SARS-CoV-2: The Virus, Its Biology and COVID-19 Disease-Counteracting Possibilities

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Abstract

Since the end of 2019, the SARS-CoV-2 virus started to spread in different countries, leading to a world-wide pandemic, with today’s infection numbers of more than 690 million and with a case fatality rate of more than 6.9 million. In addition, about 65 million patients suffer from post/long-Covid syndromes after having infections with the SARS-CoV-2 virus or variants thereof. This review highlights the biology of the virus, summarizes our knowledge of some of the viral mechanisms that counteract our immune responses, and finally also discusses the different vaccines and their specific safety profiles. Also, the possibility to fight this virus with recently available drugs (Veklury, Lagevrio and Paxlovid) will be discussed. All these data clearly argue that SARS-CoV-2 variants still exhibit a dangerous potential—although with a lower case fatality rate—and that vaccination in combination with drug intake upon infection may help to lower the risk of developing chronic or temporary autoimmune diseases.

Keywords: SARS-CoV-2; interferon pathway; immunology; vaccines; safety issues; small molecules

1. Introduction

The SARS-CoV-2 virus belongs to the α/β-Corona family of viruses (n = 7) that are associated with typical respiratory infections. While 4 out of these 7 viruses cause harmless seasonal infections, the remaining 3 have a much higher lethal potential; these were SARS-CoV-1, MERS-CoV and the recently observed SARS-CoV-2 virus. SARS-CoV-1 appeared first in China in Nov 2002 and led to the infection of nearly 8100 human beings of which 774 died of their infection (lethality 9.6%). MERS-CoV appeared about 10 years later and was transmitted from camels to humans. Infections spread all over the world over a period of nearly 6 years, with 2143 infected patients of which 750 died (lethality 34.9%). Finally, SARS-CoV-2 appeared at a market in Wuhan (China) in 2019, spread rapidly around the world with more than 690.7 million infected humans today of which 6.89 million have died so far (lethality is ~1% world-wide)1[1] (data retrieved 6.23.2023). This large number of infected people (nearly 8.5% of the world population) also led to ~10% of infected/recovered patients acquiring “Long-Covid” symptoms. Long-Covid is a synonym for a multitude of organ-specific autoimmune diseases which differ between different patients [2]. Long-Covid symptoms may encompass, e.g., cardiovascular, thromboembolic and cerebrovascular disease [3], type 2 diabetes [4] and myalgic encephalomyelitis/chronic fatigue syndrome (ME/CFS) [5]. Symptoms may last for longer than a year [6], and cases with ME/CFS are expected to be lifelong [7].

Thus, we have to realize that this zoonotic virus—although described as a “respiratory virus”—has the potential to increase dramatically the prevalence to develop autoimmune diseases. Moreover, since the necessary angiotensin-converting enzyme 2 (ACE2) receptor is expressed on endothelial cells, the blood circulation system is a major target for viral infections, resulting in an increase of thromboembolic events. ACE2 is also expressed in the duodenum and small intestine (mucus producing cells), gall bladder, kidney, testis and heart muscle, explaining the various symptoms observed in infected and post-infected patients. Most symptoms are related to inflammatory conditions and tissue destructive processes in these organs. In addition, cells of our innate and adaptive immune system, such as mast cells, neutrophils and naive B-cells also express the ACE2 surface receptor (data retrieved from [8]) and may change their normal immunological activity after infection.

In 2020, when no vaccine was available, the situation was dramatic in several countries. The mortality between different age groups differed significantly, with a lethality of nearly zero in small children (0.0221%), 4.2% in adult between 50–60 years and up to 40% in elderly patients (>80 years; data are based on the number of infected patients and case fatality rates (CFR) in Germany; data source: German Federal Statistical Office). The situation changed when the first vaccines became available in 2021, but also a second time when the far more infectious Omicron-variant out-competed rapidly the Delta-variant at the end of 2021. Unfortunately, the Omicron-variant displayed many amino acid changes (18 of which were in the ACE2 binding interface)—when comparing to the delta variant (2 in the ACE2 binding interface) [9]. Thus, many vaccinated people completely lost their neutralizing humoral immunity,
Fig. 1. Structure of the SARS-CoV-2 genome. (A,B) Cleavage specificity of the two proteases PL$_{Pro}$ and 3-CL. The proteolytic products of both polyproteins is given either by non-structural protein (NSP) number (1–16) or the corresponding protein name. PL$_{Pro}$ hydrolyses the first 3 NSPs (indicated by the light purple triangles below), while 3-CL cleaves all other NSPs (indicated by the dark purple triangles on top). NSP12 is the RNA-dependent RNA Polymerase (RdRp), and NSP13 represents a necessary RNA Helicase. (C) All NSP assemble at the endoplasmatic reticulum (ER) membrane into a functional replication machinery that allows the production of genomic RNA for packaging into the final viral particles (D, right side). The target proteins of 2 available drugs, Molnupiravir and Paxlovid, are indicated.

while still exhibiting a good T-cell immunity to cope with the new substrains (BA.1, BA.2, BA.4, BA.5). Although these Omicron-variants had a much better affinity for the ACE2 receptor resulting in a dramatic increase of infectivity (infections in Germany: 31.12.2020–1.753 million; 31.12.2021–7.150 million; 31.12.2022–37 million), infections were correlated with a much lower lethality (<0.4%; mean CFR = 0.8% in Europe with 830 million inhabitants, 262 million infected people and 2.1 million Corona-associated deaths) [10]. Thus, the virus became endemic at the end of 2022, and therefore, the year 2023 should mark the end of this world-wide pandemic caused by SARS-CoV-2.

This paper describes aspects of the biology of this virus, which in part explain the causal reasons for the onset and large heterogeneity of Long-Covid symptoms after real infections. Long-Covid describes the consequences of SARS-CoV-2 infection which are caused by inflammatory conditions and additional immunological mechanisms that all result in the onset of autoimmune diseases. These autoimmune reactions may affect all inner organs, blood vessels and nerve cells [2,3]. This is caused by certain SARS-CoV-2 proteins that inhibit immunological defense mechanisms, and by the 2 proteases which generate cellular neoantigens. The latter process results in the production of auto-antibodies which are the major cause underlying the development of Long-Covid symptoms. Moreover, we propose Omicron-associated mechanisms explaining the immune suppression sometimes seen in patients following multiple infections. These result from observations made early in 2023 that some patients display either a prolonged infection time (bacterial or viral) or recurrent infections. In addition, our current knowledge about the mRNA and vector-based vaccines will be presented, along with their safety profiles. Finally, this article reflects on the available drugs that can be used to combat SARS-CoV-2 infections and which in part are able to avoid the development of Long-Covid symptoms. Alternatively, vaccination with the available bivalent mRNA vaccines against the Omicron-variant are also a protective measure against the development of Long-Covid, with a risk reduction of ~40% [11].

2. Results

2.1 The SARS-CoV-2 Virus and Its Life Cycle

The original Wuhan strain of SARS-CoV-2 was early on described as a zoonotic virus which clusters together with the RaTG13 bat strain isolated from Rhinolophus, suggesting that RaTg13 could be the ancestral strain that circu-
Fig. 2. The life cycle of SARS-CoV-2. Endocytosis of angiotensin-converting enzyme 2 (ACE2) receptor-attached SARS-CoV-2 virus and Furin cleavage of Spike protein by TMPRSS2; uncoating and translation leads to the assembly of replication machinery at the ER after translation of orf1a and orf1b, replication of subgenomic RNA leads to all necessary proteins. (A) Expected exocytosis for Virus release. (B) Observed Virus release via the lysosomal egress pathway. The latter pathway depends on the Ras-related protein Rab-7a (RAB7; can be inhibited by the drug CID1067700) and on the ADP Ribosylation Factor Like GTPase 8B (ARL8B). Additional functions known for SARS-CoV-2 proteins are indicated on the left (information retrieved from the ViralZone 2021). The figure has been adapted from ref. [20].

SARS-CoV-2 virus exhibits a ~30,000 nucleotide long RNA genome [18] that encodes large polyproteins that are linked via a frameshift to produce 16 non-structural proteins (NSPs), including 2 proteases, a helicase (Hel) and an RNA-dependent RNA Polymerase (RdRP). The 3’-portion of the RNA genome encodes the essential proteins for virus assembly, such as the Spike protein, the membrane glycoprotein, the envelope protein as well as the nucleo-capsid protein which is necessary to protect the packaged RNA genome. Besides these major virus proteins, smaller open reading frames encode a series of accessory proteins, named orf3a/b, orf6, orf7a/b, orf8, orf9b/c and orf10, respectively. Taken together, SARS-CoV-2 virus has the capability to produce a total of 28 proteins (see Fig. 1).

2.2 Known Functions Deriving from SARS-CoV-2 Encoded Proteins

For some of these viral proteins a cellular function in infected cells is already known. About 10 NSPs/orfs are involved in inhibiting the interferon pathway [19], the two proteases (PL<sub>Pro</sub> and 3-CL) cleave the polyproteins and cellular host proteins, orf3a (Viroporin) binds to lysosomal membranes and inhibits their acidification which allows a lysosomal egress pathway (lysosomal exocytosis) [20], and in addition, inhibits autophagy by blocking the fusion of autophagosomes with lysosomes [21]. Immunosuppressive functions have also been discovered: the SARS-CoV-2-encoded M protein antagonizes the typeI/III interferon response [22], the combination of 3-CL/orf3b/N protein suppresses the innate eGAS-STING pathway [23], and the N/orf6/orf9b proteins suppress the TOM70-mediated activation of the mitochondrial antiviral signaling (MAVS) complex that activates IRF3 via the kinase Tbk1 [24]. All this appears to exhibit strong immunosuppressive functions for our innate and adaptive immune system, and immunosuppressive myeloid cell induction and T-cell depletion have been described recently [25]. The latter probably results in many individuals suffering from recurrent bacterial infections, or, pro-longed disease courses when infected with harmless viruses during the first months of 2023. Whether these observations hold true for a longer period of time has to be carefully examined.
Fig. 3. Adaptive immune response under normal (A) and hyperinflammatory condition (B) in secondary lymphatic organs (spleen or lymph nodes). (A) Immune response under physiological conditions. B-cells become activated in germinal centers by activated follicular T-helper cells (BCL6+ Tfh). This leads to a massive proliferation of B-cells and the selection of optimal B-cells after the process of hypermutation. (B) Immune response under SARS-CoV-2 infection. The germinal center reaction is compromised in COVID-19 patients that suffer from hyperinflammatory conditions. High amounts of TNFα block the differentiation of inactive follicular T-helper cells into activated follicular T-helper cells, and thus, the typical germinal center reaction is strongly compromised, if not absent. Right panel: short explanation of the observed effects in severe COVID-19 patients. The figure has been adapted from ref. [28].

One of the major dangers associated with SARS-CoV-2 infections is the fact that this virus encodes 2 viral proteases which appear to generate a massive pool of protein fragments (neoantigens) from host cellular proteins. This switches on a vicious pathway to steer autoimmune reactions [26], which is of course dependent on the affected host and their capacity to present such aberrant peptides to their immune cells. As a consequence, affected individuals experience autoimmune reactions against their own body organs. Such autoimmune reactions are then collectively termed “Long-” or “Post-Covid”, depending on how long these post-infection symptoms persist [11].

An important biological feature of SARS-CoV-2 is the final part of its replication cycle, namely how new virus particles are released from infected cells. As already mentioned above [20], the SARS-CoV-2 virus is assembled in the cytosol but leaves the infected cells by a rather unexpected route. Cell-internal lysosomes, which are usually rather important for the destruction of foreign or host proteins in acidic (pH 5) conditions, are hijacked for the egress of the virus. One of the reading frames of SARS-CoV-2, the orf3a protein, encodes a protein that inhibits the acidification of lysosomes. Therefore, the virus uses these de-acidified lysosomes as a shelter for newly produced viral particles, which are then released when these de-acidified lysosomes fuse with the outer membrane. This process is termed “lysosomal egress pathway” or “lysosomal exocytosis” (see Fig. 2, Ref. [20]). Interestingly, this pathway is dependent on RAB7 and ARL8B. Noteworthy, at least for RAB7 potential inhibitors are available [27].

This is important, because this trick bears a great danger for the infected host. Lysosomes are urgently needed in Antigen-presenting cells (APCs) after phagocytosing the virus or for B-cells after endocytosing surface antibody-bound virus. The resulting membrane vesicles (phagosome, endosome)—containing the infectious viruses—are then acidified to mature into lysosomes where all the internal protein material becomes hydrolyzed into peptides. These peptides are then used to load major histocompatibility complex (MHC) II complexes to present the foreign peptides to T-helper (T_H) cells. The fact that SARS-CoV-2 is able to deacidify lysosomes makes it nearly impossible for such cells to exhibit their natural immunological function. Thus, APCs and B-cells are hampered in their ability to present peptides, and thus, the adaptive immune system has a problem in activating T_H- and appropriate B-cells for antibody production. Since, for this reason, immune responses are limited to the innate immune system, hyperinflamma-
Fig. 4. Summary on protein-protein interactions of SARS-CoV-2-encoded proteins with host proteins and pathways. (A) Structure of SARS-CoV-2 with a total of 28 proteins. (B) Summary of all the identified protein interactions between viral proteins and host proteins of major cellular pathways. Black boxes are indicating that a specific SARS-CoV-2 protein interferes with the highlighted pathway of the infected host cell. Noteworthy, about 10 viral proteins (NSP1, NSP3, NSP12, NSP13, NSP15, orf3a, M, orf9b and orf9c) are interfering with the interferon pathway. This figure has been adapted from ref. [32].

2.3 Interference with the Host Defense Systems

A very good overview on the diverse functions of SARS-CoV-2 proteins has been published already in 2020 [32]. Intracellular interactions with host proteins have been investigated using modern mass-spectrometry. These data have also provided potential drug targets for the treatment of SARS-CoV-2. As shown in Fig. 4 (Ref. [32]), most of the 28 SARS-CoV-2 encoded proteins are able to bind to multiple host factors, and thus, may have the potential to interfere with several cellular pathways. All this cellular interference helps the virus in its own replication and affects processes such as lipid and sugar metabolism, protein stability, vesicle trafficking, mitochondrial functions and control mechanisms for inflammatory and interferon responses.

In particular, the potent inhibition of the interferon response pathway could explain severe cases of COVID-19. SARS-CoV-2 infections turn on the Interferon pathway via the RIGI/TLR3/IRF7 axis, leading to a profound type I/III interferon response to block viral replication by specific defense proteins (different IFIT proteins, ISG15 and Viperin). This usually defends against viral infections quite effectively, because all these proteins recognize viral nucleic acids, covalently modify viral proteins and target them for proteasomal destruction or synthesise unusual RNA nucleotides (e.g., 3′-deoxy-3′,4′didehydro-CTP) that can inhibit viral RNA synthesis. Of these 28 viral proteins, 10 interfere with the normal interferon pathway, indicating that shutting down this particular host defense pathway is of great importance for the lifecycle of SARS-CoV-2 and may explain retrospectively the high lethality observed for SARS-CoV-1 and MERS-CoV infected individuals.
Fig. 5. Delivery of vaccines to target cells. mRNA-based vaccines deliver a perfect mRNA to the cytosol where it could be immediately translated into Spike protein at the ER, become post-translationally glycosylated and then exported as a trimeric, receptor-like protein via the Golgi system to the outer membrane of the producing cell (secretory pathway). Vector-based vaccines are delivered first to the nucleus where they are transcribed into Spike mRNA and then exported from the nucleus to the cytosol. As already mentioned above, the Spike protein is then translated at the ER, post-translationally glycosylated inside the ER and exported via the Golgi system to the outer membrane. This allows stimulation via antigen-presenting cells of corresponding T-helper and B-cells to initiate the production of cognate antibodies. Since peptides of the foreign Spike protein are also displayed via MHC I at the outer cell surface, appropriate T-killer cells are activated as well. Thus, these novel classes of vaccines allow a profound immune stimulation without the need of any adjuvant.

Several studies have investigated the role of the interferon response in different COVID-19 patients. Based on single blood cell gene expression profiling, healthy controls can be clearly distinguished from mild, severe or critically infected patients by their gene expression profiles, the poorest interferon response being displayed in severe cases [33]. Moreover, genetic studies revealed the OAS1 gene as a restriction factor against SARS-CoV-2 infections [34,35]. The p46 variant OAS1 gene (splice variant of the original OAS1 gene, OAS1 isotype) is encoded by a 75 kb piece of DNA located on chromosome 12q24.13. This OAS1 isotype gene variant originally derived from Neanderthals and protects against severe SARS-CoV-2 infections [36–38]. Patients that bear certain SNPs or display a genetic defect in OAS1 displayed a strongly impaired or absent interferon response [39], while children with such a genetic mutation develop the multi-inflammatory syndrome in children (MIS-C) [40]. Other patients with severe outcomes after infection displayed autoantibodies against the Interferon protein [41].

Most of these poor interferon-responding patients suffered from life-threatening COVID-19 pneumonia [42] or multi-organ failure [43], which was responsible for more than 20% of COVID deaths in elderly patients [44]. In general, autoantibodies against different cytokines seem to correlate with severity of the disease course [45], validating that these pathways are crucial for viral defense. In summary, hospitalized patients with a fast type I/type III interferon response—accompanied with low levels of pro-inflammatory cytokines (IL1, IL6, TNFα)—were observed in most patients with mild COVID-19 symptoms, while a delayed or absent type I/type III interferon response—accompanied with the highest levels of pro-inflammatory cytokines (IL1, IL6, TNFα)—were observed in most patients with a severe or fatal COVID-19 disease. MIS-C in children has been associated with high levels of IL6, while the Kawasaki syndrome (KS) was associated with high levels of IL6 and IL17A [46].

2.4 Vaccination as a Strategy Against SARS-CoV-2

In order to prevent all the above-mentioned negative side effects of real SARS-CoV-2 infections on our immune system, only two possibilities exist. The first option is to vaccinate populations in order to develop immunological resistance to the virus and to prevent severe courses of disease. The second option is to take drugs that block viral replication which prevents fast spreading of the virus in infected hosts and promotes rapid clearance of the virus and virus-infected cells. The latter will be discussed in a separate chapter below.

Vaccination has been used as a strategy since the 20th century to fight and to prevent several diseases around the world. There are several vaccine developing strategies (attenuated living virus, antigen/adjuvant combinations, LPS-
toxin conjugates) that have been successfully used in the past and are currently in use to combat major diseases. Similarly, the SARS-CoV-2 pandemic that arose in the beginning of 2020 required the development of a quick vaccine strategy world-wide. Luckily, two novel strategies were available, namely the adenovirus vector-based and mRNA vaccination technology. These novel strategies were used by different companies (e.g., Biontech/Pfizer: BNT162b2; Moderna: mRNA-1273; AstraZeneca: ChAdOx1-CoV-19; Johnsen/Johnsen: Ad26.COV2.S) and ended with successful clinical trials [47–50]. These novel vaccine development strategies were in the end much faster than traditional ones. Therefore, I want to focus in this review on these two novel strategies, as both have a great potential for the future.

Adenovector-based vaccines use infectious adenoviral particles (Janssen/Johnson&Johnson: HAd26; Oxford Jenner Institute/AstraZeneca: ChAdOx1, Sputnik: HAd5 and HAd26) with the E1 gene replaced by a codon-optimized cDNA version of the Wuhan SARS-CoV-2 Spike gene. The ChAdOx1-encoded codon-optimized Spike cDNA was flanked by a cytomegalovirus (CMV) Enhancer, an intron-containing CMV Promotor and an SV40 polycl- A signal sequence. The whole cassette was located 174 bp upstream of the adenoviral pIX reading frame. The HAd26-encoded codon-optimized Spike gene is flanked by a CMV Enhancer, an intron-less CMV Promotor and an SV40 polycl-A signal sequence, located 99 bp upstream of the adenoviral pIX reading frame. Both adenoviruses infect human cells and deliver the adenovector DNA to the nucleus, with Spike mRNA transcripts then be produced and delivered to the cytosol of the cells. The mRNA is then translated at the endoplasmatic reticulum (ER) and the resulting Spike trimer is delivered via the ER-Golgi-axis as a membrane-bound, trimeric Spike protein to the outer surface of infected cells. The virus has a maintenance of 2–3 weeks and produces the recombinant Spike proteins over this time period. One of the problems with this approach is the fact, that SARS-CoV-2 is an RNA virus, with a natural life cycle only in the cytosol of infected cells. Therefore, none of the genes encoded by SARS-CoV-2 has ever been selected or optimized by nature to be expressed in the nucleus of eukaryotic cells. As a consequence, the open reading frame of the Spike gene, coding for a 1274 amino acid long Spike protein, is full of potential splice donor and acceptor signal sequences. Some of them were point-mutated by Janssen, but none of them by AstraZeneca. Therefore, alternative splice events occur, leading to shorter Spike protein variants or even fusion by splicing to the downstream located pIX gene [51,52]. None of these aberrant Spike protein variants can become a membrane-bound protein anymore, because all of these C-terminally shortened Spike protein variants lose the necessary membrane anchor, and thus, are secreted to the blood stream. Based on available RNA-Seq data (kindly provided by Prof. David A. Matthews, University of Bristol, UK), the amount of these aberrant Spike protein encoding mRNAs was estimated to be in the range of 1.5% of all Spike mRNA transcripts. Although it has not been formally proven, these shorter Spike variants—or the discovered Spike-pIX fusion protein—may be one reason for the observed thrombotic events (sinus vein thrombosis), which were observed shortly after vaccination with vector-based vaccines (see below).

Because of the associated adverse events, the Adenovirus-based vaccines were used only in the beginning of the vaccination campaign, but not used after summer 2021 for primary vaccinations in Germany. Therefore, the number of vaccinations with vector-based vaccines is quite low when compared to mRNA-based vaccines.

By contrast, mRNA vaccines used a different strategy. In vitro transcribed mRNA which was appropriately capped and poly-adenylated is packaged into lipid-nanoparticles that contain specific lipid compositions of PEG-DMG, DODMA, DSPC and Cholesterol. This particular composition guarantees the fusion of these nanoparticles with target cells upon intramuscular injection. The mRNA was synthesized with N1-methyl pseudouridin nucleotides that replaced the Uracil nucleotides to avoid immunological counterreactions. This facilitated higher concentrations of mRNA being packaged into these lipid nanoparticles (Biontech: ~30 µg; Moderna: ~50 µg). After injection, these nanoparticles fuse with target cell membranes and deliver the modified mRNA to the cytosol of these cells, where it is immediately translated at the ER, and the resulting Spike trimer is again delivered via the ER-Golgi-axis as a membrane-bound, trimeric Spike protein to the outside surface membrane of cells. The maintenance of the mRNA inside of the transduced cells is estimated to be 2–3 days (see Fig. 5).

Both approaches were game changers for the field of vaccine development. In the past, vaccine development was either based on attenuated living viruses for life-long protection, or on the use of portions of viruses or bacteria (recombinant) combined with adjuvant for short- to midterm immune protection. Vector- or mRNA-based vaccines both deliver a nucleotide-encoded antigen into human cells. By using the cell as a production unit for the protein antigen, a profound stimulation of the humoral (T_h + B) and cytotoxic axis (T_k) of our adaptive immune system can be achieved. Thus, these vaccines—all classified as “dead vaccines”—behave like “living vaccines” when viewed from the perspective of their executed immunological functions. Although longitudinal studies are necessary and yet not available due to the short observation time, one can hypothesize that persons vaccinated with either vector- or mRNA-based vaccines for several times will exhibit a very long—if not life-long—immunity against the vaccinated antigen(s).

2.5 Security Aspects of Vector- and mRNA Based Vaccines

The Federal Paul-Ehrlich Institute (PEI in Langen, Germany) has the important task to constantly evaluate the
Table 1. Evaluation of side effects and safety of 4 different vaccines that have been used in 2021 for the world-wide vaccination campaign.

<table>
<thead>
<tr>
<th>Companies</th>
<th>Biotech/Pfizer</th>
<th>Moderna</th>
<th>Oxford/AZ</th>
<th>Janssen/J&amp;J</th>
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<tr>
<td>Total vaccinations</td>
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<td>10,576,131</td>
<td>12,703,030</td>
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<tr>
<td>Complications</td>
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<td>28,289</td>
<td>46,325</td>
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<td>cases/Million</td>
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<td>550</td>
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<td>cases/Million</td>
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<tr>
<td>injections</td>
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<td>TTS*</td>
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<td>5</td>
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<td>cases/Million</td>
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<td>ITP**</td>
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<td>cases/Million</td>
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<td>injections</td>
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<td>GBS***</td>
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<td>Death cases</td>
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<td>cases/Million</td>
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<td>injections</td>
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* TTS-Incidence in Germany: 3 diagnoses/ Million individuals per year.
** ITP-Incidence in Germany: 20–40 diagnoses/ Million individuals per year.
*** Guillain-Barré-Syndrome Incidence in Europa: 8–19 diagnoses/ Million individuals per year.

benefits or risks of vaccines on the market. By law, all physicians and companies are obliged to report all cases with negative side effects to this institution. Therefore, the PEI has a large data resource to monitor side-effects and efficacy of vaccines. Based on the last report in 2021 (23.12.21; about 1 year after the vaccination campaign started in Germany), all 4 vaccines were analyzed for their potential danger. This is summarized in Table 1 where total and adjusted numbers (cases per million injections in red) are summarized for the observed vaccine-associated complications. The 4 investigated vaccines differed quite dramatically in their usage. The Biotech mRNA vaccines was used for 96.6 million, the Moderna mRNA vaccine for 10.5 million. The AstraZeneca vector-based vaccine for 12.7 million, and finally, the Janssen vector-based vaccine for 3.4 million injections. When looking at the red marked lines where all side-effects are shown in numbers per million injections, it becomes quickly clear that the normal vaccine-related side effects (headache, pain at the injection site, fatigue, fever, etc.) were lowest for the Biotech (B) product, a little more for the Moderna (M) product, highest for the AstraZeneca (AZ) vaccine and again lower for the Janssen (J) product. The reason for the increased vaccine-related side effects with the AZ product could be explained by the identified impurities in several batches of the vaccine which may explain the increased inflammatory reactions [53]. Anaphylactic reactions were rare and between 3–8 cases per million injections. Noteworthy, myo- or pericarditis cases were observed only with mRNA vaccines, and were more than doubled in the M product when compared to the B product (50 µg vs. 30 µg mRNA). Instead, cases with Thrombosis with Thrombocytopenia Syn-

drome (TTS)—including the rare sinus vein thromboses—mainly occurred with the AZ product, were less frequent with the J product and nearly absent for both mRNA vaccines. Normal Thrombosis with immune thrombocytopenia (ITP) were again increased for the AZ product, while much less for the J, B and M products. Finally, cases with Guillain-Barré-Syndrome (GBS) were slightly elevated for the AZ and J products, while nearly absent for both mRNA vaccines. Finally, the total number of deaths was low for the B, M and J products, while strongly increased for the AZ product. This was the main reason for not using the AZ product later in the year 2021, and to switch the population-wide vaccination campaign completely to mRNA vaccines. In order to evaluate the data even further, the TTS incidence is ~3 for 1 million inhabitants in Europe per year, underlining again the security of mRNA vaccines with 0.4 and 0.5 cases after 1 year of mRNA vaccinations. The ITP incidence is ~20–40 for 1 million inhabitants in Europe per year, indicating that all 4 four vaccines had lower incidences. Also, the GBS incidence is between 8–19 for 1 million inhabitants in Europe per year, indicating that only the AZ vaccine was lying exactly in this range, while all other vaccines were below this number. Thus, based on this analysis, mRNA vaccines are very safe, apart from the observed risk for myo- or pericarditic events, mostly observed in young men between 25 and 30 years of age [54]. This also supports the notion that additional vaccinations with the now available bivalent vaccines from B and M should not only protect against the Omicron-variant but could also help to lower the number of Long-Covid patients.

The displayed data were retrieved from the security report on the vaccination campaign in Germany, dated
19 patients have shown clear-cut benefits in clinical trials with COVID-19, the activity of the main protease 3-CL of SARS-CoV-2 and a mixture of Normatrelvir and Ritonavir, effectively blocks carefully restricted for treatment. By contrast, Paxlovid, a new virus mutants of these drugsoftenincreasestheprobabilityofgenerating particles. However, several studies have shown that the use of remicadinant monoclonal antibodies (cocktails) such as Casirivimab/Imdevimab (Ronapreve®; Roche/Regeneron), Regdanvimab (Regkirona®; Celltrion), Tixagevimab/Cilgavimab (Evusheld®; AstraZeneca) or Sotrovimab (Xevudy®; GSK/Vir Biotechnology) prevented severe courses in 2020/21 and certainly saved lives of high-risk patients. However, many of these recombinant antibodies became ineffective when the Omicron-variant appeared in 2022, except Sotrovimab which still appeared to have neutralizing capacity against this novel virus variant.

I also want to place the focus on some oral drugs which were introduced for the treatment of COVID-19 patients: Remdesivir (Veklury), Molnupiravir (Lagevrio) and the combination of Normatrelvir and Ritonavir (Paxlovid). Remdesivir and Molnupiravir are both nucleotide analogues that should in principle hinder the RNA dependent RNA polymerase of SARS-CoV-2 (NSP12) to replicate efficiently the viral genome and to produce new viral particles. However, several studies have shown that the use of these drugs often increases the probability of generating new virus mutants [55–58], and thus, their use should be carefully restricted for treatment. By contrast, Paxlovid, a mixture of Normatrelvir and Ritonavir, effectively blocks the activity of the main protease 3-CL of SARS-CoV-2 and has shown clear-cut benefits in clinical trials with COVID-19 patients [59–62]. The 3-CL protease is absolutely necessary for virus replication, but the half-life of the specific protease inhibitor is too low. Therefore, it has been combined with Ritonavir, a well-known drug for the treatment of HIV-1 patients. Ritonavir is by itself also a protease inhibitor, but the main function is the inhibition of the liver enzymes CYP3A4 and CYP2D6 which are responsible for metabolizing Normatrelvir. Thus, the combination of both drugs enhances the pharmacodynamic behavior of Normatrelvir and allows the use of lower doses of both inhibitors. The only reason why it is not so easy to use Paxlovid is that inhibition of CYP3A4 also has dramatic effects for many other drugs (roughly one third of all available drugs which are today on the market). SARS-CoV-2 infected individuals need to take Paxlovid for 5 days in a perfect 12 h rhythm to effectively kill all SARS-CoV-2 viruses in their body, but if these patients are already taking several other drugs on a daily basis, then their treatment requires careful medical evaluation. The list of drugs that are dependent on CYP3A4 (either for drug destruction or activation) is rather long, and therefore, many physicians hesitate to prescribe Paxlovid, although it is as yet the only effective and available drug against SARS-CoV-2 virus [63]. However, the effectiveness of Paxlovid has been shown in a retrospective clinical study of the California Department of Public Health, as yet not peer reviewed, demonstrating that bivalent vaccines and Paxlovid could avert hospitalization (7.8% vs. 11.2%) and SARS-CoV-2 related death cases (16.2% vs. 25.2%) [64]. To this end, Paxlovid is a valuable drug in the treatment of SARS-CoV-2 infections.

3. Discussion

Humanity has faced the largest world-wide pandemic of the 21th century. In 2020, we needed to learn how dangerous SARS-CoV-2 infections really were, how hospitalized SARS-CoV-2 patients had to be treated and how we could protect ourselves against airborne SARS-CoV-2 infections. But even with all these efforts, we still faced a case fatality rate of ~2.4% worldwide. Luckily, one year later we already had several vaccines available of which 4 were mainly used in the Western world. Therefore, the second year of the pandemic was accompanied by large vaccine campaigns in order to get the whole population convinced and vaccinated. Many hopes but also a much criticism was heard in 2021, because mankind has never produced, tested and legally approved new vaccines within such a short time-frame. However, all clinical studies demonstrated the efficacy of these vaccines, even after a first injection. The efficacy to combat the virus (an all subvariants thereof) increased with the second and third vaccination steps. However, at the end of 2021, a new game changer appeared in form of the Omicron-variant. Omicron appeared first in Africa, probably deriving from a single immuno-suppressed patient in which SARS-CoV-2 had evolved over months to perfectly adapt to the human host [65]. The Omicron-variant represented an immune-evasive derivative of the original Wuhan SARS-CoV-2 virus. The Omicron-variant displayed a much higher infectivity and out-competed all yet existing strains and variants within 2–3 months. Since Omicron had so much more mutations in the ACE2 binding domain (~18 mutated amino acids in the ACE-2 binding interface alone), all neutralizing antibodies failed to neutralize the virus, and thus, could not prevent novel infections. Thus, infections in 2022 occurred equally in vaccinated and not-vaccinated people, and at least 50% of the populations in different countries were re-infected at
least once. Most countries could not handle this situation anymore and things became statistically uncontrollable. At the end of 2022, most people—if not all—were vaccinated several times, infected or both. Thus, the Omicron-variant did two things: due to the high infectivity, it infected most people, but its lethality was dramatically reduced. Therefore, the CFR is now in most countries between 0.3–0.4% which is slightly higher than that for flu-infections. Thus, the SARS-CoV-2 became endemic and the SARS-CoV-2 pandemia is ending in 2023.

Despite this hopeful conclusion, we have to acknowledge a few things. First, airborne infections can be handled well by simply wearing masks, and many studies have shown this clearly [66]. As a scientist, I am quite thankful that most journals made SARS-CoV-2 studies freely available to anyone in order to speed up scientific progress. We also have realized the value of pre-print servers (like e.g., https://www.medrxiv.org/, https://www.biorxiv.org/, https://www.researchsquare.com) to rapidly spread scientific information. We should also be thankful to companies that took the efforts to produce these novel vaccines which do not need adjuvants, are dead vaccines by definition but will give a long-lasting protection against harmful infectious agents. At least the mRNA vaccines are not only very rapid in terms of development but provide also a high safety and efficacy when compared to existing vaccines. This is important because with global warming, many new viruses will come to the Western world, for which we have yet no effective vaccines available (e.g., West-Nil-virus, Krim-Kongo virus, Dengue fever, Zika virus, Chikungunya virus, etc.).

The last thing to mention is that SARS-CoV-2—regarded as a respiratory infection virus—should be classified as systemic disease causing virus. It mainly affects our blood circulation system. Blood vessels are internally lined with a thin monolayer of endothelial cells that control blood flow, blood pressure (via the smooth muscle cells behind them), blood clotting reactions and integrity of the complete circulation system. Endothelial cells are also involved in the control of the early steps of inflammatory reactions, e.g., by controlling the transmigration of white blood cells. Infection of these cells not only causes prothromboembolic situations, it also destroys the important barrier functions of the endothelial cell layer. Thus, the effects for the body are dramatic (damage, inflammation, etc.) and cause the aging of affected organs. So even if the pandemia is over, we will still face in the future the results of all this organ damage across the whole population. Also the proteases encoded by this virus, PL̂pro and 3-CL, and their effects on presenting “auto-neoantigens” to our immune system, has caused an increase in autoimmune episodes, which we termed Long- or Post-Covid (>1 year). The danger of developing autoimmune diseases is very high and today affects about 10% of SARS-CoV-2 infected patients. Vaccination studies have shown to decrease this risk for developing Long-Covid (~45%) but cannot completely avoid it. This is due to the large heterogeneity and variability of our MHC molecules which are needed for cross-talk and activation of T-cells. The individual immune system decides whether to react against a presented neoantigen—or not. The immunological heterogeneity of human individuals also explains the large variability of symptoms in Long-Covid patients, that are yet un- or badly treatable. Therefore, we will have to deal with this pandemia—or better with its long-lasting consequences—also in the future. Life expectancy has already decreased in all major countries worldwide with a mean of 1.2 years in all European countries during the last 2 years [67].

4. Conclusions

The SARS-CoV-2 pandemia has taught humanity an important lesson for our lives in modern times: viruses can easily spread around the world and can cause an enormous physic, emotional and economical damage. This pandemia has displayed the fragility of our medical systems, has produced a large number of patient deaths, has shed light on our product delivery chains and the strong influence on economic activity in the case of lock-down decisions taken by governments. We should be aware that global warming, the way we destroy nature and the way we live today, harbour a big danger for ourselves. We need to learn this lesson before the next pandemia arrives. In addition, we should be aware that not every virus is amenable to rapid and effective vaccine development (see HIV-1).

On the other hand, we have learned how basic and medical Science—in combination with the pharmaceutical industry—were able to develop and produce novel vaccines, which were then rapidly evaluated for their efficacy in clinical trials. This gives hope for the future in which similar pandemic situations may emerge. Note-worthy, these vaccines are able to avoid all the above-mentioned adverse effects of real SARS-CoV-2 infections which ended with a high rate of case fatalities. The effects on our immune system are dramatic and ~10% of all infected patients have developed Long-Covid symptoms. This is in strong contrast to the very low numbers of severe vaccination-related side effects (post-vac syndrome) which is in the range of several hundreds to thousands in Germany (~0.0015% of at least 1-time vaccinated individuals).

In contrast, we have also seen negative aspects in this pandemia: social media has been misused to spread fake news or hate-speech. In addition, social media has influenced the recognition of vaccination as a key element to combat any kind of pandemia. Based on the available data, the CFR between unvaccinated and vaccinated individuals (a factor of 15 to 20-fold) tells a clear story for most European countries [68]. Therefore, effective education of young scientists has been also recommended to block the spreading of fake news [69]. This could help in the future
to get more rapid responses of populations to viral threats and to be prepared for the next pandemic—as this is only a matter of time.

**Author Contributions**

The author confirms sole responsibility for the following: study conception and design, data collection, analysis and interpretation of results, and manuscript preparation.

**Ethics Approval and Consent to Participate**

Not applicable.

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This manuscript is dedicated to Professor Dr. Theo Dingermann on the occasion of his 75th birthday, who has been one of my esteemed scientific mentors. As the editor-in-chief of the "Pharmazeutische Zeitung" (PZ) at Avovxxa Group, he has authored numerous articles on the SARS-CoV-2 virus, the pandemic, methods of protection against airborne infections, and information about available vaccines. This contribution has been invaluable in raising awareness among the public, pharmacists, and physicians regarding the threat posed by SARS-CoV-2, as well as the efficacy and safety of mRNA vaccines, which are central to the global effort against this pandemic.

I extend my heartfelt thanks to Theo for his outstanding work in this regard.

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**Conflict of Interest**

The author declares no conflict of interest.

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