. Load all samples and calibration curve into the plate within 4 minutes.
4. Cover the plate with a plate seal and incubate for 30 minutes at room temperature (20-25°C).
5. Wash the plate 3 times (see Appendix A).
6. Add 50 μL per well of Detection Mix, cover with a new plate seal, and incubate for 30 minutes at room temperature (20-25°C).
7. Wash the plate 6 times (see Appendix A).
8. Add 50 μL per well of previously prepared substrate (see "Assay Preparation" step 6). Image plate immediately. Wait no longer than 5 minutes to commence imaging.

Note: If imaging cannot commence immediately, protect the plate from drying for up to 15 minutes by dispensing 100 μL of wash buffer into each well of the plate prior to adding the mixed substrate. When ready to image, remove the wash buffer from the plate and add the substrate.

9. Place the plate in the Q-View Imager.
10. Open Q-View Software, create or open a project, and click Acquire Image.
11. When using a Q-View Imager Pro, set the exposure time to 300 seconds.
12. When using a Q-View Imager LS, set the exposure time to 270 seconds.
13. Click the Capture Image(s) button. Users may perform Well Assignment while images are being captured.

Note: Details about these imaging steps are available in the Q-View Software Manual viewable at www.quansysbio.com/manuals or within Q-View Software under **Support > Manual**.

14. Dispose of all used and unused materials in accordance with local regulations.

ANALYZING A Q-PLEX™ IMAGE

The following summarizes a general workflow for analyzing a Q-Plex image in Q-View Software. Each of these steps is described in greater detail in the Q-View Software and Imager manuals, viewable at www.quansysbio.com/support or within Q-View Software under **Support > Manual**.

1. Acquire or import an image into Q-View Software as previously described.
2. Enter the **Software Product Code** (found on the In-Kit Certificate of Analysis) into the **Product Code** field of the software.
3. **Image Processing:** Align the plate overlay as follows:
 - a. Set the overlay: If using the **Auto-Set Plate Overlay** feature, this will occur automatically. Otherwise, go to **Overlay Options > Set Plate Overlay**.
 - b. To visualize bright or dim spots, optimize the display using **Image Options > Adjust Gamma** (does not affect the data).
 - c. Optimize overlay alignment: Go to **Overlay Options > Adjust Plate Overlay** to pivot the overlay, **Adjust Well** and **Adjust Spot** to move individual wells and spots, then **Auto-Adjust Spots** to automatically snap each circle of the overlay to the nearest spot of the image beneath.
4. **Well Assignment:** Label wells as samples, calibrators, controls, or negatives (blank), and specify their dilution factors. Use Templates to quickly assign layouts that are repeated often or export the layout as a .csv file.
5. **Data Analysis:** Once **Image Processing** and **Well Assignment** are complete, select **Data Analysis**. Click **Perform Analysis** to generate charts, concentrations, and statistics based on optimal regression settings provided by the product definition.

PERFORMANCE CHARACTERISTICS

Precision

Intra-Assay Precision – 3 samples containing and/or spiked with various concentrations of each analyte were tested 20 times on 3 plates to assess intra-assay precision. The average %CV of all samples is reported.

Inter-Assay Precision – 3 samples containing and/or spiked with various concentrations of each analyte were tested on at least 10 individual assays by at least 3 users. The average %CV of all samples is reported.

Linearity

Samples containing and/or spiked with various concentrations of each analyte were diluted with sample diluent at dilutions of 1:2, 1:4, 1:8, 1:16 and assayed. The highest, lowest, and average dilution recoveries are reported.

Analyte	Average Intra-Assay Precision (%CV)	Average Inter-Assay Precision (%CV)	Linearity Range (4 samples 1:40-1:320 dilution)	Linearity Average (4 samples 1:40-1:320 dilution)
Serotype 1	11.1%	12.3%	80%-113%	97%
Serotype 3	14.4%	18.7%	83%-118%	99%
Serotype 4	7.9%	17.0%	85%-118%	99%
Serotype 5	9.8%	9.5%	92%-119%	102%
Serotype 6B	12.2%	13.4%	88%-123%	103%
Serotype 7F	8.3%	14.7%	90%-119%	100%
Serotype 8	8.9%	18.5%	86%-110%	97%
Serotype 9N	11.0%	19.9%	91%-115%	102%
Serotype 9V	10.0%	12.9%	89%-126%	102%
Serotype 12F	9.7%	11.2%	72%-129%	93%
Serotype 14	12.0%	19.7%	82%-117%	98%

Analyte	Average Intra-Assay Precision (%CV)	Average Inter-Assay Precision (%CV)	Linearity Range (4 samples 1:40-1:320 dilution)	Linearity Average (4 samples 1:40-1:320 dilution)
Serotype 18C	9.3%	15.4%	92%-108%	100%
Serotype 19F	10.3%	14.9%	88%-125%	100%
Serotype 23F	11.7%	14.6%	83%-118%	99%
Serotype 2	12.9%	20.2%	92%-118%	101%
Serotype 10A	10.8%	17.7%	83%-137%	106%
Serotype 11A	10.8%	9.2%	84%-132%	103%
Serotype 15B	13.3%	17.8%	82%-133%	105%
Serotype 17F	9.9%	13.6%	63%-118%	98%
Serotype 19A	7.2%	20.5%	85%-121%	101%
Serotype 20	7.9%	10.8%	81%-136%	102%
Serotype 22F	11.7%	20.0%	88%-116%	107%
Serotype 33F	9.1%	12.4%	88%-121%	101%

Interference

The following substances were shown not to interfere with the assay:

Bilirubin (< 0.01 mg/mL)

Intralipid (< 5 mg/mL)

Hemoglobin (< 5 mg/mL)

Human Anti-Mouse Antibodies (HAMA) (< 0.2 ug/mL)

APPENDIX A: PLATE WASHING METHODS

Automated Wash Method

1. Use a program that will aspirate and dispense 300-400 μL wash buffer.



2. Ensure that the plate washer has a program that will not scratch the bottom of the microplate but will prevent plate drying. This is critical to prevent damage to the capture antibody arrays. The simplest method to avoid plate drying is to leave a small, uniform amount (1-3 μL) of wash in the well after the final aspiration and add the next reagent to the plate as quickly as possible.
3. Leaving a uniform amount of wash in the wells can be accomplished with an automated washer by slightly increasing the aspiration height of the washer head.

For example:

Process	Distance	Steps on a Biotek ELX-405
Aspiration Height	3.81 mm	28
Aspiration Position	1.28 mm from center	-28
Dispense Height	15.24 mm	120

No soak or shake cycles are needed

4. Connect the prepared wash buffer to your automatic plate washer.

5. Run 1-2 priming cycles to make sure that the wash buffer is running through the plate washer. When the buffer has run through the machine, the waste will be foamy.
6. In a spare microtiter plate, dispense 100 μL wash buffer, ensure that all pins dispense uniformly, then aspirate and ensure that all pins aspirate uniformly. This will ensure that all pins are functioning.
7. Prime the plate washer 1 time before the first wash step. When running the assay, perform the wash 3 or 6 times according to the protocol.
8. Tap the plate upside down on a paper towel to remove any residual wash.
9. Proceed immediately to dispense the next solution so drying does not occur.

Multichannel Pipette Wash Method

10. Just prior to washing, pour the prepared wash buffer into a trough or tray.
11. After each incubation, flick the solution out of the plate over a waste container before starting the wash protocol.
12. Using a multichannel pipette, dispense 400 μL of wash buffer into each of the wells used in the test.
13. Aggressively flick the wash buffer out over a waste container.
14. This washes the plate 1 time. When the assay procedure calls for 3 or 6 washes, repeat steps 3-4 accordingly.
15. Tap the plate upside down on a paper towel to remove any residual wash.
16. Proceed immediately to dispense the next solution so drying does not occur.

ABBREVIATED PROTOCOL

Preparation

1. Install Q-View Software (page 8).
2. Set up the imager (page 8).
3. Set up microplate washer (page 15).
4. Reconstitute and prepare reagents (page 8).

Running the Assay

5. Prepare the calibration curve using the prepared Calibrator and Sample Diluent according to the In-Kit Certificate of Analysis (page 9).
6. Prepare the samples with Sample Diluent (page 9).
7. Load the calibration curve and samples into the plate. Incubate 30 minutes at room temperature (page 10).
8. Wash the plate 3 times, add the Detection Mix, and incubate 30 minutes at room temperature (page 10).
9. Wash the plate 6 times and add the prepared Substrate (page 10).
10. Capture and analyze image of the plate (page 11).

We take great care to ensure that customers have success using our products and services. If you have further questions about running the assay, data analysis, troubleshooting, or our products or services, please contact us at 1-888-QUANSYS (1-888-782-6797) or at support@quansysbio.com.

PLATE DIAGRAM

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

[illegible]

Q-PlexTM ARRAY

For technical support, email: support@quansysbio.com

482117HU

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