Anti-Spike protein assays to determine post-vaccination antibody levels: a head-to-head comparison of five quantitative assays

Running head: Comparison of five quantitative SARS-CoV-2 antibody assays after vaccination

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Abstract (239/250)

Background
Reliable quantification of the antibody response to SARS-CoV-2 vaccination is highly relevant for identifying possible vaccine failure and estimating the time of protection. Therefore, we aimed to evaluate the performance of five different Anti-SARS-CoV-2 antibody assays regarding the quantification of anti-spike (S) antibodies induced after a single dose of BNT162b2.

Methods
Sera of n=69 SARS-CoV-2 naïve individuals 21±1 days after vaccination with BNT162b2 (Pfizer/BioNTech) were tested using the following quantitative SARS-CoV-2 antibody assays: Roche S total antibody, DiaSorin trimeric spike IgG, DiaSorin S1/S2 IgG, Abbott II IgG, and Serion/Virion IgG. Test agreement was assessed by Passing-Bablok regression. Results were further compared to the percent inhibition calculated from a surrogate virus neutralization test (sVNT) by correlation and ROC (receiver-operating-characteristics) analysis.

Results
Individual values were distributed over several orders of magnitude for all assays evaluated. Although the assays were in good overall agreement ($\rho=0.80-0.94$), Passing-Bablok regression revealed systematic and proportional differences, which could not be eliminated by converting the results to BAU/mL as suggested by the manufacturers. 7 (10%) individuals had a negative sVNT results (i.e. <30% inhibition). These samples were reliably identified by most assays and yielded low binding antibody levels (ROC-AUCs 0.84-0.93).
Conclusions

Although all assays evaluated showed good correlation, readings from different assays were not interchangeable, even when converted to BAU/mL using the WHO international standard for SARS-CoV-2 immunoglobulin. This highlights the need for further standardization of SARS-CoV-2 serology.

Keywords: SARS-CoV-2; serology; vaccination; BNT162b2; antibody testing

Abbreviations

SARS-CoV-2 Severe acute respiratory syndrome Coronavirus 2
COVID-19 Coronavirus Disease 19
NC Nucleocapsid
S Spike protein
tAb total antibody
ECLIA electrochemiluminescence immunoassay
RBD receptor binding domain
CMIA chemiluminescence microparticle assay
CLIA chemiluminescence immunoassay
ELISA enzyme-linked immunosorbent assay
BAU/mL binding antibody units per milliliter
sVNT surrogate virus neutralization test
ROC-AUC receiver-operating-characteristics area under the curve
Background

SARS-CoV-2 antibody testing played and still plays an essential role in the management of the COVID-19 pandemic (1). Detection of specific antibodies following SARS-CoV-2 infection is important at both the individual and population levels to identify those at risk of infection (2). However, now in the early vaccination era of the COVID-19 pandemic, another essential role of SARS-CoV-2 serology is added: the determination of specific antibodies after active immunization (3, 4). Although this new role has not yet been finally defined, it is already clear that antibody assays will be needed for both vaccine development and approval process and the follow-up of vaccinated individuals. Licensing studies on a scale such as Pfizer/BioNtech and Moderna vaccines will not be feasible for all vaccine candidates and the companies behind them (5). For this reason, correlates of vaccine-induced protection will have to replace, at least in part, the measurement of clinical outcomes as the sole measure. For reasons of simplicity, relatively good standardizability, and broad applicability, SARS-CoV-2 antibody tests have the best chance of becoming such substitute endpoints (6).

The first SARS-CoV-2 antibody testing systems were designed to distinguish individuals with prior COVID-19 infection from those who were still naive to this new virus (7). Therefore, these immunoassays were usually developed as qualitative rather than quantitative tests and were designed by the manufacturer to achieve the highest possible specificity and high sensitivity. High specificity was indispensable, especially at the beginning of the pandemic, because the extremely low seroprevalence rates led to many false positives and low positive predictive values even with tests having a specificity of 99% (8). In contrast, the sensitivity of SARS-CoV-2 testing was often reduced to assure the high specificities needed for these assays (9). The lower
antibody levels further aggravated suboptimal sensitivities in mild/asymptomatic infections and
during the pandemic by the natural decline in antibody levels (10-15).

Various antigens have been used for this purpose, but essentially two types can be distinguished:
nucleocapsid- and spike protein-based assays (16). Antibodies directed against SARS-CoV-2
specific nucleocapsid (NC) antigens are induced early and strongly in most infected individuals
due to the virus nucleocapsid's typical strong immunogenicity (17). Furthermore, a very high
specificity can be achieved by targeted modification of the nucleocapsid antigen so that no cross-
reactivity is observed even with closely related viruses. The discriminatory properties of such
nucleocapsid-based antibody assays can therefore be excellent (18, 19). The physiological
significance of these antibodies, on the other hand, is unclear, and these surrogate markers for a
previous infection are unlikely to be functionally relevant to confer protection or immunity. The
antibodies that react with the spike protein (S), however, act differently. At least a proportion of
these S-binding antibodies are likely to have the function of neutralizing antibodies (20). Thus, it
is not surprising that numerous studies have shown a correlation between spike protein binding
assays and various forms of functional virus neutralization assays (21-26).

In the context of SARS-CoV-2 vaccines, it is precisely these neutralizing antibodies that are of
paramount importance. The primary goal of active immunization is to induce many SARS-CoV-
2-specific neutralizing antibodies that ideally prevent the pathogen's entry and thus infection or
stop the systemic spread to prevent disease (27). The functional virus neutralization assays are
not feasible everywhere: assays with live viruses require biosafety level 3, but variants such as
pseudotyped neutralization assays are also labor-intensive and cannot be performed at high
throughput (28-30). Classical antibody assays, which measure the reactivity of antibodies in
serum/plasma with defined antigens, can be performed very rapidly and in high throughput, in contrast to neutralization tests.

Thus, anti-spike protein assays will play an important role in vaccine development, licensing, and efficacy monitoring in the future. However, these test systems must be able to reliably quantitate SARS-CoV-2 specific antibody levels, be comparable to each other, and have good to excellent agreement with the presence of neutralizing antibodies. The comparability of antibody assays is expected to be improved by the recent introduction of a first WHO International Standard for anti-SARS-CoV-2 immunoglobulin (NIBSC code 20/136) with reference to neutralizing antibodies.

In the present work, we aim to go a step further and characterize the vaccination response after the first administration of the Pfizer/BioNTech btn162b2 vaccine using five commercial quantitative anti-spike protein antibody assays (4 of them with manufacturer's correction factor for the WHO standard) in a head-to-head comparison.

Methods

Study design and participants

This prospective observational study includes sera taken from 69 individuals without a previous SARS-CoV-2 infection taken 21±1 days (mean±standard deviation) after the first dose of the Pfizer/BioNTech BNT162b2 vaccine. Further inclusion criteria were an age >18 years, whereas an insufficient amount of serum would have led to exclusion from the study. The study protocol was reviewed and approved by the Ethics Committee of the Medical University of Vienna (EK1066/2021). All participants provided written informed consent to donate blood for the evaluation of diagnostic test systems (EK404/2012).
Laboratory procedures

Serum was obtained and stored at 2 – 10°C for <7 days within the MedUni Wien Biobank, a centralized facility for the preparation and storage of biomaterial with certified quality management (ISO 9001:2015)(31). All analytical procedures were performed at the Department for Laboratory Medicine, Medical University of Vienna. The following CE-marked binding assays were applied:

The Roche Elecsys® Anti-SARS-CoV-2 S (Roche S tAb) is an electrochemiluminescence sandwich immunoassay (ECLIA) and detects total antibodies directed against the receptor-binding domain (RBD) of the viral spike (S)-protein and was measured on cobas® e801 modular analyzers (Roche Diagnostics, Rotkreuz, Switzerland). The quantification range is between 0.4 and 2500.0 U/mL. The manufacturer states intra- and interassay precision between 1 and 3%, a clinical specificity of 99.98% (99.91 – 100), and a cumulative sensitivity ≥14 days after the first positive PCR of 98.8% (98.1 – 99.3) if 0.8 U/mL is used as a cut-off.

The Abbott SARS-CoV-2 IgG II Quant-test (Abbott S IgG) is a chemiluminescence microparticle immunoassay (CMIA). It quantifies IgG-type antibodies against the RBD of the viral S-protein on an Abbott ARCHITECT platform (Abbott, Abbott Park, USA) between 21.0 and 40,000.0 AU/mL. Intra- and interassay precision ranges between 3 and 5%. According to the manufacturer, clinical specificity is 99.55% (99.15 – 99.76), and clinical sensitivity is 98.81% (93.56 – 99.94) ≥15 days after the first positive PCR at a cut-off of ≥50 AU/mL.

The DiaSorin LIAISON SARS-CoV-2 TrimericS IgG (DiaSorin TriS IgG) chemiluminescence immunoassay (CLIA) quantifies IgG antibodies against a trimeric S-protein antigen on a DiaSorin LIAISON (DiaSorin, Stillwater, USA). The quantification range is between 1.63 and 800 AU/mL. Intra- and interassay precision ranges between 0 and 5%. According to the
manufacturer, clinical specificity is 99.5% (99.0 – 99.7), and clinical sensitivity ≥15 days after
the first positive PCR is 98.7% (94.5 – 99.6) at a cut-off of ≥13 AU/mL.

The DiaSorin LIAISON SARS-CoV-2 S1/2 CLIA (DiaSorin S1/2 IgG) detects IgG antibodies
against an S1/S2 combination antigen on a DiaSorin LIAISON (DiaSorin, Stillwater, USA). The
quantification range is between 3.8 and 400.0 AU/mL. Intra- and interassay precision ranges
between 0 and 4% and, according to the manufacturer, specificity among blood donors is 98.5%
(97.5 99.2) and sensitivity is 97.4% >15 days after diagnosis at a cut-off of >15 AU/mL, whereby
results between 12.0 and 15.0 AU/mL are considered borderline.

The Virion\Serion ELISA (enzyme-linked immunosorbent assay) agile SARS-CoV-2 IgG (Serion
IgG) (Institut Virion-Serion, Wuerzburg, Germany) was analyzed on a FilterMax F5 Multiplate
Reader (Molecular Devices, San José, USA) and quantifies IgG antibodies against total S-protein
between 3 and 250 U/mL. Intra- and interassay precision ranges between 1 and 4%. According to
the manufacturer, specificity is 99.2%, and sensitivity is 96.2% at a cut-off of 15 U/mL, with
values between 10 and 15 U/mL being considered borderline results.

If applicable, binding antibody units per milliliter (BAU/mL), which are traceable to the WHO
International Standard for anti-SARS-CoV-2 immunoglobulin, were calculated by applying the
following conversion factors, as suggested by the manufacturers: Roche S tAb = \frac{U}{mL} \ast 1, Abbott S
IgG: = \frac{U}{mL} \ast \frac{1}{7}, DiaSorin TriS IgG = \frac{AU}{mL} \ast 2.6, Serion IgG = \frac{U}{mL} \ast 2.1.

We excluded prior SARS-CoV-2 infection by using the Roche Elecsys® SARS-CoV-2 ECLIA
on the cobas® e801 analyzer (Roche), which detects total antibodies to the viral nucleocapsid
antigen. These antibodies are not induced by vaccination with bnt162b2. This assay yields high
diagnostic sensitivity (90%) and specificity (99.7%) for infections that occurred at least 14 days
before blood withdrawal (7). As suggested by the manufacturer, results >1.000 COI were considered positive.

Neutralizing capacity was estimated by performing a surrogate virus neutralization test (sVNT) (GenScript, Piscataway, USA). The assay was read on a Filtermax F5 multimodal plate reader. According to the manufacturer, it shows excellent positive (100% [87.1 – 100.0]) and negative percent agreement (100.0% [95.8 – 100.0]) with conventional plaque-reduction neutralization tests (PRNT\textsubscript{50} and PRNT\textsubscript{90}). Results ≥30% are considered positive.

Statistical analysis

Continuous data are given as a median and interquartile range, categorical data as counts and percentages. Passing-Bablok regressions, Cohen’s Kappa (linear weights) and Spearman rank correlations evaluated the agreement between binding assays. The relationship between binding assays and results from the sVNT was described by quadratic curve fitting. The predictive value of binding assays regarding positivity in the sVNT was assessed by interpreting and comparing (according to DeLong) the areas under the curve (AUC) from receiver-operating-characteristics (ROC)-curves. Statistical significance was assumed if $P$ values were below 0.05. All analyses were performed using MedCalc 19.6 (MedCalc, Ostend, Belgium), and graphs were drawn using GraphPad 9 (GraphPad, La Jolla, USA).

Results

Measurement ranges differ between binding assays

29 female (42%) and 40 male (58%) participants with a median age of 42 years (29 – 51) were included. Results from the five different antibody binding assays are presented in Table 1 and Figure 1. The Abbott S IgG assay showed the highest values with a median of 1097.1 AU/mL
yielded the lowest values (63.7 AU/mL [47.8 – 87.5]), and the levels ranged from below the limit of quantification (<3.8%, 1 sample) to 148.0. Two assays, the DiaSorin TriS IgG (195.0 AU/mL [99.0 – 337.3]) and the Serion IgG (50 U/mL [30 – 89]) returned a result above the measuring range for the same donor (>800 AU/mL and >250 U/mL). However, both tests used their full available range (lowest values: 1.8 AU/mL and <3 U/mL). Roche S tAb ECLIA results came in between the other test systems (79.6 U/mL [24.7 – 142.3]), ranging from 0.4 to 508.0.

The measured values indicate that the numerical results are strongly dependent on the test system used. In the next step, we aimed to evaluate the overall agreement between the test systems.

**Agreement between results from different binding assays**

Results from the Roche S tAb assay correlated well with those of the other binding assays (Abbott S IgG $\rho=0.88$, DiaSorin TriS IgG $\rho=0.83$, DiaSorin S1/2 IgG $\rho=0.80$, Serion IgG $\rho=0.82$). However, Passing-Bablok regression revealed relevant systematic and proportional differences: Abbott S IgG $= 82.5 + 15.54*X$, DiaSorin TriS IgG $= 33.4 + 2.18*X$, DiaSorin S1/2 IgG $= 39.6 + 0.32*X$, Serion IgG $= 12.3 + 0.65*X$.

The Abbott S IgG assay correlated at $\rho=0.90$ with the remaining three test systems (DiaSorin TriS IgG and S1/2 IgG, Serion IgG). In Passing-Bablok regression, all systematic and proportional errors were statistically significant: DiaSorin TriS IgG $= 24.5 + 0.13*X$, DiaSorin S1/2 IgG $= 34.5 + 0.02*X$, Serion IgG $= 6.2 + 0.04*X$.

The DiaSorin TriS IgG assay showed an excellent correlation with the remaining two tests (DiaSorin S1/2 IgG $\rho=0.91$, Serion IgG 0.94). In the Passing-Bablok regression, nevertheless,
marked deviations became apparent: $\text{DiaSorin S1/2 IgG} = 30.5 + 0.16 \times X$, $\text{Serion IgG} = -0.0 + 0.31 \times X$.

Finally, the DiaSorin S1/2 IgG and the Serion IgG correlated at $\rho=0.91$, and the Passing-Bablok regression equation was $\text{Serion IgG} = -50.9 + 1.78 \times X$. All described relationships, as well as related residual plots, are presented in Figure 2.

Furthermore, we assessed whether the classification of results into tertiles (0 – 33.3%, 33.4 – 66.7%, 66.8 – 100%) was comparable, e.g., whether a sample yielding a result in the lowest tertile of test A was also in the lowest tertile of test B. Cohen’s kappa was between 0.60 and 0.80, indicating a good agreement, for all but for one of the ten test combinations (Roche S tAb/Serion, kappa = 0.59, see Table 2).

In conclusion, the results of the investigated test systems correlate well but are not necessarily interchangeable. Several manufacturers provided conversion factors related to the WHO International Standard for SARS-CoV-2 immunoglobulin, as described in the methods section.

Next, we wanted to clarify whether comparing converted BAU/mL instead of arbitrary values facilitates comparability.

**Associations between standardized binding assay results**

Binding antibody units per milliliter (BAU/mL) were calculated for the Abbott S IgG, the DiaSorin TriS IgG, and the Serion IgG, according to the recently proposed conversion factors. Results from the Roche S tAb ECLIA did not require conversion as indicated by the manufacturer.

As shown in Figure 3, the BAU/mL recalculation did not solve the problem of high proportional errors. The least proportional error could be observed for the relationship between Roche S tAb
Correlation of binding assay results with a surrogate neutralization assay

In a final step, the binding assays' results were compared to percent inhibition of a surrogate virus neutralization assay (sVNT). In the sVNT, the tested samples yielded median values of 63% (50 – 76), ranging from 6 to 92%. Figure 4a illustrates that all binding assays except the DiaSorin S1/2 IgG showed a quadratic relationship with the sVNT. The binding assays also differentiated those values clustered in the upper range of the sVNT. However, for the DiaSorin S1/2, the quadratic curve approached a straight line, indicating a mostly linear relationship between this binding assay and the sVNT within the observed range.

7 (10%) of the individuals yielded sVNT results below 30% inhibition, which is considered negative according to the manufacturer (Figure 1). Binding assay results were compared between positives and negatives in the sVNT by ROC-curve analysis. The resulting AUCs ranged between 0.84 and 0.93 (see Figure 4b), however, the differences between AUCs were not statistically significant. Optimal cut-offs according to Youden’s Index, as well as other cut-offs observed from the data and their respective sensitivities and specificities are given in Supplemental Table 1. For three out of the five assays (Abbott S IgG, DiaSorin S1/2 IgG, and Serion IgG), all samples with a negative sVNT result could be correctly identified with a corresponding sensitivity of approximately 60%.

Discussion

SARS-CoV-2 antibody assays become important tools to evaluate the proportion of people affected by COVID-19 and identify those who are still at infection risk. Now with the first
vaccines available a new field of use for SARS-CoV-2 antibody tests will open up. First, many
vaccinated individuals will be interested in confirming their own vaccination success based on
the detection of specific antibodies. Second, vaccination-induced antibodies may be used as
surrogate from which a protection correlate will be estimated. To date, only limited information
on the performance of quantitative SARS-CoV-2 antibody assays is available, since most
currently evaluated assays were developed in-house, as recently summarized by the CDC
COVID-19 response group (32). Only for a few commercially available quantitative CE-marked
test systems preliminary data on the performance are given in the literature (19, 22, 33-35).

Although a protection correlate for immunity in SARS-CoV-2 has not been defined yet, it is
useful to begin this important preliminary work now (6). Therefore, in the present work, we
compared different commercial SARS-CoV-2 antibody assays with spike protein reactivity using
a vaccination cohort to give a first insight into the comparability of these assays.

With regard to the numerical results, we were able to determine a broad distribution of the result
values for each individual test system, so that these were presented on a logarithmic scale. This is
in line with recently published reports, showing the antibody response after a single dose of
BNT162b2 vaccine (3, 4). Interestingly, in agreement with a study involving >500 participants in
an identical study setting, we observed very similar mean values for the measurements with the
DiaSorin S1/S2 IgG: 66.3 AU/mL versus 68.6 AU/mL (3). Therefore, it is reasonable to assume
that our cohort is representative despite the moderate number of participants. In addition, we
were able to show that the results of the different test systems varied by a factor of up to more
than 50. This leads to the initial conclusion that a direct comparability of the numerical results of
different test systems is unlikely to be given across the range of individual findings. Differences
also occurred with respect to measurement ranges, and upper measurement limits were exceeded
in 2 out of 5 systems (DiaSorin TriS IgG and Serion IgG), although the study cohort reflects the antibody response before the administration of the 2nd dose of Pfizer/BioNTech vaccine in SARS-CoV-2 naïve individuals. However, it must be mentioned that it is not yet known up to which level a differentiation of the obtained values is meaningful. A recently published paper shows that an anti-S post-vaccination titer of 61.8 AU/mL for the Abbott S IgG was associated with reinfection after a single dose of vaccination. Since this value is only just above the threshold for positivity for this specific assay (50 AU/mL) and in our cohort >95% of all observed values were far higher (5th percentile: 207.5 AU/mL), this finding is plausible and suggests vaccine failure due to very low antibody production. Alternatively, a reinfection might have been caused in these subjects by a virus variant where vaccination protection is mitigated. Nevertheless, it can be assumed that the average values of completely vaccinated persons are significantly higher than those in our collective and thus the upper measurement limits could frequently be exceeded in most assays. If clinically relevant, this could make additional dilution steps necessary, which are not yet taken into account by the manufacturers.

Despite the different levels of measurement, all systems showed good correlations with each other. When the measured values of the individual antibody tests were assigned to tertiles, good agreement was shown between the lowest third, the middle third and the highest third of the results. Thus, one individual with known immunosuppressive therapy consistently showed no formation of antibodies in all five antibody binding assays tested. With defined cut-offs for low or high vaccination titers of the different test systems, at least a partial transferability of a result from one to another test system may therefore be expected. Such transferability of results could also be anticipated via referencing the antibody assays used to an international reference standard (32). Indeed, a first WHO international SARS-CoV-2
antibody standard with the valence of 1000 BAU/mL has recently become available. This standard was used by the manufacturers for four out of five of the assays studied. However, this standardization was not introduced during the establishment of the test system, but post-hoc as a reference material to define a conversion factor of their own units in BAU/mL. It is therefore not surprising that, this subsequent correction did not reduce the existing systematic deviations (Figure 2) between the different tests. Only the Roche S tAb and Serion IgG tests were able to approximate the equivalence line, although here a very wide scattering of values around the trend lines was observed.

The in vitro binding of infection-associated antibodies to pathogen-specific antigens in an antibody test are important markers to objectify a past infection or vaccination. However, these do not necessarily say anything about the function of these antibodies (1). In SARS-CoV-2 vaccination, an important goal is to induce neutralizing antibodies that will prevent the virus from binding to the cellular receptor, the ACE2 receptor, via the surface spike protein (36, 37). Tests to neutralize live viruses can only be performed in very specialized laboratories and unfortunately, in the case of SARS-CoV-2, are not standardized, making comparability almost impossible. For this reason, we chose to use a well-characterized surrogate virus neutralization test (sVNT) as a functional reference (38–40). In this assay, a simple ELISA format is used to determine the inhibition of conjugated RBD protein by neutralizing antibodies to the plate-bound ACE2 receptor. The manufacturer suggests a threshold for positivity of 30% inhibition. For the Abbott S IgG, the DiaSorin S1/2 IgG, and the Serion IgG assay, all samples below this threshold were identified at a corresponding sensitivity of about 60%. The corresponding criteria were 3 – 17 times higher than the respective assays’ thresholds for positivity. This implies that the cut-off values given for the respective test systems are only valid for the diagnosis of a past infection, but
do not necessarily represent a threshold value for the presence of sufficient neutralizing activity. For the Roche S tAb and the DiaSorin TriS IgG assays, a single outlier avoided reaching maximum specificity before sensitivity dropped to 3 and 32%, respectively.

In conclusion, we found a good correlation between all evaluated assays, however, the values from the different test systems were not interchangeable, even when converted to BAU/mL using the WHO international standard for SARS-CoV-2 immunoglobulin. Furthermore, it should be noted that the thresholds for positivity provided by the manufacturers are of diagnostic value and are not indicative of sufficient inhibitory capacities.

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References


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Legends to Figures

Fig. 1: Results from binding assays (A) and a surrogate virus neutralization test, sVNT (B). Solid lines mark the median. The dotted line in (B) marks the manufacturer’s threshold for positivity (30%).

Fig. 2: Comparison of binding assays by linear regression (dotted lines indicate the 95% confidence interval) (A) and residual plots (B).
Fig. 3: Relationships between binding assay results and percent inhibition assayed using a surrogate virus neutralization test (threshold for positivity: 30%, dotted vertical lines). Presented are quadratic regression lines and their 95% confidence intervals.

Fig. 4: (A) Comparison of binding assay results converted to BAU/mL (binding antibody units per milliliter). Given are linear regression curves and their 95% confidence intervals. Dotted diagonal lines represent lines of equality. (B) ROC (receiver-operating-characteristics) curves of binding assays regarding the agreement with the results of a surrogate virus neutralization test (threshold for positivity: 30% inhibition).
<table>
<thead>
<tr>
<th></th>
<th>Roche S tAb</th>
<th>Abbott S IgG</th>
<th>DiaSorin TriS IgG</th>
<th>DiaSorin S1/2 IgG</th>
<th>Serion IgG</th>
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<tr>
<td>Median</td>
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<td>&lt;3.8 – 148.0</td>
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Tbl. 1: Dimensions of position and spread for 5 S-protein based SARS-CoV-2 antibody assays calculated from N=69 samples taken 21±1 days after the first shot of bnt162b2.

<table>
<thead>
<tr>
<th></th>
<th>Abbott S IgG</th>
<th>DiaSorin TriS IgG</th>
<th>DiaSorin S1/2 IgG</th>
<th>Serion IgG</th>
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Tbl. 2: Kappa (±95% confidence interval) for ten different test combinations regarding the classifications of samples into tertiles.