The genomic and clinical features of the COVID-19 Omicron variant: a narrative review [version 2; peer review: 1 not approved]

Decsa Medika Hertanto, Henry Sutanto, Maria Inge Lusida, Kuntaman Kuntaman, Djoko Santoso

1 Department of Internal Medicine, Faculty of Medicine, Airlangga University, Dr. Soetomo Teaching Hospital, Surabaya, 60286, Indonesia
2 Department of Physiology and Pharmacology, SUNY Downstate Medical Center, Brooklyn, New York, 11203, USA
3 Department of Clinical Microbiology, Faculty of Medicine, Airlangga University, Surabaya, 60132, Indonesia
4 Institute of Tropical Disease, Airlangga University, Surabaya, 60115, Indonesia

Abstract
Coronavirus disease 2019 (COVID-19) is a major cause of morbidity and mortality worldwide. Since late November 2021, the Omicron variant has emerged as the primary cause of COVID-19 and caused a huge increase in the reported incidence around the world. To date, 32-34 spike mutations have been reported to be present in the Omicron variant, 15 of which were located in the receptor-binding domain that interacts with the cell surface of the host cells, while the rest were located in the N-terminal domain and around the furin cleavage site. Recent studies have suggested that those mutations could have a major role in the transmissibility and pathogenicity of the Omicron variant. Additionally, some mutations might contribute to the change of viral tropism of this novel variant. Here, we aim to discuss the recent reports on the transmissibility and severity of the Omicron variant from both the genetic and clinical perspectives. Afterward, we also take the chance to deliver our personal view on the topic.

Keywords
COVID-19, pandemic, SARS-CoV-2, Omicron variant, emerging disease, global health, virus, genome, mutation

This article is included in the Emerging Diseases and Outbreaks gateway.
**Amendments from Version 1**

This revision was made in response to the outstanding and critical comments from Dr. Wang and Dr. Revindran-Stam. Among other things, we have added the phylogenetics of the Omicron variants, discussion about the sublineage of the Omicron variant (BA.1, BA.2, BA.3, BA.4, BA.5 and XE), added the mutations outside the S-proteins (e.g., ORF and NSP) and provided figures depicting the mutation sites, clarify the usefulness of Ct in the Omicron infection, as well as added a paragraph discussing the clinical presentations of the Omicron variant as compared to other COVID-19 variants. We believe that this revised manuscript has answered most (if not all) of the concerns raised by the esteemed Referees.

Any further responses from the reviewers can be found at the end of the article.

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**Introduction**

Coronavirus disease 2019 (COVID-19) has been a major cause of morbidity and mortality worldwide since December 2019. Up to July 2022, severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) has infected more than 570 million people and contributed to the death of more than 6.4 million individuals around the world. Intriguingly, after more than 2 years of its existence, the infection rate remains high, and the pandemic has not been resolved. Following the devastating impacts of the B.1.617.2 (Delta) variant, predominantly in health and socioeconomic sectors, there was a high expectation that the viral disease could be tamed by the rapid, collaborative and evolutionary development of COVID-19 vaccines. Indeed, the massive vaccination program in several countries has successfully reduced the fatality of the disease and together with the implementation of strict public health and social control measures (PHSCM), the infection rate could also be lowered.1-5 For example, in the United States (US), COVID-19 vaccines reduced the overall attack rate (i.e., number of new cases during specified time interval divided by the total population at start of time interval) in vaccinated individuals by 4.4% on day 300 from the start of vaccination (9% in the unvaccinated group vs. 4.6% in the vaccinated group), as well as the rate of hospitalization, intensive care unit (ICU) occupancy, incidence of major adverse events and mortality.1 Despite the reported effectivity decline of BNT162b2 and ChAdOx1 vaccines against the Delta variant (10–13% and 16% lower than B.1.1.7 (Alpha) variant, respectively),6 vaccination was proven to remain effective in reducing infection and accelerating viral clearance,7 as well as lowering mortality caused by the Delta variant.8

However, a glimpse of hope to end the pandemic was once again challenged by the presence of the new COVID-19 B.1.1.529 (a.k.a, Omicron) variant. Since its first reported appearance in South Africa in November 2021, it has rapidly taken the attention of experts around the world. A few days after its appearance, the World Health Organization (WHO) immediately classified Omicron as a Variant of Concern (VoC). Since then, it has vastly spread from Africa to Europe, then Asia, Australia and America (Figure 1). Since early January 2022, the Omicron variant has become the major COVID-19 variant in most countries (Table 1)6,7 and contributed to the rise of COVID-19 incidence from ~600,000 new cases in late November 2021 to ~3.5 million cases in late January 2022.

**The genomic basis of the Omicron variant and the predicted effects of its mutations**

**Phylogenetics of the Omicron variant**

Several studies have attempted to investigate whether there are evolutionary connections between the Omicron variant and its predecessors.8,9 For instance, using the ultrametric and metric clustering method, Kandeel et al.8 found that the Omicron variant was not closely connected with the previous SARS-CoV-2 variants, but rather formed a new monophyletic clade on its own. Similarly, another phylogenetics study by Sun et al.9 discovered that the Omicron variant was not evolutionarily derived from the Delta variant and did not share a common mutation profile with it. Of note, the Omicron variant first appeared when the Delta variant dominated the global COVID-19 infection back in late 2021, so it was hypothesized that the Delta variant evolved into the Omicron variant. Instead, they found some similarities between the Omicron and the Gamma variants,9 as also reported by Kannan et al.10 Using a different method for phylogenetics analysis (i.e., the Neighbor-joining method), Kandeel et al. revealed a connection between the Omicron variant and the Alpha variant. Furthermore, among the circulating SARS-CoV-2 variants, they found that the Alpha variant had the least nucleotide changes as compared to the Omicron variant, highlighting the close connection between the two.8

**The consequences of reported mutations on transmissibility and pathogenicity of the Omicron variant**

A study conducted based upon the genome sequencing data of 108 samples collected from patients infected with the Omicron variant revealed that this variant possessed 61-64 mutations, 54 of which were single nucleotide polymorphisms (SNPs). Of those, 34 mutations were positioned at the spike (S) proteins (Figure 2) and 32 of those spike mutations were non-synonymous, which means that the mutations alter the amino acid sequences.11 Importantly, the mutations in the Omicron variant were reported to be present in all three key regions in the spike of SARS-CoV-2: The receptor binding domain (RBD), N-terminal domain (NTD) and furin cleavage site (FCS). Overall, 15 of the spike mutations were located in the RBD, a region that is responsible for the viral attachment to the cell surface (Table 2).
Mutations in RBD

As displayed in Table 2, the N501Y mutation that converts amino acid asparagine to tyrosine at position 501 was detected in the Omicron variant. Of note, the N501Y mutation was also identified in other COVID-19 variants (e.g., C.1.2, Alpha, Beta and Gamma). This particular mutation enhanced the binding affinity of the RBD to angiotensin converting enzyme type-2 (ACE2) receptor in the surface of the host cell. As a consequence, the Omicron variant could have a stronger attachment to the host cells than some of the other COVID-19 variants, fostering its transmissibility. Meanwhile, the mutation Q498R (converting glutamine to arginine at position 498) of the RBD alone negatively affected the protein stability and binding. However, due to its known epistatic effect with N501Y, the combination of both mutations was shown to increase the affinity of the RBD to ACE2 receptor by 4-fold. In contrast with the N501Y mutation, the Q498R mutation has not been seen in other COVID-19 variants. In addition to their effect on the binding affinity to ACE2 receptor, Omicron-associated mutations in the RBD could also promote the escape from existing neutralizing antibodies. In the study done by Cao et al., the effect of Omicron-associated mutations in the RBD region on neutralizing antibodies was assessed using a high throughput yeast display screening. As the result, K417N, G446S, E484A and Q493R mutations assisted the virus to evade neutralizing antibodies, especially the ones which epitope overlapped with the ACE2 binding motif (epitope group A-D). Also, there was evidence that G339D, S371L and N440K mutations of the RBD could facilitate the virus evasion from other neutralizing antibodies (epitope group E-F). Overall, of the 247 antibodies tested, 85% were evaded by Omicron, suggesting the potentially low efficacy of preexisting neutralizing antibodies against the Omicron variant.

Mutations in NTD

Several mutations in the NTD region were shown to have significant effect on viral infectivity and the modulation of immune evasion. For example, the del69-70 boosted the infectivity of the Alpha variant through the elevation of cleaved
Table 1. Confirmed coronavirus disease 2019 (COVID-19) cases from December 1st, 2021 to February 23rd, 2022 and shares of the Omicron variant.

<table>
<thead>
<tr>
<th>20 countries with highest confirmed cases during observed period</th>
<th>Total cases within observed period (in million)</th>
<th>Relative change compared with data on December 1st, 2021</th>
<th>Shares of Omicron variant per November 29th, 2022</th>
<th>Shares of Omicron variant per February 21st, 2022</th>
</tr>
</thead>
<tbody>
<tr>
<td>World</td>
<td>165.85</td>
<td>+63%</td>
<td>0.06%</td>
<td>99.77%</td>
</tr>
<tr>
<td>United States</td>
<td>30.01</td>
<td>+62%</td>
<td>0.16%</td>
<td>99.14%</td>
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<tr>
<td>France</td>
<td>14.75</td>
<td>+190%</td>
<td>0.17%</td>
<td>99.75%</td>
</tr>
<tr>
<td>United Kingdom</td>
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<td>+82%</td>
<td>0.35%</td>
<td>95.88%</td>
</tr>
<tr>
<td>India</td>
<td>8.27</td>
<td>+24%</td>
<td>0.21%</td>
<td>99.51%</td>
</tr>
<tr>
<td>Germany</td>
<td>8.25</td>
<td>+138%</td>
<td>0.11%</td>
<td>89.91%</td>
</tr>
<tr>
<td>Italy</td>
<td>7.56</td>
<td>+150%</td>
<td>0.11%</td>
<td>89.91%</td>
</tr>
<tr>
<td>Brazil</td>
<td>6.38</td>
<td>+29%</td>
<td>0.14%</td>
<td>99.75%</td>
</tr>
<tr>
<td>Russia</td>
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<td>+64%</td>
<td>0.22%</td>
<td>99.03%</td>
</tr>
<tr>
<td>Spain</td>
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<td>0.17%</td>
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</tr>
<tr>
<td>Turkey</td>
<td>4.94</td>
<td>+56%</td>
<td>0.00%</td>
<td>97.90%</td>
</tr>
<tr>
<td>Argentina</td>
<td>3.54</td>
<td>+66%</td>
<td>0.00%</td>
<td>(Feb 7, 2022) 100.00%</td>
</tr>
<tr>
<td>Netherlands</td>
<td>3.50</td>
<td>+131%</td>
<td>1.72%</td>
<td>100.00%</td>
</tr>
<tr>
<td>Japan</td>
<td>2.97</td>
<td>+172%</td>
<td>1.61%</td>
<td>(Feb 7, 2022) 99.44%</td>
</tr>
<tr>
<td>Australia</td>
<td>2.91</td>
<td>+1,364%</td>
<td>0.44%</td>
<td>99.73%</td>
</tr>
<tr>
<td>Israel</td>
<td>2.25</td>
<td>+167%</td>
<td>0.30%</td>
<td>99.78%</td>
</tr>
<tr>
<td>Denmark</td>
<td>2.17</td>
<td>+441%</td>
<td>0.04%</td>
<td>99.92%</td>
</tr>
<tr>
<td>Portugal</td>
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<td>+179%</td>
<td>1.55%</td>
<td>99.72%</td>
</tr>
<tr>
<td>South Korea</td>
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<td>+446%</td>
<td>0.15%</td>
<td>(Feb 7, 2022) 89.86%</td>
</tr>
<tr>
<td>Poland</td>
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<td>0.00%</td>
<td>98.88%</td>
</tr>
<tr>
<td>Belgium</td>
<td>1.74</td>
<td>+98%</td>
<td>0.20%</td>
<td>99.74%</td>
</tr>
</tbody>
</table>

The data shown here was retrieved from Our World in Data and the raw data in the database was sourced from the COVID-19 Data Repository by the Center for Systems Science and Engineering (CSSE) at Johns Hopkins University and GISAID.

Figure 2. The mutation sites of the Omicron BA.1 variant. The red box depicts the mutations located in the S-proteins of the virus. In total, there are 34 mutations in the spike protein and the rest are located outside of this region (e.g., in the non-structural protein (NSP) or the open reading frame (ORF) regions). This figure was obtained from the Stanford University Coronavirus Antiviral and Resistance Database under license CC BY-SA 4.0.
spike incorporation into virions. Conversely, T95I mutation that was previously identified in the B.1.617.1 variant seems not to be substantially involved in immune evasion due to its location outside of the antigenic supersite. Interestingly, two cases of BNT162b2 and mRNA-1273 vaccines breakthrough infection were reported in COVID-19 patients harboring T95I mutation of the NTD, as well as the del142-144 mutation of the NTD and D614G mutation of the FCS, indicating the possible contribution of those mutations on viral immune escape. Meanwhile, evidence of a marked (4-16-fold) reduction of neutralizing capacity of COVID-19 convalescent sera against a recombinant vesicular stomatitis virus carrying SARS-CoV-2 spike protein with Y145D mutation was also reported.

Mutations in FCS

The FCS region has been shown to be a key part of SARS-CoV-2 pathogenesis and severity. For instance, a mutant lacking FCS (ΔPRRA) displayed a reduced replication rate in a human respiratory cell line. Meanwhile, in regard to Omicron-associated mutations, the H655Y and N679K mutations which are located proximal to the FCS, and the P681H mutation of the FCS could enhance the SARS-CoV-2 spike cleavage and increase the transmissibility of the Omicron variant. Moreover, the P681H mutation, which was previously identified in the B.1.1.7 (Alpha) variant, was demonstrated to facilitate the viral resistance against innate immunity (i.e., interferon-β) in lung epithelial cells. Yet, another study on this particular mutation did not find any notable change on the spike cleavage, viral entry or intercellular spreading. Next, the D614G mutation, a common mutation found in existing COVID-19 variants, increased viral replication in human lung epithelia and respiratory tract by fostering the virion stability and infectivity. Moreover, it increased the viral load in the upper respiratory tract, thereby promoting viral transmission. However, it is important to note that the effect of such mutations on Omicron pathogenicity may differ from what was reported in previous COVID-19 variants due to distinct interactions among variant-associated mutations.

Mutations outside of the S-region

In addition to the abovementioned mutations in the spike (S) region, several mutations were detected in the open reading frame (ORF) and non-structural protein (NSP) regions of the virus (e.g., NSP3, NSP4, NSP5, NSP6, NSP12 and NSP14). Three amino-acid deletions at L3674-, S3675- and G3676- were found in ORF1a region and could play a role in altering the cells’ ability to degrade viral components. Additionally, K856R, S2083-, L2084I, A2710T, T3255I, P3395H, L3674-, S3675-, G3676-, and I3758V were also documented in ORF1b region. Meanwhile, several mutations were detected in ORF1b region, such as P314L and I1566V; while in ORF9b (i.e., a region that is deemed to be involved in the suppression of the innate immune response to viral infection), three amino-acid deletions at E27-, N28- and A29- were

<table>
<thead>
<tr>
<th>NTD</th>
<th>RBD</th>
<th>FCS &amp; adjacent</th>
</tr>
</thead>
<tbody>
<tr>
<td>A67V</td>
<td>G339D</td>
<td>T547K</td>
</tr>
<tr>
<td>I5H681</td>
<td>S371L</td>
<td>D614G</td>
</tr>
<tr>
<td>del69-70</td>
<td>S373P</td>
<td>H655Y</td>
</tr>
<tr>
<td>T95I</td>
<td>S375F</td>
<td>N679K</td>
</tr>
<tr>
<td>GVYY142D</td>
<td>K417N</td>
<td>P681H</td>
</tr>
<tr>
<td>del142-144</td>
<td>N440K</td>
<td>N764K</td>
</tr>
<tr>
<td>Y145D</td>
<td>G446S</td>
<td>D796Y</td>
</tr>
<tr>
<td>NL211</td>
<td>S477N</td>
<td>N856K</td>
</tr>
<tr>
<td>del21I</td>
<td>T478K</td>
<td>Q954H</td>
</tr>
<tr>
<td>L212</td>
<td>E484A</td>
<td>N969K</td>
</tr>
<tr>
<td>Ins214EPE</td>
<td>Q493R</td>
<td>L981F</td>
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<td></td>
<td>G496S</td>
<td></td>
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<tr>
<td></td>
<td>Q498R</td>
<td></td>
</tr>
<tr>
<td></td>
<td>N501Y</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Y505H</td>
<td></td>
</tr>
</tbody>
</table>

*was reported by Ma et al. and was reported by the United States Centers for Disease Control and Prevention (CDC). (FCS = furin cleavage site; NTD = N-terminal domain; RBD = receptor binding domain).
present together with P10S.\textsuperscript{30,24} Covariants.org also described two other mutations in nucleocapsid: R203K and G204R, which were linked with the capability to increase viral loads.

The existing Omicron subvariants

**BA.1**

B.1.1.529.1 is the original Omicron variant first identified in Botswana, before rapidly spreading to South Africa and all over the world. It was first reported on November 24\textsuperscript{20}, 2021 by the Network for Genomics Surveillance in South Africa. As described in the previous section, this subvariant has a higher transmission rate, an increased immune evasiveness and presumably lower severity as compared to the other circulating COVID-19 variants.

**BA.2**

In a recent statement, WHO’s Technical Advisory Group on SARS-CoV-2 Virus Evolution (TAG-VE) has raised a concern regarding Omicron sublineage BA.2. For the past few months, globally, the proportion of BA.2 sublineage has been increasing relative to BA.1. BA.2 (B.1.1.529.2) was first identified among genome sequences submitted to GISAID database from the Philippines back in November 2021.\textsuperscript{25} Subsequently, the BA.2 sublineage was identified in viral genome sequences from more than 40 countries in January 2022, with Denmark, India, Sweden and Singapore having recorded the most cases. Early research showed that this sublineage was significantly more transmissible than the original B.1.1.529 variant and could cause more severe disease than BA.1. Moreover, it is also more resistant to neutralizing antibodies than BA.1. A study evaluating the response of neutralizing antibodies against BA.1 and BA.2 in 24 fully vaccinated individuals and 8 people with previous SARS-CoV-2 infection demonstrated that the median BA.2 neutralizing antibody titer was 1.3-1.4 times lower than the median BA.1 neutralizing antibody titer.\textsuperscript{26} Another study also discovered that BA.2 had a 1.4 times higher effective reproduction number and a higher replication efficacy in human nasal epithelial cells than the BA.1. Additionally, it was resistant to the BA.1-induced humoral immunity. Furthermore, the spike of BA.2 was more fusogenic than the one in BA.1, suggesting its higher pathogenicity than the original Omicron variant.\textsuperscript{27} Importantly, individuals with previous BA.1 infection could still contract the BA.2 sublineage and BA.2 was shown to be capable of inducing vaccine breakthrough.\textsuperscript{28}

BA.2 (Figure 3) shared some genomic similarities with BA.1, with 32 similar mutations presented in both subvariants. Nonetheless, 28 distinct mutations were reported in BA.2, including 4 specific mutations in the RBD (S371F, T376A, D405N and R408S) and in the NTD (T19I, del24-26, A27S and V213G) regions.\textsuperscript{26} In addition, the absence of del69-70 mutation of NTD in BA.2 could significantly lower the identifiability of this subvariant in some PCR assays.

**BA.3**

BA.3 was first documented in northwestern South Africa and constituted the combination of mutations in BA.1 and BA.2. BA.3 shared most of its mutations with BA.1 and BA.2, except for one at NSP6 (A88V). It also has 15 RBD mutations together with P10S.\textsuperscript{30,24} Covariants.org also described two other mutations in nucleocapsid: R203K and G204R, which were linked with the capability to increase viral loads.

**Figure 3. The mutation sites of the Omicron BA.2 variant.** The red box depicts the mutations located in the S-proteins of the virus. This figure was obtained from the Stanford University Coronavirus Antiviral and Resistance Database under license CC BY-SA 4.0.
mutations, but none was distinct from that of BA.1 and BA.2.\textsuperscript{28} In the population, BA.3 caused the lowest number of cases in these three lineages. It may have been due to the loss of six mutations (ins214EPE, S371L, G496S, T547K, N856K, and L981F) from BA.1 or obtaining two mutations from BA.2 (S371F and D405N). As also found in BA.2, BA.3 retained the N501Y and Q498R mutations, which enhance the binding to ACE2 receptor, and the H655Y, N679K, and P681H mutations, which increase spike cleavage and facilitate virus transmission.\textsuperscript{29}

**XE**

This subvariant is a hybrid or recombinant of BA.1 and BA.2, which was first discovered in the United Kingdom in January 2022. The XE subvariant has similar spike and structural proteins as BA.2, and obtains the 5' part of its genome from BA.1. This subvariant was found to be more transmissible than BA.1 and BA.2, making it one of the most transmissible COVID-19 variants to date. Moreover, XE subvariant had a 9.8% higher growth rate than BA.2, suggesting the potentially more infectious nature of this Omicron subvariant.\textsuperscript{30} Interestingly, despite sharing some similar mutations at NSPs with BA.1 and at other sites with BA.2, XE subvariant carries 3 specific mutations that was not seen in its predecessors: NSP3 C3241T, NSP3 V1069I and NSP12 C14599T.\textsuperscript{30}

**BA.4 and BA.5**

These two Omicron subvariants (BA.4 and BA.5) were first identified in South Africa in January and February 2022, respectively. Since then, they have spread to many countries and (re)increased the number of COVID-19 cases accordingly, creating a new COVID-19 wave in some countries. At present, there is limited information regarding the behavior of these subvariants, whether they have an enhanced transmissibility, immune evasiveness, or severity as compared to other SARS-CoV-2 variants. However, the rapid global dissemination of these subvariants may suggest that it could have a higher transmission rate than most, if not all, of the preexisting variants. Nevertheless, early results of recent studies have shed some light on these open questions. Yamasoba et al.\textsuperscript{31} explored the effect of several monoclonal antibodies on BA.4 and BA.5, and showed that BA.4 and BA.5 had a higher resistance to monoclonal antibody cilgavimab than BA.2. Another study by Cao et al.\textsuperscript{32} showed that BA.4 and BA.5 subvariants exerted stronger neutralization evasion than BA.2 against the plasma from 3-dose vaccination and from humoral immunity derived from post-vaccination BA.1 infections. Interestingly, they showed a contrasting result that cilgavimab could effectively neutralize BA.4 and BA.5. Next, using a deep-learning method, Chen et al.\textsuperscript{33} studied the variants infectivity and revealed that BA.4 and BA.5 were 36% more infectious than BA.2.

BA.4 and BA.5 sublineages contain some specific mutations, namely L452R and F486V (Figure 4), which are located in the RBD and not present in BA.1 (the original Omicron variant).\textsuperscript{34} As compared to BA.2, both the S-proteins of BA.4 and BA.5 have similarities with BA.2, except for the presence of del69-70, L452R and F486V. The F486V mutation in their S-proteins is known to be responsible for the viral infection.\textsuperscript{34} It was also reported that D405N (which is also carried by BA.2 subvariant) and BA.4/BA.5-specific L452R and F486V facilitated the neutralizing antibody evasion.\textsuperscript{32}

**The viral tropism of the Omicron variant**

The change of viral tropism in the Omicron variant is also important to scrutinize. A recent UK-based study showed that the Omicron variant was strongly associated with symptoms from upper respiratory tract more than the lower
respiratory tract.\textsuperscript{35} This could be explained by the presence of two Omicron-specific mutations: N764K and N856K. Those mutations were shown to produce cleavage sites for subtilisin-kexin isozyme-1/site-1 protease (SKI-1/S1P) serine protease predominantly situated in the upper airway. Such cleavage sites are important to cleave viral envelope glycoproteins, which modulate SARS-CoV-2 replication and pathogenesis.\textsuperscript{36} Indeed, the Omicron variant was shown to replicate faster in the bronchus than in the lung parenchyma.\textsuperscript{37} Another study also highlighted that despite the similar viral replication in human nasal epithelial cultures, the Omicron variant demonstrated lower replication in lower respiratory and pulmonary cells than the Delta variant.\textsuperscript{38} Additionally, Omicron spike protein has a lower S1/S2 cleavage efficiency than the Delta variant and Omicron tends to avoid cells expressing a high level of transmembrane protease, serine 2 (TMPRSS2). This was due to Omicron’s failure in exploiting the TMPRSS2 that promotes cell entry via plasma membrane fusion. As a consequence, the cell entry of the Omicron variant was largely mediated through the endocytic pathway.\textsuperscript{39,40} The fact that SKI-1/S1P was also present in pulmonary macrophages\textsuperscript{41} could suggest the potential consequences of N764K and N856K mutations on the host innate immunity. Nevertheless, more studies are needed to confirm this notion.

What can we tell from the Ct value of patients with the Omicron variant?

Real-time reverse transcription polymerase chain reaction (rtRT-PCR) or quantitative RT-PCR (qRT-PCR) is a gold-standard diagnostic tool for identifying SARS-CoV-2, the causative agent of COVID-19. This method usually detects \( \geq 2 \) genes of SARS-CoV-2, including ORF1ab/RdRp, N and E.\textsuperscript{40} At present, due to the increasing incidence of Omicron variant infection, S gene with S gene target failure (SGTF) is frequently used as an indicator for screening of the Omicron variant. Most reported Omicron variant sequences include a deletion in the S gene, which can cause an SGTF in some PCR assays.

Cycle threshold (Ct) is the thermal cycle number at which the amplified DNA that shows as a fluorescent signal exceeds and thus passes the threshold for positivity. A higher Ct value means that the tool requires more copy numbers to reach the positivity threshold, indicating a lower viral concentration in the specimens. Conversely, the lower the Ct level the greater the amount of identified target ribonucleic acid (RNA) in the sample. A recent study assessing the real-time (RT) qPCR data from 10,324 specimens and comparing 97 Omicron against 107 Delta variant infections reported that the Ct values were higher for Omicron infections than for Delta infections (Omicron: Ct 23.3 vs. Delta: Ct 20.5), suggesting a lower peak viral RNA of the Omicron variant. Moreover, the clearance phase for Omicron was shorter while the clearance rate was similar with the Delta variant.\textsuperscript{41} Similarly, studies conducted in France also reported a significantly higher Ct value for Omicron infection than the non-Omicron variants.\textsuperscript{42,43} These findings suggest that the high transmissibility of the Omicron variant might not be due to a high viral load in the upper respiratory tract, despite the presence of D614G mutation which was known to increase viral replication in the upper respiratory tissue.\textsuperscript{23} Indeed, a study comparing 2,001 Alpha (median Ct = 22.0), 792 Delta (median Ct = 19.7), and 1,935 Omicron variant (median Ct = 20.8) samples reported that the Omicron infection did not have higher viral loads than those with Delta when stratified by the major PCR platforms used and by symptomatic vs. asymptomatic status. Additionally, consistent with prior studies, the study displayed higher viral loads in symptomatic patients, as compared to the asymptomatic ones. Overall, the study suggested that the rapid dissemination of the Omicron variant was not attributed to higher nasal viral loads as compared to prior variants.\textsuperscript{44} Thus, at present, the exact cause of a higher transmissibility in Omicron variant infection than other variants, including Delta, remains to be elucidated.

Clinical features of the Omicron variant

A recent observational study conducted in Texas, US, from late December 2021 to early January 2022 reported that the Omicron variant displayed some different clinical patterns than its predecessors.\textsuperscript{45} Compared to patients infected by Alpha and Delta variants, these patients were younger and predominantly female. The number of patients requiring hospitalization was significantly lower in Omicron than in Alpha and Delta variant infections (19.8% vs. 54.6% and 43.1%, respectively). Of those who were hospitalized, the average length of stay was significantly shorter in Omicron than Alpha and Delta (3.2 days vs. 5.1 days and 5.4 days, respectively). Moderate to severe cases (e.g., number of patients requiring extracorporeal membrane oxygenation [ECMO], mechanical ventilation and high-flow oxygenation) and mortality were also lower in Omicron infection. As expected, while Alpha and Delta variants affected mostly the unvaccinated individuals, the Omicron variant proportionally infected both the unvaccinated (44.1%) and vaccinated people (55.9%).\textsuperscript{45} These findings and comparable results from other studies\textsuperscript{46,47} confirm the hypothesis that the Omicron variant causes less severe disease, resulted in a lower hospitalization rate. Importantly, the fact that Omicron caused an increased mRNA vaccine (i.e., BNT162b2 and mRNA-1273) breakthrough incidence\textsuperscript{45} needs to be swiftly responded to by speeding up the vaccination booster campaign. The third vaccine (booster) dose has been shown to rescue and broaden the viral neutralization.\textsuperscript{48–51} A study comparing the mRNA vaccines effectiveness on 16,087 Omicron versus 4,261 Delta cases revealed that the second dose vaccine effectiveness against the Delta variant reduced from 89% to 80% after 240 days. Interestingly, against the Omicron variant, the second dose vaccine effectiveness was only 36% at day 7-59 and
completely diminished after 180 days. However, the third dose (booster) vaccination recovered the vaccine effectiveness against the Delta and Omicron variants to 97% and 61% after day 7, respectively. Therefore, booster vaccination is expected to be beneficial to tackle this emerging COVID-19 variant.

Table 3 lists the reported clinical symptoms associated with Omicron variant infection. In most studies, upper respiratory symptoms (e.g., runny or stuffy nose, sneezing, cough and sore throat) dominated, while the lower respiratory complaint (e.g., shortness of breath) was identified in less than 20% of patients infected by the Omicron variant. Non-respiratory symptoms, such as fatigue, headache, muscle pain and fever, were also seen in some patients, although the number varies between studies. Importantly, COVID-19-pathognomonic symptoms, such as anosmia and ageusia, were only observed in a limited number of patients across studies (less than 25%). Overall, this observation is consistent with the abovementioned viral tropism of the Omicron variant.

Overall, the symptoms of the Omicron variant infection were different than those caused by other COVID-19 VoCs infection. For example, the ZOE COVID study in the United Kingdom reported that in the Delta variant infection, the loss of smell or anosmia was more prevalent (52.7% vs. 16.7%) and sore throat was less common (60.8% vs. 70.5%) than in the Omicron infection. Additionally, the duration of sickness was shorter in the Omicron infection, as well as the rate of hospitalization. Whilst, the Alpha variant that was phylogenetically connected with the Omicron variant shared many similar symptoms, including fatigue, headache, runny nose, sneezing, cough and sore throat. However, anosmia and dysosmia that were common in the Alpha variant were not frequent in the Omicron variant infection. Another study investigating the clinical characteristics of the Gamma variant infection in 313 healthcare workers in Brazil showed that common cold, headache, cough and sore throat were the most common reported symptoms of the Gamma infection, while anosmia and ageusia were only reported in less than 25% of the sampled individuals. This clinical presentation is very similar with the clinical characteristics presented by the Omicron variant, possibly due to their close phylogenetics connections.

---

### Table 3. Reported clinical symptoms of the Omicron variant.

<table>
<thead>
<tr>
<th>Symptoms</th>
<th>Brandal et al.</th>
<th>Young et al.</th>
<th>CDC Team</th>
<th>Maisa et al.</th>
<th>Menni et al.</th>
<th>Hajjo et al.</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Norway N = 81</td>
<td>Singapore N = 87</td>
<td>USA N = 43</td>
<td>France N = 277</td>
<td>UK N = 4,990</td>
<td>Jordan N = 500</td>
<td>N = 5,978</td>
</tr>
<tr>
<td>Sneezing</td>
<td>35 (43%)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>3,143 (63%)</td>
<td>-</td>
<td>3,178 (63%)</td>
</tr>
<tr>
<td>Runny or stuffy nose</td>
<td>63 (78%)</td>
<td>30 (35%)</td>
<td>22 (59%)</td>
<td>74 (27%)</td>
<td>3,818 (77%)</td>
<td>33%</td>
<td>~4,172 (70%)</td>
</tr>
<tr>
<td>Loss of smell/anosmia</td>
<td>10 (12%)</td>
<td>3 (3%)</td>
<td>3 (8%)</td>
<td>23 (8%)</td>
<td>~17%</td>
<td>1%</td>
<td>~892 (15%)</td>
</tr>
<tr>
<td>Loss of taste/ageusia</td>
<td>19 (23%)</td>
<td>-</td>
<td>-</td>
<td>25 (9%)</td>
<td>-</td>
<td>-</td>
<td>44 (12%)</td>
</tr>
<tr>
<td>Cough</td>
<td>67 (83%)</td>
<td>39 (45%)</td>
<td>33 (89%)</td>
<td>143 (52%)</td>
<td>2,486 (50%)</td>
<td>47%</td>
<td>~3,003 (50%)</td>
</tr>
<tr>
<td>Hoarseness</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>2,145 (43%)</td>
<td>9%</td>
<td>~2,190 (40%)</td>
</tr>
<tr>
<td>Sore throat</td>
<td>58 (72%)</td>
<td>40 (46%)</td>
<td>-</td>
<td>88 (32%)</td>
<td>3,517 (71%)</td>
<td>45%</td>
<td>~3,928 (66%)</td>
</tr>
<tr>
<td>Shortness of breath</td>
<td>10 (12%)</td>
<td>-</td>
<td>6 (16%)</td>
<td>31 (11%)</td>
<td>~5%</td>
<td>-</td>
<td>~297 (6%)</td>
</tr>
<tr>
<td>Reduced appetite</td>
<td>27 (33%)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>~25%</td>
<td>-</td>
<td>~1,275 (25%)</td>
</tr>
<tr>
<td>Nausea or vomiting</td>
<td>-</td>
<td>-</td>
<td>8 (22%)</td>
<td>20 (7%)</td>
<td>~18%</td>
<td>-</td>
<td>~927 (17%)</td>
</tr>
<tr>
<td>Abdominal pain</td>
<td>5 (6%)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>~18%</td>
<td>-</td>
<td>~904 (18%)</td>
</tr>
<tr>
<td>Diarrhea</td>
<td>-</td>
<td>5 (6%)</td>
<td>4 (11%)</td>
<td>17 (6%)</td>
<td>~19%</td>
<td>-</td>
<td>~975 (18%)</td>
</tr>
<tr>
<td>Fatigue/lethargy</td>
<td>60 (74%)</td>
<td>-</td>
<td>24 (65%)</td>
<td>158 (57%)</td>
<td>-</td>
<td>32%</td>
<td>~402 (45%)</td>
</tr>
<tr>
<td>Headache</td>
<td>55 (68%)</td>
<td>-</td>
<td>-</td>
<td>121 (44%)</td>
<td>3,729 (75%)</td>
<td>13%</td>
<td>~3,970 (68%)</td>
</tr>
<tr>
<td>Muscle pain</td>
<td>47 (58%)</td>
<td>-</td>
<td>-</td>
<td>107 (39%)</td>
<td>~30%</td>
<td>29%</td>
<td>~1,796 (31%)</td>
</tr>
<tr>
<td>Fever</td>
<td>44 (54%)</td>
<td>24 (28%)</td>
<td>14 (38%)</td>
<td>169 (61%)</td>
<td>~35%</td>
<td>48%</td>
<td>~2,238 (37%)</td>
</tr>
<tr>
<td>Asymptomatic</td>
<td>-</td>
<td>20 (23%)</td>
<td>3 (7%)</td>
<td>-</td>
<td>31%</td>
<td>-</td>
<td>~178 (28%)</td>
</tr>
</tbody>
</table>

*Multiple symptoms could be reported by one person (UK = United Kingdom; USA = United States of America).
It is also crucial to acknowledge that although studies have shown compelling evidence of a less severe disease and a lower hospitalization due to Omicron infection, the high transmissibility of this variant might still impact the healthcare system readiness and increase the absolute number of hospitalization. These conditions would cause a severe delay on the non-COVID-19 patient care and might result in bigger indirect consequences than we expected. Also, the effect of the Omicron variant in immunocompromised individuals, and patients with concomitant diseases and comorbidities (e.g., diabetes mellitus) is still unknown. In general, these individuals have an impaired immune response against infections, a condition that favors Omicron and other COVID-19 variants. Nonetheless, the effect of Omicron on this special population is yet to be explored.

Regardless, implementing an effective PHSCM could still be beneficial to limit person-to-person transmission of Omicron variant. The application of well-fitting mask, imposing strict physical distancing, cough etiquette and hand hygiene, as well as avoiding crowds remain vital. The WHO has already advised to enhance surveillance with rapid testing, cluster investigations, contact tracing and quarantine, as well as case isolation to cut the chain of transmission. Only by a collaborative effort, the COVID-19 pandemic could be brought to an end.

Data availability
No data are associated with this article.

Acknowledgements
An earlier version of this article can be found on https://www.preprints.org/ (doi: 10.20944/preprints202202.0224.v1).

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Leyi Wang
Veterinary Diagnostic Laboratory and Department of Veterinary Clinical Medicine, College of Veterinary Medicine, University of Illinois at Urbana-Champaign, Urbana, IL, USA

Vanessa Revindran-Stam
Veterinary Diagnostic Laboratory, College of Veterinary Medicine, University of Illinois at Urbana-Champaign, Urbana, IL, USA

The genomic and clinical features of the COVID-19 Omicron variant: a narrative review. (Hertanto et al.)

Summary:

The ongoing COVID-19 pandemic is a major cause of worldwide mortality and morbidity. The review manuscript by Hertanto et al. looks at the Omicron variant that first emerged in South Africa in November of 2021 and how it became a Variant of Concern relatively quickly. This review paper is split into two parts, the first covers the genomic basis of the Omicron variant and the second part looks at the clinical features of the same variant.

Major comments:

What we find lacking in this review is that authors fail to introduce the readers to any basic background of Omicron. There are phylogenetic studies that point to the suggestion that the Omicron variant likely diverged early on from other SARS-CoV-2 strains. Examination of the whole genome has indicated that the Omicron variant can be classified into two different lineages, BA.1 and BA.2. Maybe more could have been presented on this earlier rather than at the end in the highlights section of the paper. If this is a narrative review, the hope is that an overall picture of the subject is presented. We find that this is lacking in this narrative.

The major focus of the section on the genomic bases of the Omicron variant is on the mutations that are present in the 3 key regions of the spike of SARS-CoV-2: the receptor binding domain, the N-terminal domain and the furin cleavage site. There are more than just the mutations found in
these three regions that account for the mutations present in the Omicron variant. The authors do not cover these mutations or their effect.

The first sentence in Mutations in RBD section: "As displayed in Table 2, the N501Y mutation that converts amino acid asparagine to tyrosine at position 501 was identified in the Omicron variant", is slightly misleading. On first read, it implies that this mutation was identified first in the Omicron variant. This mutation has been identified in other variants, and this is represented five sentences later in the same paragraph.

On the section titled: What can we tell from the Ct value of patients with the Omicron variant? : While the authors present that Omicron variant samples had a higher Ct value compared to other variant samples, there is nothing notable to take away from this section. The authors do not highlight any key point as to what having a higher Ct value means for the Omicron variant.

**Minor Comments:**

Table 2: Reported mutations in S-protein of the Omicron variant. This table was not helpful and did not contribute anything to the review. All the mutations identified in the Omicron variant should have been presented, not just the ones in the RBD, NTD and Furin Cleavage site. Legend for Table 2 mentions the mutations that are potentially suitable for Omicron screening but no where in the body of the text is there mention of using the mutations for screening purposes.

Clinical features of the Omicron variant:

For this section of the paper, the authors highlight the different clinical patterns observed in the Omicron variant compared to its predecessors. This section is written and presented very well. Being able to read some comparisons between the Omicron, Alpha and Delta variant was a plus and helped accomplish the author's goal. With the presentation of the Omicron came asymptomatic and symptomatic variant infection. It would be a nice touch if the authors mentioned something notable about this occurrence.

The data presented in Table 3 might have been better served if presented in graph/chart form. It would have had a better impact if that data was presented differently.

**Summary:**

Overall, this review is missing in many areas. There is no new scientific information that is presented. There are other reviews, (1: Omicron genetic and clinical peculiarities that may overturn SARS-CoV-2 pandemic: a literature review; 2: Literature review of Omicron: A grim reality amidst COVID-19) that give a good understanding of the Omicron variant and the impact on the COVID-19 pandemic.

**References:**


References

Is the topic of the review discussed comprehensively in the context of the current literature?
Partly

Are all factual statements correct and adequately supported by citations?
Partly

Is the review written in accessible language?
Yes

Are the conclusions drawn appropriate in the context of the current research literature?
Partly

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Virology, Molecular Diagnostics, SARS-CoV-2

We confirm that we have read this submission and believe that we have an appropriate level of expertise to state that we do not consider it to be of an acceptable scientific standard, for reasons outlined above.
Author Response 25 Jul 2022

Djoko Santoso, Airlangga University, Dr. Soetomo Teaching Hospital, Surabaya, Indonesia

The ongoing COVID-19 pandemic is a major cause of worldwide mortality and morbidity. The review manuscript by Hertanto et al. looks at the Omicron variant that first emerged in South Africa in November of 2021 and how it became a Variant of Concern relatively quickly. This review paper is split into two parts, the first covers the genomic basis of the Omicron variant and the second part looks at the clinical features of the same variant.

First, we would like to thank the Referee(s) for the time and willingness to review our manuscript, as well as for the critical assessment, which we believe has improved our manuscript significantly. Here, we address the comments and concerns raised by the Referee(s) point-by-point:

**Major comments:**

What we find lacking in this review is that authors fail to introduce the readers to any basic background of Omicron. There are phylogenetic studies that point to the suggestion that the Omicron variant likely diverged early on from other SARS-CoV-2 strains. Examination of the whole genome has indicated that the Omicron variant can be classified into two different lineages, BA.1 and BA.2. Maybe more could have been presented on this earlier rather than at the end in the highlights section of the paper. If this is a narrative review, the hope is that an overall picture of the subject is presented. We find that this is lacking in this narrative.

We thank the Referee(s) for these excellent and useful suggestions. Previously, we did not include the phylogenetics of the Omicron variant because at the time of writing, it was not yet clear whether it had its own clade or shared some connections with the preexisting COVID-19 variants. In this revised version, we added a paragraph discussing the phylogenetics of the Omicron variant reported by previous studies.

Similarly, regarding the BA.1 and BA.2 classifications, at the time of the writing (January 2022), BA.1 was the dominating subvariant, whereas BA.2 just began to rise. As the consequence, we provided a warning at the end of the original manuscript to highlight the potential hazard of this BA.2 subvariant. At present (6 months later), we clearly see that BA.2 took over the domination from BA.1 and as suggested, we have now added some discussion about the existing Omicron subvariants / sublineages (not only BA.1 and BA.2, but also BA.3, BA.4, BA.5 and XE subvariants).

In general, we fully agree that ideally, in our narrative review, we should be able to capture the overall picture of the subject of interest, which is Omicron variant in our case. However, the rapid development and progression of the disease, as well as the rapid changing information about this variant make it difficult to cover everything simultaneously. Employing the unique feature of F1000 Research (which allow us to submit updated versions of our manuscript), we hope to be able to keep this review live and updated, should we find new information regarding the Omicron variant in the future.
The major focus of the section on the genomic bases of the Omicron variant is on the mutations that are present in the 3 key regions of the spike of SARS-CoV-2: the receptor binding domain, the N-terminal domain and the furin cleavage site. There are more than just the mutations found in these three regions that account for the mutations present in the Omicron variant. The authors do not cover these mutations or their effect.

Again, thank you for this meticulous observation. Indeed, we originally focused only on the spike protein mutations because the spike region was the most studied part of the virus (due to its importance in the binding to ACE2 receptor and cell invasion) and the S-protein mutations were the ones that were heavily linked with the clinical features of the SARS-CoV-2 variants, including the Omicron variant. As reported by early papers about this variant, the mutations in RBD, NTD and FCS could be associated with the transmission, pathogenicity and immune response of the virus. However, we agree with the Referee(s) that several other mutations are now discovered outside of this region, including the ones in the NSP and ORF regions. Therefore, we have added a paragraph discussing this aspect as well in our revised manuscript.

The first sentence in Mutations in RBD section: "As displayed in Table 2, the N501Y mutation that converts amino acid asparagine to tyrosine at position 501 was identified in the Omicron variant", is slightly misleading. On first read, it implies that this mutation was identified first in the Omicron variant. This mutation has been identified in other variants, and this is represented five sentences later in the same paragraph.

Thanks for this correction regarding the semantics of “identify”. Since the Referee(s) feel that the word “identified” could be misleading, we want to avoid similar misunderstanding in the future. Therefore, we have replaced “identified” with “detected” in the revised manuscript and introduce the associated information “Of note, the N501Y mutation was also identified in other COVID-19 variants (e.g., C.1.2, Alpha, Beta and Gamma)” right after the sentence.

On the section titled: What can we tell from the Ct value of patients with the Omicron variant? : While the authors present that Omicron variant samples had a higher Ct value compared to other variant samples, there is nothing notable to take away from this section. The authors do not highlight any key point as to what having a higher Ct value means for the Omicron variant.

Thank you for the critics regarding this Ct section. With all due respect, allow us to disagree with the Referee(s) on this particular point. We believe that there are some important insights can be obtained from this section that the Ct value of the Omicron variant was higher than Delta, despite the higher transmissibility of the Omicron variant. A higher Ct value indicates lower viral loads in the sample taken (normally from the upper respiratory tract) and this was contradictory with the rapid spreading of the Omicron variant. Taking into account the results of previous studies, we suggest that the Omicron variant might be using another way to increase its transmissibility, rather than increasing the viral loads in the upper airways. This was also confirmed by a recent study by Laitman and colleagues.
which concluded that the rapid dissemination of the Omicron variant was not attributed to higher nasal viral loads as compared to prior variants.

Minor Comments:
Table 2: Reported mutations in S-protein of the Omicron variant. This table was not helpful and did not contribute anything to the review. All the mutations identified in the Omicron variant should have been presented, not just the ones in the RBD, NTD and Furin Cleavage site. Legend for Table 2 mentions the mutations that are potentially suitable for Omicron screening but no where in the body of the text is there mention of using the mutations for screening purposes.

As mentioned above, we previously focused on the S-protein mutations because they were more heavily studied than other mutations outside of the spike region. In response to the Referee(s) comment, we now added new figures displaying the site of the mutations outside and inside of the spike region that we obtained from The Stanford University Coronavirus Antiviral and Resistance Database. We hope that these figures are informative and suffice. We believe that research is still ongoing, thus it could be that some newly discovered mutations associated with the Omicron variant will be subsequently reported in the future.

Regarding the legend for Table 2, we have now removed the text regarding “...potentially suitable for Omicron screening” since it is not relevant anymore to be included.

Clinical features of the Omicron variant:
For this section of the paper, the authors highlight the different clinical patterns observed in the Omicron variant compared to its predecessors. This section is written and presented very well. Being able to read some comparisons between the Omicron, Alpha and Delta variant was a plus and helped accomplish the author's goal. With the presentation of the Omicron came asymptomatic and symptomatic variant infection. It would be a nice touch if the authors mentioned something notable about this occurrence.

This is a very interesting and useful comment. We have now added a new paragraph comparing the reported clinical characteristics of the Omicron vs. Delta vs. Alpha vs. Gamma variants. Of note, COVID-19 symptoms are heterogeneous and could be influenced by gender, race, ethnic, environmental factors, comorbidities etc., so head-to-head comparisons between two or more distinct studies have to be carefully interpreted.

Regarding the asymptomatic vs. symptomatic, as per the definition, there is no symptom reported in the “asymptomatic” individuals, therefore we could not compare it with the clinical characteristics of the symptomatic patients. However, the study by Laitman and colleagues for example, revealed that the Ct value in the symptomatic individuals were lower than the asymptomatic ones (which we believe might not be too surprising that higher viral loads may induce overt disease). Additionally, the fact that some patients were asymptomatic or mildly affected by the virus could be associated with an increasing herd immunity in the population. Nevertheless, a quantitative calculation of herd immunity is needed to confirm the notion.
The data presented in Table 3 might have been better served if presented in graph/chart form. It would have had a better impact if that data was presented differently.

We sincerely thank the Referee(s) for this suggestion. However, we believe that a table is more appropriate to display the exact number and percentage, as done by the original studies. To provide a clearer overall data regarding the most reported symptoms of the Omicron variant, we now calculated and added the approximate mean percentage from 6 studies that we included in this manuscript.

Summary:
Overall, this review is missing in many areas. There is no new scientific information that is presented. There are other reviews, (1: Omicron genetic and clinical peculiarities that may overturn SARS-CoV-2 pandemic: a literature review; 2: Literature review of Omicron: A grim reality amidst COVID-19) that give a good understanding of the Omicron variant and the impact on the COVID-19 pandemic.

We wholeheartedly appreciate the efforts made by the Referee(s) and we fully agree that our narrative review is still far from perfection. We are confident that following the revision, our manuscript has improved significantly and we hope that it can be approved by the Referee(s).

Indeed, the nature of a narrative review does not allow us to add new scientific information. Per definition (https://libguides.csu.edu.au/review/Traditional), a narrative or traditional literature review is a comprehensive, critical and objective analysis of the current knowledge on a topic. Therefore, most journals prohibit adding original data into a narrative review. This is clearly different than a meta-analysis that uses a more quantitative approach on a more specific and focused topic of interest.

The abovementioned reviews about the Omicron variants are nice examples and we believe that our revised manuscript can complement those reviews.

**Competing Interests:** None
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