Humidified Warmed CO₂ Treatment Therapy Strategies Can Save Lives With Mitigation and Suppression of SARS-CoV-2 Infection: An Evidence Review

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The coronavirus disease (COVID-19) outbreak has presented enormous challenges for healthcare, societal, and economic systems worldwide. There is an urgent global need for a universal vaccine to cover all SARS-CoV-2 mutant strains to stop the current COVID-19 pandemic and the threat of an inevitable second wave of coronavirus. Carbon dioxide is safe and superior antimicrobial, which suggests it should be effective against coronaviruses and mutants thereof. Depending on the therapeutic regime, CO₂ could also ameliorate other COVID-19 symptoms as it has also been reported to have antioxidant, anti-inflammation, anti-cytokine effects, and to stimulate the human immune system. Moreover, CO₂ has beneficial effects on respiratory physiology, cardiovascular health, and human nervous systems. This article reviews the rationale of early treatment by inhaling safe doses of warmed humidified CO₂ gas, either alone or as a carrier gas to deliver other inhaled drugs may help save lives by suppressing SARS-CoV-2 infections and excessive inflammatory responses. We suggest testing this somewhat counter-intuitive, but low tech and safe intervention for its suitability as a preventive measure and treatment against COVID-19. Overall, development and evaluation of this therapy now may provide a safe and economical tool for use not only during the current pandemic but also for any future outbreaks of respiratory diseases and related conditions.

Keywords: anti-COVID-19, antiviral, anti-cytokine storm, improve COVID-19 symptoms, carrier gas composition, enhancer antiviral, protect and improve organs function, suppression COVID-19 pandemic

BACKGROUND

The coronavirus disease (COVID-19) outbreak has presented enormous challenges for healthcare systems worldwide and caused terrible societal and economic impacts. There is also an urgent need to address health inequality in treating the current COVID-19 pandemic. Even now, scientists are racing to unravel sometimes conflicting information to understand the source, diagnose, and find effective treatments for SARS-CoV-2, and to conduct clinical trials of antiviral drugs and vaccines. Other COVID-19 mysteries include the appearance of new symptoms, the relation of silent hypoxia and sudden deaths, spikes insignificant vessel blockages, and increased risks of clotting (1). The
Precise control of the unique properties and intervention parameters of warmed humidified CO₂ gas make it a promising anti-COVID-19 therapy for mitigation and suppression of SARS-CoV-2 infection.

The virus is now known to be able to target a wide variety of cells throughout the human body through ACE2 and TMPRSS2 receptors (2) and is believed to have caused a spike in a rare syndrome: “multi-system inflammatory state requiring intensive care” in children. Furthermore, the mode of transmission and the extent of environmental contamination is yet unknown. While the virus may not technically be airborne, it is definitely borne in the air as aerosols (3).

One of the most critical unanswered questions is why some COVID-19 patients develop severe disease, while others do not? Does the answer hidden in the origin and continuing evolution of SARS-CoV-2 virus mutation into mild and wild different strains (4)? Alternatively, does the answer depend on the two phases of the individual human body immune responses; a protective phase and a damaging phase due to inflammation-cytokine storms (5)? Other questions include whether bacterial co-infections such as bacterial pneumonia and sepsis with antibiotic resistance lead to increased COVID-19 disease severity and mortality (6) and how long it will take to create an effective vaccine. Potential SARS-CoV-2 vaccines have a variety of approaches that depend on viral life cycles (7), and it is estimated that a vaccine will either arrive in 1 or 2 years or will never arrive at all. Even if the vaccine trials are successful, will the new vaccine cover all SARS-CoV-2 mutant strains, and give full immunity to everyone with no issues when translation to clinical practice? Can we produce enough, how much will it cost and who will pay (a particularly important issue in developing countries)? Can the new vaccine stop the threat of a second inevitable wave of coronavirus, or other pandemic viruses emerging to produce a similar situation in the future?

Gas therapy is a highly effective viral inactivation strategy. Carbon monoxide (CO) gas is very flammable and highly poisonous and referred to as the “Silent Killer,” because it binds to the parts of human blood that carry oxygen molecules, so it chemically blocks the body and organs from getting the needed oxygen. However, CO gas has also been shown to have antimicrobial and antiviral activities against infected cells (8), and two clinical trials (NCT02425579, NCT03799874) have demonstrated that the administration of low concentrations of CO is well-tolerated and safe in patients with sepsis-induced ARDS (9, 10). Similarly, while high concentrations of inhaled ozone (O₃) can damage the lungs, cause chest pain, coughing, shortness of breath, throat irritation, and worsen chronic respiratory diseases such as asthma as well as compromise the ability of the body to fight respiratory infections (11), ozone gas therapy has been demonstrated to inactivate airborne viruses (12) and could inactivate the SARS-CoV-2 virus through oxidizing the sulphydryl groups in cysteine of the virus-cell (13). There are also at least four ongoing clinical trials (NCT04290871 - NCT04306393 - NCT04305457 - NCT04290858) testing the use of inhaled nitric oxide (NO) gas for patients with COVID-19 (14), as increasing airway NO levels via gas inhalation or precursor molecules may improve oxygenation in COVID-19 subjects (15). As with the other gases, there is another side to NO, which can be harmful due to the formation of highly toxic and irritating nitrogen dioxide (NO₂) gas and methemoglobinemia (16).

THE HYPOTHESIS AND EVIDENCE

Carbon dioxide (CO₂) is a fundamental biological gas and has been used for medical purposes for over a century due to its unique properties (Figure 1). Carbon dioxide gas is natural, biocompatible, chemically stable, and safer than any other medical gases (NO, O₃, or CO). It has been shown to possess antioxidant and anti-inflammatory properties, to improve blood oxygenation and enhance oxygen delivery to organs, to protect and improve lung function, to function as a carrier, or enhancer gas for drug delivery by rapid and direct open airway inhalation with easy administration in home, GP, emergency unit, and
ICU settings. These unique biological, physical, and medical properties of \( \text{CO}_2 \) make it a promising anti-COVID-19 therapy for mitigation and suppression of SARS-CoV-2 infection. Our hypothesis depends on inhaling precise doses of humidified and warmed \( \text{CO}_2 \) medical gas, either alone or as a composite carrier gas with other COVID-19 inhaler medications (bronchodilators, antivirals, antibiotics, or anti-cytokine agents), to disinfect the SARS-CoV-2 virus inside the infected human lung, as a preventative measure to stop coronavirus infection spreading, and to improve the treatment of mild, moderate, and severe COVID-19 symptoms. The following benefits and evidence of using medical carbon dioxide gas support the hypothesis.

**Universal Virucidal and Antimicrobial Activity**

Direct inactivation technologies have several limitations against the current virus. Moist, warm \( \text{CO}_2 \) gas could become a competitive disinfection technology. Carbon dioxide gas is an antiviral, antibacterial, and anti-infection agent effective not only on solid surfaces but also in aqueous solutions and water treatment settings (17). Heated, un-pressurized carbon dioxide bubbled through wastewater or aqueous media effectively destroys both waterborne bacteria and viruses (18). Moreover, supercritical \( \text{CO}_2 \) can in-activate and eliminate coronaviruses from an animal, human tissues and solid surfaces (19–21). Supercritical \( \text{CO}_2 \) offers a novel, user-friendly process to sterilize acellular tissue, such as lung matrices, for use in tissue and organ engineering (22). \( \text{CO}_2 \) can also enhance the effect of some other antibacterial agents, further improving the protection imparted (17). When breathing is impaired, \( \text{CO}_2 \)-levels in the human body drop, which creates a favorable environment for bacterial growth and a higher risk of infection. Pure \( \text{CO}_2 \) significantly decreased the growth rate of most viruses and bacteria at body temperature; this inhibitory effect of \( \text{CO}_2 \) increased exponentially with time (23). This phenomenon could be attributed to unravels the secret of structure and function of the Endothelial Surface Layer (ESL) (24–27). As the venous ESL is probably comprised of nanobubbles of \( \text{CO}_2 \), generated from tissue metabolism, that presumably kills the viruses and bacteria exiting to the blood flow on the way to leaving via the lungs (27). Even though
the mechanism of inactivation of microorganisms by CO$_2$ is
not yet resolved, there are a number of hypotheses that have
been proposed to explain the unique disinfection action of CO$_2$
gas (28).

CO$_2$ gas is far superior to other similar gases, with much
higher viral inactivation rates at lower temperatures (18–100°C)
without the need for pressurization (18, 29). CO$_2$ interacts
with water moisture to generate carbonic acid (pH 4.18), a reduced
pH could affect virus and microbial cell inactivation, as lipid
membrane stability is disrupted and permeability to carbon
dioxide increases (30, 31). However, a reduction in the pH of
the medium is not sufficient to account for the antimicrobial
action of CO$_2$, since it shows a specific inhibitory effect which
is greater than that of the other acids used to lower the pH of
media (hydrochloric acid, phosphoric acid, etc.) (32). These acids
do not penetrate the microbial cells as easily as carbon dioxide
(33). Cheng et al. believe that CO$_2$ molecules could enter virus
capsids much more easily than H$^+$ and inactivate the virus (34).
CO$_2$–protein binding could also damage the capsid, inactivating
the virus. Both mechanisms may be active during dense phase
carbon dioxide treatment (DPCD) which has also been shown
to effectively inactivate viruses (31). The warm atmospheric
pressure CO$_2$ gas during DPCD is suggested to have high viral
inactivation effect by penetrating the virus capsid due to the
high density of CO$_2$ with a high interfacial area (α) produced by
the continuous CO$_2$-moist contact surface area (29). Following
this; CO$_2$ can bind inside the capsid proteins through acid/base
interactions (35), producing the high virus inactivation rates
(18). Also, when compared with other gases (Air, O$_2$, N$_2$, and
Argon), CO$_2$ gas has the highest inactivated viruses and bacteria
rates in different NaCl solutions, even at ambient temperatures
and normal atmospheric pressure (18). Recently, Edwards et al.
demonstrate the effectiveness of aerosol administration of nasal
saline comprising calcium and sodium salts diminishes exhaled
particles and acts as a new natural defense against airborne
pathogens in the human airways (36). Moreover, Zare and his
teamwork report that spraying micron-sized water droplets can
act as an effective disinfectant by causing inactivation of over 98%
of the bacteria. They propose that the combined action of reactive
oxygen species present in micron-size water droplets (but not in
bulk water) along with the droplet surface charge is responsible
for the observed bactericidal activity (37). The efficiency of CO$_2$
technology will require adjustment and control of the mechanical
and dynamic behavior of moist CO$_2$ bubbles and properties such
as temperature, flow and density rates, pressure, electrolyte pH,
bubble size and thickness, surfaces area, and duration. All of these
factors contribute to the observed fast microbial death (38).

Safe and Tolerance for Human Clinical Trials and Treatment
Carbon dioxide (CO$_2$) gas is natural, inexpensive, non-toxic at
low concentrations (5,000 ppm), non-flammable, and readily
available in high purity from a variety of sources. When CO$_2$ gas
dissolves in water, it exists in chemical equilibrium with carbonic
acid (pH = 4.18) which plays an essential role in the bicarbonate
buffer system used to maintain acid-base homeostasis in the
human body. The duration and concentration of carbon dioxide
inhalation may be the key to the effective and protective role of
CO$_2$ gas therapy. A recent study investigated that pre-treatment
by CO$_2$ inhalation for 10 min, but not for 60 min, could improve
lipopolysaccharide LPS-induced lung injury (39). A pre-clinical
sheep model used perflubron combined with 12% CO$_2$ to re-
open constricted airways treatment for severe acute asthma (40).
As a reference, OSHA has set a CO$_2$ permissible exposure limit
(PEL) of 5,000 ppm over 8 h and 30,000 ppm over 10 min. This
compares favorably to CO$_2$ gas at 50 ppm, NO gas at 25 ppm,
and O$_3$ gas at 0.10 ppm for 8 h. Humans can tolerate up to
10% CO$_2$ before severe adverse effects are encountered (41)
although CO$_2$ tolerance decreases with age (p < 0.0001) (42).
Two clinical trials (NCT02616770 & NCT02334553) showed that
perflubron carried in gas with ascending doses of carbon dioxide
(4, 8, and 12% CO$_2$) administered to healthy subjects was safe
and effective in subjects with mild asthma (43, 44), while other
ongoing clinical trials (NCT03903913) are testing the safety and
tolerability the same formulation in subjects with cystic fibrosis.
Moreover, CO$_2$ concentrations of up to 35% have been applied in
other clinical trial study used “CO$_2$ inhalation challenge model”
through a protected inhalation system to measure the anxiolytic
and panicylic effects of new test compounds (45, 46).

Suppressing Cytokine Storm
Evidence is accumulating inferring that a subcategory of
patients with acute COVID-19 might experience cytokine storm
syndrome (47). CO$_2$ gas is one of the potential treatment
strategies to dampen an overactive immune system and to
quell a cytokine storm (48, 49). Many researchers have
reported that the presence of CO$_2$ reduces the production
of proinflammatory cytokines such as tumor necrosis factor-
alpha (TNF-α) and interleukins 1 and 6 (IL-1 and IL-
6, respectively), suggesting that the gas temporarily inhibits
macrophage activity via a mechanism that could be associated
with the reduction of the local or systemic pH (50–54). Carbon
dioxide gas can also affect the production of pro-and
anti-inflammatory cytokines in endotoxin-stimulated human
whole blood cultures under hypercapnic, normocapnic, and
hypocapnic conditions (55). In another study, CO$_2$ was shown
to differentially affect the cytokine release of macrophage
subpopulations exclusively via alteration of extracellular pH.
Decreasing the extracellular pH to 6.5 mimicked the effects
of CO$_2$ and a decrease to 5.5 suppressed IL-6 release in cell
lines (53).

Inhaled Carrier Gas Delivery System
CO$_2$ gas has unique safety, chemical stability, biocompatibility,
and properties as well as a higher density than oxygen,
high solubility in tissue and blood and high tolerance in
vascular system (56). CO$_2$ itself is a respiratory stimuli,
enhances mucus clearance, and seems to be a bronchodilator
by general induction of smooth muscle relaxation (57).
Additionally, warmed and humidified CO$_2$ insufflation leads
to an improved body core temperature (BCT) maintenance,
a reduction of the inflammatory and cytokine responses (58,
59) and improved quality of postoperative course, compared
with standard insufflation (60, 61). Also, it can reduce intraoperative hypothermia, coagulation dysfunction, early postoperative cough pain, days to first flatus and solid food intake, and the length of hospital stays (62). In recent years, CO₂-based technologies have accordingly gained considerable interest in the pharmaceutical industry. CO₂ bubble-generating carrier systems can be used to locally accumulate a drug at diseased tissue, reducing side effects on the healthy tissue and improving their therapeutic effectiveness (63). CO₂ may also be used as an enhancer and carrier gas for delivery of effective medical agents into a surgical wound (64) or respiratory diseases such as severe acute asthma and cystic fibrosis (40, 43, 44).

**Clinical Usage and Medical Purposes**

Medical carbon dioxide has been used as a pure gas or in specialized mixtures with other gases in anesthesia, as an insufflation gas for minimally invasive surgery (65), and in carboxytherapy (66). It can be used in the expansion of blood vessels to increase carbon dioxide level after rapid breathing, and to stimulate breathing after a period of non-breathing (67). Transdermal carbon dioxide gas therapy is widespread and uses carbon dioxide gas at high humidity, to increase tissue blood flow. Tissue oxygenation generates new blood vessels, and well-oxygenated tissues improve the effectiveness of antibiotic therapy. This is complemented by the antioxidant effect of CO₂ itself, which reduces oxidative stress in open surgery (68), and improves wound healing (69).

**Benefits of Hypercapnic Therapy**

Hypercapnic therapy (elevated CO₂ levels) has beneficial effects on the physiology of the respiratory, cardiovascular, and nervous system. In human critical care, hypercapnic acidosis (HCA) is frequently acceptable and improves innate immune function, resistance to infection, and protects and improves lung functions in patients with advanced lung disease. However, all these benefits require careful consideration of when and for how long hypercapnia will be applied. Hypercapnic acidosis, but not buffered hypercapnia, was reported to reduce the severity of sepsis-induced lung injury (70). Recent studies suggest that HCA is protective in the earlier phases of bacterial pneumonia-induced sepsis, just as HCA is protective in preclinical models of early and prolonged systemic sepsis (71). Also, CO₂ gas in therapeutic hypercapnia and other forms of acidosis techniques is an excellent antioxidant and anti-inflammatory agent (72). Hypercapnic acidosis was associated with benefits on lung and distant organs in several disease models, apart from the reduction of ventilation parameters.
such as ventilator-induced lung injury (73), acute respiratory distress syndrome (ARDS) (74), ischemia-reperfusion injury (75) and sepsis (76), therapeutic hypercapnia through inspired carbon dioxide attenuated lung injury, as measured by gas exchange, reduced cytokine release, lung oedema formation, and histological lung injury. Hypercapnic acidosis improves ventilation-perfusion matching that also improves gas exchange (77), prevents oedema formation (78), clears the alveolar fluid in pulmonary oedema (79), maintains the integrity of the blood-brain-barrier and reduces neurologic deficits after trauma (80). HCA also reduces the oxidative stress that contributes to pathologic thick mucus gel formation in the lung (81, 82). It is hoped that hypercapnia therapy may offer real benefits, but well-planned and executed clinical studies will be required.

Recent COVID-19 Contradictory Studies

The partial pressure of CO$_2$ in the atmosphere varies between 0.03 and 0.06% (83) but forms a high proportion (12.5–13.5%) with water vapor (1.3%) of the mainstream cigarette smoke (84). Recent studies have discovered the unusually low prevalence of current smoking was observed among hospitalized COVID-19 patients compared to the expected prevalence based on smoking prevalence in China. This preliminary analysis does not support the argument that current smoking is a risk factor for hospitalization for COVID-19, and might even suggest a protective role (85). Other cross-sectional studies in both COVID-19 out- and in-patients strongly suggests that daily smokers have a very much lower probability of developing symptomatic or severe SARS-CoV-2 infection as compared to the general population (86, 87). However, on the other hand, researchers at Baylor College of Medicine, the University of South Carolina and other institutions have identified tobacco smoking as a potential risk factor for infection of the COVID-19 virus, due to increasing the expression of ACE2, the receptor of SARS-CoV-2, in the lungs (88, 89). These two contradictory studies support our hypothesis of moist warm CO$_2$ gas resulted from cigarettes smoking could kill the SARS-CoV-2 viruses inside the infected lungs of smoker patients, and that leads to decreasing the infected COVID-19 patient from the smoker, not the nicotinic or other outcomes of mainstream cigarette.

TESTING THE HYPOTHESIS (A): PRECLINICAL STUDY AND INACTIVATION MECHANISMS

Herein, we recommend preclinical studies to optimize the relation between disinfection efficacy and toxicity level of warm humidified CO$_2$ gas while considering other related parameters to discover the possible mechanism of action of disinfection by CO$_2$ gas. The temperature inside healthy lungs is around 37°C, the pH is between 7.38 and 7.42, and the relative humidity ranges from 30 to 70%. It is essential to keep humidity stable as too high humidity provides optimal conditions for microbial growth, and low humidity and dry air can dry mucous membranes and make them more susceptible to infection (90). The SARS-CoV-2 virus is highly stable at 4°C, but it is very sensitive to heat. It is remarkably stable in a wide range of pH values (pH 3–10) at room temperature (22°C) (91, 92). However, the stability of SARS-CoV-2 under different environmental conditions of temperature, pressure, relative humidity, and pH with biological tissue and barriers require further investigation.

TESTING THE HYPOTHESIS (B): CLINICAL EVALUATION AND IMPLICATIONS

Whilst the properties and clinical applications of CO$_2$ have been known for many decades; parameters must be systematically studied before it can be used in a new clinical setting.

(I) Healthy, Non-symptomatic, Mild, and Moderate Care Levels

Optimizing the balance between disinfection efficacy and toxicity of humidified warmed CO$_2$ gas considering other parameters (temperature, relative humidity, pressure flow and density rates, electrolyte pH, bubble size and thickness, surfaces area, and duration) will be key. Different regimes will be needed to protect healthy and non-symptomatic patients and improve the condition of those suffering mild and moderate COVID-19 symptoms. Multiple-ascending dose studies in which subjects with mild to moderate COVID-19 will be enrolled [CO$_2$ max 14%, tolerance decreases with age ($p < 0.0001$)] (42). The suggested study could consist of a screening period, a run-in, dosing and evaluation periods, and a follow-up period. The dosing and evaluation period of the study could divide into three connected components. First, a dose-escalation study—This segment of the treatment period is designed to assess the safety and tolerability of escalating doses of medical CO$_2$ gas (2–4%) in a healthy volunteer (Figure 2), and (4, 8, 12, and 14%) in those with mild-moderate COVID-19 symptoms (Figure 3). Second, a daily dosing study - This segment of the treatment period is designed to assess the short term (5 days) safety and tolerability of 1–2 times daily administrations of a fixed dose of medical CO$_2$ gas in healthy volunteers, and 2–3 times daily administration of a fixed dose of medical CO$_2$ gas in patients with mild-moderate COVID-19. Third, a drug delivery study - This segment of the treatment period is designed to assess the safety, efficacy, enhancing, and tolerability of humidified warmed CO$_2$ gas (2–14%) composed with other inhaled medication such as an antiviral (Remdesivir or IFN-β SNG001), short-acting bronchodilator, antibiotic, anti-inflammation. The recommended clinical trial study may well-include placebo-control, humidified warmed CO$_2$ gas (2–14%), and humidified warmed CO$_2$ gas (2–14%) composed with other inhaled medication. Administration can be achieved through using simple comprised cartridge MDI puffer, portable nebulizer, or circularize II high-efficiency aerosol drug delivery system nebulizer in a negative pressure environment. Direct air/oxygen inhalation for a few minutes can be used to recover patients to
baseline carbon dioxide levels. A safety monitoring committee must also review the results from each cohort before deciding continuation of the study at the next prescribed dose level, based on consideration of the clinical significance of safety and tolerability parameters.

(II) Severe Care Level

The damage mechanisms of SARS-CoV-2 are still unclear, with severe COVID-19 cases complicated by high mortality rates due to compromised immune function and a high probability of antibiotic-resistant secondary infections. Most severe COVID-19 cases are associated with respiratory failure, with many already suffering from internal hypercapnia acidosis (with humidity levels near 100%) that disrupt not only cardiac and neurological functions but also immune system function by suppressing both innate and adaptive immune responses to viral and bacterial proliferation and infection (54, 93–96). This dysfunction of the immune system with increasing SARS-CoV-2 infection can lead to an overreaction of the immune system (cytokine storm), during which white blood cells are misdirected to attack and inflame even healthy tissue, leading to failure of the lungs, heart, liver, intestines, kidneys, and genitals (Multiple Organ Dysfunction Syndrome, MODS). This may, in turn, lead to the lungs shutting down (Acute Respiratory Distress Syndrome, ARDS), which makes absorption of oxygen difficult. Most deaths due to COVID-19 are due to respiratory failure. To save the lives of severely affected patients, we must first stop the causes of SARS-CoV-2 infection and prevent secondary infections. However, due to the absence of a specific COVID-19 antiviral treatment, most severe COVID-19 patients are admitted to the intensive care unit to fight the symptoms, aiming to lower the mortality rate through intensive monitoring and supportive organ function treatments by anti-cytokine medications with artificial blood purification system machines (97). Herein, we cautiously suggest that external artificial hypercapnia acidosis (warmed humidified CO$_2$ 15–25%) could be applied to disinfect and stabilize the lungs of SARS-CoV-2-infected patients and prevent secondary infections (Figure 4). However, it should only be considered for severely affected patients if they are already connected to life support and artificial blood purification through mechanical means, and a controlled gas mixture consisting of 25% CO$_2$ and 75% O$_2$ is delivered through a protected inhalation system while monitoring a wide range of physiological parameters, and administering supportive organ function treatments.

CONCLUSION AND EXPECTING OUTCOMES

There is an urgent global need for a universal vaccine to cover all SARS-CoV-2 mutant strains to stop the threat of an inevitable second wave of coronavirus. Currently, there are hundreds of clinical trials, but not yet any approved antiviral drugs specific for the treatment of COVID-19. The physical, biological, and medical properties of CO$_2$ gas suggest that humidified warmed CO$_2$ gas possesses multiple bioactivities and offer a new concept to SARS-CoV-2 viral disinfection and COVID-19 treatment. This inexpensive and broadly applicable therapy could lead to a massive reduction in the global number of infected, especially when used as a carrier for delivery of other inhaled drugs and creates new possibilities for mitigation and suppression of any COVID-19 second wave, or indeed any new future respiratory viral pandemic. In the future, more bioactive properties of CO$_2$ could be identified, and their mechanisms of action investigated. We believe well-designed clinical trials of CO$_2$ and its various bioactive properties are warranted to examine its efficacy against these diseases in human beings. It is hoped that...
this hypothesis will serve as a stimulus for further investigation into this issue.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/supplementary material, further inquiries can be directed to the corresponding author/s.

AUTHOR CONTRIBUTIONS

All authors contributed to the article and approved the submitted version. AE-B: conceptualization, methodology, writing—original draft, and writing—review and editing. EB: writing—review and editing. MG and KH: conceptualization and writing—review and editing.

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