Review article

Understanding the implications of SARS-CoV-2 re-infections on immune response milieu, laboratory tests and control measures against COVID-19

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ABSTRACT

Several months after the emergence of severe acute respiratory syndrome coronavirus-2 (SARS-CoV-2), cases of re-infection after recovery were reported. The extent and duration of protective immunity after SARS-CoV-2 infection is not fully understood. As such, the possibility of re-infection with SARS-CoV-2. Furthermore, cases of re-infection were mainly due to different variants or mutant SARS-CoV-2. Following the fast and pandemic-scale spread of COVID-19, mutations in SARS-CoV-2 have raised new diagnostic challenges which include the redesign of the oligonucleotide sequences used in RT-PCR assays to avoid potential primer-sample mismatches, and decrease sensitivities. Since the initial wave of the pandemic, some regions had experienced fresh outbreaks, predisposing people to be susceptible to SARS-CoV-2 re-infection. Hence, this article sought to offer detailed biology of SARS-CoV-2 re-infections and their implications on immune response milieu, diagnostic laboratory tests and control measures against COVID-19.

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1. Introduction

The news of COVID-19 re-infection after months of recovery in a male patient showed how the immune response should work. This suggested that the immune system through its memory keeping abilities might have remembered its previous encounter with SARS-CoV-2 and swung into action, preventing the re-infection before it could do much damage [1]. On the contrary, more severe symptoms of re-infection cases were reported by public-health workers in Nevada [2]. This had left scientist and researchers with the questions of the possibility of the immune system failing to protect against the virus and also leaving the system more prone to SARS-CoV-2 viral attack. Duelling anecdotes are common in the see-saw world of the COVID-19 pandemic, and a firm conclusion cannot be drawn about long-term immune responses to SARS-CoV-2 from just a few cases [2].

It was believed for months that the second infection (re-infection) was merely a continuation of the first, but recent findings in the disparity in variants between the sequencing of the viral genome of first and second infections from Hong Kong and Nevada teams respectively seemingly rule out the initial belief [1, 2, 3]. It is also worthy of note that a general conclusion cannot be drawn from only two sets of cases as reported by Hong Kong and Nevada teams, and it is still unclear how frequently re-infections can occur. With over 90 million known coronavirus infections worldwide so far, a few re-infections might not be a cause to worry. More information on the prevalence of re-infection is needed. Since the initial wave of the pandemic, some regions had experienced fresh outbreaks, predisposing people to be susceptible to SARS-CoV-2 re-infection. In the Hong Kong re-infection case study, it was reported to have occurred after he had travelled to Spain and was screened for SARS-CoV-2 at the airport on his return to Hong Kong. Also, following the relieve from the first wave of the pandemic, scientists in public-health laboratories are beginning to find their feet again, expanding their horizon of epidemic surveillance in areas of tracking re-infections, protocols that can rapidly sequence large numbers of viral genomes from positive SARS-CoV-2 tests. All of these will make it easier to find and verify re-infections in the near future.

Cases with possible re-infection with SARS-CoV-2 have been recently reported in different parts of the world [4]. In many of these instances, it is difficult to differentiate a diagnostic true re-infection or a positive Polymerase Chain Reaction (PCR) as a result of the body forming a memory cell of a previous episode of an infection.

Cases of prolonged PCR-positive result had been reported among some individuals who have recovered from the SARS-CoV-2 infection [5]. The duration of viral RNA detection has been shown to vary. In some instances, viral RNA is detected 104 days after the onset of symptoms from upper respiratory samples [6, 7, 8]. More so, intermittent negative PCR tests have been reported in some patients, especially when SARS-CoV-2 concentration in the specimen becomes relatively low or undetected by the PCR test [4].

It is noteworthy that the detection of SARS-CoV-2 RNA does not always represent viable infectious virus in a patient. Additional challenges of lack of testing facilities and genetic sequencing can also lead to the error of classifying suspected cases as ‘confirmed’ re-infections. This is further complicated by the lack of established protocol and criteria for the identification of re-infections. Consequently, there is a need for additional tests to confirm for the viability of the virus and test results must be interpreted alongside the clinical and epidemiological presentation of individual patients.

Recent published data describing re-infections based on genetic sequencing as confirmation of second infections with SARS-CoV-2, following a first confirmed infection will provide insight into the features and recurrence of re-infection is pivotal, as it will influence our understanding of acquired immune response following initial SARS-CoV-2 infection [1]. Furthermore, media reports of cases in the Netherlands, Spain and several additional cases globally that are under investigation may also be of importance [9, 10, 11, 12]. Hence, this article sought to offer detailed biology of SARS-CoV-2 re-infections and their implications on immune response milieu, diagnostic laboratory tests and control measures against COVID-19.

2. SARS-CoV-2 mutation mechanisms

To understand how SARS-CoV-2 enters the cell, there is a need to discuss the virus mutation mechanisms. Generally, SARS-CoV-2 enters the cell using the human angiotensin-converting enzyme (ACE)-2 and human proteases as an entry activator [13]. This entry is through its receptor-binding domain (RBD) proteolytically activated by human proteases [14], and mediated by a viral spike protein which is a glycoprotein with two domains S1 and S2. The domain S1 initiates infection by interacting with host receptor ACE-2 and inducing conformation changes, spike protein S2 that acts as a class 1 viral fusion protein which mediates virion fusion to the cell membrane [15].

A study conducted in the United States of America showed that SARS-CoV-2 spike putative PPC site mutation affected its cleavage to S1 and S2, as the mutant SARS-CoV-2 S-protein was no longer cleaved. This could result to poor PPC motif fusion of SARS-CoV-2 during viral packaging, a step critical for viral entry into HeLa cells, Calu-3 cells, and MRC-5 cells [13].

As SARS-CoV-2 continues to adapt to its human host, a mutation known as D614G makes was seen in the virus’s spike protein. While the D614G mutation forms a part of a clade known as the G clade. A clade is like a lineage in a phylogenetic tree [15]. Clade G is mostly prevalent in Europe and it carries the D614G in the virus spike protein [16]. Korber et al. [15] found that the G clade varied from the original form from Wuhan and is linked to 3 other mutations indicating that the D614G is more transmissible than the original Wuhan form. Aside from this, there are ~8000 reported single nucleotide polymorphisms in the SARS-CoV-2 genomes than could result to changes in its infectivity. However, the D614G pattern is consistent throughout the globe with very rare exceptions. In another study, it was proved that the dominant form of SARS-CoV-2 is the variant carrying the D614G spike mutation, which enables the primary interaction of the virus with the host cell. Although, other non-mutated spike protein also interacts with the host protein [17]. Mutations occur within coronaviruses in three mutually inclusive processes. First, during viral replication some mutation may arise due to a copying error. However, this can be reduced in SARS-CoV-2 because coronavirus polymerase enzyme contains a proofreading mechanism [18, 19]. Second, coinfection of same host due to a recombination between two viral lineages may trigger genomic variability [20]. Third, mutations acquired through evolution can undergo induction by host RNA editing mechanisms which is associated with innate immune response [21, 22, 23]. Currently, there is no evidence that recurrent mutations in SARS-CoV-2 can lead to increased viral transmission. A recent study done in the United Kingdom showed that recurrent mutations in SARS-CoV-2 are not associated with increased viral transmission [23]. Instead, it is triggered by host immunity through RNA editing systems which are expected to be selectively neutral. Conversely, these mutations can be used to monitor the spread of SARS-CoV-2 [24].

One study by Becerra-Flores and Cardozo [25] showed that the G614 variant in spike had higher infectiousness and spread more rapidly than D614. This finding has an important implication for vaccine development and immunotherapeutic intervention. Vaccine platforms that elicit broadly neutralizing antibodies against both D614G and D614 should be considered. Furthermore, there are concerns that COVID-19 patients carrying D614G mutant might not respond to transfusion of neutralizing antibody (Nabs) from a donor with an S protein variant. Despite the significance of the immunotherapeutic use of Nabs in some COVID-19 patients, they could potentially trigger immunopathogenic processes in COVID-19 patients with dissimilar viral genome content or enhanced infectivity [25]. Interestingly, the risk of being hospitalised with G614 spike-pseudotyped viruses was less significant. The study found that there was no significant impact on disease severity, but G614 is
associated with higher case fatality rates (proportion of people who die from COVID-19 among all diagnosed individuals whether hospitalized or not) across countries [25]. In terms of mortality data, a national study done in China demonstrated that deaths from COVID-19 in countries out of China was three times higher (15.2%) compared to 5.6% in China [26]. This could be due to the emergence of viral mutations and evolution capability of SARS-CoV-2 over time. However, it remains unclear if the different case fatality rate reported across countries may be the import of clade’s differences in virulence as observed by a study in the United Sates [27].

SARS-CoV-2 mutations are in the S protein (nt23403), RNA polymerase (nt14408), RdRp (nt14408), and nucleoprotein (nt28881) [28, 29]. A study done in China showed that mutations could occur in various regions of the SARS-CoV-2 virus like the ORF1ab, S, ORF3a, ORF8, and N regions. The study showed a mutation rate of 30.53% and 29.47% in ORF8 and ORF1a, respectively [29]. Similarly, in an evaluation study of emerging SARS-CoV-2 mutation hotspots across 4 geographic areas, Pachetti et al. [29] showed that mutations located at positions 2891, 3036, 14408, 23403, and 28881 were mainly observed in Europe, while those located at 17746, 17857 and 18060 were present in North America. This finding suggests a differential pattern of mutation across the countries due to founder effect.

Mutation of SARS-CoV-2 occurs in distinct patterns across countries. As demonstrated by data from CoV_GLUE S-D614G and nsp-12-P323L, all continents had the two main mutations that determined the virus clade G except for three cases in Asia. However, D614G in the S protein was often found co-evolving with the P323L mutation in the nsp12 protein and is found in 2342 and 2318 samples, respectively. Both mutations – D614G and P323L were reported in Switzerland, Spain, Italy, and France. Furthermore, ORF8-L84S is the third most frequent mutation that determines the virus clade S found. This was found in 740 sequences reported in 71 European cases reported by Coppée et al. [30]. Meanwhile, association of ORF8 mutation in S clade with mutations in ORF3a, nsp4, and the N proteins of SARS-CoV-2 has been demonstrated by Lorusso et al [31]. This has implication for possible reinfection potential. A study demonstrated that SARS-CoV-2 has mutated in such a manner that facilitated its rapid transmission all over the world [32]. Virus mutation may well pose a huge challenge to COVID-19 vaccine – nevertheless this can be prevented. If SARS-CoV-2 mutates as result of its reaction to a COVID vaccine, there are various paths it might assume. Evolution- and mutation-proof vaccines provides protection against all circulating strains, so that entrants of new strains are covered by the vaccine. To achieve this, an extra effort will be needed during vaccine clinical trials. For instance, by testing upper respiratory swab samples of people who have received the experimental vaccine, researchers can tell the virus suppressed level in subjects. Furthermore, by analyzing the whole genome of SARS-CoV-2 in vaccinated people, it might be possible to study the evolutionary escape process (if present). Blood samples from vaccines can be used to work out the number of sites on the SARS-CoV-2 that are being attacked by vaccine-induced immunity. These put together can provide detailed information about the mutation and evolution processes taking place in SARS-CoV-2 so that adjustments be provided in an updated vaccine candidate during further re-designing and development stages.

3. Biology of SARS-CoV-2 re-infection

Early experimental study showed that re-infection by coronaviruses is possible. However, little was known about the possibility of re-infection by SARS-CoV-2 since the COVID-19 pandemic is still in its early phase [33]. A study on rhesus macaques showed that re-infection could not occur after challenging the monkeys with the same dose of SARS-CoV-2 strain as the first infection [34]. However, recent findings have shown that re-infection is possible, specifically with a different strain, with confirmed cases of re-infection in Hong Kong [1] and Nevada [3]. To et al. [1] and Edridge et al. [33] suggested that natural infection may be responsible for re-infection in coronaviruses (HCoV-NL63, HCoV-229e, HCoV-OC43 and HCoV-HKU1), and this could be a general occurrence for all coronaviruses, including SARS-CoV-2.

Differences were observed in the genomes of confirmed re-infection cases of SARS-CoV-2. The differences between the initial and subsequent infection of the viral genomes are attributed to the clade/lineage, the number of single nucleotide variants (SNVs), the difference in the amino acids and number of dinucleotide multi-nucleotide variant (MNV) [1, 3]. Although the re-infection case in Nevada, USA belongs to the same clade as the first infection, clade 20C, there are some genomic differences as obtained from the genomic sequence analysis (Table 1).

According to Jain et al. [35], clade 20C possess genomic variants in the regions C14408T, A23403G, C1059T, and G25563T. However, viral particle B has an additional variant in the region C3037T [3]. The mutation in the viral genome A is nonsense; however, viral genome B included a synonymous mutation in the C3037T region [27]. While the A23403G occurred in the S region, the other mutation occurred in the ORF1ab region, which is the longest ORF in the SARS-CoV-2 genome [36]. The extra change in the S region could be responsible for the severity in the re-infection case in the Nevada case.

According to the To et al. [1], they highlighted that the viral genome in the first episode of infection in the re-infection case in Hong Kong was of the GISAID Clade V. Next strain clade 19A, Pangolin lineage B.2 and with a probability of 0.99; while the re-infection viral genome was of the GISAID Clade G, Next strain 20A, Pangolin lineage B.1.79 with a probability of 0.70. Specifically, lineage B.1 belongs to variant G614 which is widely distributed globally with substitution in the regions C241T, C3037T, C14408T and G23403A; while lineage B.2 belongs to the variant V351 [37]. Furthermore, these two viral genomes differ from one another by changes in the amino acids in the spike protein, accessory proteins (ORF3a, ORF8 and ORF10), nucleoprotein, membrane protein and non-structural proteins (NSP3, NSP5, NSP6, NSP12) [1]. The genome of the first virus is related to the Clade GR obtained in England between March and April 2020 [1], with varying nucleotide and amino acids mutations (GISAID Database). Analysis of 10,022 samples to understand the genomic variability of SARS-CoV-2 also showed that G614 variant had been the most common variant since the onset of the pandemic in December 2019 (Koyama, 2020 #462) [36]. Evidence has shown that G614 has a higher titre of viral particles in upper respiratory tract specimens [15, 37], but it is associated with lowered RT-PCR cycle thresholds and not necessarily increased diseases severity [15].

The re-infection case in Belgium showed that the first infection belongs to the lineage B.1.1 while the second infection belong to lineage A, with eleven genomic mutations identified in the two strains [38]. According to Gupta et al. [39] there were some genetic variations in the case of re-infection among the two healthcare workers identified in India. Using WGS, there were six non-synonymous (NS) variations in patient 1 when the two virus strains were compared. Virus strains in patient 2 showed nine NS variations between the two strains in the different genomic regions. Most of these variations were observed between the orf1b and envelope protein region.

4. Immune response during SARS-CoV-2 infection

The defensive characteristics of antibodies following SARS-CoV-2 infection is not still been fully understood. Nevertheless, regular identification of antibodies and quantification of their titres could provide information about antiviral protection associate produced by antibodies over time [40]. The binding of IgG antibodies with SARS-CoV-2 have been reported to occur in people over a period of time during the day 10 and day 21 after infection [41, 42]. One of the available studies showed that more patients (>91%) developed IgG seropositivity due to primary SARS-CoV-2 infection [43]. There are still needs to determine long-term survival of antibodies to SARS-CoV-2, but it is known that antibody levels against some coronaviruses decline with time (the range is 12–52 weeks counting from the
Table 1. Studies that reported cases of SARS-CoV-2 re-infection.

| Country (Citation) | Age/gender/general health condition | Period between episodes (RT-PCR positive outcomes) | No. of cases | Key Clinical findings | Scheduling of RT-PCR and Ct figures | Sequencing | Mutation | Immunoglobulin testing |
|-------------------|-----------------------------------|-----------------------------------------------|---------|----------------------|-----------------------------------|-----------|-----------|---------------------|
| Hong Kong, China (To et al) [1] | 33-year-old immunocompetent male | 142 days | 1 | First episode: dry cough, fever, headache. Second episode: no symptoms | First episode: positive outcome 3 days after symptom onset with Ct of 30.5. Second episode: positive outcome 1–3 days (Ct 26–28) & 5 days (Ct 32) post symptom onset. | 1st and 2nd viral genomes from dissimilar lineages and differentiated by 24 nucleotides. First episode: Next strain 19A/GISAID V/Rambout clade B.2 (Hong Kong). Second episode: Next strain 20A/GISAID G/Rambout B.1.79 (Spain). | Amino acid variations in Spike protein (N-terminal domain, upstream helix, subdomain 2), nucleoprotein, non-structural proteins (NSP3, NSP5-6, NSP12), accessory proteins (ORF3a, ORF8, ORF16). | First episode: negative for IgG 10 days after symptom commenced. Second episode: negative for IgG 1–3 days following hospitalization with a reactive outcome on the 5th day. |
| Washoe, Nevada, USA (Tillet et al) [3] | 25-year-old immunocompetent male | 48 days | 1 | First episode: less severe symptoms (dry cough, sore throat, diarrhea, headache). Second episode: more severe symptoms (pyrexia, headache, dry cough, dizziness, hypoxia) with stronger immune response. | First episode: positive outcome on the 24th day following the commencement of symptoms (Ct 35.2). Second episode: positive outcome on the 6th day following the commencement of symptoms (Ct 35.3). | 1st and 2nd viral genomes originated from a common lineage (Next strain 20C) and differentiated by 7 nucleotides. | SNVs (25563G>T, 3037C>T, 14408C>T, 19132A>G) which results in 4 amino acid changes [NSP12 (P323L), NSP13 (S485L), SP (D614G), ORF3a (Q57H)]. Second episode: 10 SNPs (C1457T, C8782T, C17531C, C17747T, A17858G, C18060T, C18877T, C23403G, G25563T) which results in 5 amino acid changes [NSP2 (R218C), NSP3 (I432T, P504L, Y541C), ORF8 (L84S)]. | First episode: no immunoglobulin test was done. Second episode: reactive for IgG/IgM on day 7 post symptom onset. |
| Belgium (Van Elslande et al.) [38] | 51-year-old immunocompetent female on corticosteroid for asthma management | 93 days | 1 | First episode: pyrexia, migraine, dry cough, dyspnea, chest pain. Second episode: migraine, dry cough, fatigue. | First episode: N1-gene (Ct 25.6). Second episode: N1-gene (Ct 32.6). | 1st and 2nd viral genomes from dissimilar lineages and differentiated by 11 nucleotides. First episode: Rambout clade B.1.1. Second episode: Rambout clade A. | Amino acid variations in Spike protein [G24043A, A23873G, C24726T], nucleoprotein [A28881G, A28882G, C28883G], accessory proteins [ORF1a (C3037T, C8782T, C11654T), ORF1b (T14408C, T17427G)]. | First episode: no immunoglobulin test was done. Second episode: reactive for IgG on day 7 with a value of 1/34 and for neutralizing antibodies on the 6th week with a value of 1/320 following symptom onset. |
| Ecuador (Prado-Vivar et al.) [52] | 46-year-old immunocompetent male | 63 days | 1 | First episode: less severe symptoms (migraine, drowsiness). Second episode: more severe symptoms (pyrexia, dry cough, dyspnea, sore throat). | First episode: positive outcome on the 11th day following the commencement of symptoms (Ct 36.85, ORF3a gene). Second episode: positive outcome on the 4th day following the commencement of symptoms. | No common mutations between the viral sequences of both first and second episodes. First episode: 8 SNPs (C2113T, C3037T, C7765T, C14408T, C17531C, C18877T, A23403G, G25563T) which results in 4 amino acid changes [NSP2 (R218C), NSP3 (I432T, P504L, Y541C), ORF8 (L84S)]. Second episode: 10 SNPs (C1457T, C8782T, C17531C, C17747T, A17858G, C18060T, C18877T, A24694T, T28144C) which result in 5 amino acid changes [NSP2 (R218C), NSP3 (I432T, P504L, Y541C), ORF8 (L84S)]. | First episode: negative for IgG 4 days following symptom onset. Second episode: positive for IgG on the 30th day with a value of 34.1 following symptom onset. |
| India (Gupta et al.) [39] | 25-year-old immunocompetent male | 108 days | 2 | First & second episodes: no symptoms. | First episode: positive outcome (Ct 36). Second episode: positive outcome (Ct 16.6). | 1st and 2nd viral genomes with 9 distinctive variants. | Synonymous mutations: C241T, C6445T, G11383A, T11408C, C18877T, C25207T, C26735T. Nonsynonymous mutations: T1947C, G17584T, A24694T, C23403G, C23934T, G25563T. | First and second episodes: no immunoglobulin test was done. |

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### Table 1 (continued)

| Abbreviations | CRP: C-reactive protein; SNV: Single nucleotide variants; SNP: Single nucleotide polymorphism; NSP: non-structural protein; SP: Spike protein; NS: Nonsynonymous |
|---------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

- **25-year-old female**: First episode: positive outcome (Ct 28.16). Second episode: positive outcome (Ct 16.92).

- **1 and 2 viral genomes with synonymous mutations**:
  - C241T
  - C18877T
  - C23929T
  - C26735T
  - C29215T

- **1 and 2 viral genomes with nonsynonymous mutations**:
  - C3267T
  - C13730T
  - C14408T
  - G17584T
  - G19109T
  - T22882G
  - A23403G
  - G24794A
  - G25563T

**Abbreviations**
- CRP: C-reactive protein
- SNV: Single nucleotide variants
- SNP: Single nucleotide polymorphism
- NSP: non-structural protein
- SP: Spike protein
- NS: Nonsynonymous

**Inception of symptoms**
- It has been shown that the antibody level of SARS-CoV-2 can last up to 94 days [4] after infection, with recent studies showing that antibody levels rise within 21–28 days, then remain unwavering for about 120 days [45]. However, this significantly varies with the severity of COVID-19 [46] considering that the magnitude and timing of antibody levels and the effect of cellular immunity have not been adequately studied in large groups of people.

**Several reports have shown that the antibody response levels and severity of the disease appears to be correlated** [47] and there are suggestions that SARS-CoV-2 antibody-associated defence could not be long-lasting in asymptomatic persons [48]. Wang et al. [49] noted a significantly lower antibody response in asymptomatic patients with the low-grade disease, together with IgM lower response and antiviral neutralizing rates, in comparison with the patients that have severe COVID-19. These results put together, showed that after the initial SARS-CoV-2 infection, many patients appear to have elevated antiviral response, just that there might be a decline in the defence mechanism over time. This seems to be expected from people with underlying severe disease that are less in number; and could be the situation described some studies.

**The present pandemic of COVID-19 contains a wide range of cases ranging from very serious or terminally ill patients to undiagnosed (asymptomatic) cases, forming a huge number. In most severe cases, immune dysregulation is assumed to carry out a key role. A recent study conducted by researchers at the Sanford Burnham Prebys Medical Discovery Institute and the University of California in San Francisco reported that an encoded protein via the sub-genomic sequence ORF9c of SARS-CoV-2 stands liable for the little antiviral reaction in infected persons permitting active viral replication in the host [50]. This can lead to a successful reticence on the immune system against viral infection.**

**In almost all investigated individuals, anti-SARS-CoV-2 IgG have consistently been detected at the end of the follow-up period (up to 94 days) and a neutralising antibody response is developed by more than 90% of individuals who have been infected [4]. In some animal models, a previous SARS-CoV-2 infection has been reported to offer protection against subsequent challenge in rhesus macaques [51]. There is a paucity of data on the duration of immunity after SARS-CoV-2 infection. In addition, there is no categorical evidence on the role of humoral immune response in the clearance of SARS-CoV-2 infection.**

**Although re-infections are possible, the specific circumstances, associated symptoms and disease progression, as well as the overall extent, are yet to be explicitly researched and comprehended. According to To et al [1], their case study did not have any detectable antibodies as at re-infection, but developed detectable neutralising antibodies; however, after the infection episode. According to case reports by Tillett et al [3] and Van Elslande et al [38], the antibody status were not measured after the patient’s first infection, but following their second infection, antibody responses were observed.**

**According to a case reported by Prado-Vivar et al [52], antibodies were not detected as at the time of the first episode of infection, even though the measurement was only done four days after the onset of symptoms; antibodies were present, however, after the second episode of infection [52]. At the moment, there is dearth of information on the role of antibodies and level of neutralising antibodies, as well as the durability of the antibodies levels following SARS-CoV-2 infection. There is need for investigations using larger population groups to properly define these. In the described re-infection cases, the virus isolates were confirmed to contain different mutations. This confirmed infections by new SARS-CoV-2 variants in the patients. For proper understanding of the possibility of re-infections and possible immune response escape, the size of mutations as well as the exact points of the mutations in the genome might be helpful in biomedical intervention against the COVID-19 pandemic. There is need, also, for investigations that will evaluate the chances of common mutations in the viral genomes from re-infected
patients that could shed light on the virus’s re-infection ability. Additionally, it is also pertinent to elucidate the level of divergence that a SARS-CoV-2 isolate needs in order to be able to reinfect a previously infected person. Furthermore, the role of cellular immunity in the prevention of COVID-19 re-infection needs to be investigated. Available data have shown that persons that got re-infected with SARS-CoV-2 had previous mild symptom from 1st infection [24]. However, the severity of a second infection varies among patients. The reasons for this still remain enigma and require further investigations. In a bid to provide an explanation to this phenomenon, it was opined that asymptomatic patients and those with mild SARS-CoV-2 infection evoke weaker immune response to SARS-CoV-2 (Figure 1) and this might be the reason they tend to be susceptible for reinfection. The few cases of reinfection identified so far had been patients that had shown symptoms, which means asymptomatic re-infected patient could be easily missed [24].

4.1. B cells and interferon response in COVID-19

Type I interferons (IFNs) are immunoregulatory cytokines that play important role in immune response to viral infections. Studies have shown that Type I interferons (IFNs) affects both innate and adaptive immune responses [53, 54]. They enhance B cells response to viral infection, cytotoxic production and neutralizing antibodies production. However, unregulated IFNs can lead to autoantibody production [53]. The study by Bastard et al. showed that autoantibodies generated by B

![Diagram](image1.png)

**Figure 1.** Diagram depicting differences in sensitivity of RT-PCR and serological diagnostic test to SARS-CoV-2 at different stages of infection. RT-PCR is sensitive at the early stage of infection but becomes less sensitive over time due to low level of RNA. This may result in false negative result. If the infection re-emerged it can be wrongly assumed to be reinfection. On the other hand, serological tests are less sensitive at the early stage of infection but sensitivity increases towards the advance and recovery stages. However, antibody titre starts to decline 1–2 months after the acute infection.

![Diagram](image2.png)

**Figure 2.** Antibody response during COVID-19 severe cases.
cells in 10% of patient with life-threatening COVID-19 pneumonia targeted type I IFNs [54]. These autoantibodies counteract the ability of IFNs to block SARS-CoV-2 infection thereby, making the host more susceptible to viral infection and in turn more severe symptoms [54].

Additionally, immunological retention is an important aspect of robust immunity against SARS-CoV-2 infection. Based on previous reports that indicated COVID-19 patients to have exhibited varied immune responses, a recent study provided details on the use of cell-based immune indicators for improved patient outcomes in convalescent plasma specimens [55].

Basically, response to viral antigens by human B cells is by secretion of germline or near-germline antibodies from plasmablasts on the exterior of the follicles [55]. Assoon as the T cells bind to the CD40 surface markers, resulting to the release of specific cytokines, B cells go into a process referred to as class switching. Consequently, they will now be found inside the germinal cores within various lymphoid organs and develop functionally. This brings about the production of long-lived plasma cells and memory B cells both of which can respond to a recurrent issue with similar or another antigen [55].

Plasmablast increase in the early B cell response has been proposed to result to poor patient outcomes. Contrarily, memory B cells produced post SARS-CoV-2 infection confer strong specific immunity. They are both seen in the classical CD27+ class-switched type, activated CD24-form and the natural CD27+ type, which is similar to an innate immune cell, with both IgG and IgM markers.

A new study discovered that higher number of memory B cells may indicate an efficacious response to an acute infection and aid in understanding T cell response. Both switched and unswitched memory B cells are associated with a shorter length of symptoms [56]. The degree of IgM + memory is highly associated with anti-RBD IgG1 antibody response. Hence, it seems like some COVID-19 patients do present a memory response that is protective, either before or after the production of the IgM + memory cells [56].

Certain B cell memory cross-reacts with past coronavirus infections. Nevertheless, the anti-RBD IgG1 response is proportional to the IgM + memory cells, since the IgM + memory cells do not produce switched immunoglobulins. Thus, researchers concluded that, coronavirus infections possibly produce a good amount of IgM + cells, some of which go into germinal cores and change to IgG1 production. This capacity of IgM + memory cells to transfer to germinal core following activation is very common, and might be beneficial in induction of immunity to pathogens like the coronaviruses with various extremely similar strains [55].

An apparently illogical discovery was that T bet + B cell occurrence did not correlate with that of resting memory B cells, since the former is important in B Cell immunity formation. One justification may be that they are not a crucial aspect of recovery, but are limited to acute and chronic viral infection. This part will require additional studies to explain the processes involved [56, 57, 58, 59]. Alternative explanation for the increased convalescent rates with higher amount of memory B cells is that these cells form part of a bigger pool, resulting to better T cell facilitation of B cell germinal center responses in unexposed individuals during primary infection. This elucidation is plausible since memory production is strictly connected to the production of specific antibodies against the responsible agent [56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66].

4.2. Antibody response in Severe/ICU/deceased patients with COVID-19

Majority of SARS-CoV-2 infected persons develop antibodies to the viral S and N proteins, which are mostly used as the antigens in COVID-19 serological assays. The S- protein is an essential target for broadly neutralizing antibodies against SARS-CoV-2, as they prevent viral entry into susceptible host cells [67]. Available data on the role of anti-SARS-CoV-2 in viral clearance, modulation of COVID-19 severity, and the durability of humoral immune responses after primary SARS-CoV-2 infection is limited. Furthermore, an improved understanding of humoral immune response to SARS-CoV-2 is crucial to direct strategies useful in vaccination production and immunotherapeutic (such as the use of neutralizing antibodies or convalescent plasma).

Clinical findings on the longevity of anti-SARS-CoV-2 titers are not in full conformity with one another, with some reports showing swift waning of anti-SARS-CoV-2 IgG about 90 days following after infection [48]. However, others studies reported stable anti-SARS-CoV-2 IgG titers detected several months after infection [68]. Sera anti-SARS-CoV-2 responses appear to rise in patients with severe COVID-19 compared to those asymptomatic or with mild symptoms. This observation has raised concerns about the effectiveness of humoral immune response against SARS-CoV-2 infection (Figure 2). However, a new finding suggested that the quality of anti-SARS-CoV-2 IgG and neutralizing antibody rather than quantity predicts the clinical outcome and prognosis of COVID-19 [69]. The study applied a panel of sero-diagnostic assays on COVID-19 patients who have convalesced or died due to COVID-19 [69].

Serum anti-SARS-CoV-2 IgM and IgG are mainly detected between 1- and 2-weeks post onset of symptoms (Figure 2). Whereas, SARS-CoV-2 RNA levels decline as the neutralizing antibody titters rise. Essentially, increased NAb have been demonstrated in patients with severe COVID-19. However, little is known about the involvement of humoral immune responses on COVID-19 induced-lung diseases. Of note, humoral immunity and B-cell responses to SARS-CoV-2 infection are short lived. This suggests that immunity following SARS-CoV-2 infection may drastically decline 12-24 months after primary infection [70].

Anti-SARS-CoV-2 IgM in severely infected COVID-19 patients in the ICU gets activated at the early days (1st week) post infection continues to rise until it peaks at the end of 3rd week. By the end of the 5th week, IgM levels begins to decline with a concomitant maintenance (plateau) of anti-SARS-IgG level after it got peaked at the 3rd week. The IgG persists in serum and also detectable for about 12–24 months post infection. Patients with more severe illness show higher antibody titers compared to those with milder illness. IgM and IgG levels reached their peak at weeks 4 and 6 respectively. Outpatients had lower IgM and IgG levels than inpatients. Dead patients had highest antibody level. However, antibody response in severe COVID-19 vary in individuals depending on prevailing co-morbidity and other clinical conditions of the patients.

5. Laboratory diagnosis of SARS-CoV-2 re-infection

To differentiate between SARS-CoV-2 positive cases, especially with prolonged viral shedding, from cases with true re-infection, epidemiological and virological data from every infection incidence needs thorough assessment.

COVID-19 compatible symptoms in a person that tested positive to SARS-CoV-2 need to be assessed, and a swab taken for laboratory analysis. Some other respiratory viruses, such as seasonal influenza, causes COVID-19 similar symptoms and should be considered as differential diagnoses. Outlined below are the main criteria that should be fulfilled to identify true re-infection in combination with individual overall clinical assessment [43].

a. Laboratory confirmation of two different strains of infection (the incident is determined or supported by phylogenetic and epidemiological data) with timely classified illness/infection episodes (least estimated period)

b. Additional investigation of suspected or confirmed/probable re-infections, to further validate that re-infection occurred and document patients’ characteristics after exposure in the two infection episodes will provide better insight into the causes of re-infection. Such knowledge could further provide guidance on public health interventions.
6. Criteria for SARS-CoV-2 re-infection diagnosis

False positive SARS-CoV-2 RT-PCR can occur due to error during pre-analytical and analytical phases of testing, especially during sample collection and testing. Contamination during analysis can also lead to false positive result and it was stated that test procedure with high sensitivity and specificity could lead to a low positive predictive value in areas with low SARS-CoV-2 infections [43].

The period between the first infection and supposed re-infection might play a role in determining true infection. Due to decline in antibody levels with subsequent waning immunity, infection between a confirmed RT-PCR negative test after infection and another positive result could be considered as a re-infection. This is however, dependent on the longer time lapse between these two events. In contrast, another infection after a confirmed RT-PCR negative result with a short time-lapse will probably be the detection of the residual viral particle as supposed to a re-infection.

Presence of viral RNA fragments in the absence of viable virus can bring about positive RT-PCR result. This false positivity could be ruled out by:

a. Virus culture: A positive culture will indicate a viable virus and a true infection is of a different strain. However, a negative culture result will indicate a non-viable viral RNA shedding which cannot be ascribed to an ongoing infection [43].

b. Viral load quantification using the cycle of threshold (CT) value of PCR has shown to correlate with the viability of the virus. According to a recent pre-published study, a viral load of 6.610 RNA copies/mL could make the probability of detecting the virus less than 5%. However, this method could not be relied upon as it has not been established and validated for this purpose [43].

c. Sequencing and phylogenetic analysis using whole-genome sequencing (WGS) could be a tool to evaluate the probability of a second infection as re-infection. However, care has to be taken when using this method as there is the plausibility of virus mutation within the host, in the case of one strain infection, and simultaneous infection with two different virus strains.

SARS-CoV-2 can infect a host alongside other seasonal coronaviruses, with re-infection with β-CoV hCoV-OC43 reported in some studies 90 days after the initial COVID-19 infection [70]. Modelling has shown that there is an average of 45 weeks of protective immunity against hCoV-OC43 and hCoV-HKU1 before re-infection can occur [70]. Nevertheless, re-infection with other coronaviruses can occur with stable and high antibody titres. In a study where 133,266 laboratory-confirmed cases were evaluated with 243 positive swabs after 45 days of their first SARS-CoV-2 infection, it was found that 54 of these cases had re-infection considering their Ct values or symptoms of COVID-19 [71]. However, these cases cannot be determined to be true re-infection because neither WGS nor viral culture was used to identify these cases. Yet, the risk and incidence of re-infection was estimated to be 0.04% (95% CI: 0.03–0.05%) and 1.09 (95% CI: 0.84–1.42) per 10,000 persons respectively [71].

The media is inundated with potential cases of re-infection which are under investigation; however, six cases of confirmed SARS-CoV-2 re-infection have been reported to date. Although, re-infection is a rare scenario coupled with non-traceability of samples from the first episode...
and previously positive laboratory results in some settings, increase in testing of symptomatic and asymptomatic persons will increase the likelihood of identifying re-infection cases. This could help in understanding the factors that favour re-infection.

In a recent review study to understand the infectiousness of SARS-CoV-2, five of the re-infection cases were included [4]. No onward transmission to close contacts was observed in the cases of re-infection. Also, there is limited scientific evidence to support the infectiousness of a re-infected person and based on the small sample size of the re-infected cases with non-employability of WGS to decipher the phylogeny, these cases might not be actual cases of re-infection. However, symptomatic and asymptomatic re-infected individuals should be managed as if it is the first infection while ensuring COVID-19 universal precautions.

7. Clinical and laboratory features of SARS-CoV-2 re-infection cases

In August, 2020, the previous study reported, based on whole-genome sequencing and serological data, the evidence of re-infection in a 33-year-old Hong Kong resident with dissimilar viral strain from the first infection which was 142 days apart (Table 1). A following first episode of the symptomatic viral infection, clinical hallmarks of the infection which include dry cough, sore throat, pyrexia and headache lasted for 72 h with detectable neutralising immunoglobulin which began to wane within 8 weeks as these symptoms were resolved with therapeutic intervention. The resolution of the clinical manifestations was further marked by the absence of detectable viral nucleic acid for two consecutive PCR testing using samples from the throat and the nasopharyngeal tract. The waning effect of these immunoglobulins was predicated on predisposing the patient to the second wave of infection by a different viral strain from a dissimilar lineage with 24 nucleotide variants compared to the first strain and was observed to be characterised by no clinical manifestations which is associated with low antibody titters, slightly raised C-reactive protein and a relatively elevated viral load with progressive decrease over a period. The evidence for re-infection of these different strains was further consolidated by the phylogenetic tracing of the first strain from Hong Kong while the second strain was contracted from Spain were [1].

Four months earlier, similar viral hallmarks (dry cough, nausea, sore throat, diarrhoea and headache) were reported in a 25-year-old resident male of Washoe in Nevada, USA. The immunocompetent patient, who had no history of underlying pathology, was observed with symptoms which resolved with steady improvement between April and May 2020 during the period of isolation. However, towards the end of the following month, the symptoms resurfaced with evidence of atypical pneumonia as revealed by chest radiography. Further observation revealed that the patient presented with hypoxia and breathing discomfort. During the two clinical presentations by the patient, serological markers (IgM/IgG) were detected, and nucleic acid amplification was performed using nasopharyngeal swabs to confirmed the COVID-19 status. Genomic analysis of the samples collected for the positive results of April 18, 2020 (first instance of infection) and June 5, 2020 (second instance of infection) which revealed two negative COVID-19 outcomes between these periods, revealed that a significant difference in the nucleotide sequence of the viral isolates collected during these two waves of infection. The second wave of infection was observed to be characterised with a more severe form of the viral infection with a more robust immune response compared to the first wave of infection. The computed genetic discordance disclosed value of 83.6 as opposed to 23.1, which serves as the natural rate of substitution. The findings revealed that the patient contracted two genetically different viruses on two distinct occasions which clearly defeats the possibility for herd immunity [3].

Similar to the reported cases in Hong Kong and the United States, the third evidence of re-infection was reported with a first wave of infection in March 2020 in a 51-year-old woman from Belgium with similar clinical presentations (pyrexia, muscular weakness, dyspnea, dry cough, headache, chest pain, loss of smell and taste) of symptomatic re-infection no less than 3 months and 3 days following a moderate COVID-19 re-infection. Unlike previous cases, although she was reported to be immunocompetent, she had a respiratory disorder (asthma) which was managed using corticosteroids. Apart from slight elevations in the liver enzyme profile, biochemical and haematological analysis revealed no obvious signs of pathology with a 94% oxygen saturation. After a three months period, her symptoms were observed to relapse with similar complaints of the first wave of infection (dry cough, headache, rhinitis and muscle weakness) and no migration history. Although the severity of this second episode of infection was milder compared to the first episode, it resolved within a shorter period (7 days) and both samples from the first and second episodes tested positive for immunological markers (IgM/IgG). Genetic discordance was observed between the viral isolates from the first and second episodes of infection based on 11 nucleotide variants which further buttressed the evidence for infections with two genetically distinct viral strains [38].

Another case of re-infection was also reported in Ecuador which was in contrast to the Nevada case but similarly to the cases of Hong Kong and Belgium, the genetic confirmation test revealed two waves of infection (in May and in July 2020 respectively) which had a second episode of infection that was much more severe with a surge in antibody titters (IgM/IgG) compared to the first episode of infection which were four weeks apart during when the COVID-19 status of the patient was confirmed to be negative using RT-PCR [52].

In the case of India which involved reported evidence of asymptomatic re-infection in two healthcare professionals (male and female aged 25 and 28 respectively) unlike previous reports, RT-PCR confirmed their two episodes (5th and 17th of May 2020; and 21st August and 5th September 2020) of positive COVID-19 status with three months apart during when they were confirmed negative on 13th and 27th May 2020 respectively. Genomic and phylogenetic analysis was used to confirm the genetically dissimilar viral strains which ruled out any suspicion for viral shedding or reactivation [39]. More recently on 25th September, 2020, Mahallawi reported another case of SARS-CoV-2 reinfection in Saudi Arabia [40]. This was case of a 31-year-old man that presented with myalgia, headache (with no other symptoms) and was not suffering from any chronic disease.

8. SARS-CoV-2 mutations and consequences on laboratory tests

Following the fast and pandemic-scale spread of COVID-19, mutations in SARS-CoV-2 has raised new diagnostic challenges which include the redesign of the oligonucleotide sequences used in RT-qPCR assays to avoid potential primer–sample mismatches, and decrease sensitivities [72]. Reportedly, as at 30th March 2020, of all high coverage of 1825 SARS-CoV-2 genome sequences deposited in the Global Initiative on Sharing All Influenza Data (GISAID) database [73]. About 79% (26 of 33) of the primer binding sites used in the RT-qPCR assays were mutated in at least one genome [73]. Of significance was the GGG substitution to AAC at the beginning of the binding site of the forward primer in the gene that encodes for the nucleocapsid phosphoprotein. The AAC variant was found in 14% (258 of 1825) of the genomes isolated and sequenced in 24 different countries.

Despite the possibility of sequencing errors, the consistent detection of some specific variants, there is need to continue to optimise the oligonucleotides used in assays being developed. The global sharing of SARS-CoV-2 genomes and the frequent updating of reports on sequence analysis that are available on the GISAID4 website will help facilitate oligonucleotide optimisation.

Since the emergence of SARS-CoV-2 continuous efforts have been made to map its genetic diversity and identify variants/mutants that have a selective advantage [74]. Some key variations of interest include changes in immune targets, such as the spike glycoprotein; changes in
primer binding and probe-binding sites, which can reduce the sensitivity of diagnostic tests; and genetic variations that might affect transmissibility and virulence [75, 76, 77].

For instance, the Cobas® system (Roche) uses a dual target assay to detect SARS-CoV-2, with qRT-PCRs targeting both the ORF1ab region and the E–gene [78]. During SARS-CoV-2 testing, Artesi et al. [78] found out that most SARS-CoV-2 test were negative at the E-gene and positive for ORF1ab region. However, the Cobas Roche system classified the diagnostic result as Subsequent genomic sequencing showed mutation that interfered with E-gene during qRT-PCR [78]. However, the C > U transition at position 26,340 of the SARS-CoV-2 genome during sequencing could be responsible for negative E-gene RT-PCR. Available evidence suggests that there was heavily interference most likely at the E-gene domain [79].

Indeed, mutation in SARS-CoV-2 genes used in COVID-19 Diagnostics have occurred (Table 2). These mutations can affect the accuracy of RT-PCR test by producing false negative result in a single qRT-PCR primer/ probe assay [78]. Thus, during SARS-CoV-2 assay, two or more positions of the viral genome should be targeted, as mutations in all targeted genes in very unlikely for now. However, there is the need for continuous global molecular surveillance of SARS-CoV-2 to promptly detect future mutation that could affect RT-PCR test results.

In another instance, Sun et al. [79] reported a 12-hp deletion on E gene on mutant and wild-type SARS-CoV-2 strains isolated from the same clinical sample. Furthermore, Ziegler et al. [80] found that a single nucleotide polymorphism (SNP) in the E gene SARS-CoV-2 from a patient interfered with detection in a widely used RT-PCR assay. Consequently, this underscores the necessity of targeting two independent essential genes of SARS-CoV-2 for reliable detection.

Getting an adequate epidemiological data about the global COVID-19 pandemic requires accurate RT-PCR testing to identify SARS-CoV-2 infected individuals. Artesi et al. [78] further identified a C-to-U transition at position 26,340 of the SARS-CoV-2 genome which is associated with failure of the cobas® SARS-CoV-2 E-gene qRT-PCR in eight patients. This work highlights the necessity of monitoring SARS-CoV-2 for the emergence of SNPs especially in the case reinfection which might negatively affect RT-PCR assays used for COVID-19 diagnostics.

9. Implications of SARS-CoV-2 re-infections on pandemic control measures

Prior to the discovery of the ability of SARS-CoV-2 to be involved in a number of re-infection cases, there have been several notions that individuals who recovered from COVID-19 infection develop immunity and are protected against viral re-infection due to the presence of immunological response by serum neutralising immunoglobulins. The notion which was held by certain governments on the grounds that patients who recovered from COVID-19 were fortified against re-infection and could afford to travel as well as resume work during the pandemic was however, debunked by WHO on the basis of the lack of substantial evidence to confirm the relationship between neutralising antibodies and immunity against re-infection [80]. Despite this development, several studies have demonstrated that the viral RNA can replicate and diminish in numerical strength in body fluids of convalescent infected individuals for up to 12 weeks [81]. However, researchers before the advent of the documentation of the first COVID-19 re-infection case could not understand whether these group of infected individuals are experiencing either viral shedding or re-infection by a different SARS-CoV-2 strain due to the lack of evidence of COVID-19 genome sequencing to differentiate these instances. Unlike viral shedding which does not correlate with infectivity and rarely occurs for up to 4 weeks [82, 83, 84], re-infection refers to cases where an individual who was previously infected with the virus and underwent recovery, tests positive a second time for a mutated strain of the same viral pathogen.

The first of the cases of infection were reported in Hong Kong [1], which was followed by Nevada in the United States [85], Beijing in China [86], Japan [87] and Mumbai in India [88]. Most of these cases had mild symptoms, including sore throat or dry cough between 2-5 days of their first infection before recovery. Their second infection was characterised by a mutated strain which intensified the pyrexia, headache and myalgia. These latest reports have laid to rest the persistent queries and concerns on the possibility of SARS-CoV-2 to reinfect the human biological system, frequency of re-infection, severity of the second compared to the first infection and the possible impact of subsequent infections on preventive measures which involve the natural immunity (personal measure), surrounding (environmental measure) and vaccines (organisational measure).

9.1. Personal measures

Personal measures are aimed at depleting the risk of transmission of SARS-CoV-2 infection through human-to-human contact and interaction in order to persuade and avoid mounting overwhelming challenge on an individual's adaptive immunity. These personal measures range from hand hygiene, physical distancing to the proper handling, usage and disposal of personal protective equipment (PPE).

Despite the advocacy to adhere to these personal precautionary measures, the rebellious quest to return to "normal life" without adhering to these safety precautions is evident through global anti-face mask protests in Spain [89], Rome [90] and the UK [91]. These actions which have become a setback against tackling the pandemic have the potential to induce subsequent waves of re-infection which interferes with adaptive immunity and ensures that long-term immunological response by the body to the virus is compromised.

Adaptive immunity mainly involves the synthesis and defensive roles of cytotoxic T cells (killer or CD8+ T cells) and B cells (immunoglobulin-producing plasma cells) which are responsible in ameliorating the severity of COVID-19 re-infection [1].

Following the viral invasion of the biological system, IgM is released within a week to two weeks which is mobilised to tackle the infection before gradually waning in a matter of months. Within 2–3 weeks following recovery, IgG antibodies are released. Previous studies which conducted whole-genome sequencing in four health professionals in Mumbai, India and in a 33-year-old resident of Hong Kong to confirm re-infection reported the absence of detectable IgG antibodies which could be associated with their mild illness following their first infection. The levels of the immunoglobulins in COVID-19 asymptomatic individuals with their mild symptoms have been observed to be lower compared to their counterparts with severe cases [92]. By implication, the rapid waning levels of immunoglobulins expose these asymptomatic patients to be more prone to a second wave of infection which may present more severe hallmarks of the disease compared to those who are critically ill if the personal safety measures are not in place. A previous study revealed that anti-SARS-CoV mounted against the spike protein was correlated with a severe form of acute pulmonary injury [92].

At the time of re-infection, the 33-year-old resident of Hong Kong had no detectable IgG but developed detectable antibody following 5 days post hospitalisation [1]. The absence of detectable neutralising antibodies as observed both in individuals who recovered from COVID-19 infection [93] and asymptomatic patient [48] does not necessarily imply that the biological system of these patients did not mount immunological response but could actually be that the waning effect of the antibodies secreted in response to the virus was below the limit of detection of the assays used for the investigation of anti-SARS-CoV-2 in body fluids collected within 5-6 weeks after the onset of clinical manifestations. This further implies that the low antibody titers in these patients can predispose them to re-infection which is possible within 4 and a half months following the first episode of symptomatic viral infection. Hence it becomes clear that herd immunity by natural infection is not sufficient to shield the body against the virus [1].
Apart from the B cells, T cell immunity has been revealed to sustain the defensive momentum by inducing long-term immunity which mainly targets the structural proteins of the virus [94, 95, 96, 97]. While T helper cells were demonstrated to be instrumental in targeting both structural (membrane, spike and nucleoprotein) as well non-structural (nsp3, nsp4) proteins and ORF8, cytotoxic T cells were shown to target ORF8, ORF3a and nsp6. These cells have been detected in patients who recovered from COVID-19 infection following several months post initial infection [98]. The T helper cell response is mainly elicited in response to mutation at amino acid 222 of the spike protein [99].

In order to guarantee its survival against the immunological response of the biological system, generational strains of the virus have developed ways of evading the immune system by undergoing over 30 mutations within 8 genomes and resulting in cases of a second episode of infection which either present with mild or severe symptoms compared to the initial episode of infection. These cases of re-infection have been confirmed using whole-genome analysis to indicate that the viral strains which induce the first and second episodes of infections originate from dissimilar lineages with up to 24 nucleotide variants, hence both strains vary from each other. Unlike the first episode, the second episode of infection is characterised with raised levels of C-reactive protein (biomarker for acute infection), relatively elevated viral load and seroconversion of IgG against SARS-CoV-2.

In order to evade the attack of neutralising antibodies against its spike protein [100], viral mutations of its spike protein amino acids emerged through natural selection, features of recent epidemiology and random genetic drift. This led to different amino acid residues (L18F, Q780E, D614G and A222V) of the spike proteins between the viral strains responsible for the first and second episodes of infection. Previous studies have provided evidence that reveals that samples from 614G infections have raised levels of viral RNA and produced antibody titers in pseudo-viruses from in vitro experiments [101, 102]. Although these studies established the relationship between D614G and replication, it remains unclear whether these mutations are responsible for re-infection and if individuals who contract the infection for a second time with a viral strain that has amino acid variations in its spike protein are able to transmit the virus.

9.2. Environmental measures

These measures aimed at reducing the risk of viral transmission to the individual through contact with inanimate and contaminated objects [103]. The survival duration of SARS-CoV-2 on these objects is dependent on several factors which include the type/nature of the surface and specific viral strain in addition to relative humidity and temperature.

9.3. Organisational measures

These measures are aimed at minimising the possibility of exposure and transmission of SARS-CoV-2 through non-pharmaceutical and pharmaceutical approaches. Several concerns have been raised on the possibility of amino acid variants of the spike protein to aid the virus in evading vaccine-induced immunity. However, this may be unlikely as these mutations (e.g., D614G) are not localised within the receptor-binding domain (RBD) of the viral spike protein but situated between each of the spike protomer that stabilises the mature trimeric form located on the viral surface through the hydrogen bonding. As a result of this, the amino acid changes do not influence the immunogenicity of the RBD receptors, which are thought to be vital for immunoglobulin neutralisation. However, it remains unclear the specific impact of these mutations on the spike protein’s role in viral entry and fusion with the host’s angiotensin-converting enzyme receptor as well as the influence of the mutation on the therapeutic entry inhibitors.

Because more clarity is required on the role of these mutations during natural infection by SAR-CoV-2, it is recommended to consider the existence of these mutations during vaccine design and administration [104]. In order to prevent subsequent cases of re-infection, it is imperative that vaccination studies should not just be restricted to those who never had the infection but also for those who have recovered from the infection. Furthermore, booster doses of these vaccines should be considered since these therapeutic agents may not be able to provide lifelong defence against SARS-CoV-2 through a sustained immunological response to the virus [1].

10. Conclusion

Patients with SARS-CoV-2 re-infections are highly unlikely to be due to natural viral evolution and the implications are that SARS-CoV-2 can adapt with enough genetic agility to avoid an innate immune response in a manner to re-establish detectable levels of infection in an individual. Hence, this underscores the need for further investigations to provide more robust genomics and immunology data and their correlates with transmissibility and viral shedding. These could help to predict the likelihood of SARS-CoV-2 re-infection and the potential of re-infected symptomatic and asymptomatic persons to transmit the SARS-CoV-2 at a second instance. There is a need to have a globally accepted and adhered standardize and robust laboratory criteria and case definitions for SARS-CoV-2 re-infection to avoid mis-diagnosis, especially in developing countries.

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