Article

Possible correlation of electrochemiluminescence based numerical cut off index value with concentration of anti-SARS-CoV-2 antibody: Is it worth reporting?

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Abstract

Background: Many laboratories are reporting a numerical cut-off index value (COI) for most anti-SARS-CoV-2 qualitative tests. These numerical values in patients’ report ultimately created great confusion in the public and physicians, therefore this study was designed to evaluate the correlation of electrochemiluminescence (ECLIA) based numerical COI values with quantitative ELISA of anti-SARS-CoV-2 antibody.

Design and Methods: Two hundred and twenty-eight (228) recovered COVID-19 patients were included; their serum samples were analyzed by quantitative ELISA and ECLIA for anti-SARS-CoV-2 antibodies.

Results: One hundred and seventy-three (75.8%) patients tested positive by ECLIA and ELISA assay and thirty-seven (6.2%) were tested negative by both methods. A weak positive correlation (r=0.37) was found between numerical COI value of ECLIA with ELISA concentration, which was statistically significant with p<0.001. All values were dispersed on scatter plot and there was no significant linear relationship between ECLIA and ELISA assay.

Conclusions: As both testing techniques are base upon the same immunological phenomena of detecting antibodies against nucleocapsid protein. We suggest that COI values are not meant to describe the immunity level of the individuals thus the physicians should not consider it as a quantitative value for antibody levels in COVID-19 patients.

Introduction

On March 11, 2020 the World Health Organization (WHO) declared COVID-19 a global pandemic.1 The causative agent of COVID-19 is a corona virus named as SARS-CoV-2 and it belongs to a family of viruses that may cause respiratory symptoms ranging from common cold to severe pneumonia. The host or infected individual in response to the infection produces specific antibodies including IgM, IgG and IgA.2,3 The detection of these antibodies indicates that the individual has been exposed to the virus. During the recent pandemic, several diagnostics companies introduced testing kits for detection of these antibodies, to estimate the percentage of the population previously infected with the virus information needed to devise strategies for community surveillance in order to protect the public’s health.4 Many tests based on different techniques are readily available in the market to detect the antibody. The two most commonly used techniques to detect antibodies against SARS-CoV-2 that entered the market included Electrochemiluminescence Immunoassay (ECLIA) and Enzyme-linked Immunosorbent assay (ELISA). ECLIA is a qualitative assay used to find the patients who had been exposed to the virus while utilizing cut off index (COI) as a reference to give a positive or negative result. The test targets antibodies including IgG, IgM and IgA against nucleocapsid of SARS-CoV-2. On the other hand, ELISA is a quantitative test intended to define the titers of IgG in patients who had been previously exposed.5 In case of SARS-CoV-2, it is still unknown if the presence of IgG confers any protection against re-infection. As the number of COVID cases increased, there was emergence of different techniques to estimate the exposure of the population to the causative agent and to detect the development of corresponding antibodies. New techniques were developed and marketed to diagnose the disease in acute phase as well as to show the percentage of the population that had been exposed to the virus.6 On the other hand, there were techniques which were intended to show the quantitative values of one specific types of antibody. Of these testing systems, ECLIA based testing system and ELISA based Testing system have been used by many laboratories in Pakistan.7

Both testing systems have been claimed to have high sensitivity and specificity,8 but COI is mistakenly considered as a quantitative variable to define the overall anti- SARS-CoV-2 antibodies in recovered patients. The objective of the current study was to record any possible correlation of ECLIA based numerical COI values provided by Roche Diagnostics (Rotkreuz, Switzerland) testing system with quantitative ELISA provided by AESKU Diagnostics (AESKU Diagnostics; Wendelsheim, Germany) for anti- SARS-CoV-2 antibody detection.

Significance for public health

Many laboratories in Pakistan are reporting numerical COI values, which ultimately created great confusion among the patients and physicians. These values are, used indiscriminately and wrongly compared to other testing systems which were in general intended to be used for quantitative analysis of the antibodies developed in the persons exposed to the virus. There had been obvious misunderstanding in the public including the healthcare sector when these different techniques were used indiscriminately without a proper orientation towards the utility and limitations of a given testing system. As a result, the different numerical COI values which were included in the laboratory reports of the test created a great havoc and raised suspicions about the certainty of the diagnostic techniques. This correlation is important because this number game has been talk of the town and lay person uses them to get the idea of one’s own immunity status.
Design and Methods

This prospective study was performed from 5th June to 30th October 2020 at National Institute of Blood Disease and Bone Marrow Transplantation (NIBD) Hospital, Karachi, Pakistan. The adult fully recovered corona survivors (NIBD) Hospital, Karachi, Pakistan. The adult fully recovered corona survivors of either gender aged 18 to 60 years were recruited after two weeks of negative PCR. All subjects were healthy with no known co-morbidities. The study was approved by institution’s ethical review committee. Three to five ml blood sample was taken in EDTA tube for anti-SARS-CoV-2 antibody analysis by both ELISA and ECLIA. For quantitative ELISA AESKULISA® SARS-CoV-2 NP IgG (AESKU Diagnostics) was used to create scatter plots, which gives slope of 0.69 (95% CI -5.83-4.78) and did not address the confusion existent in the public and physicians that whether higher COI values meant a higher humoral immunity level (Ig level) or not. Although manufacturer of ECLIA had clearly described the quantitative nature of the testing system, the included numerical values or COI reporting by laboratories were creating confusion. Physicians were confused as well and were considering these COI values comparable to Ig levels reported by quantitative ELISAs. Every participant of the study was tested by two antibody-based testing systems i.e. Eclays® Roche testing system using ECLIA technique and Quantitative ELISA by AESKULISA® diagnostics testing system. We found lack of any significant quantitative correlation between these two. The results suggest that the COI values are not meant to describe the immunity level of the individuals. The correlation of electrochemiluminescence based numerical COI of anti-SARS-CoV-2 antibody with actual concentration is not reported in literature.

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Sittings et al. in 2020 determined IgG levels by using ELISA technique as the serum was collected after 28 days of infection, they did not perform IgM levels on ELISA as they collected samples after 4 weeks of illness and its already known that IgM levels markedly reduced after 4 weeks of infection.9 The positivity rate of IgM antibody was only 60%, with a marked reduction in antibody levels 4 weeks after onset of illness.10 Anti-SARS-CoV-2 S-specific IgG antibodies were identifiable from day 7 onwards, peaking at approximately day 25 Serum IgG antibodies were still maintained at a high level after 4 weeks of infection.11 Hou et al also reported IgM levels increased during the first week after SARS-CoV-2 infection, peaked 2 weeks and then reduced to near background levels in most patients.12 There were 8.34% individuals who were tested positive only by ECLIA system and negative by ELISA based system, which may be due to IgM detected by Roche and not.

Statistical analysis

Data was analyzed by EP evaluator (vers. 10.0). Mean and Standard deviation for quantitative variables were calculated along with frequencies for all categorical variables. The mean values for ECLIA anti-SARS-CoV-2 were calculated by using numerical cut of index (COI) value. Scatter plots were created by using test method as ECLIA on Y-axis and reference method as ELISA on X-axis to see the relationship between COI and concentration. Correlation coefficient (r value) was calculated by Pearson correlation. Cohen’s Kappa was calculated to see the agreement, Cohen’s Kappa >75% considered as high agreement. Furthermore, t-test was applied to compare the means of ECLIA and ELISA and p value of less than 0.05 was taken as statistically significant.

Results

Two hundred and twenty-eight (228) COVID-19 survivors were included in the study. The mean age of subjects was 36.6±11.6 years. There were 192 (84.2%) males and 62 (15.8%) females. The mean anti-SARS-CoV-2 antibody by ECLIA was 30.38±31.04COI and by ELISA was 44.35±38.49U/ml. The mean anti-SARS-CoV-2 antibody level was higher by ELISA and there was a significant statistical difference between these two means with p value <0.000. Approximately, 75.8% (173) individuals tested positive either by ECLIA and ELISA or both whereas 16.2% (37) tested negative by both methods. About 8.34% (18) individuals tested positive by ECLIA and tested negative by ELISA assay. Antibody results obtained by ECLIA and ELISA of all patients were used to create scatter plot, which gives slope of 0.69 (95% CI -0.60-0.78), intercept of -0.53 (95% CI -5.83-4.78) and did not show any significant linear relationship between ECLIA and ELISA.

Discussion

The current study was inspired by the curiosity to understand the numerical COI values given by any ECLIA testing system to address the confusion existent in the public and physicians that
detected by ELISA although this could not be proven because of the combo nature of the Roche ECLIA testing technique. We strongly suggested that ECLIA being a qualitative test should be reported as positive or negative. No numerical values for qualitative assays should be documented in lab reports to avoid confusion in physicians as well as nonclinical individuals. This suggestion is in line with the manufacturer’s claim whose intention was to give only the qualitative results, but confusion was created because of the associated numerical values. As these COI values lack any relationship to the antibody/immunity status of an individual. This practice has been adopted by our laboratory although some other laboratories in our country are still issuing reports with numerical values included thus creating confusion patients and physicians.

Conclusions

The results of this study suggest a lack of significant quantitative correlation between these two testing systems. Although greater sample size may be required to find any possible correlation, but it can be strongly suggested comparing these two systems. Additionally studies are required to explain the meaning of numerical values generated by the Roche system and their correlation with the immunity status/antibody titers in a given individual.

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Dedication: To my Mother (Late Zarina Ramzan Ali), Father (Late Ramzan Ali) and Brother (Late Rehman Ali).

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