N-Acetylcysteine as Adjuvant Therapy for COVID-19 – A Perspective on the Current State of the Evidence

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Abstract: The looming severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) has caused a long-lasting pandemic of coronavirus disease 2019 (COVID-19) around the globe with substantial morbidity and mortality. N-acetylcysteine, being a nutraceutical precursor of an important antioxidant glutathione, can perform several biological functions in mammals and microbes. It has consequently garnered a growing interest as a potential adjunctive therapy for coronavirus disease. Here, we review evidence concerning the effects of N-acetylcysteine in respiratory viral infections based on currently available in vitro, in vivo, and human clinical investigations. The repurposing of a known drug such as N-acetylcysteine may significantly hasten the deployment of a novel approach for COVID-19. Since the drug candidate has already been translated into the clinic for several decades, its established pharmacological properties and safety and side-effect profiles expedite preclinical and clinical assessment for the treatment of COVID-19. In vitro data have depicted that N-acetylcysteine increases antioxidant capacity, interferes with virus replication, and suppresses expression of pro-inflammatory cytokines in cells infected with influenza viruses or respiratory syncytial virus. Furthermore, findings from in vivo studies have displayed that, by virtue of immune modulation and anti-inflammatory mechanism, N-acetylcysteine reduces the mortality rate in influenza-infected mice animal models. The promising in vitro and in vivo results have prompted the initiation of human subject research for the treatment of COVID-19, including severe pneumonia and acute respiratory distress syndrome. Albeit some evidence of benefits has been observed in clinical outcomes of patients, precision nanoparticle design of N-acetylcysteine may allow for greater therapeutic efficacy.

Keywords: N-acetylcysteine, SARS-CoV-2; COVID-19, coronavirus, repurposing approved drugs, engineering nanoparticles, virus infected cells, respiratory viral diseases, antioxidant, glutathione, T lymphocytes, immune modulating activity, anti-inflammatory response, antiviral effect, clinical translation

Introduction

The acute respiratory disease COVID-19 caused by the novel coronavirus, SARS-CoV-2 (severe acute respiratory syndrome coronavirus 2), has spawned a global pandemic with untold morbidity and mortality, accompanied by devastating disruption to all facets of society, economy, and health care system.1,2 SARS-CoV-2 is a single-stranded, positive-sense RNA virus that was initially identified in Wuhan city in China in December 2019 from an outbreak of pneumonia cases in connection with Huanan Seafood Wholesale Market.3 It is closely related to other...
tremendously pathogenic beta-coronaviruses that have emanated in this century, namely severe acute respiratory syndrome coronavirus 1 (SARS-CoV-1) and Middle East respiratory syndrome coronavirus (MERS-CoV). Unlike SARS-CoV-1 and MERS-CoV that exhibit only limited human-to-human transmissions, a person being infected with SARS-CoV-2 who is just mildly ill or even asymptomatic can spread the disease to an average of two or three others, resulting in an exponential rate of increase in infection cases. SARS-CoV-2 virus has engendered 10-fold higher in the number of cases than the 2003 SARS epidemic in a quarter of the time. The rapid transmission of this highly pathogenic virus has warranted a pressing global need for the instantaneous development and deployment of therapeutic approaches and preventive measures against the disease.

Common hematological manifestations of COVID-19 infection include lymphocytopenia associated with intensification of the inflammatory process and direct infection of lymphocytes and destruction of lymphoid organs, increased ferritin levels owing to inflammation, and a higher rate of erythrocyte sedimentation in severe disease. Blood group A and males are more likely to become infected. Male sex, older age, and the presence of comorbidities are correlated with increased risk of COVID-19-related mortality. The development of thrombocytopenia may occur in severe disease due to reduced platelet production and increased destruction or consumption of platelets. Elevated D-dimer levels concomitant with high levels of fibrin degradation products and low antithrombin activity render COVID-19 patients to be at risk of hypercoagulability and thrombotic complications. Neutrophil-to-lymphocyte ratio and plasma D-dimer concentrations are relatively easy to quantify and possess clinical value for disease prognosis. High levels of proinflammatory cytokines and chemokines, including interleukin (IL)-6, IL-2, IL-10, tumor necrosis factor alpha (TNF-α), and interferon gamma (IFN-γ) may cause multiorgan damage as well as cardiovascular complications. Cardiac arrhythmia has also been reported and is associated with a cytokine storm-triggered systemic hyperinflammatory state and immune response that may cause injury to cardiac monocytes, resulting in myocardial dysfunction and the ensuing development of arrhythmia. Similarly, infection of alveolar pneumocytes cells by SARS-CoV-2 virus triggers the initiation of systemic inflammation and elevated immunoreactivity that potentiate T-cell and macrophage activation infiltrating infected myocardial tissues, leading to cardiovascular damage and myocarditis. Additionally, atherosclerotic plaques can be destabilized by systemic inflammatory response which happens simultaneously with pro-inflammatory and pro-oxidative effects of SARS-CoV-2, thereby giving rise to acute coronary syndrome and ischemic heart disease among COVID-19 patients.

Previous research with coronaviruses using both in vitro and in vivo experimental designs has contributed to a valuable guiding foundation for elucidating therapeutic strategies for the treatment of COVID-19. On top of understanding the microbial pathogenesis and the molecular and cellular mechanisms of disease biology, activity of a therapeutic agent in a translational research, be it either in vitro or in vivo, is critical to proffer advancement to first-in-human clinical trials based on laboratory findings of pharmacology, toxicology, and immunology. For COVID-19 repurposing of an existing clinically approved drug, it is pivotal to demonstrate its antiviral, anti-inflammatory, and related effects against SARS-CoV-2 in cell-based systems in vitro. The assessment of drug potency may be impacted by the type of the virus (eg, full-length wildtype, reporter viruses, or sub-genomic replicons, etc.) and the cell culture utilized (eg, Vero E6, Huh7, FRhK, or human airway epithelial cells), thus, necessitating high quality and standardized cellular assays, or at least with robust and universally accepted control groups.

Recent randomized controlled trials depict that repurposed antiviral drugs such as remdesivir, lopinavir, and interferon beta-1α regimens have small or null effect on hospitalized patients with COVID-19, as determined by outcomes such as overall mortality, initiation of mechanical ventilation, and duration of hospitalization. Combination treatment of remdesivir with anti-inflammatory drug baricitinib is associated with shorter time to recovery and accelerated improvement in clinical status, notably among those receiving high-flow oxygen or non-invasive ventilation. Treatment with dexamethasone reduces 28-day mortality in patients who are receiving invasive mechanical ventilation or oxygen without invasive mechanical ventilation, however, no discernible benefit and the possibility of causing harm have been found among those who are not receiving respiratory support. Neutralizing antibody bamlanivimab results in fewer hospitalizations and a lower symptom burden. Safe and effective vaccines that can confer significant protection against COVID-19 infection in real-world settings encompass BNT162b2 [Pfizer-BioNTech], mRNA-1273
There is a growing body of evidence that highlights the intrinsic antimicrobial and antibiofilm activities in many respiratory pathogens, including Escherichia coli, Haemophilus influenzae, Haemophilus parainfluenzae, and Klebsiella pneumoniae. Pseudomonas aeruginosa, Staphylococcus aureus, Acinetobacter baumannii, and other Gram-negative bacteria such as Haemophilus influenzae, Pseudomonas aeruginosa, and Klebsiella pneumoniae. Pseudomonas aeruginosa, Staphylococcus aureus, Acinetobacter baumannii, and other Gram-negative bacteria such as Haemophilus influenzae, Pseudomonas aeruginosa, and Klebsiella pneumoniae.

The overwhelming impact of the COVID-19 crisis has driven the push for reimagining and repositioning of previously approved medical treatments for other indications to speed up the discovery and development of safe and efficacious agents to enlarge the alternatives for adjunctive treatment or prevention of progression into severe COVID-19 illness. From the clinical front, it is presently worrisome to have no effective antimicrobial agents to treat the infected individuals and, optimally, eliminate viral shedding and the ensuing transmission cascades.

N-acetylcysteine is a mucolytic drug which exhibits antioxidant and anti-inflammatory effects. The compound has been available in clinical practice for several decades to treat various medical conditions, including bronchitis, acute respiratory distress syndrome, paracetamol intoxication, chemotherapy-related toxicity, doxorubicin cardiotoxicity, heavy metal intoxication, ischemia-reperfusion cardiac injury, human immunodeficiency virus infection or acquired immunodeficiency syndrome, and neuropsychiatric disorders. N-acetylcysteine is also marketed as a dietary supplement that is suggested to possess antioxidant and hepatic-protecting effects. The antioxidant characteristic of N-acetylcysteine has been ascribable to its reactivity with •OH, CO₃₂⁻, •NO₂, and thiyl radicals, the ability to repair oxidative damaged key cellular molecules, and activity as a precursor for biosynthesis of glutathione. There is a growing body of evidence that highlights the intrinsic antimicrobial and antibiofilm activities in many respiratory pathogens, including Escherichia coli, Pseudomonas aeruginosa, Staphylococcus aureus, Acinetobacter baumannii, and Klebsiella pneumoniae. High concentrations of N-acetylcysteine do not carry the risk of adverse interactions with most commonly used antibiotics and can exert intrinsic antimicrobial activity against Haemophilus influenzae. Being a precursor for glutathione biosynthesis which is crucial to redox regulation and control, N-acetylcysteine is often prescribed as a mucolytic agent in conjunction with antibiotic treatment in respiratory tract infections to improve the outcomes of the course of therapy.

N-acetylcysteine has recently been suggested as an adjunctive therapy to the standard care for SARS-CoV-2 infection considering the favorable risk and benefit ratio and its effects on synthesizing glutathione, improving immune function, and modulating inflammatory response. It achieves the therapeutic effects through two main activities: 1) mucolytic action conferred by the free sulfhydryl group which reduces disulfide bonds in the cross-linked mucus glycoproteins matrix, thus decreasing the viscosity of mucus; 2) antioxidant activity attributable to a direct interaction with free radicals, an indirect effect as a precursor to cysteine which is required for glutathione biosynthesis, and a replenishment of thiol pools that is central to redox regulation and control. In light of these properties, we hypothesize that N-acetylcysteine plays a role in the treatment of COVID-19 infection by the following postulated mechanisms of action (Figure 1):

1. Envelope (E) and spike (S) proteins have a triple cysteine structural motif located directly after the E protein’s transmembrane domain (NH₂…L-Cys-A-Y-Cys-Cys-N…-COOH) and a similar motif located in the carboxy terminus of the S protein (NH₂…S-Cys-G-S-Cys-Cys-K…-COOH). The position, orientation, and composition of these two motifs may serve as a center for the structural link between the E and S proteins which is mediated by the formation of disulfide bonds between the corresponding cysteine residues. Previous studies have indicated that the entry of viral glycoprotein is affected by thiol-disulfide balance within the viral surface and the cell-surface of the host. Any perturbations in the thiol-disulfide interchange equilibrium would deter the entry of the virus into host cells. Cleavage of disulfide bridges by N-acetylcysteine disrupts the structural components of the interacting proteins, thereby impairing receptor binding affinity and infectivity.

2. N-acetylcysteine is a chemical reducer of disulfide bonds via its free sulfhydryl groups may interact with the extracellular disulfide bridges of angiotensin II receptor, alter its tertiary structure, and inhibit the binding of angiotensin II to its surface receptors (AT₁a receptors) with subsequent attenuation of signal transduction and cell action. The AT₁a receptors possess two sets of disulfide bridges at the extracellular domain of the receptors: C18-C274 and C101-C180. N-acetylcysteine can reduce the disulfide bonds in a dose-dependent manner, decreasing angiotensin II and increasing angiotensin 1–7 (a biologically active peptide exerting many opposing effects on the renin-angiotensin system). From the clinical front, it is presently worrisome to have no effective antimicrobial agents to treat the infected individuals and, optimally, eliminate viral shedding and the ensuing transmission cascades.

The overwhelming impact of the COVID-19 crisis has driven the push for reimagining and repositioning of previously approved medical treatments for other indications to speed up the discovery and development of safe and efficacious agents to enlarge the alternatives for adjunctive treatment or prevention of progression into severe COVID-19 illness.
actions to angiotensin II), thus protecting against lung inflammation and fibrosis.\(^{43}\)

3. The sulfhydryl group of N-acetylcysteine inhibits angiotensin converting enzyme, reducing production of angiotensin II.\(^{44}\) In human lungs, angiotensin converting enzyme is expressed in lower lungs on type I and II alveolar epithelial cells. Following infection, viral entry begins with the attachment of spike (S) protein expressed on the viral envelope to angiotensin converting enzyme on the alveolar surface. Hence, N-acetylcysteine may prevent viral entry by limiting viral protein angiotensin converting enzyme interaction and internalization of the receptor-ligand complex.\(^{45}\) It also protects against oxidative stress and prevents glycosylation of proteins which may confer protection against respiratory disease syndrome and lung failure.

4. The antioxidant effect of N-acetylcysteine ameliorates oxidative stress and inflammatory response in COVID-19.\(^{46}\) It amplifies the signaling functions of toll-like receptor 7 protein and mitochondrial antiviral-signaling protein for boosting type 1 interferon production.\(^{47}\) Type I interferon functions to induce expression of various interferon-stimulated genes that exert antiviral activities to host cells.\(^{48}\)

5. The receptor for advanced glycation end products (RAGE) and its ligands have a crucial role in the pathogenesis of COVID-19 pneumonia and acute respiratory distress syndrome as well as lung inflammation. Circulating levels of soluble RAGE
(sRAGE, a decoy receptor) are positively associated with acute respiratory distress syndrome severity and mortality risk, whereas reduction in circulating levels of sRAGE drop results in disease resolution. Advanced glycation end products are formed by a reaction of the dicarbonyl compounds methylglyoxal and glyoxal with amino acids in proteins during glycolysis. Methylglyoxal and methylglyoxal-derived AGE can further activate inflammatory cells by binding to RAGE. N-acetylcysteine induces endogenous glutathione and hydrogen sulfide synthesis, thus attenuating methylglyoxal-induced protein glycation and additional glycosylation events in SARS-CoV-2 which may then inhibit the virus’s infectivity and associated pathologies.

6. N-acetylcysteine inhibits NF-κB activation by suppressing TNF-induced IkB kinases, followed by impediment of proteasome-dependent degradation. This prevents translocation of NF-κB from cytoplasm to the nucleus and block expression of pro-inflammatory cytokines and chemokines which have been correlated with severity and lethality in various acute respiratory viral infections, including Influenza A H5N1, highly pathogenic H1N1, SARS-CoV, MERS-CoV, and SARS-CoV-2.

Experimental in vitro Models
In vitro studies show that N-acetylcysteine holds therapeutic promise in numerous viral respiratory tract infections.

Influenza Viruses
In H5N1-infected A549 cells, N-acetylcysteine was found to attenuate H5N1-induced cytopathogenic effects, apoptosis, and virus yields. In addition, it decreased pro-inflammatory cytokine levels of CXCL8, CXCL10, CCL5, and interleukin-6. It was posited that the antiviral and anti-inflammatory effects of N-acetylcysteine were mediated by the inhibition of the oxidative metabolic pathway including transcription factor NF-κB and mitogen activated protein kinase p38. In another study, N-acetylcysteine was demonstrated to have immune-modulating properties by increasing influenza virus-specific cytotoxic T-lymphocyte clones and interferon-γ generation.

In an in vitro A549 model infected with influenza (strains A and B), N-acetylcysteine was shown to inhibit expression and release of MUC5AC, IL6, and TNF-α, reduce intracellular hydrogen peroxide level, restore intracellular pool of total thiols, diminish NF-κB translocation to the cell nucleus, attenuate activation of mitogen activated protein kinase p38, and prevent replication of the viruses. N-acetylcysteine also prevented apoptosis in H1N1-infected cell lines.

In Vero cells (ATCC CCL-81) infected with porcine H1N1 strain, N-acetylcysteine was reported to exert a dose-dependent inhibition on virus replication. However, the inhibitory effect of N-acetylcysteine was significantly less than that observed with H5N1, suggesting the susceptibility is strain-dependent.

Respiratory Syncytial Virus
In in vitro modeling of RSV infection carried out in cultures of primary normal human bronchial epithelial cells, N-acetylcysteine significantly inhibited viral infection, re-established the number of beating cells, restored the virus-induced decrease in expression of β-tubulin and genes involved in ciliagenesis such as DNAI2 and FOXJ1, inhibited IL-13, MUC5AC, and GOB5 upregulation, inhibited ICAM1 expression, increased heme-oxygenase 1 expression which correlated with the restoration of cellular antioxidant capacity, and enhanced intracellular hydrogen peroxide and glutathione concentrations.

In another experiment, N-acetylcysteine significantly reduced MUC5AC expression in RSV-infected A549 cells. It also diminished induction of TNF-α and IL-6, decreased NF-κB nuclear translocation and p38 mitogen activated protein kinase phosphorylation, abolished intracellular hydrogen peroxide production, restored intracellular thiol levels, and reduced virus titre.

Likewise, RSV infection of A549 epithelial cell lines showed the inhibitory effects of N-acetylcysteine on virus-induced chemokine expression and DNA-binding activity of NF-κB subunits of p50 and p65.

Experimental in vivo Models
In vivo studies depict some beneficial effects of N-acetylcysteine in numerous viral respiratory tract infections.

Influenza Viruses
In mice following intranasal infection with a lethal dose of influenza A virus A/PR/8, the use of N-acetylcysteine
increased survival rate from 58% (ribavirin monotherapy) to 92% (N-acetylcysteine and ribavirin), suggesting the antioxidant’s role in improving host defense mechanism and preventing pathogenesis of lung inflammation.61,62 Further experiment depicted that survival rates of mice were 20% in the N-acetylcysteine group, 60% in the oseltamivir group, and 100% in combination treatment of N-acetylcysteine and oseltamivir.63 In these three models, the mice were infected with 2–3 LD50 of influenza virus and N-acetylcysteine was administered as a single daily dose of 1,000 mg/kg.61–63 N-acetylcysteine increased survival by contributing to host defense mechanisms or by direct antioxidant effect against oxidative stress associated with viral infection.63 On the contrary, mice infected intranasally with a deadly dose of porcine A/swine/Iowa/4 (H1N1) influenza strain showed no significant difference in percent survival and mean survival time between N-acetylcysteine and control groups, showing that different virus strains resulted in different susceptibilities to N-acetylcysteine.58 The porcine A/swine/Iowa/4 (H1N1) strain was found to be more resistant to N-acetylcysteine than the human strain A/PR/8 (H1N1) deployed in other animal models.61–63 The lack of treatment effect was possibly due to the virus phenotypic susceptibility to N-acetylcysteine and the viral pathotype.64

Findings from H9N2 swine influenza virus-infected murine model indicated that N-acetylcysteine ameliorated pulmonary edema and inflammatory response, reduced myeloperoxidase activity in the lungs, decreased total and differential cell counts, neutrophils, macrophages, IL-6, IL-1β, TNF-α, and CXCL-10 in the bronchoalveolar lavage fluid, and inhibited protein expression of TLR4 and its mRNA in the lungs.65

In an experiment of H3N2-infected mice, N-acetylcysteine decreased pulmonary responsiveness and increased immune cytokine expression of IFN-γ in comparison to saline controls, with no effect on virus titers and expression of IL-4, IL-13, and IL-12p40.66

**Clinical Trials**

Currently, there have been several clinical studies to measure the effects of N-acetylcysteine on viral respiratory tract infections. A randomized controlled trial demonstrated long-term N-acetylcysteine regimen resulted in significantly lower episodes of influenza and influenza-like illness, disease severity, and duration of confinement to bed due to respiratory diseases. Only 25% of the virus-infected subjects in the N-acetylcysteine arm developed symptoms compared to 79% receiving placebo.67 Benefits were also documented in patients diagnosed with community acquired pneumonia whose plasma levels of malondialdehyde (oxidative stress parameter) and TNF-α (inflammatory mediator) were decreased and total antioxidant capacity was increased significantly among those treated with N-acetylcysteine.68 Moreover, N-acetylcysteine resulted in a lower rate of ventilator-associated pneumonia compared with placebo. Patients treated with N-acetylcysteine had significantly shorter length of stay in the ICU, reduced time to hospital discharge, and a higher rate of complete recovery.69 A clinical case study revealed that combination therapy of oseltamivir and N-acetylcysteine improved clinical outcomes in a woman with H1N1 influenza pneumonia, with clearance of pulmonary infiltrates, decreased requirement for oxygen supplementation, and lower plasma level of C-reactive protein.70

Nonetheless, the majority of human studies utilizing N-acetylcysteine in COVID-19 infection have been small in scale, non-randomized, and lack appropriate comparison groups (Table 1). In a single-center, randomized, double-blind, placebo-controlled trial conducted at the Emergency Department of a hospital in Brazil, 140 patients with severe COVID-19 infection and oxyhemoglobin saturation of less than 94% or respiratory rate higher than 24 breaths/minute were randomized to receive N-acetylcysteine 21,000 mg (approximately 300 mg/kg) for 20 hours or dextrose 5% (placebo). Of the intention-to-treat population comprising 135 patients, 16 patients (24%) receiving placebo required endotracheal intubation and invasive mechanical ventilation compared to 14 patients (21%) in the N-acetylcysteine group ($P=0.675$). No differences were noted in the duration of mechanical ventilation, death rate, rate of ICU admission, duration of stay in the ICU, and hospital stay.71

A single-center, randomized, double-blind, placebo-controlled trial evaluating a combined metabolic cofactors supplementation encompassing L-serine, N-acetylcysteine, nicotinamide riboside, and L-carnitine tartrate in 309 adult patients with laboratory-confirmed COVID-19 infection reported that mean recovery time in the intervention group was significantly shorter compared to the placebo group (5.7 vs 9.2 days, $P<0.0001$). In the intervention group, serum alanine aminotransferase ($P=0.032$), lactate dehydrogenase ($P<0.0001$), and creatinine levels ($P<0.0001$) were significantly lower on Day 14 compared to Day 0. Moreover, patients receiving the metabolic
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<tr>
<td>de Alencar et al (2020), Brazil</td>
<td>Single center, double-blind, randomized, placebo-controlled trial, Phase 4</td>
<td>Patients aged 18 years or older diagnosed with severe COVID-19 (suspected or confirmed) with oxyhemoglobin saturation of less than 94% while breathing ambient air and respiratory rate higher than 24 breaths/min.</td>
<td>140</td>
<td>21 g of N-acetylcysteine (300 mg/kg) was administered intravenously in 2 divided doses: 14 g in the first 4 hours and 7 g in the next 16 hours (n=70).</td>
<td>Dextrose 5% in water was administered intravenously (n=70).</td>
<td>No adverse event was noted in patients who received N-acetylcysteine. All patients tolerated the drug and the volume well.</td>
<td>SARS-CoV-2 was confirmed in 63 (94.0%) in the control group and 65 (95.6%) in the intervention group. No significant differences were observed in rates of mortality (13.4% vs 13.2%), ICU admission (43.3% vs 47.1%), ICU stay (9 vs 8 days), hospital stay (11 vs 10 days), and invasive mechanical ventilation use (23.9% vs 20.6%) between the intervention and control groups.</td>
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<td>Puyo et al (2020), United States</td>
<td>Case study</td>
<td>A 54-year-old Caucasian male patient with hypertension, hyperlipidemia, and obesity, who was tested positive for COVID-19 11 days prior to admission.</td>
<td>1</td>
<td>Hydroxychloroquine 400 mg was given as a single oral dose and N-acetylcysteine was administered intravenously at 75 mg/kg over 4 hours, then 35 mg/kg over 16 hours, followed by 17 mg/kg over 24 hours on Day 2. Prophylactic anticoagulation was started with subcutaneous heparin 5,000 units every 8 hours. An additional 200 mg dose of hydroxychloroquine was given on Day 2.</td>
<td>No control arm.</td>
<td>The patient tolerated hydroxychloroquine and N-acetylcysteine well.</td>
<td>The combination therapy resulted in progressive clinical improvement and a significant decrease of inflammatory markers such as ferritin levels, C-reactive protein, and lactic acid. However, the patient developed pulmonary embolism and deep vein thrombosis. Following thrombolysis and heparinization, his clinical condition continued a positive trend until discharge.</td>
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<tr>
<td>Liu et al (2020), China</td>
<td>Case study</td>
<td>A 64-year-old Chinese male with an anastomotic fistula after radical treatment of esophageal cancer and right-side encapsulated pyopneumothorax was confirmed for COVID-19, presenting with dry cough and shortness of breath and dyspnea on the day of admission.</td>
<td>1</td>
<td>Repeated bedside bronchoscopy with 10–15 g/time of N-acetylcysteine nebulized inhalation solution lavage was given in combination with routine nebulization and sputum suction airway management.</td>
<td>No control arm.</td>
<td>No adverse event was reported.</td>
<td>The patient’s hypercapnia was significantly improved and disengaged from mechanical ventilation intermittently. He was discharged after 6.5 weeks of hospital stay.</td>
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<td>Ibrahim et al (2020), United States</td>
<td>Prospective observational study</td>
<td>A 44-year-old male with G6PD deficiency was confirmed for COVID-19, presenting with fever, cough, and shortness of breath 5 days prior to admission. In addition, nine other COVID-19 patients without G6PD deficiency requiring veno-venous extracorporeal membrane oxygenator.</td>
<td>10</td>
<td>In the patient with G6PD deficiency, 30 g of N-acetylcysteine was administered intravenously in three divided doses over 24 hours. It was then discontinued for 7 days. Intravenous N-acetylcysteine was re-started at 600 mg every 12 hours for 1 week and was subsequently withheld for 10 days. IV NAC was started again until discharge. For the other nine patients, intravenous N-acetylcysteine of dose ranged from 600 mg every 12 hours for 4–9 days or 20 g over 24 hours was given.</td>
<td>No control arm.</td>
<td>No adverse event was reported.</td>
<td>In a G6PD deficient patient, N-acetylcysteine elicited improvements in hemolysis indices (direct bilirubin), liver enzymes (ALT and AST), and inflammatory markers (C-reactive protein and ferritin), decreased dependence on respirator and veno-venous extracorporeal membrane oxygenator, and reduced neutrophil to lymphocyte ratio. In non-G6PD deficient patients, significant overall reductions in inflammatory markers (C-reactive protein and ferritin) were observed with the use of N-acetylcysteine.</td>
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<td>Altay et al (2020), Turkey</td>
<td>Single center, double-blind, randomized, open-label, placebo-controlled trial, Phase 2</td>
<td>Adults aged 18 years or older with confirmed COVID-19 within 24 hours and had a stable clinical course</td>
<td>Hydroxychloroquine of an initial dose of 2×400 mg orally, followed by 400 mg/day (2x200 mg) for 5 days. Subsequently, a combined metabolic cofactors supplementation (CMCS) comprising L-carnitine tartrate 7.46 g/day, N-acetylcysteine 5.1 g/day, Nicotinamide riboside 2 g/day plus Serine 24.7 g/day was given orally twice a day for 14 days (n=71).</td>
<td>Only mild adverse events occurred in two CMCS-treated patients. Both patients experienced a mild rash on the upper body and they completed the study.</td>
<td>Patients treated with CMCS had significantly shorter time to full recovery compared with placebo (6.6 vs 9.3 days, P=0.0001). Furthermore, there was a significant reduction in liver enzymes (ALT and AST) and lactate dehydrogenase levels on day 14 in the CMCS group. There were no significant differences on serum levels of neutrophil, lymphocyte, white blood cell, and platelets between CMCS and placebo groups.</td>
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<td>Bhattacharya et al (2020), India</td>
<td>Single center, observational, retrospective cohort study</td>
<td>Adults aged 18 years or older with confirmed COVID-19 and were hospitalized</td>
<td>Patients received standard care plus Ivermectin single dose, Atorvastatin 10 mg daily, and intravenous N-acetylcysteine (n=2).</td>
<td>No adverse drug reaction occurred in any of the study subjects.</td>
<td>144 patients were discharged with a mean hospital stay of 12 days. The study concluded that triple combination of Ivermectin, Atorvastatin, and N-acetylcysteine had no case fatality rate and adverse effect, thus can be considered as adjunct treatment in coronavirus disease.</td>
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<td>Hernández et al (2020), Spain</td>
<td>Multicenter, retrospective, observational cohort study</td>
<td>Adults aged 18 years or older with positive SARS-CoV-2 diagnostic test, COVID-19 symptoms ≥5 days prior to diagnosis of SARS-CoV-2, past medical record of visits due to COVID-19 disease, or at least 1 month of follow-up of COVID-19 symptoms, whichever occurs earlier.</td>
<td>40</td>
<td>Patients received both standard care and ImmunoFormulation consisting of transfer factors (oligo- and polypeptides from porcine spleen, ultrafiltered at &lt;10 kDa – Imuno TF®) 100 mg, 800 mg anti-inflammatory natural blend (Uncaria tomentosa, Endopleura uchi and Haematococcus pluvialis - MiodesinTM), 60 mg zinc orotate, 48 mg selenium yeast (equivalent to 96 μg of Se), 20,000 IU cholecalciferol, 300 mg ascorbic acid, 480 mg ferulic acid, 90 mg resveratrol, 800 mg spirulina, 560 mg N-acetylcysteine, 610 mg glucosamine sulphate potassium chloride, and 400 mg maltodextrin-stabilized orthosilicic acid (equivalent to 6 mg of Si – SiliciuMax®) taken 3 times daily (n=20).</td>
<td>Patients received standard care only (n=20).</td>
<td>None of the patients on ImmunoFormulation experienced an adverse drug reaction.</td>
<td>90.0% of patients receiving ImmunoFormulation recovered compared to 47.4% in the control group (P=0.0057). Irrespective of disease severity, the ImmunoFormulation cohort showed a significantly shorter mean duration to recovery compared to controls (Severe: 16.0 vs 25.4 days, Moderate: 17.6 vs 28.0 days, Mild: 11.2 vs 28.0 days). In terms of symptomatology, the use of ImmunoFormulation led to significantly shorter duration of fever (2.3 vs 21.8 days), dry cough (4.4 vs 24.0 days), dyspnea (3.7 vs 20.0 days), headache (2.0 vs 26.5 days), diarrhea (5.3 vs 25.3 days), and generalized body weakness (1.9 vs 23.3 days).</td>
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cofactors supplementation showed a greater magnitude of decrease in plasma level of downregulated cytokines compared with placebo group on Day 14. The plasma levels of cytokines associated with inflammatory reactions (CSF-1, IL-15RA, IL18, MCP-1, and TNF) were significantly downregulated in the intervention group compared to the placebo group on Day 14. Only two patients in the intervention group (2.8%) reported a similar adverse event of mild rash on the upper part of the body.72

A case report of a 54-year-old male patient with pre-existing hypertension, hyperlipidemia, and obesity who was tested positive for COVID-19 11 days prior to admission to Holy Family Hospital in Massachusetts showed that combination therapy of hydroxychloroquine and N-acetylcysteine led to a progressive clinical improvement and a decrease in inflammatory markers for inflammation, for instance, C-reactive protein and lactic acid. The patient was mechanically ventilated for only 3 days, was then transferred from the ICU to the general ward on Day 7, and was finally discharged from the hospital on Day 12 with stable vital signs and normalized laboratory results.73

In another case report, a 64-year-old man presented with acute respiratory distress syndrome due to COVID-19 severe pneumonia and secondary bacterial lung infection was admitted to Taihe Hospital in Shiyan, China and was given repeated bedside bronchoscopy at intervals of 1 day or 2 days with a large dose of 10,000–15,000 mg/time of N-acetylcysteine nebulized inhalation solution lavage combined with routine nebulization and sputum suction airway management. The patient’s refractory hypercapnia was gradually improved and was cured and discharged after 46 days of hospitalization.74

Similar clinical benefit was noted in a case study of a 44-year-old man presented to the emergency department of NYU Langone, New York, with a previously diagnosed G6PD deficiency and a current COVID-19 infection with a redispersed risk of hemolysis. The use of intravenous N-acetylcysteine at 30,000 mg in three divided doses over 24 hours was associated with an immediate improvement in hemolysis indices (direct bilirubin levels), liver enzyme levels (alanine aminotransferase and aspartate aminotransferase), resolution of hemolysis as quantified by a sustained reduction in total and direct bilirubin and an elevated haptoglobin, and a sustained decrease in neutrophil to lymphocyte ratio which was an inflammatory predictor. Furthermore, patient’s oxygenation continued to improve and his veno-venous extracorporeal membrane oxygenator was discontinued after 2 weeks. It was observed that the combination of corticosteroids and intravenous N-acetylcysteine resulted in a marked reduction in inflammatory markers (C-reactive protein and ferritin) and the patient was finally discharged home after 41 days of hospital stay. Following these encouraging outcomes, a cohort of nine COVID-19 patients without G6PD deficiency were prescribed with intravenous N-acetylcysteine and demonstrated a significant overall reduction in inflammatory markers (C-reactive protein and ferritin), whereas a rebound inflammation was noted in six patients following discontinuation of the therapy. The median C-reactive protein level during IV N-acetylcysteine administration was 55 mg/dL which was considerably lower compared to the time before administration (143 mg/dL) or after N-acetylcysteine discontinuation (69 mg/dL). N-acetylcysteine was found to mitigate COVID-19-associated cytokine storm, elicit progressive clinical improvement, and facilitate hospital discharge readiness.75

A retrospective case series study in the inpatient department of Medical College Kolkata, India revealed that two out of 148 patients with pre-existing comorbidities and laboratory-confirmed COVID-19 infection were treated with triple combination therapy of Ivermectin, Atorvastatin, and N-acetylcysteine, and the regimen was found to be a useful adjunct to standard of care (oxygenation, restricted fluid therapy, anticoagulation, and corticosteroid). It had no apparent adverse effects and could potentially reduce mortality in patients with COVID-19.76

A multicenter, retrospective, cohort study in Clinic Bascoy and Clinica Arvila Magna, Spain assessed the effectiveness of an ImmunoFormulation containing N-acetylcysteine versus standard care in 40 patients with laboratory-confirmed COVID-19 infection. Ninety percent of patients in the ImmunoFormulation cohort recovered from the infection as compared to 47% in the control cohort (P=0.006). The mean recovery time for the ImmunoFormulation cohort was significantly shorter in comparison to the control cohort for patients with mild symptoms (11.22 vs 28.00 days), moderate symptoms (17.57 vs 28.00 days), and severe symptoms (16.00 vs 25.42 days). The duration of symptoms was also significantly shorter in the ImmunoFormulation cohort compared to the control group, notably for fever (2.25 vs 21.78 days), dry cough (4.38 vs 24.00 days), dyspnea (3.67 vs 20.00 days), headache (2.00 vs 26.50 days), diarrhea (5.25 vs 25.25 days), and weakness (1.92 vs 23.30 days) (all P<0.05). The use of ImmunoFormulation was associated with resolution of all clinical symptoms within 2–5 days,
except for the loss of taste or smell (19.73 days) that was recognized as a long-term complication of COVID-19. No patient experienced any adverse drug reactions. The ImmunoFormulation containing 560 mg N-acetylcysteine daily was an effective adjuvant therapy on evolution of symptomatology in COVID-19 patients.77

Notwithstanding the inconsistencies on the degree of efficacy of N-acetylcysteine as an adjunct in COVID-19 infection, all reported studies support the apparent safety profile of N-acetylcysteine irrespective of the dose and route of administration. There are many more clinical studies that are currently underway to generate the necessary evidence of its efficacy to inform future patient care and clinical action (Table 2).78–83

Discussion

There has been no in vitro or in vivo research that specifically examines the effect of N-acetylcysteine on COVID-19 infection. The potential use of N-acetylcysteine in COVID-19 is largely inferred from previous research on other viruses such as influenza and respiratory syncytial virus. Promising results have been noted in the ability of N-acetylcysteine to synthesize glutathione, improve T lymphocyte proliferative response, and modulate the inflammatory pathway. Glutathione precursors such as N-acetylcysteine are suggested as a potential therapeutic approach for blocking NF-κB activation and addressing inflammatory pathway. Glutathione precursors such as N-acetylcysteine are suggested as a potential therapeutic approach for blocking NF-κB activation and addressing cytokine storm syndrome and respiratory distress in patients suffering from COVID-19 pneumonia.84 In plasma, N-acetylcysteine reacts with cystine, reducing it to cysteine and yielding diacetylcysteine and N-acetylcysteine-cysteine via redox exchange reactions, and subsequently enters human erythrocytes and sustains glutathione synthesis.85 Additional research has illuminated the action of N-acetylcysteine on T-cell proliferation and IL-2 secretion which implies intracellular thiols regulate selective signaling pathways for a novel target of immunoregulation.86 Furthermore, in vitro treatment of murine T-cells with N-acetylcysteine induces reactive oxygen species scavenging and initiates NFAT expression and nuclear translocation. Addition of N-acetylcysteine to Cd4cre-Gclc T-cells increases Myc expression and CD98 production. Indeed, intrinsic Gclc expression in T-cells is necessary for antigen-specific immunity to virus infections. The antioxidative glutathione pathway is found to be central to metabolic integration and reprogramming in inflammatory responses mediated by T-cells.87 In this regard, high-dose N-acetylcysteine has been exemplified to replenish depleted pulmonary glutathione concentrations and yield concomitant favorable effects on lung function.88 The therapy also increases glutathione levels in peripheral blood T lymphocytes and disrupts the mammalian target of rapamycin activation in chronic inflammatory disease.89 Given that severe COVID-19 possesses shareable dominant risk factors with idiopathic pulmonary fibrosis, the effectiveness of N-acetylcysteine for slowing the rate of deterioration of vital capacity and single-breath lung diffusion capacity for carbon monoxide presents a rational option for treating patients infected with COVID-19.88,90 Therefore, it is conjectured that high dose N-acetylcysteine can enhance innate and adaptive immunity by elevating stores of glutathione levels in T lymphocytes, along with modulating immune-system responses to alleviate the degree of severity of COVID-19 infection and thus improve patient outcomes. Future in vitro studies to test candidate therapeutic compound in COVID-19 should deploy standardized assays, for example, Vero cells that can competently replicate and isolate the virus readily, whereas in vivo studies should utilize transgenic mice and Syrian hamsters, or cats and ferrets if opting for larger experimental animals.91

Of note, one of the important regulators of inflammation in COVID-19 is IL-6.92 Elevations in serum IL-6 concentrations have been identified under critical conditions such as sepsis, acute respiratory distress syndrome, and COVID-19.93,94 IL-6 is crucial for innate and adaptive immunity, efficient pathogen clearance, and physiological functions such as regulation of acute-phase response, hematopoiesis, lipid homeostasis, metabolism, and neural development.95 In essence, anti-inflammatory, pro-resolution, and antimicrobial activities of IL-6 are facilitated by classical signalling, whilst pro-inflammatory activities of IL-6 are facilitated by trans-signalling.96 Targeting the pathological effects of IL-6-mediated inflammation should avoid the unintentional concurrent abolition of its anti-inflammatory and pro-resolution functions.93 Blockade of IL-6 and the consequent reduction of downstream effects on inflammation and the innate immune response may have beneficial effects on clinical outcomes in patients with COVID-19, including those with acute hypoxemic respiratory failure.97 Contemporary in vitro and in vivo evidence indicating a potential effect of N-acetylcysteine in IL-6 inhibition may stimulate further research to understand how the drug affects disease outcomes and maximize its benefits with concomitant...
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<tr>
<th>Study, Year, Country</th>
<th>Study Design, Phase</th>
<th>Patient Characteristics</th>
<th>Number of Patients</th>
<th>Intervention</th>
<th>Control</th>
<th>Safety Outcome</th>
<th>Outcome Measures</th>
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<tr>
<td>Vardhana et al (2020), United States</td>
<td>Single center, non-randomized, open-label, parallel-group trial, Phase 2</td>
<td>Adults aged 18 years or older with documented COVID-19 infection.</td>
<td>84</td>
<td>Patients will receive N-acetylcysteine IV 6 g/day in addition to supportive and/or COVID-19 directed treatments at the discretion of the treating physician. Treatment interruptions for up to 48 hours are permissible if there is a clinical indication to hold the study drug. Patients can restart drug if they have been off drug for less than 48 hours.</td>
<td>No control arm.</td>
<td>-</td>
<td>Primary outcome measures include number of patients who are successfully extubated and/or transferred out of critical care due to clinical improvement and number of patients who are discharged from the hospital due to clinical improvement. The study is scheduled to complete in May 2022. ClinicalTrials.gov Identifier: NCT04374461.</td>
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<td>Lai-Becker et al (2020), United States</td>
<td>Multicenter, randomized, open-label, parallel-group, controlled trial, Phase 4</td>
<td>Adults aged 18 years or older with known or suspect COVID-19 disease.</td>
<td>200</td>
<td>Inpatients will receive N-acetylcysteine 25 mg/kg orally (rounded up to the nearest 600 mg) 4 hourly until discharge or N-acetylcysteine 1,200 mg twice daily for a week post-discharge. Outpatients will receive N-acetylcysteine 2,400 mg orally for a week, then 1,200 mg orally twice daily for 2 weeks.</td>
<td>Patients will not receive N-acetylcysteine.</td>
<td>-</td>
<td>Primary outcome measures include respiratory rate, hospital length of stay, need for mechanical ventilation, length of time intubated, outpatients on N-acetylcysteine needing admission to the hospital, and recovery disposition. The study is scheduled to complete in May 2021. ClinicalTrials.gov Identifier: NCT04374461.</td>
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<td>Alamdari et al (2020), Iran</td>
<td>Single center, randomized, parallel-group, controlled trial, Phase 1</td>
<td>Adults aged 18 to 90 years with confirmed COVID-19 disease, admitted to ICU, and required intubation and mechanical ventilation (PaO₂/FiO₂ &lt; 100–200).</td>
<td>20</td>
<td>Patients will be treated with mixture of Methylene blue, Vitamin C, N-acetylcysteine.</td>
<td>Patients will receive standard medical therapy (supportive therapy).</td>
<td>-</td>
<td>Primary outcome measures include mortality rate, improvement in PaO₂/FiO₂ ratio, duration of hospital stay, duration of ICU stay, need for vasopressor, days free of dialysis, C-reactive proteins, and white blood cell count. The study is scheduled to complete in September 2020. ClinicalTrials.gov Identifier: NCT04370288.</td>
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<td>Olagunju et al (2020), Nigeria</td>
<td>Multicenter, randomized, parallel-group, controlled trial, Phase 4</td>
<td>Adults aged 18 to 75 years with COVID-19 infection confirmed ≤2 days before randomization, currently hospitalized and requiring medical care and had a peripheral capillary oxygen saturation (SpO2) &lt;94% on room air at screening.</td>
<td>90</td>
<td>Patients will receive standard care and daily antioxidant supplement composed of two proprietary formulations that include reduced glutathione, N-acetylcysteine, superoxide dismutase, and bovine lactoferrin and immunoglobulins.</td>
<td>Patients will receive standard care only.</td>
<td>-</td>
<td>Primary outcome measures include time to clinical improvement and proportion of patients with SARS-CoV-2 polymerase chain reaction negative result at Day 14. The study is scheduled to complete in February 2021. ClinicalTrials.gov Identifier: NCT04466657.</td>
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<td>Alhawassi et al (2020), Saudi Arabia</td>
<td>Multicenter, randomized, parallel-group, double-blinded, placebo-controlled trial, Phase 3</td>
<td>Adults aged 18 years or older hospitalized with confirmed COVID-19 infection and were given oxygen supplementation.</td>
<td>1,180</td>
<td>Patients will receive N-acetylcysteine 150 mg/kg every 12 hours for 14 days orally or intravenously diluted in 200 mL diluent (Dextrose 5% in Normal Saline).</td>
<td>Patients will receive matching placebo administered in the same schedule and volume as N-acetylcysteine.</td>
<td>-</td>
<td>Primary outcome measure includes time to recovery. The study is scheduled to complete in August 2021. ClinicalTrials.gov Identifier: NCT04455243.</td>
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<td>O’Connell et al (2020), United States</td>
<td>Single center, randomized, open-label, parallel-group, controlled trial, Phase 1</td>
<td>Adults aged 18 years or older with confirmed COVID-19 infection who have not been admitted to the hospital prior to study enrollment.</td>
<td>42</td>
<td>Patients will receive oral N-acetylcysteine 600–1,800 mg 3-times daily alone or oral N-acetylcysteine 600–1,800 mg 3-times daily plus oral Famotidine 20–80 mg 3-times daily.</td>
<td>No control arm.</td>
<td>Primary outcome measure was number of participants with treatment-related adverse events.</td>
<td>Efficacy outcome measures include rate of hospitalization and time to symptom resolution. The study is scheduled to complete in August 2021. ClinicalTrials.gov Identifier: NCT04545008.</td>
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pharmacotherapy to selectively inhibit the trans-signalling pathway through which IL-6 signals.

Oral and intravenous N-acetylcysteine regimens are associated with adverse events such as diarrhea, nausea, vomiting, and mild-to-moderate hypersensitivity reactions such as urticaria, rash, angioedema, and wheezing.\textsuperscript{98,99} Such adverse events are dose-related and antihistamines are effective for treating and preventing recurrence of the anaphylactic reactions. Urticaria should be treated with antihistamine and N-acetylcysteine can be continued after symptom resolution. Angioedema and respiratory symptoms require the administration of antihistamine, corticosteroid, or symptomatic management.\textsuperscript{100} In such cases, N-acetylcysteine should be discontinued, and if necessary, it can be re-started 1 hour after the administration of antihistamine in the absence of signs and symptoms of anaphylaxis.\textsuperscript{101,102} Patients with a history of atopy (asthma and allergy) are recommended to receive prophylactic antihistamine 15 minutes prior to N-acetylcysteine administration to eliminate the excess risk of adverse reactions.\textsuperscript{102,103}

At present, there are six registered clinical trials evaluating the potential therapeutic value of N-acetylcysteine against COVID-19.\textsuperscript{78–83} The findings will build on in vitro and early in vivo evidence for the use of N-acetylcysteine as an adjuvant therapy for COVID-19. Similar to clinical trials on other infectious respiratory illnesses, study endpoints should appropriately encompass changes in SARS-CoV-2 associated blood cytokine levels, change in peripheral-blood lymphocyte count and activation, improvement in clinical status, hospital length of stay, time to recovery, safety outcomes, and overall mortality. While SARS-CoV-2 variants appear to occur sporadically in different geographical locations across the globe, they do not culminate in more severe diseases than the ancestral strain, albeit altered virus virulence, pathogenesis, and transmissibility have been documented.\textsuperscript{104,105} Future trials in patients infected with mutated SARS-CoV-2 strains may help to confirm or refute the possibility of a treatment benefit with N-acetylcysteine.

It is noteworthy that molecular attraction forces between chitin found on the exoskeleton of mites on human skin (Demodecidae or Pyemotidae species) and lipids present on the viral envelope cause interactions between arthropod and coronavirus which play a major role in the transmission of SARS-CoV-2.\textsuperscript{106} Natural small molecules, namely cyclodextrins and phytosterols, may hinder viral lipid-dependent attachment to host cells and thus reduce infectivity of the virus.\textsuperscript{107} Nowadays, these natural compounds exist in various forms of dietary supplements.\textsuperscript{107} Their potential to exhibit antiviral effects invigorates additional research to create novel biomedical applications for the treatment and prevention of COVID-19. This also implies the possibility that N-acetylcysteine has a role in inhibiting arthropod-coronavirus interaction and producing antimicrobial effect on endosymbionts of Demodex folliculorum.\textsuperscript{108} Ivermectin is a broad-spectrum anti-parasitic drug which elicits anti-inflammatory and acarcidal actions against Demodex mites,\textsuperscript{109,110} in addition to anti-viral activity against a range of viruses.\textsuperscript{111} Possible mechanisms include downregulation of the expression of pro-inflammatory genes, including those of IL-8, TNF-α, and cathelicidin LL-37 and downregulation of the ACE-2 receptor and viral entry into the cells of the respiratory epithelium and olfactory bulb.\textsuperscript{112} As a free radical scavenger, N-acetylcysteine can prevent ivermectin-induced cell death due to reactive oxygen species generation,\textsuperscript{113} suggestive of potential benefits with their combined use. However, current clinical findings do not support the use of ivermectin for the treatment of COVID-19 as it does not significantly improve the time to resolution of symptoms.\textsuperscript{114}

Patients with COVID-19 pneumonia can present with blood coagulation abnormalities, commonly manifested by elevated levels of fibrinogen and D-dimer in tandem with mild thrombocytopenia.\textsuperscript{115,116} Rise in D-dimer levels has been linked to poorer prognosis and higher mortality rate.\textsuperscript{117,118} D-dimer levels, lung inflammation, and pulmonary hemorrhage are influenced by neutrophil elastase activity.\textsuperscript{119,120} As such, suppression of elastase and neutrophil activation may be helpful in hemorrhagic or thrombotic complications associated with COVID-19.\textsuperscript{121} N-acetylcysteine has been found to inhibit release of elastase and modulate neutrophil activity when used at high concentrations.\textsuperscript{122} In neutrophilic airway inflammation in cystic fibrosis, high-dose N-acetylcysteine decreases the neutrophil burden in airways and the number of airway neutrophils actively releasing elastase-rich granules.\textsuperscript{123} Treatment with N-acetylcysteine can also attenuate lung emphysema induced by elastase as depicted in amelioration of airspace enlargements, partial recovery of expiratory flows, and normalization of lung collagen content.\textsuperscript{124} This evidence sheds light on the possible role of N-acetylcysteine on mediating inflammation-mediated lung injury and abnormalities of blood coagulation in severe COVID-19.
Novel formulations have recently been discovered to overcome the low bioavailability and short plasma half-life of N-acetylcysteine.\textsuperscript{125} Loading of N-acetylcysteine into lipid-based and polymer-based nanoparticles can improve muco-penetrating properties in pulmonary and oral delivery and hence the drug efficacy.\textsuperscript{126,127} A detailed in vivo and in vitro investigation is warranted to derive their applicability and efficacy in clinical settings.

As the most commonly approved nanomedicines by the US FDA,\textsuperscript{128} lipid-based nanoparticles afford many advantages such as relatively simple formulation, self-assembled structure, biocompatibility, drug load capacity, and adjustable physicochemical properties to suit biological characteristics.\textsuperscript{129} However, the lipid nanoparticle system is limited by low physicochemical stability and low-to-moderate encapsulation efficiency for N-acetylcysteine.\textsuperscript{130} A liposomal formulation of co-encapsulated azithromycin and N-acetylcysteine has been demonstrated to have a synergistic effect against bacterial strains.\textsuperscript{131} The high cellular membrane permeability and targeting accuracy associated with such a delivery system illuminates a potential implication for the co-formulation strategy of N-acetylcysteine with antiviral drug.

Recent evidence shows that polymer-based nanoparticles like N-acetylcysteine-loaded poly(lactic-co-glycolic acid) have desirable efficacy in preventing acute lung injury by mitigating the effects of reactive oxygen species and inflammation. The formulation delivers N-acetylcysteine directly to the lungs with increased pulmonary deposition and higher pulmonary concentrations with a lower dose of N-acetylcysteine.\textsuperscript{132} Polymer-based nanoparticle enhances pharmacokinetic and pharmacodynamics properties of encapsulated drug via prolonged drug release and particle retention.\textsuperscript{133,134} While poly(lactic-co-glycolic acid) is one of the most commonly used biodegradable polymers, its poor drug loading requires higher polymer load to facilitate dose delivery.\textsuperscript{135} Polymer-based nanoparticles are biodegradable, water soluble, biocompatible, biomimetic, and stable during storage. The surfaces can be easily modified for additional targeting. Disadvantages comprise higher risk of particle aggregation and toxicity.\textsuperscript{127}

To date, N-acetylcysteine has been used intranasally for the treatment of non-allergic chronic rhinitis with goblet cell metaplasia in which significant reduction in neutrophils, lymphocytes, goblet cells, bacterial count, turbinate hypertrophy, nasal symptoms, and rhinorrhea have been observed. The benefits are attributed to N-acetylcysteine’s mucolytic activity, the ability to restore surface ciliary activity at the nasal epithelium, and modulation of inflammatory response which is central to the immune defence mechanism of nasal mucosa.\textsuperscript{136} Likewise, acute recurrent rhinosinusitis treated with nasal douche consisting of flunisolide (corticosteroid) and N-acetylcysteine has been associated with a less stuffy nose, reduction in severity of symptoms and rhinosinus signs as assessed by endoscopy, improved cytoplasmic grading of neutrophil and eosinophil cells, less exacerbations, and increased mucociliary motility.\textsuperscript{137} In addition, topical intranasal drug combining tiaminoheptane sulphate (vasoconstrictor) and N-acetylcysteine exhibits a rapid decongestant effect with a significant decrease of resistance and increase of inspiratory flow.\textsuperscript{138} Topical delivery of N-acetylcysteine in the nostrils has also been investigated in ragweed sensitive patients and discovered that late phase allergic response mediated nasal symptoms can be reduced.\textsuperscript{139} In vivo studies have demonstrated nasal application of N-acetylcysteine in rats can reduce goblet cell loss and inflammation as well as promote wound healing of nasal mucosa,\textsuperscript{140} and combination of N-acetylcysteine and non-ionic surfactant (polyoxyethylene (C25) lauryl ether) can improve nasal bioavailability as a result of mucolytic activity of N-acetylcysteine in decreasing mucus viscosity.\textsuperscript{141} Accruing evidence suggests nasal route drug administration may be a potentially attractive strategy for N-acetylcysteine on top of the highly vascularized nasal cavity and large surface area for drug absorption.\textsuperscript{142}

Conclusions
The COVID-19 pandemic has highlighted the critical need for new drugs to complement existing therapies. In view of the widespread recognition of the safety and efficacy of N-acetylcysteine in numerous diseases over several decades, the aroused scientific interest has prompted the evaluation of its efficacy in COVID-19 clinical trials. As such, it opens a window for drug discovery with additional advantages, comprising the known pharmacological and human safety profiles. Advances in nanoparticle design are foreseen to have an impact on optimizing drug delivery and targeted activity of N-acetylcysteine in coronavirus disease. Currently, there is some evidence supporting the use of N-acetylcysteine as an adjunctive therapy for COVID-19. Further studies are warranted to design a formulation with
increased bioavailability or target-oriented delivery, ascertain optimal dosage and route of administration, and conduct well-controlled, adequately powered, randomized clinical trials to determine the safety and efficacy of N-acetylcysteine in patients with COVID-19.

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**References**


