Review Article

A review on SARS-CoV2 drug regimens inferring plausible mechanisms to impede the viral propagation and cytokine storm

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ABSTRACT

A sharp outbreak of pandemic severe acute respiratory syndrome coronavirus 2 (SARS-CoV2) in early 2020, has shaken up the whole global health care system and economy with its rapid chain-of-infection and high mortality index. This article emphasizes on the immunopathology of virus and current drug therapies. We have put down a compendious postulation on different drug actions vis-à-vis their reversing strategies for the pertinent diseases. A critical analysis has been made on the cytokine storm and focusing on to the interplays among lymphocytes and other innate immune cells (NK cells, macrophages, dendritic cells etc.) with secreted cytokines as well as on to the drug effectiveness by tallying with the contexts of SARS-CoV and Middle East respiratory syndrome coronavirus infections. The hustle and bustle is ongoing for repurposing existing drugs, designed for other infectious microbes. Tackling the adversities in anti-viral treatment has become too challenging to stop this contagion. Thereafter, the scientific front liners are on trials of hundreds of drugs by making comparisons to other contagious epidemics. Current review summarizes different drug actions on viral propagation and host immunomodulation. Our conjecture will intrigue to re-evaluate the mostly used drugs at present state.

Keywords: SARS-CoV2, Cytokine storm, Anti-viral drugs, Remdesivir, Favipiravir, Immunopathology

INTRODUCTION

In 21st century, the deadly outbreak of newly emerged coronavirus disease 2019 (COVID-19) caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV2) is the third most lethal pathogenic human corona virus, after SARS-CoV and MERS-CoV in 2003 and 2012, respectively. The outbreak was first reported on December 31, 2019 in Wuhan City, Hubei Province, China. Originating from China, till now it has outspread nearly 215 countries including all major continents. On 26th January 2021, globally above 98 million confirmed cases, and more than 2.1 million confirmed fatalities have been recorded (fatality rate of 2.15% worldwide). Fever (83-98%), cough (59-82%), shortness of breath (19-55%), and muscle ache (11-44%) are the usual symptoms of SARS-CoV2 infection but some patients may develop sore throat, rhinorrhea, headache and fever, indicating as critical symptoms. When SARS-CoV2 progresses from severe to critical, patients may develop severe cytokine storm, secondary acute respiratory distress syndrome, followed by the shock, tissue perfusion disorders, and even multi-organ failure. Due to the high infectivity, mortality rate and the absence of definitive treatment protocols or therapeutic agents or vaccines, the urgency of this outbreak lead to the use of broad-spectrum clinically available non-specific antiviral drugs and non-viral drugs. In spite of significant efforts for the development of vaccines and therapeutic drugs none of them have reached to bedside. The vaccines which are
came to light with promise and now under clinical trials but warrant more time. In some countries, several cocktail drugs or different combinational therapies are being used for the treatment of the SARS-CoV2 infected patients. Existing drugs that were previously designed for other specific viral infections are used to treat SARS-CoV2 infection, and most of these agents were tested for their safety. Several clinical trials are in a row with different drugs, cocktails, and combinational therapies. These drugs either can be directly target the viral infection response pathways or boost host innate antiviral immune response or alleviate damage induced by dysregulated inflammatory responses. In the current context, we review the detail immunopathology of the infection with special emphasis to cytokine storm and the mechanism of different non-specific antiviral drugs used against SARS-CoV2 infection for the better therapeutic interventions against SARS-CoV2 infection. The review is also illustrating the world wide current status of the potential drugs used so far against the infection.

PATHOLOGICAL ASPECTS OF SARS-COV2

Pathogenesis

Recent understanding of the pathogenesis of SARS-CoV-2 infection is still limited. Before 2019, there were six CoVs (HCoV-229E, HCoV-OC43, HCoV-NL63 and HCoV-HKU1) that could infect humans and cause respiratory disease i.e., the “common cold”. Estimated 80% of the SARS-CoV2 infection is asymptomatic or mild, while remaining is severe or critical in condition. SARS-CoV2 can infect lower respiratory tract and causes severe pneumonia, which is similar to SARS-CoV and MERS-CoV, including fever, dry cough, dyspnea, myalgia, fatigue, normal or decreased leukocyte counts. Further SARS-CoV2 infection leads to lung inflammation, acute respiratory distress syndrome (ADRS), cardiac and renal injury and multi organ failure. Critical correlation with SARS-CoV and MERS-CoV infection and information available for SARS-CoV2 infections so far will facilitate better clinical perception of the current pandemic.

Immunopathology of SARS-COV2 infection

SARS-CoV2 is mainly characterized by fever, pneumonia, lymphopenia, exhausted lymphocytes, lung inflammation and cytokine storm. SARS-CoV2 progress to ADRS approximately 8-9 days after symptom onset. The immunopathology of SARS-CoV2 infection closely resembles with SARS-CoV and MERS-CoV infection, which results an aggressive inflammatory responses and damage to the airways like alveolar cells, ciliated and goblet cells in the airways. Disease severity in patients is not only due to the viral infection but also the host response. The disease severity increase with age is also similar with SARS-CoV and MERS-CoV. Improved insight into immunopathological change in patient is important for better clinical management. The SARS-CoV2 spike protein (S) is 20-30 amino acids longer than SARS-CoV and binds to the ACE-2 receptor over a host cell which includes human airway epithelial cells, goblet cells, type 2 alveolar epithelial cells, vascular endothelial cells, macrophages in the lung, cardiac cells and intestinal epithelium, all of which highly express the transmembrane ACE-2 receptor. The host response and clearance of viral infections heavily depends on type I interferon (IFN1) expression and programming the immune cells into an “anti-viral state”.

Immune cells first identify the viral RNA through virus derived pathogen associated molecular patterns (PAMPs) which activate pattern recognition receptors (PRRs) resulting in the production of IFN1 and expression of IFN-stimulated genes (ISGs) that target viral life cycle, including virus binding to attachment receptors, virus entry, RNA synthesis, progeny virion assembly and egress. As SARS-CoV, SARS-CoV2 can also be detected by endosomal RNA PRRs, toll-like receptors (TLR)-3,7,8, retinoic acid-inducible gene I (RIG-I) and melanoma differentiation- associated protein 5 (MDA5) (Figure 1). RIG-I and MDA5 can sense viral molecules in the cytoplasm. Activated RIG-I (and MDA5) exposes its caspase activation and recruitment domains (CARD), which bind to the mitochondrial antiviral signaling (MAVS) protein to activate tank binding kinase 1 (TBK1) and I-kappa-B kinase ε (IKKe), leading to activation of transcription factors interferon regulatory factor (IRF) 3 and 7 results to production of IFN1 and as well as NFkB (Figure 1). The activated RIG-I binds to MAVS, leading to its prion like aggregation, where receptor-interacting serine-threonine kinase 1 (RIP1) and Fas-associated death domain (FADD) are involved for the initiation of NFkB pathway. Activation and nuclear translocation of interferon regulatory factor (IRF) -3, -7 and NFkB which leads to expression of IFN1 and pro-inflammatory cytokines (IL-1, IL-6, TNF-α) (Figure 1).

Similar to SARS-CoV, SARS-CoV2 may alter ubiquitination and degradation of RIG-I/ MDA5 and also inhibits activation of MAVS. SARS-CoV2 may also inhibit the TNF receptor-associated factors (TRAF) 3 and 6 in response to TLR3, 7/8 ligation and NFkB signaling pathway. SARS-CoV2 can counteract IFN1 signaling through the inhibition of STAT family transcription factor phosphorylation (Figure 1). The suppression of innate immune mechanisms in infected epithelial cells, monocytes and macrophages allow SARS-CoV2 to proliferate without activating the innate anti-viral response mechanism (Figure 1). Infected cells undergo cell death and released virus particles trigger innate inflammatory mechanisms through their recognition by PRRs on innate immune cells which result activation and expression of pro-inflammatory cytokines (including IL-1β, IL-6, TNF-α, etc.). T lymphocytes play a crucial role in this anti-viral response, including CD4+ T cell derived cytokines, CD8+ T cell mediated cytotoxicity, and B cell activation resulting in antibody production (Figure 1).
The tissue injuries caused by the virus induce the production of pro-inflammatory cytokines, macrophages and granulocytes. Numerous studies have described abnormal levels of the cytokines and chemokines (IL-1, IL-2, IL-4, IL-6, IL-7, IL-10, IL-12, IL-13, IL-17, M-CSF, G-CSF, GM-CSF, IP-10, IFN-γ, MCP-1, MIP-1-α, hepatocyte growth factor (HGF), TNF-α, and vascular endothelial growth factor (VEGF), CXCL10, CXCL8, CXCL9, CCL2, CCL3, CCL5) present in SARS-CoV2 infected patients. These results in the ‘cytokine storm’ termed as ‘macrophage activation syndrome’ or ‘secondary hemophagocytic lymphohistiocytosis (HLH)’ which leads to further tissue damage and multi-organ failure, including ADRS (Figure 1B). At this juncture overreaction from the immune system can endanger a patient's life. Chemokines, opsonize professional microbe like neutrophils at the site of infections. Higher IL-1α and IL 4 levels in severe SARS-CoV2 patients are strongly associated with lung injury.

Cytokines such as interleukin 1b, interleukin 6 and tumor necrosis factor steer neutrophils and other immunocytes from blood capillaries to the site of infection. This recruited monocytes, macrophages and neutrophils to the site of infection, which exhibited strong and poorly controlled inflammatory responses, resulting the tissue damage and systemic inflammation and contribute to morbidity. A number of systemic inflammatory conditions and uncontrolled activation of immune responses is not limited to the innate mechanisms. As a result of pro-inflammatory cytokine expression and the presence of nuclear antigens, adaptive immune cells become activated and trigger a “second wave” of inflammation. Finally, severely ill SARS-CoV2 infected patients are experiencing lymphopenia and atrophy of the lymph nodes and spleen. Primary and secondary forms of HLH and associated cytokine storm result in inflammatory cell death and hypo-cellularity of lymphatic organs. IL-6 in patients show accelerated inflammatory process, contributing to the cytokine storm and worsening the prognosis which is associated with cardiac damage in these patients.

Elevated liver enzymes and creatinine seen in some SARS-CoV2 infected patients without respiratory failure, suggesting that the inflammatory cytokine storm is the cause of damage to extra-pulmonary tissues and organs. These cytokines can augment heartbeat, elevate body temperature, trigger blood clots that trap the pathogen and modulate body temperature, fever, weight loss and other physiological responses that have evolved to kill the virus. However, a crucial role seems to be played by IL-6, whose increased levels in the serum have been correlated with respiratory failure, ARDS, and adverse clinical outcomes.

**TREATMENT WITH DRUGS**

The rapidity of pandemic SARS-CoV2 infection with concordant severe mortality rate gets going for an antidote or an authentic therapeutic strategy to stop this chain-infection. In 2003, the minimal severity of SARS-CoV and in 2012, the epidemic emergence of MERS-CoV had already led the scientific experts in generating vaccines which might be effective for SARS-CoV2 in the context of their sameness. Taxonomic parities with related virions and the symptomatic mimicries of SARS-CoV2 with other contagious diseases clued the clinicians for miscellaneous drug usage. Manifold methodologies of a particular type of drug or concoctions of drugs have been used by the medical front-liners, unintendedly. Amongst all extensively used drugs, those which have shown substantial results are enlisted here with plausible mechanism of actions.

**Molecular mechanism of viral RNA-dependent RNA polymerase inhibitors**

Remdesivir, Favipiravir and Ribavirin act as nucleotide analogs to inhibit the viral RNA-dependent RNA polymerase (RdRp) during viral replication through competitive inhibition and/or chain termination (Figure 2).
Remdesivir (RDV, GS-5734)

It is a novel antiviral nucleotide analogue developed by Gilead sciences, for the treatment of Ebola virus disease and Marburg virus infections. It has broad-spectrum in vitro activity against different RNA viruses such as Ebola virus, MERS-CoV, SARS-CoV, Nipah virus and Hendra virus. The anti-MERS-CoV activity of RDV is likely through premature termination of viral RNA transcription and has shown prophylactic and therapeutic efficacy in nonclinical models of these two coronaviruses.16

Experiments on enzyme kinetics revealed that whenever RDV, the adenosine tri-phosphate (ATP) analogue, was added into growing RNA chain, the reaction stopped after a certain time period as this inhibitor caused chain termination after adding three or more ribo-nucleotides into it.17 Quantitative analyses have determined earlier that the Michaelis Menten parameters (V_max/K_m) in this competitive inhibition by RDV, is highly significant to understand its efficiency of incorporation rate into viral genome, in comparison to competitor substrate ATP.17 The lastly added additional three nucleotides on growing chain following RDV help it from the exonucleolytic digestion (3´ to 5´) to which it is more sensitive.15 So, it could be a novel strategy for repurposing RDV against RNA virus like SARS-CoV2.

Favipiravir (FAV, T-705)

Favipiravir (6-fluoro-3-hydroxy-2-pyrazinecarboxamide, Fujifilm Toyama Chemical Co. Ltd, Tokyo, Japan) is selectively potent RdRp inhibitor against all serotypes and strains of influenza A, B and C virus and many other RNA viruses. Favipiravir performs by getting transformed into its most active form, Favipiravir-ribofuranosyl-5´-triphosphate (FRTP), after entering into the cell.18 A hypothesis on the functional property of FRTP explained that it might be mis-incorporated in a nascent growing RNA chain, otherwise the RdRp could be blocked (FRTP; IC50 of 0.341 μmol/l for influenza viruses) by binding to this guanosine triphosphate (GTP) analogue during viral RNA replication and transcription.18 Indeed, this binding site of FRTP on RdRpo is conserved in RNA viruses and it is unable to block human DNA polymerases (α, β and γ) up to IC50 of 1000μmol/l and slightly blocks human RNA polymerase II, at an IC50 of 905μmol/l.18

FRTP introduces lethal mutagenesis of various nucleoprotein clones by generating identifiable transition mutations (G to A and C to T or C to U) and transversions with almost no viable drug resistant mutants.18 Another line of evidence has shown that the single molecular addition of FRTP only can prevent the RNA chain extension and double incorporation leads to complete stoppage of any further extension.19 Thereafter, this mechanism of action might delineate favipiravir as a serious virucidal substance against SARS-CoV2.

Ribavirin (RBV)

Ribavirin (tribavirin) is used against respiratory syncytial virus, hepatitis C, lassa fever, Crimean-Congo hemorrhagic fever, Venezuelan hemorrhagic fever and Hanta virus. It is on the world health organization's list of essential medicines as the safest and most effective medicine for human health and was patented in 1971 and approved for medical use in 1986. Ribavirin, a guanosine analogue, inhibits ionosine monophosphate dehydrogenase (IMPDH) by getting into singly phosphorylated form i.e. ribavirin monophosphate (RMP) and then to triple phosphorylated state: RTP. RTP inhibits IMPDH driven GTP synthesis and that low levels of the cellular GTP, accelerates the incorporation of Ribavirin by viral RdRp, as a mutagenic GTP-analogue into the growing RNA. Scarcity in guanosine also may hamper with 5´-guanosine cap formation, and naturally this uncapped viral RNA is prone to degradation by RNase. Some studies indicated that the Th1 cell polarization is enhanced by Ribavirin during T-cell regulated anti-viral response.20 By inhibiting RdRp, Ribavirin may destabilize the nascent viral RNA and thus it might block the chain extension which needs further studies emphasizing on drug action.

Molecular mechanism of viral entry inhibitor/immunomodulator

Hydroxychloroquine (HCQ) and chloroquine (CQ); HCQ and CQ both are potent anti-malarial drugs and show immunomodulatory effects in autoimmune conditions (rheumatoid arthritis and systemic lupus erythematosus) by mediating anti-inflammatory responses. CQ was first developed in 1934 and HCQ in 1955.

Glycosylation inhibition might be a major ruse of CQ for blocking the pre-entry step of virus into host cell. Interplay between CQ and glycosyl-transferases may occur within human cells as CQ has shown to inhibit quinine reductase 2, a structural neighbor of UDP-N-acetyl-glucosamine 2-epimerases, involved in sialic acid biosynthesis. The sialic acids are present at the extremity of ACE2 (expressed in lung, heart, kidney and intestine).21 CQ derivatives by reducing N-terminal glycosylation on ACE2 receptor may change the property of ACE2’s ligand binding site for S1 spike protein on virus envelope and consequently the viral ingestion might become hindered (Figure 2).21

HCQ and CQ are accumulated in lysosomes by lysosomotropism and interfere in its enzymatic activity which rapidly onset the immunomodulatory or anti-inflammatory consequences (Figure 3). CQ derivatives briskly increase the organellar pH of lysosome just after its entry into the alveolar type II cells which lead to inhibition of hydrolytic enzymes and normal degradation of cargo, hindering MHC-II mediated presentation.22 Sudden rise in pH impairs processing of endosomal TLRs (TLR7, TLR8, TLR9). CQ and HCQ interfere with
TLR7-RNA interactions by directly binding to nucleic acid and can inhibit RNA-mediated TLR7 signaling.24 (Figure 3).25 Hereafter this hampered TLR signaling might be a cause of drug effectiveness in a COVID-19 patient.25 HCQ/HCQ blocks the binding of cytosolic RNA with cyclic-GMP-AMP synthase (cGAS) (Figure 3).26 Cytosolic cGAS-STING complex captures cytosolic RNA and then converts the transcription of IFN1 gene (IFNβ, IL-6, IL-12 etc.).26 SARS-CoV RNA is able to bind with cGAS and induces cGAS-STING pathway and SARS-CoV2 is robustly responsive to IFN1, thereafter, this CQ/HCQ might abrogate the pro-inflammatory cytokine release due to anti-viral response.21,25

Jakinibs (JAK-inhibitors) are the type of drug that performs in auto-immune state of immune response by blocking the JAK activation by trans-phosphorylation on heterodimeric interferon receptor (IFNAR) in virus infected cells (Figure 3).27 Baricitinib and ruxolitinib are clinically moderately used jakinibs in SARS-CoV2 mediated viral infection.28 In normal antiviral response of natural killer (NK) cells and macrophages, pro-inflammatory cytokines (IL-1, IL-6 etc.) are released and myeloid cells (including dendritic cells) overproduces IFN1s which lead to activation of regulatory T cells and B cells.29 These two drugs are known for their first-generation competitive inhibition reaction kinetics where they compete with ATP to bind at the JH1 Tyrosine kinase domain on JAK molecule in its active form.29 In downstream signaling of JAK-STAT pathway, the active phosphorylated dimer of STAT molecule regulates the IFN gene signature which promotes the IFN auto-inflammatory loop (Figure 3).29

**Molecular mechanism of different immune-suppressive drugs**

Jakinibs (Baricitinib and Ruxolitinib); baricitinib was first used in February, 2017, in European union as a second-line medication for moderate to severe rheumatoid arthritis in adults. Later on April, 2018, this drug got an approval from FDA advisory committee. It acts as an inhibitor of JAK1 and JAK2 janus kinase subtypes. Ruxolitinib also acts as an inhibitor of JAK1 and JAK2 and is used for the treatment of myelofibrosis and polycythemia vera. In November 2011, it was approved by USFDA for the usage in intermediate to high risk myelofibrosis.

The Jakinibs act by blocking the trans-phosphorylation of JAKs on IFNAR which leads to inactivation of STAT molecules. As a result the IFN auto-inflammatory loop becomes hampered.27 Hereafter, by dampening the pro-inflammatory cytokine synthesis jakinibs might block the anti-viral response in a COVID-19 patient.

**Tocilizumab (TCZ)**

Tocilizumab (atiluzumab) is a humanized monoclonal antibody against interleukin-6 receptor (IL-6R) and was developed by Hoffman–La Roche and Chugai. It is used as an immunosuppressive agent for treatment of mainly
rheumatoid arthritis and systemic juvenile idiopathic arthritis and later approved for Castleman’s disease, neumyelitis optica, giant cell arteritis, cytokine release syndrome like autoimmune disorders.

Tocilizumab has shown noticeable effectiveness against SARS-COV2.30 Strategy for this drug is to block the IL-6R from binding with IL-6, through competitive inhibition (Figure 3).30 Tocilizumab binds to soluble and membrane bound IL-6R and then blocks the cis and trans signaling by glycoprotein 130 (gp130) subunits.30 Hence, the SARS-CoV2 induced cytokine storm might be shattered to some extent, although there is not sufficient information on how this drug functions to block the infection.

Molecular mechanism of neuraminidase inhibitor

Viral neuraminidases are a category of enzyme found on the capsid surface of Influenza viruses. They cleave the terminal sialic acid moiety on host membrane glycoprotein to get escaped from the budding state in host cell to a free virion for further propagation.

Oseltamivir (OTV, GS-4104)

Oseltamivir (Tamiflu) is an orally administered antiviral drug used for the treatment of influenza A and B.

It is used as a prodrug (phosphate) of neuraminidase inhibitors against influenza like symptoms and after administration it is converted into active carboxylate form by hepatic esterases in host body.31 The active form binds to and inhibits the active site of the neuraminidases enzymes which exist on all influenza viruses and crucial for the release of the progeny virus from infected cells (Figure 2).31 In case of SARS-CoV2, this neuraminidase glycoprotein inhibitor might have subtle approach of inhibition towards virus S1 (spike) protein which has high sequence similarity with H1N1 neuraminidase segment and NS1/NS2 virulence factors.32 This could be the mechanism of oseltamivir in restricting SARS-CoV2 propagation within host.

Molecular mechanism of hemagglutinin inhibitor

Nitazoxanide (NZX)

Nitazoxanide or 2-(acetyloxy)-N-(5-nitro-2-thiazolyl) benzamide which was first synthesized in the early 1970s as an oral anti-parasitic agent, has been extensively commercialized in India as a broad-spectrum anti-parasitic drug for the treatment of intestinal infections by Cryptosporidium parvum and Giardia lamblia. It is also effective for anti-helminth, anti-protozoan, anti-viral and anti-cancer treatments. This drug is effective in blocking the maturation of viral hemagglutinin at its post-translational stage (Figure 2).33 SARS-CoV2 has a hemagglutinin esterase protein in its surface and this drug may block the packaging of this structural protein inside infected host cell. Therefore, with this defective hemagglutinin protein, the virus could not be able to get its infectious form back.

Molecular mechanism of nuclear transport inhibitor

Ivermectin (IVT)

Ivermectin is an anti-parasitic drug for head lice, scabies, river blindness, strongyloidiasis, trichuriasis, ascariasis and lymphatic filariasis. In vitro studies of Ivermectin on SARS-CoV2 infection have shown satisfactory evidences on blocking the enhanced antiviral responses.34 The molecular pathway of blocking relies upon Importin α/β1 mediated cytoplasmic transportation of molecular cargo into nucleus through nuclear pore complex (Figure 2).35,36 Ivermectin binds to Importin heterodimer and may prevent its signal-dependent nucleo-cytoplasmic shuttling of viral nucleocapsid protein cargo and simultaneously might block the host cell’s antiviral response.37

Molecular mechanism of viral protease inhibitors

Lopinavir (LPV)/ritonavir (RTV)

LPV-RTV is the most clinically favored antiretroviral, dual-drug regimen for SARS-CoV2 treatment, used all over the world and was first developed as an inhibitor of HIV-1 protease.38, 39 LPV is a novel protease inhibitor which is derived from RTV and is same in their mechanism of action.40,41 SARS-CoV2 main protease, 3CLPRO plays a major role in proteolytic cleavage of replicase polyproteins which is definite important in forming the core structure of Replication Transcription Complex (RTC) during viral replication (Figure 2).42,43 The molecular interactions like hydrogen bond, electrostatic and van der Waals interactions might occur in between the active site of SARS-CoV2 3CL conserved domain and the drug which may hamper the proteolysis of replicase polyproteins.42 Subsequently, the viral maturation may become arrested and the nascent virion might lose its infectivity.

OTHER TREATMENT STRATEGIES

Convalescent plasma therapy and monoclonal antibody therapy are other two prominent therapies against SARS-CoV-2. In the first case, the serum was separated from the collected blood sample of a person who had just got recovery from infection and the serum which contains antibodies was injected into a newly infected person to directly activate the immune response.44,45 It has been reported that this plasma transfusion might be beneficial to critically ill COVID19 patients.46 Existing risk factors of serum disease and antibody-dependent enhancement of infection, associated with sera transfusion enhanced the urgency of monoclonal antibody therapy.47 As SARS-CoV-2 enters into the host cell by using the ACE2
receptor for entry and the TMPRSS2 serine protease for S protein priming, an anti-S1 human monoclonal antibody has shown its competency against the virus. Research reports have declared that monoclonal antibodies, such as CR3022, 47D11 (human) can be used as potential therapeutics to stop the course of virus or to immunize an uninfected host that is supposed to get viral exposure.

CONCLUSION

On the cusp of 2020, we, the humankind start to grapple with a stronghold global force, namely COVID-19. This pandemic has emerged as a myriad challenge. Global pharmacopoeia with flimsy pretext of general anti-viral drugs exaggerates more insight into drug administration on COVID-19 patients. A groundswell of suspicion has developed regarding SARS-CoV2 pathogenicity, embodied with multiple organ failure. This all-hands-on-deck demands a streamlined technology to countermand this pandemic viral surge, immediately. Keeping abreast of diversified strategies for repurposing drug applications, here, we have reviewed some specific ones as per their worldwide consistent usage, for last few months since SARS-CoV2 emergence. These forefront drug therapies are merely scratching in unfolding the COVID19-enigma. Amongst all, remdesivir, favipiravir and tocilizumab are indefinitely close to combat the chain-infection. To date no such WHO-approved or comprehensible drug has been reported, although it is undeniable that different countries are “on track” of trials for forthcoming solution. Since the heady first days of SARS-CoV2, the Stepford experts are interweaving brew of drugs, technologies in search of an epitome one against it to hold up our indigenous territories of life on healthy mother earth.

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