

Analysis of SARS-CoV-2 Antibodies in COVID-19 Convalescent Plasma using a Coronavirus Antigen Microarray

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Abstract

The current practice for diagnosis of COVID-19, based on SARS-CoV-2 PCR testing of pharyngeal or respiratory specimens in a symptomatic patient at high epidemiologic risk, likely underestimates the true prevalence of infection. Serologic methods can more accurately estimate the disease burden by detecting infections missed by the limited testing performed to date. Here, we describe the validation of a coronavirus antigen microarray containing immunologically significant antigens from SARS-CoV-2, in addition to SARS-CoV, MERS-CoV, common human coronavirus strains, and other common respiratory viruses. A comparison of antibody profiles detected on the array from control sera collected prior to the SARS-CoV-2 pandemic versus convalescent blood specimens from virologically confirmed COVID-19 cases demonstrates complete discrimination of these two groups. This array can be used as a diagnostic tool, as an epidemiologic tool to more accurately estimate the disease burden of COVID-19, and as a research tool to correlate antibody responses with clinical outcomes.

Background

COVID-19 caused by the SARS-CoV-2 virus is a worldwide pandemic with significant morbidity and mortality estimates from 1-4% of confirmed cases¹. The current case definition for confirmed SARS-CoV-2 infection relies on PCR-positive pharyngeal or respiratory specimens, with testing largely determined by presence of fever or respiratory symptoms in an individual at high epidemiologic risk. However, this case definition likely underestimates true prevalence, as individuals who develop subclinical infection that does not produce fever or respiratory symptoms are unlikely to be tested, and testing by PCR of pharyngeal or respiratory specimens is only around 60-80% sensitive depending on sampling location and technique and the patient's viral load². Widespread testing within the United States is also severely limited by the lack of available testing kits and testing capacity limitations of available public and private laboratories. Therefore, the true prevalence of SARS-CoV-2 infection is likely much higher than currently reported case numbers would indicate.

Serology can play an important role in defining the true prevalence of COVID-19, particularly for subclinical infection². Early studies of serology demonstrate high sensitivity to detect confirmed SARS-CoV-2 infection, with antibodies to virus detected approximately 1 to 2 weeks after symptom onset³. Unlike PCR positivity, SARS-CoV-2 antibodies are detectable throughout the disease course and persist indefinitely⁴. Multiple serologic tests have been developed for COVID-19⁵ including a recently FDA-approved lateral flow assay. However, these tests are limited to detection of antibodies against one or two antigens, and cross-reactivity with antibodies to other human coronaviruses that are present in all adults⁶ is currently unknown. Prior use of serology for detection of emerging coronaviruses focused on antibodies against the spike (S) protein, particularly the S1 domain, and the nucleocapsid (N) protein⁷. However, the optimal set of antigens to detect strain-specific coronavirus antibodies remains unknown.

Protein microarray technology can be used to detect antibodies of multiple isotypes against hundreds of antigens in a high throughput manner^{8,9} so is well suited to serologic surveillance studies. This technology, which has previously been applied to other emerging coronaviruses¹⁰, is based on detection of binding antibodies, which are well-correlated with neutralizing antibodies¹¹ but do not require viral culture in biosafety level 3 facilities. Recently, our group developed a coronavirus antigen microarray (CoVAM) that includes antigens from SARS-CoV-2 and tested it on human sera collected prior to the pandemic to demonstrate low cross-reactivity with antibodies from human coronaviruses that cause the common cold, particularly for the S1 domain². Here, we further validate this methodology using convalescent blood specimens from COVID-19 cases confirmed by positive SARS-CoV-2 PCR.

Methodology

Specimen Collection

The SARS-CoV-2 convalescent blood specimens from nasopharyngeal SARS-CoV-2 PCR-positive individuals were collected from different sources to increase the number of positive specimens available for analysis. Two sera and plasma samples were obtained from acute COVID-19 patients from the Oregon Health Sciences University Hospital

(OHSU), Portland, OR. These were sourced from discarded clinical laboratory specimens exempted from informed consent and IRB approval under condition of patient anonymity. Four plasma samples were obtained from outpatients of the University Hospital Basel, University of Basel, Basel, Switzerland. These patients were screened in accordance with Swiss regulations on blood donation and approved as plasma donors according to the Blood Transfusion Service of the Swiss Red Cross with informed consent. These donors were diagnosed with COVID-19 based on SARS-CoV-2 positive nasopharyngeal swab PCR tests. At time of plasma donation, each had two negative nasopharyngeal swab SARS-CoV-2 PCR- tests and negative SARS-CoV-2 PCR tests in blood, and they were qualified as plasma donors. Plasma was collected from these convalescent donors at the Regional Blood Transfusion Service of the Swiss Red Cross in accordance with national regulations. One convalescent plasma was isolated from a large-volume apheresis collection following standard protocol from a documented recovered COVID-19 blood donor who was more than 28 days post symptomatic.

The negative control sera used in this study were collected between November 2018 and May 2019 for a larger study where residents of a college resident community in the Eastern United States were monitored prospectively to identify acute respiratory infection (ARI) cases using questionnaires and RT-qPCR, so as to characterize contagious phenotypes including social connections, built environment, and immunologic phenotypes¹². A total of 144 de-identified blood specimens were tested on the CoV antigen microarray. Electronic informed consents including future research use authorization was obtained under protocols approved by the Institutional Review Boards (IRBs) of the University of Maryland and the Department of Navy Human Research Protections Office.

Coronavirus Antigen Microarray

The coronavirus antigen microarray used in this investigation includes 67 antigens across subtypes expressed in either baculovirus or HEK-293 cells (Table 1). These antigens were provided by Sino Inc. (Wayne, PA) as either catalog products or custom synthesis service products. The antigens were printed onto microarrays, probed with human sera, and analyzed as previously described^{9,13,14}.

Briefly, lyophilized antigens were reconstituted to a concentration of 0.1 mg/mL in phosphate-buffered saline (PBS) with 0.001% Tween-20 (T-PBS) and then printed onto nitrocellulose-coated slides from Grace Bio Labs (GBL, Bend, OR) using an OmniGrid 100 microarray printer (GeneMachines). The microarray slides were probed with human sera diluted 1:100 in 1x GVS Fast Blocking Buffer (Fischer Scientific) overnight at 4°C, washed with T-TBS buffer (20 mM Tris-HCl, 150 mM NaCl, 0.05% Tween-20 in ddH₂O adjusted to pH 7.5 and filtered) 3 times for 5 minutes each, labeled with secondary antibodies to human IgA and IgG conjugated to quantum dot fluorophores for 2 hours at room temperature, and then washed with T-TBS 3 times for 5 minutes each and dried. The slides were imaged using ArrayCam imager (Grace Bio Labs, Bend, OR) to measure background-subtracted median spot fluorescence. Non-specific binding of secondary antibodies was subtracted using saline control. Mean fluorescence of the 4 replicate spots for each antigen was used for analysis.

Statistical Analyses

The mean fluorescence intensity (MFI) of each antigen was determined by the average of the median fluorescence signal of four replicate spots. The fluorescence signal for each spot was determined by its signal intensity subtracted by the background fluorescence. Antigens containing a human Fc tag were removed from the analysis, as the secondary antibodies used for quantification are known to bind to human Fc; non-human Fc tag did not interfere with the assay. All statistical analyses were conducted using R version 3.6.3 (R Foundation for Statistical Computing, Vienna, Austria).

MFI was normalized using the *normalize.quantiles.use.target* function from the *proprocessCore* package (version 1.48.0). As a target for normalization, a vector containing the median MFI for IgG or IgA was constructed. Descriptive statistics were used to summarize the IgA and IgG reactivity measured as MFI. Wilcoxon Rank Sum tests with $p < 0.05$ corrected for multiple comparisons were used to compare the mean differences between groups.

Receiver Operating Characteristic Area Under the Curve (ROC AUC) values for each antigen were calculated by comparing positive and negative specimens using the *pROC* package (version 1.16.2). Antigens were ranked based on their ROC AUC values, and high performing antigens with ROC AUC ≥ 0.95 were identified. Data visualization was performed using the *ggplot2* package (version 3.3.0).

Results

Discrimination of SARS-CoV-2 Convalescent Plasma using Coronavirus Antigen Microarray

The coronavirus antigen microarray was used to detect IgG and IgA antibodies against a panel of antigens, including coronavirus spike protein (S) as separated receptor-binding (RBD), S1, and S2 domains or whole protein (S1+S2) and nucleocapsid protein (NP), from multiple coronaviruses including SARS-CoV-2, SARS-CoV, MERS-CoV, and the four common cold coronaviruses (HKU1, OC43, NL63, 229E) as listed in Table 1. To determine the antibody profile of SARS-CoV-2 infection, the differential reactivity to these antigens was evaluated for SARS-CoV-2 convalescent plasma from PCR-positive individuals (positive group) and sera collected prior to the COVID-19 pandemic from naïve individuals (negative group) as shown in Figure 1.

The positive group demonstrates high IgG reactivity to SARS-CoV-2 NP, S2, and S1+S2 antigens and moderate IgG reactivity to SARS-CoV-2 S1 and RBD antigens, while the negative group demonstrates low IgG cross-reactivity to SARS-CoV-2 S1+S2 and no cross-reactivity to other SARS-CoV-2 antigens (Figure 2). The positive group also demonstrates high IgG cross-reactivity with SARS-CoV NP and MERS-CoV S2 and S1+S2 antigens, while the negative group demonstrates low cross-reactivity with MERS-CoV S1+S2 and no cross-reactivity with other SARS-CoV and MERS-CoV antigens. The two groups do not differ significantly in reactivity to common cold coronaviruses and other seasonally circulating respiratory viruses.

Similar trends are observed for IgA but with lower reactivity overall (Figure 3). The positive group again demonstrates high IgA reactivity to SARS-CoV-2 NP, S2, and

S1+S2 and moderate IgA reactivity to SARS-CoV-2 S1 with high IgA cross-reactivity to SARS-CoV NP, while the negative group demonstrates low IgA cross-reactivity to all SARS-CoV-2, SARS-CoV, and MERS-CoV antigens.

Selection of High-Performing Antigens to Detect SARS-CoV-2 Infection

Each coronavirus antigen was evaluated for performance in discriminating the positive group from the negative group across a full range of assay cutoff values to generate Receiver Operating Characteristic (ROC) curves for which Area Under Curve (ROC AUC) was measured (Figure 4). High-performing antigens for detection of IgG or IgA defined by ROC AUC ≥ 0.95 included all SARS-CoV-2 antigens and MERS-CoV S2 for IgG and SARS-CoV-2 S2 and S1+S2 for IgA (Table 2). Each of these antigens discriminated between the positive group and the negative group with high significance (Figure 5).

Discussion

This study reveals several insights into the antibody response to SARS-CoV-2 infection. The antibody profiles of naïve individuals include high IgG reactivity to common cold coronaviruses with low-level cross-reactivity with S2 domains from SARS-CoV-2 and other epidemic coronaviruses, which is not surprising given the high degree of sequence homology and previously observed serologic cross-reactivity¹⁵ between S2 domains of betacoronaviruses, a group that includes SARS-CoV-2, SARS-CoV, MERS, and common cold coronaviruses HKU1 and OC43. However, naïve individuals do not show cross-reactivity to other SARS-CoV-2 antigens. Even for the nucleocapsid protein, which also has high sequence homology between betacoronaviruses, cross-reactivity is only seen between SARS-CoV-2 and SARS-CoV and not with MERS-CoV or common cold coronaviruses. In addition, the quantitative difference between high antibody reactivity to SARS-CoV-2 S2 in the positive group and low-level antibody cross-reactivity in the negative group is large enough that these antigens still discriminate these groups with high significance.

This study also informs antigen selection and design for population surveillance and clinical diagnostic assays and vaccine development. The observation that naïve individuals with antibodies to common cold coronaviruses do not show cross-reactivity to SARS-CoV-2 nucleocapsid protein dispels concerns that the high sequence homology of this protein across betacoronaviruses would impair its performance as a diagnostic or vaccine antigen. The low-level antibody cross-reactivity of naïve individuals for SARS-CoV-2 spike protein containing S2 domain may not preclude its use as a diagnostic antigen given large quantitative difference in antibody reactivity between positive and negative groups, but this cross-reactivity may influence response to vaccination with spike protein antigens containing the S2 domain.

The coronavirus antigen microarray can be useful both as an epidemiologic tool and as a research tool. The high throughput detection of SARS-CoV-2-specific antibody profiles that reliably distinguish COVID-19 cases from negative controls can be applied to large-scale population surveillance studies for a more accurate estimation of the true prevalence of disease than can be achieved with symptom-based PCR testing. In

addition, detection of these antibodies in SARS-CoV-2 convalescent plasma donations can provide validation prior to clinical use for passive immunization. The variation in the SARS-CoV-2 antibody profiles among acute and convalescent donors suggests that epitope characterization of convalescent donor plasma will be informative for evaluation of passive immune therapy efficacy in COVID-19 patients. The central role of inflammation in the pathogenesis of severe COVID-19¹⁶ can be more closely studied by analyzing both strain-specific and cross-reactive antibody responses, particularly to test hypotheses regarding antibody-dependent enhancement with critical implications for vaccine development¹⁷.

Conclusions

A coronavirus antigen microarray containing a panel of antigens from SARS-CoV-2 in addition to other human coronaviruses was able to reliably distinguish convalescent plasma of PCR-positive COVID-19 cases from negative control sera collected prior to the pandemic by detecting both strain-specific and cross-reactive antibodies. Further studies are needed to apply this methodology to large-scale serologic surveillance studies and to correlate specific antibody responses with clinical outcomes.

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Author Disclosures

The coronavirus antigen microarray is intellectual property of the Regents of the University of California that is licensed for commercialization to Nanommune Inc. (Irvine, CA), a private company for which Philip L. Felgner is the largest shareholder and several co-authors (de Assis, Jain, Nakajima, Jasinskis, Obiero, Davies, and Khan) also own shares. Nanommune Inc. has a business partnership with Sino Biological Inc. (Beijing, China) which expressed and purified the antigens used in this study. The convalescent plasma used in this study was collected for clinical use by independent blood centers using licensed plasma or platelet processing systems manufactured by Cerus Corporation, for which multiple authors (Corash, Bagri) are shareholders and employees. Manuel Battegay, Andreas Buser and Andreas Holbro are employees of the University of Basel and have no conflicts of interest.

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Figures and Tables

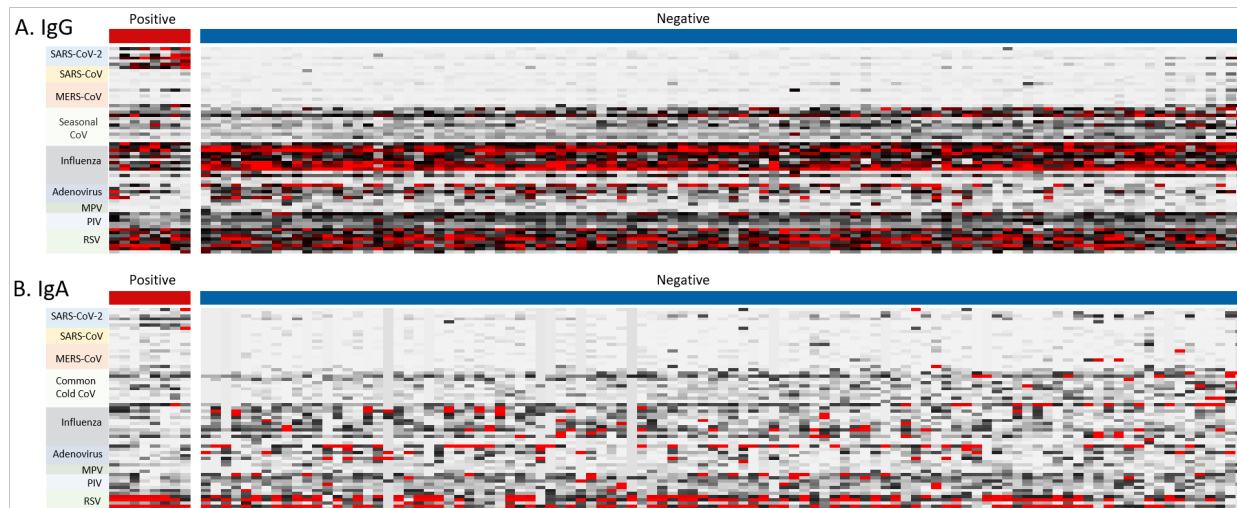


Figure 1. Heatmap for coronavirus antigen microarray. The heatmap shows IgG (A) and IgA (B) reactivity measured as mean fluorescence intensity across four replicates, against each antigen organized into rows color coded by virus, for sera organized into columns classified as positive (convalescent from PCR-positive individuals) or negative (prior to pandemic from naïve individuals). Reactivity is represented by color (white = low, black = mid, red = high).

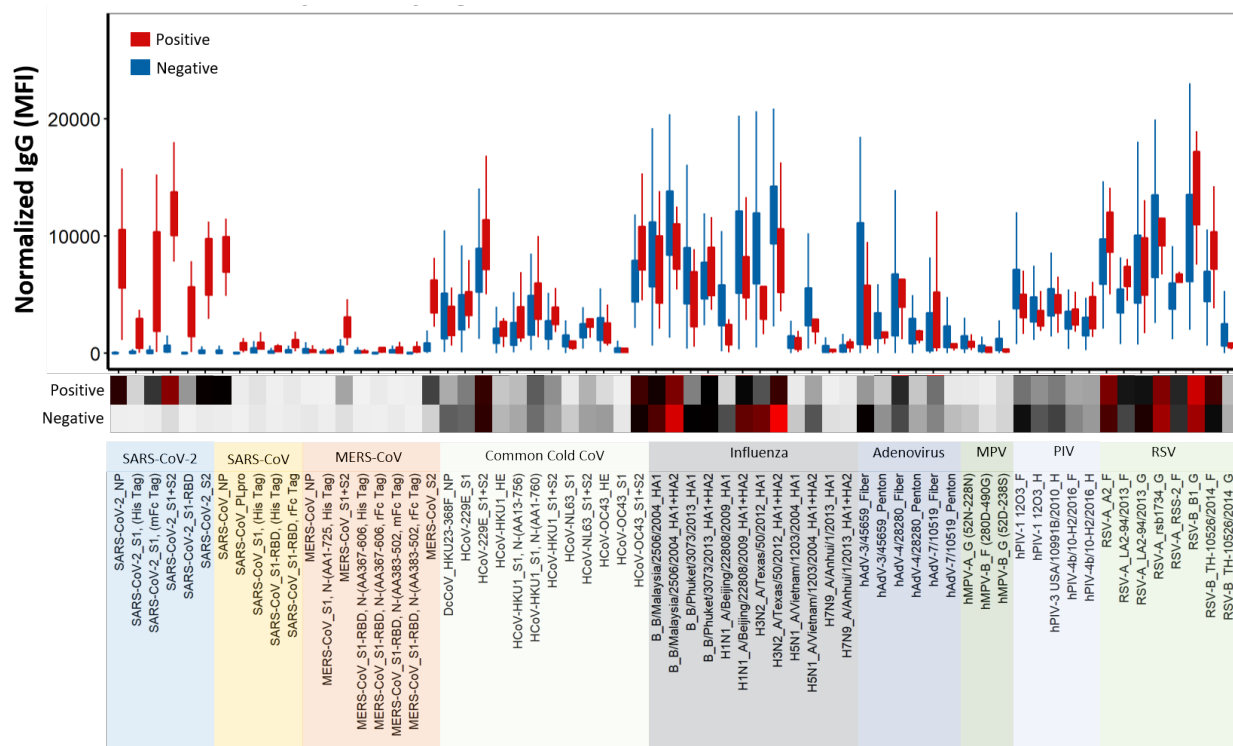


Figure 2. Normalized IgG reactivity of positive and negative sera on coronavirus antigen microarray. The plot show IgG reactivity against each antigen measured as mean fluorescence intensity (MFI) with full range (bars) and interquartile range (boxes) for convalescent sera from PCR-positive individuals (positive, red) and sera from naïve individuals prior to pandemic (negative, blue). Below the plot, the heatmap shows average reactivity for each group (white = low, black = mid, red = high). The antigen labels are color coded for respiratory virus group.

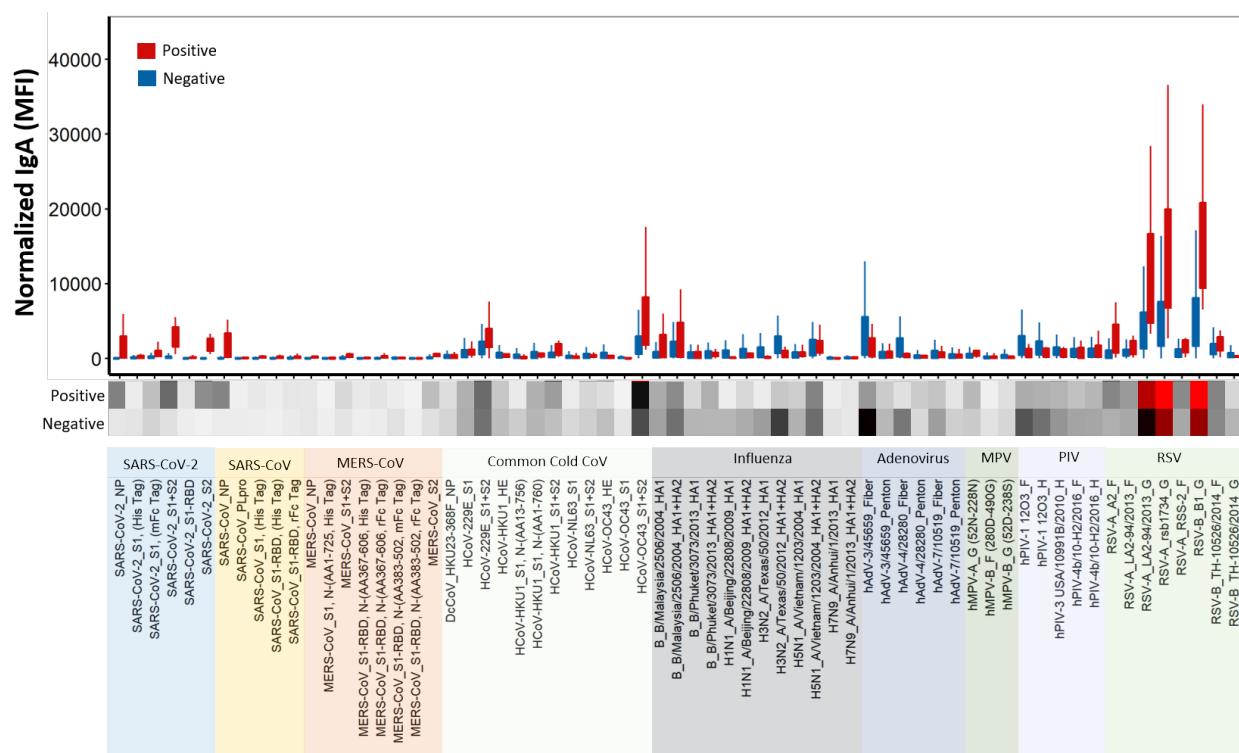


Figure 3. Normalized IgA reactivity of positive and negative sera on coronavirus antigen microarray. The plot show IgG reactivity against each antigen measured as mean fluorescence intensity (MFI) with full range (bars) and interquartile range (boxes) for convalescent sera from PCR-positive individuals (positive, red) and sera from naïve individuals prior to pandemic (negative, blue). Below the plot, the heatmap shows average reactivity for each group (white = low, black = mid, red = high). The antigen labels are color coded for respiratory virus group.

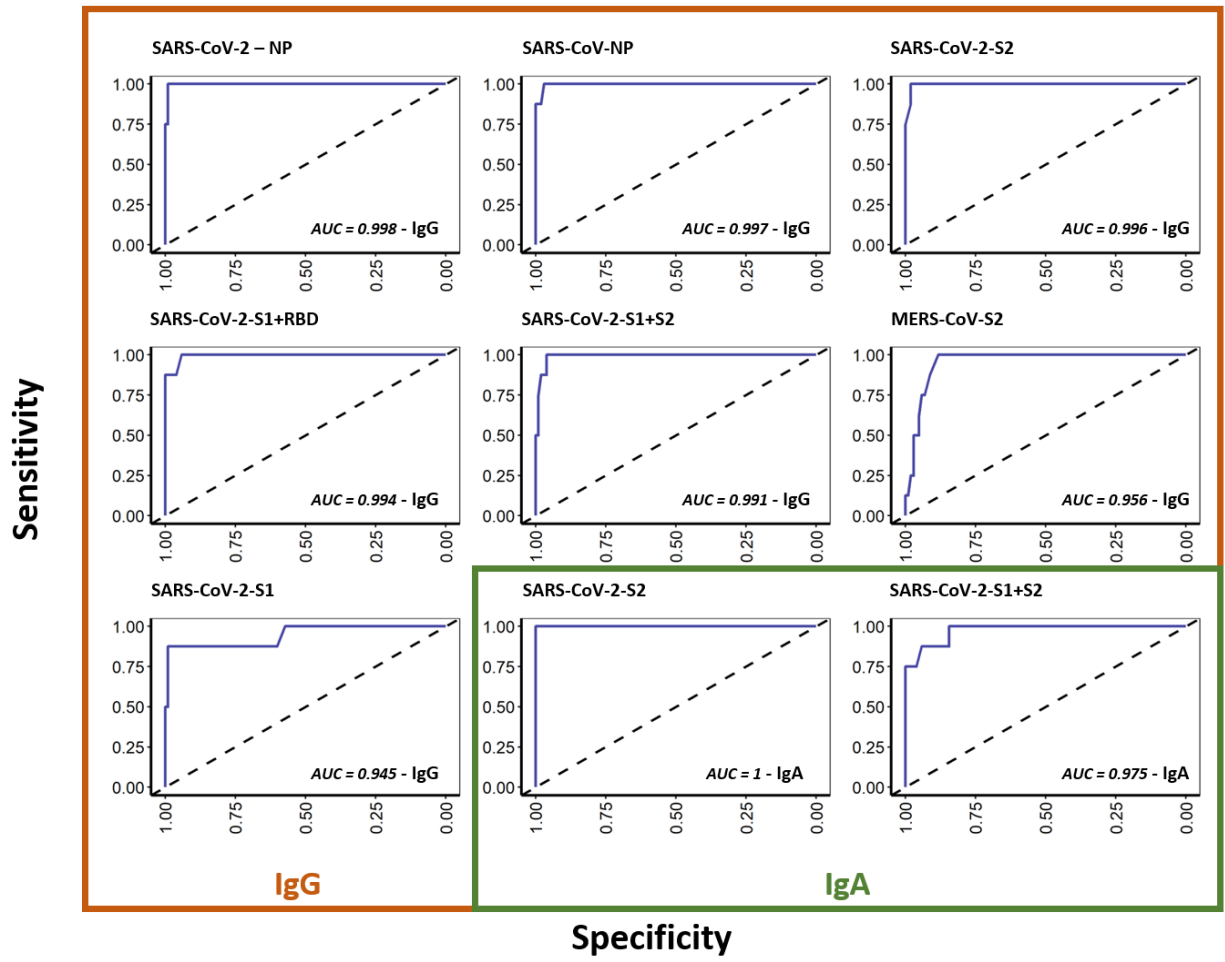


Figure 4. ROC curves for high-performing antigens. ROC curves showing sensitivity versus specificity for discrimination of positive and negative sera were derived for each individual high performing antigen (ROC AUC \geq 0.95) for both IgG and IgA (solid blue line) and compared to no discrimination (ROC AUC = 0.5, dashed black line).

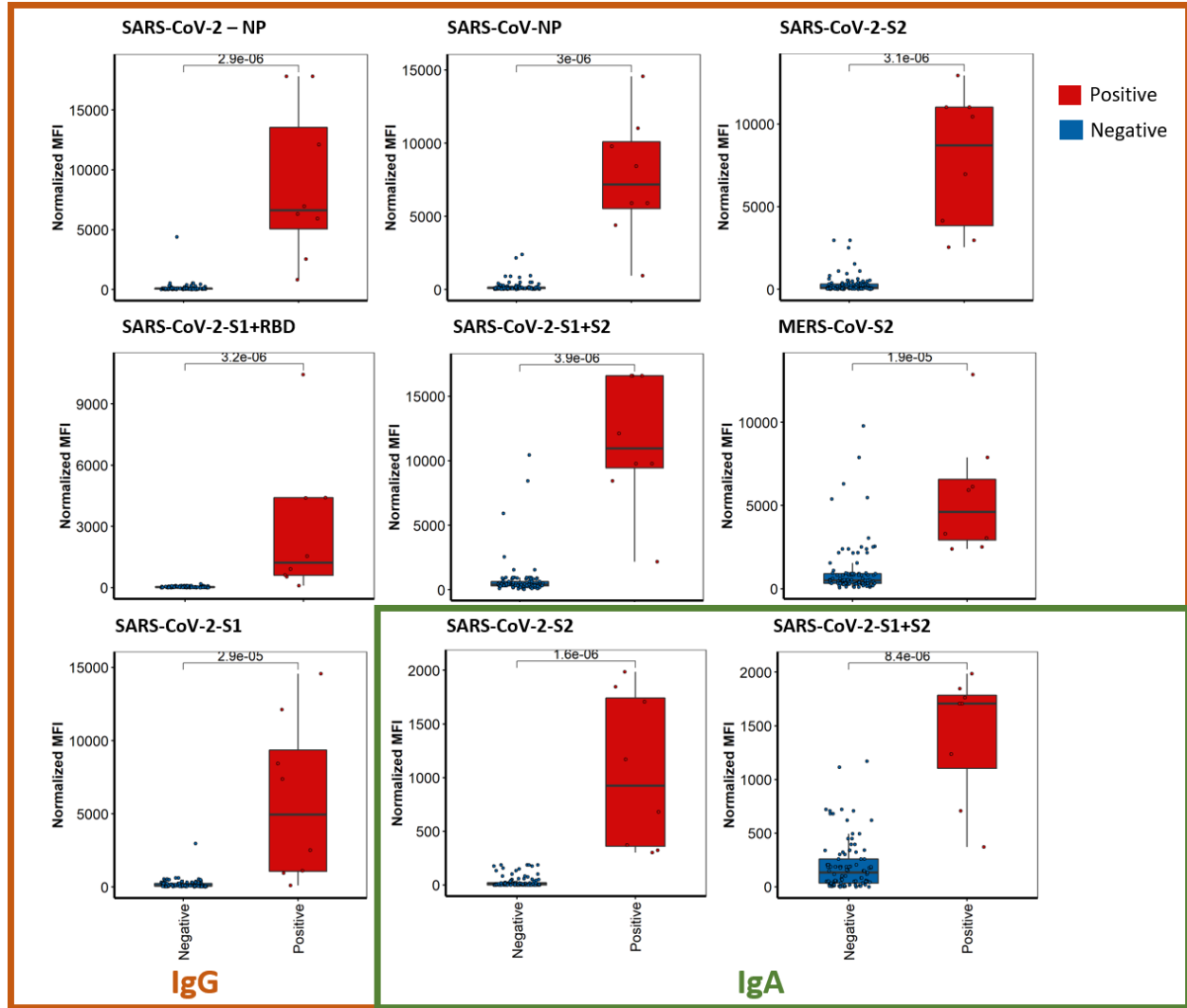


Figure 5. Normalized antibody reactivity of positive and negative sera for high-performing antigens. IgG and IgA reactivity against each high-performing antigen (ROC AUC ≥ 0.95) measured as mean fluorescence intensity (MFI) for convalescent sera from PCR-positive individuals (positive, red) and sera from naïve individuals prior to pandemic (negative, blue) are shown as box plots, including full range (bars), interquartile range (boxes), median (black line), and individual sera (dots) with p-values for each antigen calculated by Wilcoxon Rank Sum test.

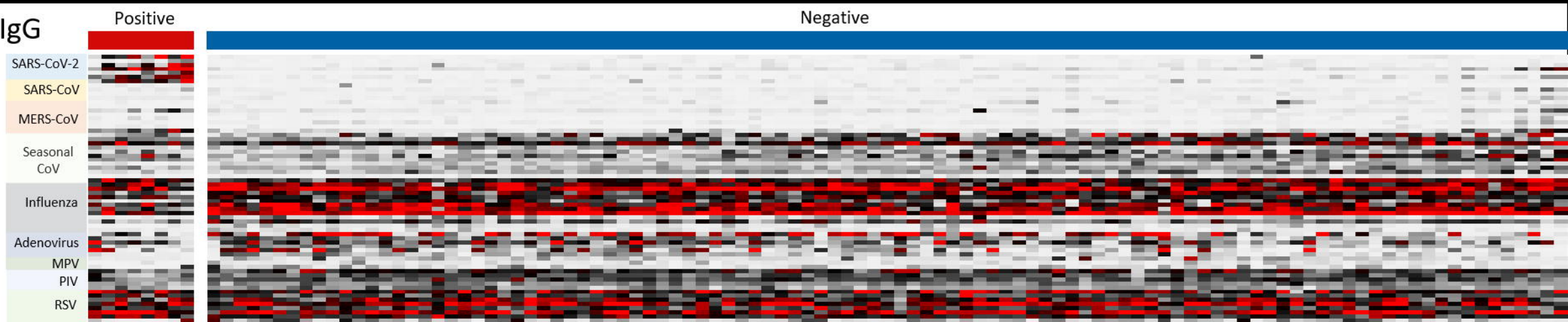
Virus	Subtype	Strain	Protein	GenBank	Expression	Construct	Source	Cat. No.
CoV	Beta	SARS-CoV-2	NP		Baculovirus	N-(AA)-His-C	Sino	40588-V08B
CoV	Beta	SARS-CoV-2	S1-RBD		HEK293	N-(AA)-mFc-C	Sino	40592-V05H
CoV	Beta	SARS-CoV-2	S1		HEK293	N-(AA)-His-C	Sino	40591-V08H
CoV	Beta	SARS-CoV-2	S1		HEK293	N-(AA)-mFc-C	Sino	40591-V02H
CoV	Beta	SARS-CoV-2	S1		HEK293	N-(AA)-Fc-C	Sino	40591-V05H1
CoV	Beta	SARS-CoV-2	S2		Baculovirus	N-(AA)-His-C	Sino	40590-V08B
CoV	Beta	SARS-CoV-2	S1+S2		Baculovirus	N-(AA)-His-C	Sino	40589-V08B1
CoV	Beta	SARS	PLpro	AAX16193.1	E. coli	N-(AA1541-1859)-His-C	Sino	40524-V08E
CoV	Beta	SARS	S1-RBD	AAX16192.1	Baculovirus	N-(AA306-527)-Fc-C	Sino	40150-V31B2
CoV	Beta	SARS	S1-RBD	AAX16192.1	Baculovirus	N-(AA306-527)-His-C	Sino	40150-V08B2
CoV	Beta	SARS	S1	AAX16192.1	Baculovirus	N-(AA1-667)-His-C	Sino	40150-V08B1
CoV	Beta	SARS	NP	NP_828858.1	Baculovirus	N-(AA1-422)-His-C	Sino	40143-V08B
CoV	Beta	MERS	NP	AFS88943.1	Baculovirus	N-(AA1-413)-His-C	Sino	40068-V08B
CoV	Beta	MERS	S1-RBD	AFS88936.1	Baculovirus	N-(AA383-502)-Fc-C	Sino	40071-V05B
CoV	Beta	MERS	S1-RBD	AFS88936.1	Baculovirus	N-(AA383-502)-rFc-C	Sino	40071-V31B
CoV	Beta	MERS	S1-RBD	AFS88936.1	Baculovirus	N-(AA367-606)-rFc-C	Sino	40071-V31B1
CoV	Beta	MERS	S1-RBD	AFS88936.1	Baculovirus	N-(AA367-606)-His-C	Sino	40071-V08B1
CoV	Beta	MERS	S1	AFS88936.1	HEK293	N-(AA1-725)-His-C	Sino	40069-V08H
CoV	Beta	MERS	S1	AFS88936.1	Baculovirus	N-(AA1-725)-His-C	Sino	40069-V08B1
CoV	Beta	MERS	S1+S2	AFS88936.1	Baculovirus	N-(AA1-1297)-His-C	Sino	40069-V08B
CoV	Beta	MERS	S2	AFS88936.1	Baculovirus	N-(AA726-1296)-His-C	Sino	40070-V08B
CoV	Alpha	NL63	S1	A0A1L2YV18	HEK293	N-(AA19-717)-His-C	Sino	40600-V08H
CoV	Alpha	NL63	S1+S2	A0A1L2YV18	Baculovirus	N-(AA19-1296)-His-C	Sino	40604-V08B
CoV	Alpha	229E	S1	A0A1L7B942	HEK293	N-(AA16-536)-His-C	Sino	40601-v08H
CoV	Alpha	229E	S1+S2	A0A1L7B942	Baculovirus	N-(AA16-1115)-His-C	Sino	40605-V08B
CoV	Beta	HKU1	S1	YP_173238.1	HEK293	N-(AA1-760)-His-C	Sino	40021-V08H
CoV	Beta	HKU1	S1	Q0ZME7	HEK293	N-(AA13-756)-His-C	Sino	40602-V08H
CoV	Beta	HKU1	S1+S2	Q0ZME7	Baculovirus	N-(AA13-1295)-His-C	Sino	40606-V08B
CoV	Beta	HKU1	HE	Q0ZME7	HEK293	N-(AA16-394)-His-C	Sino	Custom
CoV	Beta	HKU23-368F	NP	AHN64796.1	HEK293	N-(AA1-448)-His-C	Sino	40458-V08B
CoV	Beta	OC43	S1	AVR40344.1	HEK293	N-(AA13-533)-His-C	Sino	Custom
CoV	Beta	OC43	S1+S2	AVR40344.1	Baculovirus	N-(AA13-1304)-His-C	Sino	40607-V08B
CoV	Beta	OC43	HE	ATN39879.2	HEK293	N-(AA16-394)-His-C	Sino	40603-V08H
RSV	A	LA2-94/2013	F	A0A023RA53	Baculovirus	N-(AA1-526)-His-C	Sino	Custom
RSV	A	LA2-94/2013	G	A0A076FRQ0	HEK293	N-(AA64-321)-His-C	Sino	Custom
RSV	A	A2	F		Baculovirus	N-(AA1-529)-His-C	Sino	11049-V08B
RSV	A	rsb1734	G		HEK293	N-(AA66-297)-His-C	Sino	11070-V08H
RSV	A	RSS-2	F		Baculovirus	N-(AA1-529)-His-C	Sino	40037-V08B
RSV	B	TH-10526/2014	F	K7WL19	Baculovirus	N-(AA1-525)-His-C	Sino	Custom
RSV	B	TH-10526/2014	G	A0A142MLK4	HEK293	N-(AA64-310)-His-C	Sino	Custom
RSV	B	B1	G		HEK293	N-(AA67-299)-His-C	Sino	13029-V08H
MPV	A	PER/CFI0320/2010/A	G		HEK293	52N-228N-His	Sino	Custom
MPV	B	PER/CFI0466/2010/B	G		HEK293	52D-238S-His	Sino	Custom
MPV	B	PER/CFI0320/2010/A	F		HEK293	280D-490G-His	Sino	Custom
PIV	1	12O3	F	A0A1V0E1X5	Baculovirus	N-(AA22-497)-His-C	Sino	Custom
PIV	1	12O3	H	A0A1B2CW87	Baculovirus	N-His-(AA60-575)-C	Sino	Custom
PIV	3	USA/10991B/2010	H	T1UD13	Baculovirus	N-His-(AA55-575)-C	Sino	Custom
PIV	4	hPIV-4b/10-H2/2016	F	A0A1V0E1N6	Baculovirus	N-(AA22-486)-His-C	Sino	Custom
PIV	4	hPIV-4b/10-H2/2016	H	A0A1V0E1N4	Baculovirus	N-His-(AA48-575)-C	Sino	Custom
Adeno	3	hAdV-3/45659	Fiber	P04501	E. coli	N-His-[Prot]-C	Sino	Custom
Adeno	3	hAdV-3/45659	Penton	Q2Y0H9	Baculovirus	N-His-[Prot]-C	Sino	Custom
Adeno	4	hAdV-4/28280	Fiber	P36844	Baculovirus	N-[Prot]-His-C	Sino	Custom
Adeno	4	hAdV-4/28280	Penton	Q2KSF3	Baculovirus	N-[Prot]-His-C	Sino	Custom
Adeno	7	Adeno7_10519	Fiber	P15141	Baculovirus	N-His-[Prot]-C	Sino	Custom
Adeno	7	Adeno7_10519	Penton	Q2KS58	Baculovirus	N-[Prot]-His-C	Sino	Custom
Flu	H1N1	A/Beijing/22808/2009	HA1	ADD64203.1	HEK293	N-(AA1-344)-His-C	Sino	40035-V08H1
Flu	H1N1	A/Beijing/22808/2009	HA1+HA2	ADD64203.1	HEK293	N-(AA1-529)-His-C	Sino	40035-V08H
Flu	H3N2	A/Texas/50/2012	HA1	AGL07159.1	HEK293	N-(AA1-345)-His-C	Sino	40354-V08H1
Flu	H3N2	A/Texas/50/2012	HA1+HA2	AGL07159.1	Baculovirus	N-(AA1-530)-His-C	Sino	40354-V08B
Flu	B	B/Malaysia/2506/2004	HA1	CO05957.1	HEK293	N-(AA1-362)-His-C	Sino	11716-V08H1
Flu	B	B/Malaysia/2506/2004	HA1+HA2	CO05957.1	HEK293	N-(AA1-556)-His-C	Sino	11716-V08H
Flu	B	B/Phuket/3073/2013	HA1	EPI529345	HEK293	N-(AA1-361)-His-C	Sino	40498-V08H1
Flu	B	B/Phuket/3073/2013	HA1+HA2	EPI529345	Baculovirus	N-(AA1-547)-His-C	Sino	40498-V08B
Flu	H5N1	A/Vietnam/1203/2004	HA1	AAW80717.1	HEK293	(AA1-342)-mFcg1-His	Sino	10003-V06H1
Flu	H5N1	A/Vietnam/1203/2004	HA1+HA2	AAW80717.1	HEK293	(AA1-531)-mFcg1-His	Sino	10003-V06H3
Flu	H7N9	A/Anhui/1/2013	HA1	AGJ51953.1	HEK293	N-(AA1-338)-His-C	Sino	40103-V08H1
Flu	H7N9	A/Anhui/1/2013	HA1+HA2	AGJ51953.1	HEK293	N-(AA1-524)-His-C	Sino	40103-V08H

Table 1. Content of coronavirus antigen microarray. The virus group, subtype, and strain, protein, GenBank identification where available, expression system, gene construct, and vendor source and catalog number are shown for each antigen.

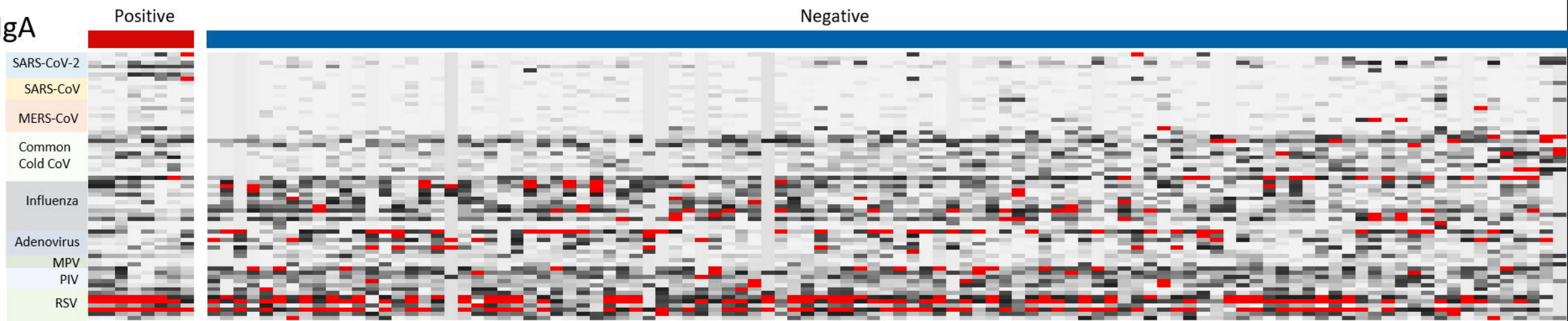
IgG Rank	Epitope	AUC	IgA Rank	Antigen	AUC
1	SARS-CoV-2_NP	0.998	1	SARS-CoV-2_S2	1
2	SARS-CoV_NP	0.997	2	SARS-CoV-2_S1+S2	0.975
3	SARS-CoV-2_S2	0.996	3	MERS-CoV_S1+S2	0.791
4	SARS-CoV-2_S1-RBD	0.994	4	SARS-CoV-2_NP	0.753
5	SARS-CoV-2_S1+S2	0.991	5	MERS-CoV_S2	0.752
6	MERS-CoV_S2	0.956	6	MERS-CoV_S1-RBD, N-(AA383-502, rFc Tag)	0.751
7	SARS-CoV-2_S1, (mFc Tag)	0.945	7	SARS-CoV-2_S1-RBD	0.735
8	MERS-CoV_S1-RBD, N-(AA367-606, His Tag)	0.894	8	MERS-CoV_S1, N-(AA1-725, His Tag)	0.734
9	MERS-CoV_S1, N-(AA1-725, His Tag)	0.845	9	SARS-CoV-2_S1, (mFc Tag)	0.72
10	MERS-CoV_S1+S2	0.836	10	SARS-CoV_NP	0.714
11	MERS-CoV_NP	0.829	11	SARS-CoV_S1-RBD, rFc Tag	0.684
12	SARS-CoV_S1-RBD, rFc Tag	0.763	12	MERS-CoV_S1-RBD, N-(AA367-606, rFc Tag)	0.65
13	MERS-CoV_S1-RBD, N-(AA383-502, mFc Tag)	0.758	13	MERS-CoV_S1-RBD, N-(AA367-606, His Tag)	0.632
14	SARS-CoV_PLpro	0.736	14	SARS-CoV-2_S1, (His Tag)	0.631
15	SARS-CoV-2_S1, (His Tag)	0.712	15	SARS-CoV_PLpro	0.594
16	MERS-CoV_S1-RBD, N-(AA367-606, rFc Tag)	0.709	16	SARS-CoV_S1, (His Tag)	0.577
17	MERS-CoV_S1-RBD, N-(AA383-502, rFc Tag)	0.669	17	MERS-CoV_NP	0.559
18	SARS-CoV_S1, (His Tag)	0.576	18	SARS-CoV_S1-RBD, (His Tag)	0.535
19	SARS-CoV_S1-RBD, (His Tag)	0.536	19	MERS-CoV_S1-RBD, N-(AA383-502, mFc Tag)	0.533

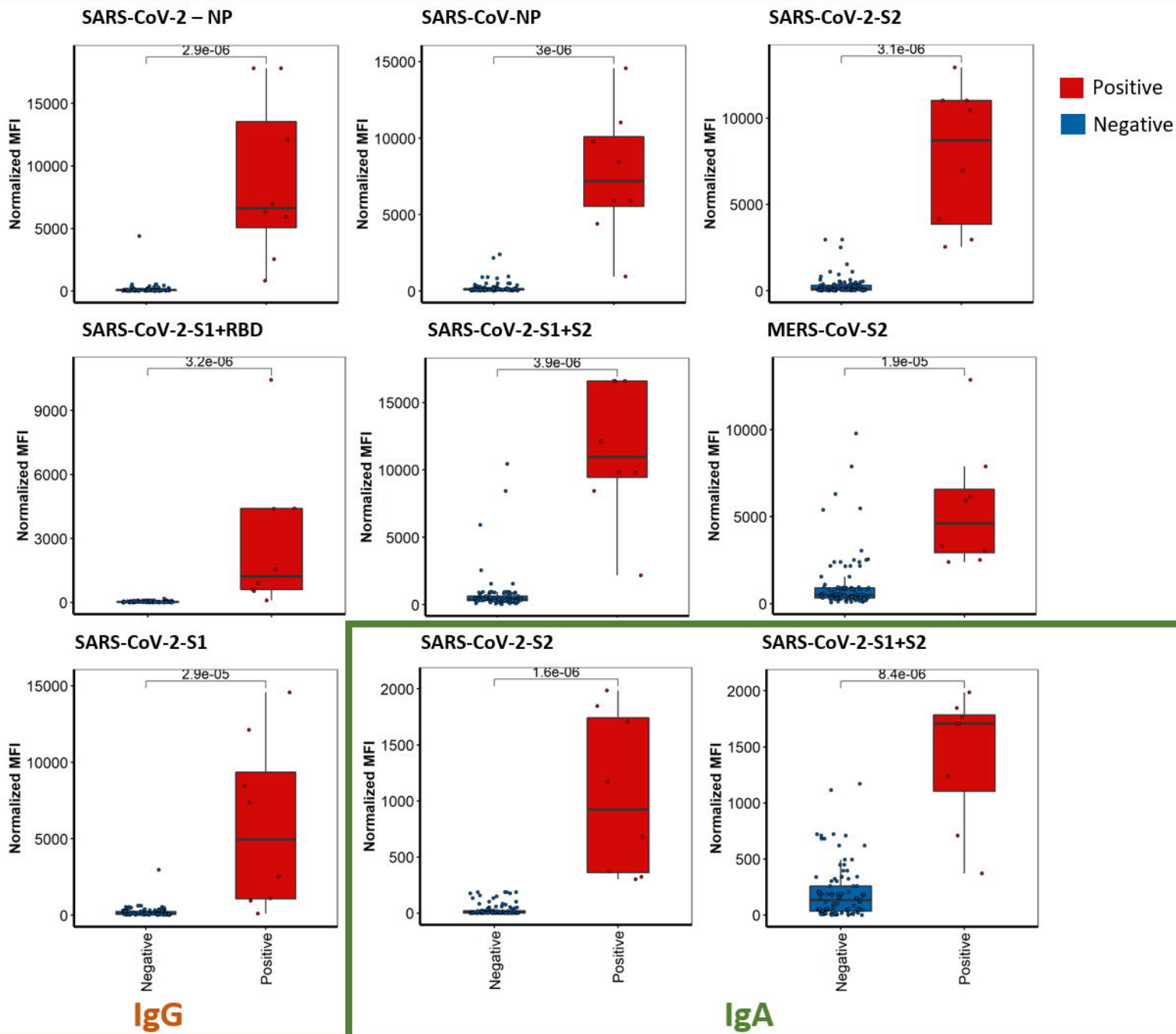
Table 2. Receiver Operating Characteristic Area Under Curve (ROC AUC) for SARS-CoV-2, SARS-CoV, and MERS-CoV antigens. ROC AUC values for discrimination of positive and negative sera were derived for each individual antigen for both IgG and IgA and ranked, and high-performing antigens with ROC AUC ≥ 0.95 are indicated above the lines.

A. IgG



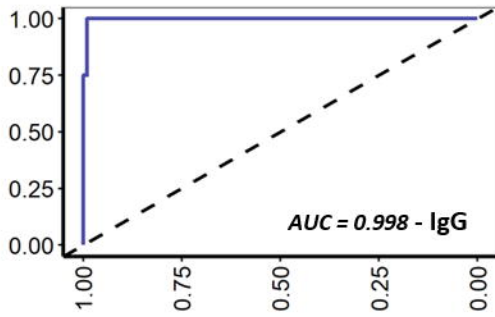
B. IgA



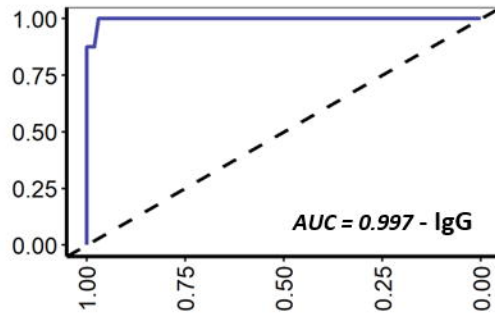


Sensitivity

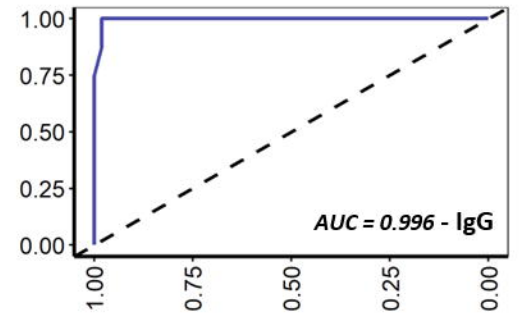
SARS-CoV-2 – NP



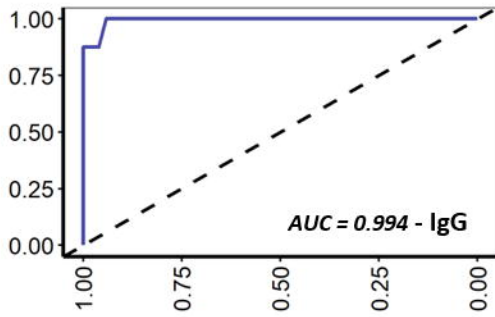
SARS-CoV-NP



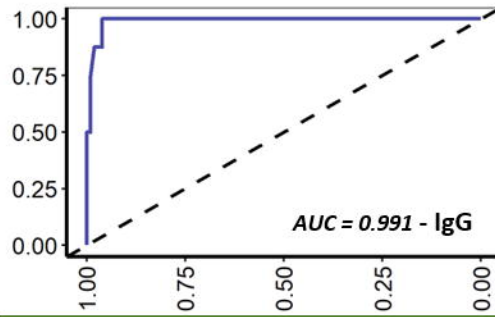
SARS-CoV-2-S2



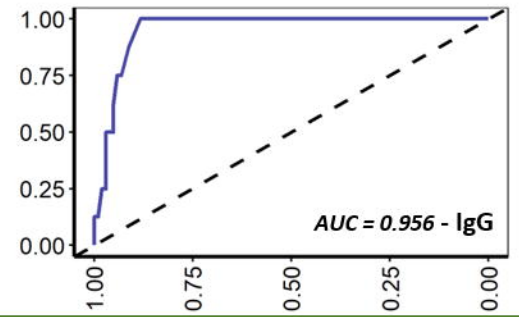
SARS-CoV-2-S1+RBD



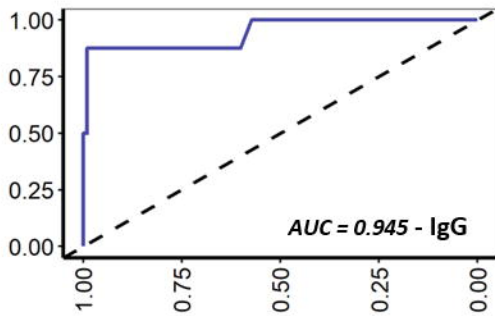
SARS-CoV-2-S1+S2



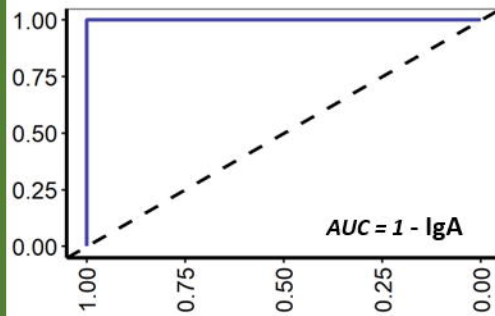
MERS-CoV-S2



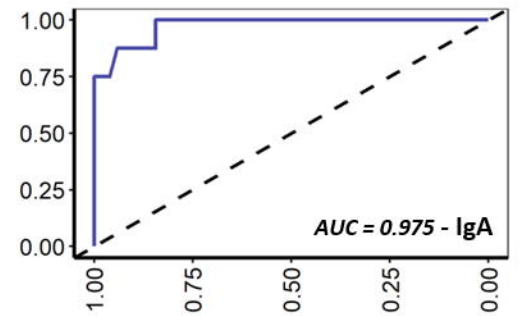
SARS-CoV-2-S1



SARS-CoV-2-S2



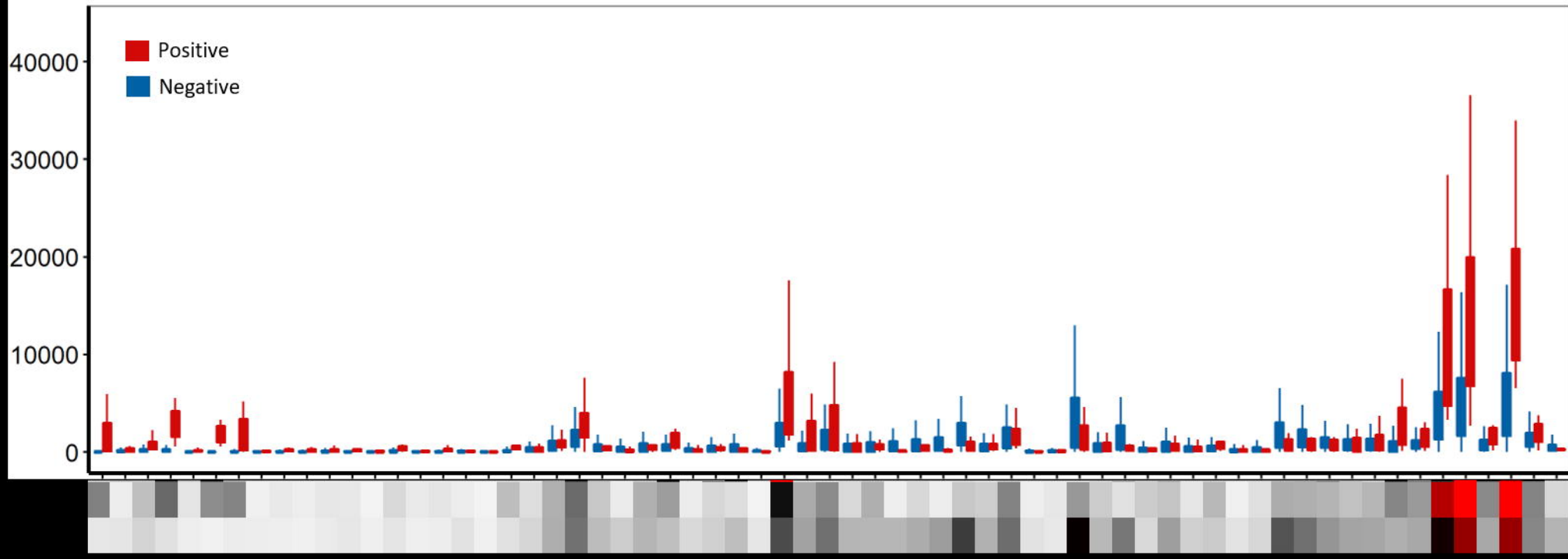
SARS-CoV-2-S1+S2



IgG

IgA

Specificity



SARS-CoV-2	SARS-CoV	MERS-CoV	Common Cold CoV	Influenza	Adenovirus	MPV	PIV	RSV
SARS-CoV-2_NP SARS-CoV-2_S1, (His Tag) SARS-CoV-2_S1, (mFc Tag) SARS-CoV-2_S1+S2 SARS-CoV-2_S1-RBD SARS-CoV-2_S2	SARS-CoV_NP SARS-CoV_PLpro SARS-CoV_S1, (His Tag) SARS-CoV_S1-RBD, (His Tag) SARS-CoV_S1-RBD, rFc Tag	MERS-CoV_NP MERS-CoV_S1, N-(AA1-725, His Tag) MERS-CoV_S1+S2 MERS-CoV_S1-RBD, N-(AA367-606, His Tag) MERS-CoV_S1-RBD, N-(AA367-606, rFc Tag) MERS-CoV_S1-RBD, N-(AA383-502, mFc Tag) MERS-CoV_S1-RBD, N-(AA383-502, rFc Tag) MERS-CoV_S2	DcCoV_HKU23-368F_NP HCoV-229E_S1 HCoV-229E_S1+S2 HCoV-HKU1_HE HCoV-HKU1_S1, N-(AA13-756) HCoV-HKU1_S1, N-(AA1-760) HCoV-HKU1_S1+S2 HCoV-NL63_S1 HCoV-NL63_S1+S2 HCoV-OC43_HE HCoV-OC43_S1 HCoV-OC43_S1+S2	B_B/Malaysia/2506/2004_HA1 B_B/Malaysia/2506/2004_HA1+HA2 B_B/Phuket/3073/2013_HA1 B_B/Phuket/3073/2013_HA1+HA2 H1N1_A/Beijing/22808/2009_HA1 H1N1_A/Beijing/22808/2009_HA1+HA2 H1N1_A/Beijing/22808/2009_HA1+HA2 H3N2_A/Texas/50/2012_HA1 H3N2_A/Texas/50/2012_HA1+HA2 H5N1_A/Vietnam/1203/2004_HA1 H5N1_A/Vietnam/1203/2004_HA1+HA2 H7N9_A/Anhui/1/2013_HA1 H7N9_A/Anhui/1/2013_HA1+HA2	hAdV-3/45659_Fiber hAdV-3/45659_Penton hAdV-4/28280_Fiber hAdV-4/28280_Penton hAdV-7/10519_Fiber hAdV-7/10519_Penton	hMPV-A_G (52N-228N) hMPV-B_F (280D-490G) hMPV-B_G (52D-238S)	hPIV-1 12O3_F hPIV-1 12O3_H hPIV-3 USA/10991B/2010_H hPIV-4b/10-H2/2016_F hPIV-4b/10-H2/2016_H	RSV-A_A2_F RSV-A_LA2-94/2013_F RSV-A_LA2-94/2013_G RSV-A_rsb1734_G RSV-A_RSS-2_F RSV-B_B1_G RSV-B_TH-10526/2014_F RSV-B_TH-10526/2014_G

