

# **Long-term carbon dioxide toxicity and climate change: a major unapprehended risk for human health.**

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## **Abstract**

As atmospheric levels of carbon dioxide continue to escalate and drive climate change, the safe level for breathing is not clear. The toxicity of CO<sub>2</sub> has been defined for short-duration exposures at high concentrations but it is unknown what levels will compromise human health when individuals are perpetually exposed for their lifetime. There is now substantial evidence that permanent exposure, to CO<sub>2</sub> levels predicted in the future, will have significant effects on humans. Blood CO<sub>2</sub> levels in populations are increasing and unhealthy concentrations have been measured from people in common indoor environments where reduced thinking ability, altered brainwaves and health symptoms have been observed at levels of CO<sub>2</sub> above 600 ppm for relatively short-term exposures. Although humans and animals are able to deal with elevated levels of CO<sub>2</sub> in the short-term due to compensation mechanisms in the body, the eventual failure of these may have severe consequences in a perpetual environment of elevated CO<sub>2</sub>. Protein malfunctions in cells due to elevated CO<sub>2</sub> and associated low pH has the potential to cause threats to life including cancer, neurological disorders, lung disease, diabetes, etc. In particular excess CO<sub>2</sub> causes the overexpression of carbonic anhydrase, the enzyme that catalyses CO<sub>2</sub> in the body, causes calcification in the kidneys, arteries and tissues, along with other diseases and this may be an existential threat. Although there is very low awareness of this risk and its proximity is yet to be adequately defined, it is likely that human physical and mental health will be affected in the near-future with the progressive severity dependant on CO<sub>2</sub> emissions.

## 1. Introduction

An axiom of modern science, as quoted from TS Huxley, is “do not pretend that conclusions are certain which are not demonstrated or demonstrable”. Carbon dioxide is one of the most frequently overlooked of all toxic gases. Even to refer to CO<sub>2</sub> as a toxic gas is a surprise to many safety professionals (Henderson 2006). In indoor environments CO<sub>2</sub> concentration is often elevated relative to ambient outdoor levels due to the fact that the exhaled breath from humans contains high CO<sub>2</sub> (about 4%) and ventilation may not be adequate to prevent the resulting increase in CO<sub>2</sub>. Despite the possible adverse effects on health where many people occupy buildings or vehicles, there is very little awareness of this issue in the general community.

The average ambient concentration of CO<sub>2</sub> (in fresh air) has been rapidly increasing and is recently around 410 ppm (Scripps Institution of Oceanography 2020; Schmidt 2020) (Figure 1). This increase is due to humanity’s activities, largely resulting from the burning of fossil fuels (Eggleton 2013).

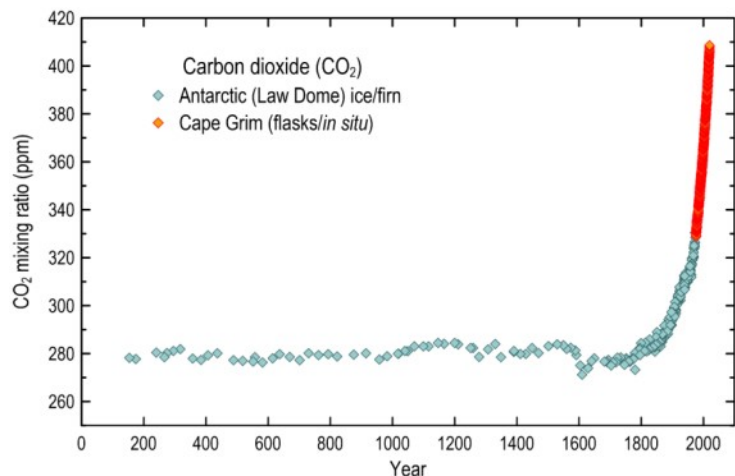


Figure 1. The atmospheric carbon dioxide concentrations (in ppm) over the last 2000 years, based on measurements of air trapped in Antarctic ice, shown in blue-grey diamonds, and the modern Cape Grim, Tasmania direct air measurements, shown in orange. (From Schmidt 2020).

Very early primate ancestors of humans were evolving around 23 million years ago. Throughout all of the ensuing period of human evolution levels of CO<sub>2</sub> in the ambient atmosphere remained relatively stable at below or close to 300 parts per million (ppm), this being derived from a combination of studies of relict features including air trapped in ice cores (Schmidt 2020), the composition of fossil plankton (Zachos et al. 2001) and Carbon-13 (<sup>13</sup>C) content in fossil plant material (Cui et al. 2020). Since about 1820, CO<sub>2</sub> levels have increased rapidly and are now above 415 ppm (Figure 1). This is a potentially catastrophic problem for many species of animals, including humans, for a number of reasons. The most well publicised issue is that of climate change. The mechanisms and history of global warming associated with CO<sub>2</sub> increase are well understood and the increase in atmospheric energy gradients will produce more extreme temperatures and weather events. To many people, climate change itself may not appear to be catastrophic – for example it might be possible to escape the effects of even a 5 degree C increase this century by moving to a cooler and safer geographic location. However, it is possible that humans have overlooked the more direct and immediate toxicity aspect of increasing atmospheric CO<sub>2</sub>. The earth’s atmosphere has

already reached CO<sub>2</sub> levels that are outside the range breathed by humans throughout their evolution and these new levels are relatively evenly distributed around the globe (see Figure 2). As well, in earlier pre-primate epochs, the effect of elevated atmospheric CO<sub>2</sub> causing physiological stress, has been postulated to be a cause of mass extinction events (Knoll et al. 1996).

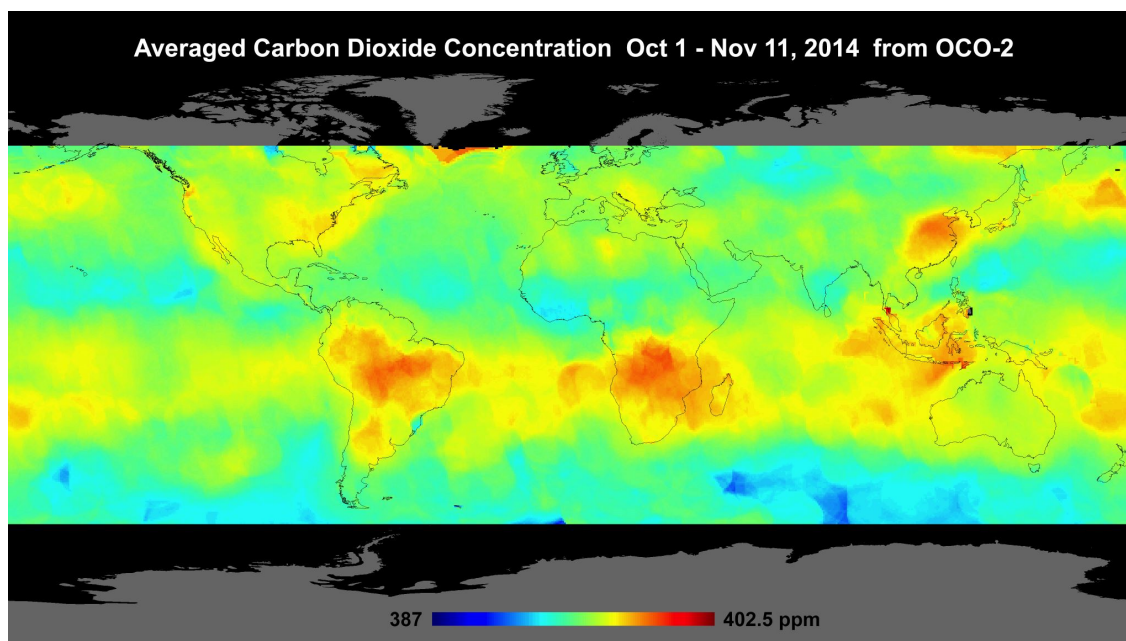


Figure 2. Global atmospheric carbon dioxide concentrations from Oct. 1 through Nov. 11, 2014 as recorded by NASA's Orbiting Carbon Observatory-2. Carbon dioxide concentrations are highest above northern Australia, southern Africa and eastern Brazil. Analysis of the African data showed that the high levels at the time were largely driven by the burning of savannas and forests. Elevated and more persistent carbon dioxide levels can also be seen above industrialized Northern Hemisphere regions in China, Europe and North America. Source: <https://www.nasa.gov/jpl/oco2/pia18934>.

We know that breathing CO<sub>2</sub> is toxic to humans when levels are high with numerous deaths reported based on occupational exposure (Scott et al. 2009). Although the CO<sub>2</sub> exposure limit for an 8-hour working day has been set at 5,000 ppm (OSHA 2012), this limit was decided in 1946 and based on relatively short-term observations of fit and healthy submariners (Scott et al. 2009). The safe level for lifetime exposure may be significantly lower than this and a number of researchers suggest there could be toxicity effects at CO<sub>2</sub> levels predicted in the near future with ongoing anthropogenic emissions (Portner et al. 2004; Robertson 2006; Ezraty et al. 2011; Antic 2012; McNeil and Sasse 2016; Karnauskas et al. 2020). So the question is: how long will it take, at present and future rates of increase, to reach levels that will impact on human health (no matter where you live) over a lifetime? To answer this question, the safe level of CO<sub>2</sub>, for continuous breathing in humans, needs to be determined. This paper is an attempt to evaluate available knowledge and to examine the likely and possible risks (for the near to medium-term future).

## 2. The role of carbon dioxide in breathing

Breathing is one part of physiological respiration and is required to sustain life (Raven et al. 2007). Aerobic organisms like birds, mammals, and reptiles, require oxygen to release energy by cellular respiration, through the metabolism of molecules such as glucose. During aerobic

respiration, glucose is broken down by oxygen to release energy, while carbon dioxide and water are the by-products of the reaction. Breathing delivers oxygen to where it is needed in the body and removes carbon dioxide thereby exchanging oxygen and carbon dioxide between the body and the environment. Carbon dioxide (CO<sub>2</sub>) is mostly a waste product and needs to be removed from our body. CO<sub>2</sub> from respiring tissues enters the blood plasma and diffuses into the red cells, where it is rapidly hydrated to H<sup>+</sup> and bicarbonate (HCO<sub>3</sub><sup>-</sup>) by the enzyme carbonic anhydrase (CA) (Arlot et al. 1985; Adeva-Andany et al. 2014). This enzyme enables the breakdown of CO<sub>2</sub> which returns to the plasma as bicarbonate and is then transported to the lungs (Adeva-Andany et al. 2014). When the bicarbonate reaches the lungs, CA in the alveoli catalyses the reverse reaction generating water and carbon dioxide which is exhaled as a gas. CA thus allows a large pool of otherwise slowly reacting plasma HCO<sub>3</sub><sup>-</sup> to be utilized in CO<sub>2</sub> excretion (Arlot et al. 1985).

There is an optimal range for the concentrations of CO<sub>2</sub> in the air we breathe. Too little can mean that breathing is too slow and not enough oxygen is brought into the body (Patton and Thibodeau 2009). Too much can compromise our ability to remove CO<sub>2</sub> (from our bodies) as a waste product. So what are the effects of too much CO<sub>2</sub> and what is the level that can cause health problems (in humans)?

### **3. Health effects from short term exposure to high levels of CO<sub>2</sub>**

Breathing too much CO<sub>2</sub> results in high levels of CO<sub>2</sub> in the blood (hypercapnia) associated with a decrease in blood pH (increased acidity) resulting in a condition known as acidosis. The decreases in blood and tissue pH produce effects on the respiratory, cardiovascular, and central nervous systems (CNS) (Eckenhoff and Longnecker 1995). Changes in pH act directly and indirectly on those systems producing effects such as tremor, headache, hyperventilation, visual impairment, and CNS impairment. In terms of worker safety, the US Occupational Safety and Health Administration has set a permissible exposure limit (PEL) for CO<sub>2</sub> of 5,000 parts per million (ppm) (or 0.5 %) over an 8-hour work day (OSHA 2012). They report that exposure to levels of CO<sub>2</sub> above this can cause problems with concentration, an increased heart rate, breathing issues, headaches and dizziness.

Exposures to 1-5 % CO<sub>2</sub> for short-term periods have been documented to produce symptoms on humans and animals such as dyspnea (shortness of breath), modified breathing, acidosis, tremor, intercostal pain, headaches, visual impairment, lung damage, increased blood pressure, bone degradation, reduced fertility, alterations to urine and blood chemistry as well as erratic behaviour (Halperin 2007; Rice 2004; Guais et al. 2011; Schaefer et al. 1963; Yang et al. 1997). These levels of CO<sub>2</sub> also induce panic attacks, interrupt the processes of metabolic enzymes and disrupt normal cell division processes (Colasanti et al. 2008; Guais et al. 2011; Abolhassani et al. 2009).

Health risks continue to escalate, with progressively higher CO<sub>2</sub> concentrations causing more severe reactions and faster responses. A value of 40,000 ppm is considered immediately dangerous to life and health given that a 30-minute exposure to 50,000 ppm produces intoxication, and concentrations around 70,000 ppm produce unconsciousness (NIOSH 1996). Additionally, acute toxicity data show the lethal concentration for CO<sub>2</sub> is 90,000 ppm (9%) for a 5-minute exposure.

There are indoor situations where exhaled human breath and restricted air flow can produce

extreme and dangerous levels of CO<sub>2</sub>. For example infant deaths have been associated with levels of up to 8% (80,000 ppm) CO<sub>2</sub> for an infant covered by blankets (Campbell et al. 1996).

#### **4. Physiological compensation for elevated CO<sub>2</sub>**

For understanding the long-term effects of breathing sustained elevated CO<sub>2</sub>, it is important to consider compensation mechanisms in the body, that regulate for increased CO<sub>2</sub> and acidity in the blood, and how these change over time with persistent exposure. The lowering of blood pH triggers various compensatory mechanisms, including pH buffering systems in the blood, increased breathing to reduce excess CO<sub>2</sub> in the bloodstream, increased excretion of acid by the kidneys to restore acid-base balance, and nervous system stimulation to counteract the direct effects of pH changes on heart contractility and vasodilation (widening of the blood vessels) (Burton 1978; Eckenhoff and Longnecker 1995). In respiratory acidosis, for a period the kidneys retain bicarbonate helping to normalise the pH of the blood as it passes through them. This occurs within 6 to 8 hours of exposure but achieves full effect only after a few days. With continued high levels of CO<sub>2</sub> in the blood, metabolic acidosis occurs and the kidneys do not respond in producing bicarbonate (Schaefer et al 1979a). After this the body uses the bones to help regulate the acid levels in the blood. Bicarbonate and a positive ion (Ca<sup>2+</sup>, K<sup>+</sup>, Na<sup>+</sup>) are exchanged for H<sup>+</sup>. The kidneys are involved in a wider array of physiological compensation responses to CO<sub>2</sub> induced pH imbalance (acidosis). The kidney tubule recovers filtered bicarbonate or secretes bicarbonate into the urine to help maintain acid-base balance in the blood and this again involves the CA enzyme (Adeva-Andany 2014).

#### **5. Health effects at common indoor CO<sub>2</sub> concentrations**

##### **5.1 Classrooms**

There is a large volume of recent literature that has documented the occurrence and levels of CO<sub>2</sub> in classrooms across the world including kindergartens, day-care centres, primary schools, high schools and universities (Bako-Biro et al 2011; Widory and Javoy 2003; Kukadia et al. 2005; Dijken et al. 2005; Branco et al. 2015; Heudorf et al. 2009; Santamouris et al. 2008; Ferreira and Cardoso 2014; Gaihre et al. 2014; Jurado, et al. 2014; Lee and Chang 2000; Muscatiello et al 2015; Carreiro-Martins et al. 2014). There is general agreement that the levels of CO<sub>2</sub> in 20-50% of classrooms commonly exceed 1,000 ppm and are often much higher, sometimes reaching levels as high as 6000 ppm for extended periods. A number of studies have identified CO<sub>2</sub> associated symptoms and respiratory diseases such as sneezing, rales, wheezing, rhinitis, and asthma (Carreiro-Martins et al. 2014; Ferreira and Cardoso 2014). Other symptoms; i.e. cough, headache, and irritation of mucous membranes, were also identified (Ferreira and Cardoso 2014). Lack of concentration was associated with CO<sub>2</sub> levels above 1000 ppm. Gaihre et al. (2014) found that CO<sub>2</sub> concentrations exceeding 1000 ppm is associated with reduced school attendance. Teachers also report neuro-physiologic symptoms (i.e., headache, fatigue, difficulty concentrating) at CO<sub>2</sub> levels greater than 1000 ppm (Muscatiello et al. 2015).

##### **5.2 Offices**

Offices have levels of CO<sub>2</sub> similar to classrooms depending on the number or density of workers and the types of ventilation systems (Lu et al. 2015; Tsai et al. 2012, Seppanen et al. 1999). These studies

have found strong evidence of the relationship between CO<sub>2</sub> levels in offices and Sick Building Syndrome (SBS) health effects such as headaches, dizziness, fatigue, respiratory tract symptoms, eye symptoms, nasal and mucous membrane symptoms (Seppanen et al. 1999; Lu et al. 2015; Tsai et al. 2012; Vehviläinen et al. 2016; MacNaughton et al. 2016). Seppanen et al. (1999) conducted a review of available literature and were careful to eliminate other confounding airborne building contaminants. The reviewed studies included over 30,000 human subjects, and they concluded that the risk of SBS symptoms decreased significantly with carbon dioxide concentrations below 800 ppm. Whether CO<sub>2</sub> itself is responsible for the health symptoms is still a subject of debate since historically it has been assumed, despite lack of direct evidence, that other airborne contaminants are the cause (Zhang et al. 2017).

More recently a number of studies have demonstrated that CO<sub>2</sub> has direct impacts on human physiology at levels commonly found in indoor environments (Azuma et al. 2018). Symptoms such as fatigue and drowsiness caused directly by CO<sub>2</sub> have been demonstrated by the use of electroencephalogram (EEG) techniques (Snow et al. 2018; Pang et al. 2020). In a study of office workers, a 20% increase in blood CO<sub>2</sub>, to significantly above normal levels, was measured along with sleepiness, headaches, heart rate variation and poor concentration in air that averaged 2,800 ppm CO<sub>2</sub> (Vehviläinen et al. 2016) while lowered heart rate and arousal level (fatigue) is clear at 4000 ppm (Xia et al. 2020). Measurements of end-tidal PCO<sub>2</sub> show a significant increase of CO<sub>2</sub> in human blood during tests conducted on astronauts after 4 months continually exposed to about 5,000 ppm (Hughson et al. 2016). Increases in blood CO<sub>2</sub> were associated with restricted lung function at levels between 2,000 and 3000 ppm CO<sub>2</sub> (Shriram et al. 2019). Zheutlin et al. 2014 used statistical data to determine an increasing trend in the average levels of CO<sub>2</sub> in the blood for a national sample of 5,000 people from 1999 to 2012. Heart rate variation at 2700 ppm is confirmed by Snow et al. (2019) for 10 minute exposure. MacNaughton et al. (2016) found that a 1,000 ppm increase in CO<sub>2</sub> from background levels was associated with a 2.3 bpm increase in heart rate after adjusting for potential confounders. Another older study (Goromosov 1968) reported harmful physiological effects on humans at only 1,000 ppm CO<sub>2</sub> with changes in respiration, circulation, and cerebral electrical activity. These physiological effects are being observed at much lower levels of CO<sub>2</sub> than previously anticipated (Azuma et al. 2018).

### 5.3 Vehicles

Although rarely studied for health effects, vehicles can often contain even higher levels of CO<sub>2</sub> particularly where there are multiple passengers for relatively long journey times. CO<sub>2</sub> levels can build up to 5,000 ppm after less than an hour of driving with two people in a car with only internal air (Gładyszewska-Fiedoruk 2011). With five people in a car with recirculated air levels of CO<sub>2</sub> can exceed 10,000 ppm (1%) after only 28 minutes, this being a level that is known to result in respiratory acidosis (Constantin et al. 2016). Buses with high numbers of passengers consistently reach average CO<sub>2</sub> concentrations of > 2500 ppm (Chiu et al 2015). Airliners can contain levels of around 2000 ppm for the duration of the flight (Gładyszewska-Fiedoruk 2012). Measurements on an Italian submarine showed a steady increase to 5000 ppm CO<sub>2</sub> after 2 hours of being submerged (Ferrari et al. 2005). Extremely high CO<sub>2</sub> concentrations (10,000-20,000 ppm) are commonly found inside motorcycle helmets in both stationary and moving situations (Bruhwiler et al. 2005).

#### 5.4 Cognitive impairment and anxiety in elevated CO<sub>2</sub> environments

There is now a significant body of research studies involving the detrimental effect of CO<sub>2</sub> on learning and cognitive abilities in humans at common indoor concentrations (Du et al. 2020; Lee et al. 2022). Testing of students has found that CO<sub>2</sub> can negatively affect attention, memory, concentration and learning ability impacting on academic performance (Bako-Biro et al. 2011; Coley et al. 2007). Several recent university studies of cognitive effects of CO<sub>2</sub> have been notable in their strong research design (Satish et al 2012; Allen et al 2016; Allen et al 2018; Scully et al 2019) with the testing environments injected with pure CO<sub>2</sub> meaning that the analysis of CO<sub>2</sub> effects was not confounded by the presence of other substances. These studies showed that low level CO<sub>2</sub> (between 950 ppm and 2500 ppm CO<sub>2</sub>) affected the cognitive abilities of students, information professionals and pilots in the indoor environment. Satish et al. (2012) tested only variations in CO<sub>2</sub> over periods of 2.5 hours of exposure. For seven of nine scales of decision-making performance (basic activity, applied activity, task orientation, initiative, information usage, breadth of approach, and basic strategy), performance was significantly impaired in a dose-response manner with higher CO<sub>2</sub> levels. For example, compared with mean raw scores at 600 ppm CO<sub>2</sub>, mean raw scores at 1,000 ppm CO<sub>2</sub> were 11–23% lower, and at 2,500 ppm CO<sub>2</sub> were 44–94% lower. As part of a larger study that included volatile organic compounds (VOCs), Allen et al. (2016) found that, after CO<sub>2</sub> was independently modified (from a baseline of 480–600 ppm) for individual 8-hour exposures, cognitive function scores were 15% lower at 950 ppm and 50% lower at 1400 ppm. This study used similar methodology to score cognitive function and the results largely repeated the findings of the earlier work (Satish et al 2012). However, one difference was that, at 1500 ppm CO<sub>2</sub>, even focussed activity was found to have declined (Allen et al 2016). In a study of pilots' performance, Allen et al. (2018) found that negative impacts on cognitive function were observed between 700 ppm and 1500 ppm CO<sub>2</sub>. Another study found similar negative effects on human cognitive abilities, in experiments involving 140-minute sessions, as well as increased fatigue at levels of 3000 ppm CO<sub>2</sub> compared with 600 ppm (Kajtar and Herczeg 2012). This study also measured some physiological parameters with heart rate analysis suggesting significantly increased mental effort at 3000–4000 ppm.

These studies are further supported by the finding of CO<sub>2</sub>-induced changes in human brainwaves, measured by electroencephalography (EEG), that have known associations with mental impairment (Lee et al. 2022). EEG combined with cognitive tests provided evidence of brainwave changes in frontal, parietal and occipital lobes that showed the negative impact of 1000 ppm and 2500 ppm CO<sub>2</sub> on working memory, mental workload and visual concentration after only 15 minutes of exposure. This is physiological evidence for the effect of CO<sub>2</sub> on cognitive decline.

Cognitive and neurological effects are also observed in animal studies. Mice exposed from birth to 1,000 ppm CO<sub>2</sub> for 38 days had decreased Insulin-like Growth Factor-1 (IGF-1) which resulted in greater anxiety and reduced cognitive function (Kiray 2014). Neurons were reduced in number and were malformed at this CO<sub>2</sub> level for several areas of the brain, with the largest effect for those areas associated with learning and memory. Mice exposed to only 890 ppm from preconception to 3 months of age showed increased corticosterone levels in the blood and brain (Wyrwoll et al 2022). Chronic elevation of brain corticoids associates with cognitive impairment and could harm the developing brain (Stolp 2022).

## 5.5 Sleep impacts

Sleep quality is reduced at 1900 ppm CO<sub>2</sub> when compared to 800 ppm in both subjective indices and electroencephalogram (EEG) measurements of brain activity (Xu et al. 2020). When CO<sub>2</sub> in arterial blood rises to a certain extent, it will produce carbon dioxide narcosis respiratory inhibition, weakened respiration, and hypoxia of human cells resulting in poor sleep quality (Xu et al. 2020).

Another example is people affected by sleep disordered breathing (SDB). Brillante et al (2012) found that the development of nocturnal hypercapnia in normal indoor CO<sub>2</sub> air concentration, quantitated by a large difference in carbon dioxide in the blood between morning and evening, predicted increased mortality in SDB patients. This is a result of the lack of efficacy of an individual's respiratory regulatory system in sleep for maintaining normal blood gas tensions (Brillante et al. 2012). As CO<sub>2</sub> levels in the atmosphere increase into the future, the impacts on and number of affected individuals will logically increase.

## 6. Health effects from long term exposure to elevated CO<sub>2</sub> levels below 1%

Where indoor levels of CO<sub>2</sub> are relatively high and affecting health, it is generally possible to obtain relief by going outdoors. However this may not be the case in a climate change future where ambient CO<sub>2</sub> is persistently high and effects of continuous long-term exposure must be considered. There have been very few studies related to long-term exposure at lower CO<sub>2</sub> levels, elevated above ambient, perhaps for logistical reasons since it is difficult to arrange an experiment for relevantly long timeframes. We are looking for information on the effect on humans of CO<sub>2</sub> levels at 1,000 ppm or less – noting that this is the level that some feasible models predict could be reached in the ambient atmosphere in less than 100 years (Smith and Woodward 2014). Given the lack of research at these CO<sub>2</sub> levels, it seems reasonable to examine the research available for medium-term studies on levels of CO<sub>2</sub> less than 10,000 ppm (1%). Table 1 provides a summary of health effects, found in the published literature and discussed in this paper, resulting from breathing CO<sub>2</sub> at levels at or below 1%.

Table1. Documented health effects from breathing CO<sub>2</sub> at concentrations at or below 1%.

CO <sub>2</sub> Level	Health effect	Exposure	Source
10,000 ppm (1%)	Kidney calcification, decreased bone formation and increased bone resorption in guinea pigs	6 weeks	Schaefer et al., 1979a
10,000 ppm (1%)	Cognitive impairment, increased diastolic blood pressure	75 min	Tu et al. 2020
8500 ppm	Increased lung dead space volume	20 days	Rice 2004
7000 ppm (0.7%)	35% increase in cerebral blood flow (implications for cognitive effects seen in other studies)	23 days	Sliwka et al. 1998
5000-6600 ppm	Headaches, lethargy, moodiness, mental slowness, emotional irritation, sleep disruption	Short-term	Chronin et al. 2012; Law et al. 2010



5000 ppm	Kidney calcification, bone degradation in guinea pigs	8 weeks	Schaefer et al 1979b
5000 ppm	Elevated blood CO <sub>2</sub> levels in astronauts	4 months	Hughson et al. 2016
5000 ppm	Current allowable levels for continuous exposure in submarines and spacecraft	Operational continuous	Halperin et al. 2007; Chronin et al 2012
5000 ppm	Permissible exposure limit (PEL) for a work day	8 hours	OSHA 2012
5000 ppm	Vascular and neurological changes	1.5 minutes	Bright et al. 2020
4000 ppm	Increase in heart rate, lowered arousal level/ increased fatigue and sleepiness	17 minutes	Xia et al. 2020
3500 ppm	Cognitive impairment, reduced human vigilance (increased sleepiness)	4 hours	Pang et al. 2020, Zhang et al. 2020
3000 ppm	Cognitive impairment, anxiety, neural damage, oxidative stress in mice	38 days	Kiray et al. 2014
3000 ppm	Impaired visual attention, decision making and executive ability	Short-term	Cao et.al. 2022
3000 ppm	Systemic inflammation and physiological stress in rodents	9-13 days	Beheshti et al. 2018
2700 ppm	Drowsiness measured by EEG	10 min	Snow et al. 2018
2700 ppm	Increase in heart rate	10 min	Snow et al. 2019
2000-4000 ppm	Unhealthy blood CO <sub>2</sub> levels - 15% above normal range, sleepiness, headaches, heart rate variations	4 hours	Vehviläinen et al. 2016
2000-4000 ppm	Inflammation and vascular damage in mice	2 hours	Thom et al.2017
2000-3000 ppm	Restrictive lung behaviour and elevated blood CO <sub>2</sub>	3 hours	Shriram et al. 2019
2000 ppm	Kidney effects in animals (likely calcification) - incomplete study	Chronic studies	Schaefer 1982
1900 ppm	Reduced sleep quality, drowsiness	8 hours	Xu et al. 2020
1400-3000 ppm	Significant impairment of cognitive function including fatigue	2.5 to 8 hours	Satish et al 2012; Allen et al 2016; Kajtar & Herczeg 2012
1200 ppm	Reduced cognitive function	2.5 hours	Scully et al. 2019
1000 ppm	Harmful changes in respiration, circulation, and the cerebral cortex	A short time	Goromosov 1968

1000 ppm	Oxidative stress and damage to DNA in bacteria (implications for cancer diseases in humans)	3 hours	Ezraty et al. 2011
1000 ppm	Cognitive impairment, anxiety, neural damage, oxidative stress in mice	38 days	Kiray et al. 2014
1000 ppm	Level associated with respiratory diseases, headache, fatigue, difficulty concentrating in classrooms	Short-term	Carreiro-Martins et al. 2014; Ferreira and Cardoso 2014; Seppanen et al. 1999
1000 ppm	EEG changes in brainwaves associated with mental impairment	15 min	Lee et. al. 2022
950-1400 ppm	Health symptoms (respiratory, skin, eyes, headaches, cognitive, dizziness, sensory), increase in heart rate	30 min	MacNaughton et al. 2016
950-1000 ppm	Moderate impairment of cognitive function	2.5 to 8 hours	Satish et al 2012; Allen et al 2016; Allen et al 2018
890 ppm	Impaired lung function in new-born female mice, slightly lower blood pH. Reduced growth, hyperactivity, increased stress hormone associated with cognitive impairment	3 months	Larcombe et al. 2021; Wyrwoll et al. 2022
800 ppm	Level associated with Sick Building Syndrome - headaches, dizziness, fatigue, respiratory tract, eye, nasal and mucous membrane symptoms	Short-term	Seppanen et al. 1999; Lu et al. 2015; Tsai et al. 2012
700-900 ppm	Gastropods: reduced growth, reduced food absorption and oxygen uptake, reproductive failure	6-10 months	Navarro et al. 2022; Mardones et al. 2022
700 ppm	Modification of behaviour, stress hormone and respiratory muscle structure in rats	15 days, 6hrs/day	Martrette et al. 2017
420 ppm	Current average outdoor air concentration – no clear effect, possible increase in disease and anxiety	Lifetime	NOAA Global Monitoring Laboratory
280-300 ppm	Pre-industrial outdoor level from about 1820 to at least 25 million years ago - no effect	Lifetime	Beerling and Royer 2011; Zachos 2001.

## 6.1 Spacecraft

A good information source may be the safety guideline documents for activities where humans are required to remain in enclosed spaces for long periods such as spacecraft and submarines. NASA sought to determine the safe levels for long-term exposure to CO<sub>2</sub>, but found little research focused on levels below 10,000 ppm CO<sub>2</sub>; as such, there was no definitive study available to guide standards (Cronyn et al. 2012). They set the maximum allowable CO<sub>2</sub> concentration limits, for long term (1,000 day) habitation of submarines and spacecraft, at 5000 ppm (James and Macatangay 2009).

International Space Station (ISS) crew members have repeatedly reported symptoms associated with

acute CO<sub>2</sub> exposure at levels of 5,000 to 6,600 ppm CO<sub>2</sub> (see Table 1). The most commonly reported symptom was headache; other symptoms reported included lethargy, mental slowness, emotional irritation, and sleep disruption (Law et al. 2010). For space flight, Cronyn et al. (2012) identified three potential areas of operational impact of elevated CO<sub>2</sub>: renal calculi (kidney calcification) and bone reabsorption; cerebral blood flow; and mission performance. With no definitive research to provide insight into these areas, further evaluation was recommended to examine the effects on human subjects of various low-to-moderate CO<sub>2</sub> concentrations (from ambient levels up to 1%). Consequently, flight rules have been employed to reduce CO<sub>2</sub> limits in the ISS to about 3 mm Hg (4,000 ppm) (Ryder et al. 2017).

## 6.2 Submarines

Studies of CO<sub>2</sub> effects on humans in enclosed submarines have been reviewed by the US government (Halperin 2007) although most of these studies are for high (> 1%) CO<sub>2</sub> levels at relatively short exposure durations. At these levels (>1%), many of the debilitating and acute symptoms described above were noted. Current safe levels for continuous exposure in submarines were deemed to be around 5,000 ppm CO<sub>2</sub>. This level is set arbitrarily at one-third of the level where there were obvious signs of health problems (James and Macatangay 2009). It was also noted that if problems are observed, a submarine can surface so that its occupants can be exposed to the ambient atmosphere. Halperin (2007) reports that exposures to CO<sub>2</sub> levels as low as 7,000 ppm can lower blood pH by up to 0.05 units and induce renal (kidney) compensation in healthy subjects. This compensation occurs over a variable period of time, but effects of lowered pH on clinical status or performance have not been reported either experimentally or operationally. Given that kidney compensation cannot occur indefinitely, there is some doubt about whether submariners could sustain the “safe” level of 5,000 ppm CO<sub>2</sub> if they spent years exposed to it.

## 7. Protein malfunctions caused by elevated CO<sub>2</sub> in the body

Carbon dioxide is a fundamental physiological gas known to profoundly influence the behaviour and health of millions of species within the plant and animal kingdoms (Phelan et al 2021). It is known that CO<sub>2</sub> plays a major role in sensing and signalling on a cellular level although the physiological response is complex and not well understood. Research has found that increases in CO<sub>2</sub> resulting in reduced pH alters mitochondrial metabolism reducing oxygen consumption and releasing intracellular calcium stores (Phelan et al 2021).

The endoplasmic reticulum (ER) is a large dynamic structure within cells that serves many roles including protein synthesis and calcium storage. Elevated CO<sub>2</sub> levels cause protein malfunctions by altering ER folding machinery. These malfunctions are associated with ER stress triggering maladaptive responses and effecting a range of diseases e.g., chronic lung disease and reduced immunity, tissue and organ malfunction (Kryvenko and Vadasz 2021)

### 7.1 Elevated CO<sub>2</sub> causes carbonic anhydrase enzyme dysfunction and related diseases

Carbonic anhydrase (CA) enzymes are widely expressed in the nervous system and tissues throughout the body where they play important physiological roles (Aspatwar et al 2021). The reaction that CA catalyses, the reversible hydration of CO<sub>2</sub> to bicarbonate and a proton, lies at the

heart of a wide range of vital physiological processes (Chegwiddden and Carter 2021; Aspatwar et al 2021). Abnormal regulation of carbonic anhydrase-related proteins causes a range of diseases conditions including cancer, Alzheimer's, cardiovascular disease, diabetes, glaucoma, epilepsy, stroke, bipolar disorder etc. The use of CA isozyme inhibitor drugs for treating these diseases is now a major science (Celebioglu et al 2021; Lemon et al 2021). However, CA inhibitors can cause CO<sub>2</sub> retention and result in cellular malfunctions (Phelan et al 2021). The progressive increase in CA disfunction due to increasing atmospheric CO<sub>2</sub> will result in worsening outcomes for the diseases mentioned above.

## 7.2 CO<sub>2</sub> induced deposits of calcium in the body

Carbonic anhydrase (CA) enzymes participate in metabolic reactions that convert CO<sub>2</sub> and result in the precipitation of calcium carbonate (Adeva-Andany et al. 2015; Kim et al. 2012; Tan et al. 2018). CA is implicated in calcification of human tissues, including bone and soft-tissue calcification (Adeva-Andany et al. 2015). The enzyme may be also involved in bile and kidney stone formation and carcinoma-associated micro-calcifications. The molecular mechanisms regulating the development of calcification in human tissues and arteries are similar to those that regulate physiological mineralization in bone tissue, being poorly understood (Adeva-Andany et al. 2015). Carbon dioxide conversion by the CA enzyme provides bicarbonate and hydrogen ions that fuel the uptake of ionized calcium which is then deposited in the body tissues as calcium carbonate. With elevated CO<sub>2</sub> in cells there is increased release of calcium from the ER (Kryvenko and Vadasz 2021).

Kidney calcification is known to occur with longer term exposure to elevated CO<sub>2</sub> levels (Rice 2004; Schaefer et al., 1979a). A similar causal link between the activity of CA enzyme, which is mainly responsible for the reversible breakdown of CO<sub>2</sub>, and calcium deposits has also been established for arteries (Adeva-Andany et al. 2014). As part of a US Navy experimental program in the 1960's and 1970's investigating impacts of long-term CO<sub>2</sub> exposure, Schaefer et al (1979b) found that, in a study of guinea pigs in an enclosed environment breathing 5,000 ppm CO<sub>2</sub> for 8 weeks, the kidneys started to calcify along with bone degradation (see Table 1). Schaefer (1982) also indicated that preliminary experiments had found kidney calcification effects in animal studies for CO<sub>2</sub> concentration as low as 2,000 ppm. Although these studies did not identify a mechanism, they established the casual link between CO<sub>2</sub> and kidney calcification. More recent studies have found that tissue calcification is promoted where CA is overexpressed due to increased CO<sub>2</sub> in the body (Song et al 2021; Phelan et al 2021). Increased CA activity is also linked to cancer where the enzyme helps create a hostile, low pH environment suitable for cancers to flourish in (Hulikova et al. 2014; Logozzi et al. 2019; Di Fiore et al. 2020).

## 8. Other important physiological CO<sub>2</sub> effects on health

### 8.1 Impaired growth and muscle development

Female mice exposed to 890 ppm CO<sub>2</sub> for 3 months from pre-pregnancy showed reduced growth (Wyrwoll et al 2022) and a range of lung and respiratory impairments including lower lung compliance (Larcombe et al. 2021). Prolonged exposure to CO<sub>2</sub> on behaviour, hormone secretion and respiratory muscles in young female rats exposed at 700 ppm CO<sub>2</sub> during 6 hours per day for 15

days (Martrette et al. 2017) (see Table 1). CO<sub>2</sub> exposure, though not continuous, produced significant disturbances in behaviour and was accompanied by increased plasma levels of corticosterone, suggesting that prolonged exposure to CO<sub>2</sub> was stressful producing anxiety. Increased corticosterone associated with hyperactivity in mice was also found from long-term exposure to 890 ppm CO<sub>2</sub> (Wyrwoll et al 2022).

## 8.2 Increased cerebral blood flow

Cerebral blood flow (CBF) effects from breathing CO<sub>2</sub> are a significant issue for humans. As CO<sub>2</sub> in the blood increases, CBF increases to effectively wash out CO<sub>2</sub> from brain tissue and helps regulate central pH (Ainslie and Duffin, 2009). In a 23-day experiment on humans, Sliwka et al. (1998) found that cerebral blood flow is increased in the presence of 7,000 ppm (0.7%) CO<sub>2</sub> by as much as 35% (see Table 1) and that CBF remained elevated until the end of the evaluation period, 2 weeks after the exposure. The impacts of persistent increase in CBF are unclear although there may be a risk of raised intracranial pressure (ICP) which can compress and damage delicate brain tissue. There is also evidence that the CBF response to increased CO<sub>2</sub> is impaired in Alzheimer's patients and that this is linked to the decline in cognitive abilities (Glodzik et al 2013) which will worsen as CO<sub>2</sub> in the atmosphere increases.

## 8.3 Oxidative Stress

In humans, carbon dioxide is also known to play a role in oxidative stress caused by reactive oxygen species (ROS) (Ezraty et al. 2011; Kiray et al. 2014). ROS are produced by aerobic metabolism of molecular oxygen and play a major role in various clinical conditions including malignant diseases, diabetes, atherosclerosis, chronic inflammation and neurological disorders such as Parkinson's and Alzheimer's diseases (Waris and Ahsan 2006). In particular, oxidative damage to cellular DNA can lead to mutations resulting in the initiation and progression of cancer. Ezraty et al (2011) demonstrated that current atmospheric CO<sub>2</sub> levels play a role in oxidative stress and that increasing CO<sub>2</sub> levels between 400 and 1,000 ppm exacerbated oxidative stress and damage to DNA in bacteria. Kiray et al. (2014) concluded that oxidative stress and oxidative damage to brain tissue in mice is associated with low Insulin-like Growth Factor 1 (IGF-1) levels in mice impacting the growth of bones and tissues. Increased CO<sub>2</sub> promotes the production of ROS leading to greater incidence of cancers and other diseases including the promotion of virus activity (Waris and Ahsan 2006). Ezraty et al (2011) concluded that with higher atmospheric CO<sub>2</sub> concentrations, this exacerbation might be of great ecological concern with important implications for life on Earth.

## 8.4 Inflammation

Inflammation is a serious illness that is known to be caused by elevated CO<sub>2</sub> exposure in humans and animals (Thom et al. 2018; Beheshti et al. 2018; Zappulla 2008; Jacobson et al. 2019). CO<sub>2</sub> increases result in higher levels of Interleukin, a protein involved in regulating immune responses, which causes inflammation and vascular damage in mice (Thom et al. 2017). Rodents exposed to 3,000 ppm CO<sub>2</sub> in spacecraft experiments for 9-13 days showed evidence of inflammation and physiological stress (Beheshti et al. 2018).

## 8.5 Stroke

Another study has shown that increased CO<sub>2</sub> in the blood of patients can increase the severity of Subarachnoid haemorrhage; a life-threatening form of stroke, due to the dilatation of arterial cerebral vessels (Reiff et al 2020).

## 9. Discussion

Evidence reviewed in this paper suggests that there is a direct risk to the human species posed by the breathing of ambient atmospheric CO<sub>2</sub> concentrations that are rapidly increasing. The level of ambient atmospheric CO<sub>2</sub> that provides unacceptable risk and the exact effect on physiology are not clearly determined. If this level is reached in the near future, the global human society should be concerned. Some climate models suggest that atmospheric CO<sub>2</sub> levels could be as high as 1,000 ppm in this century. This is completely unknown for the whole primate evolutionary lineage which has only experienced levels well below and recently up to the current level of around 410 ppm.

Given the seriousness of the risks identified in this paper an important question has to be: why has there been no societal discussion of this issue uniquely or in relation to climate change? With the startling rapid rise of CO<sub>2</sub>; the permanent changing of our breathable air composition, one would surely expect the question of whether we can physiologically deal with future projected levels of CO<sub>2</sub>, to at least be examined. However, it appears that the issue is not even thought of. There is an assumption even among scientists that if this was a real issue it would have already been researched, i.e., we would know about it, though this perception is not scientific and precludes novel investigation. It appears that the issue of long-term CO<sub>2</sub> elevated toxicity has never been significantly investigated since we have never been in this situation before. Our body's CO<sub>2</sub> compensation system, as discussed earlier, has been perfectly adequate to deal with a stable level of atmospheric CO<sub>2</sub>. It is also possible that climate change has become the main focus of rising CO<sub>2</sub> levels and there is a lack of vested interest amongst climate scientists about the potential dangers of long-term exposure to elevated CO<sub>2</sub>. The most recent IPCC report on the health impacts of climate change didn't discuss the issue at all (IPCC 2022). This is surprising since the previous IPCC report (Smith and Woodward 2014) described the findings of Satich et al. (2012) as a reported "reduction in mental performance at 1,000 ppm CO<sub>2</sub> and above, within the range that all of humanity would experience in some extreme climate scenarios by 2100". Since the earlier report there have been many more studies of cognitive decline (Du et al. 2020) and the damaging effects of elevated CO<sub>2</sub> on physiology (this paper). It is therefore disturbing and shocking that the latest IPCC report failed to recognise this serious and fundamental aspect of climate change. CO<sub>2</sub> toxicity at elevated levels is a discipline of environmental medicine which has not focussed on the potential problem because chronic impacts of increasing environmental CO<sub>2</sub> have not yet been recognised. This may also help to explain why there are very few researchers involved at this stage and public awareness is close to non-existent.

There are few long-term physiological studies of human exposure to 1,000 -2,000 ppm CO<sub>2</sub> or less. However, there are short-term exposure studies describing disease symptoms and physiological effects as well as reduced cognitive ability in humans at levels around 800 - 950 ppm CO<sub>2</sub>; these are CO<sub>2</sub> levels that are typically present in offices, classrooms and apartments (Gall et al. 2016). It

appears that many of the physiological effects of CO<sub>2</sub> are due to the stimulation of the autonomic nervous system resulting in elevated blood pressure, respiration, and heart rate (MacNaughton et al. 2016) and this is also associated with a decline in cognitive ability due to increased Cerebral Blood Flow (CBF) with resulting effects on central nervous system and brain cortical function (Satish et al 2012; Glodzik et al 2013; Bright et al. 2020). The effect on cortical function is supported by a study of infants that showed an inverse relationship between blood CO<sub>2</sub> and electrocortical activity (Wikstrom et al. 2011). Long-term exposure to environmentally relevant levels of CO<sub>2</sub> leads to increases in the levels of CO<sub>2</sub> in human blood (Zheutlin et al. 2014; Hughson et al. 2016; Vehviläinen et al. 2016). This is retention of CO<sub>2</sub> in the human body at greater than normal levels and disturbingly the levels of CO<sub>2</sub> in our blood will continue to increase as atmospheric levels rise.

Increased CO<sub>2</sub> in the blood also affects protein behaviour causing inflammation (Thom et al. 2018), calcification (Schaefer 1982) of body tissue and a range of demonstrated pH-related protein malfunctions (Duarte et al. 2020), both with potentially serious outcomes. Reduced pH due to CO<sub>2</sub> has been found to alter mitochondrial metabolism affecting oxygen consumption (Phelan et al. 2021). Elevated CO<sub>2</sub> causing protein malfunctions can trigger diseases including cancer, neurological disorders, lung disease, reduced immunity and organ failure (Duarte et al 2020; Kryvenko and Vadász 2021). Perhaps the most significant protein malfunction is the overexpression of the CA enzyme due to increased CO<sub>2</sub> in the body. This widely distributed enzyme plays a vital role allowing for respiration and the processing of CO<sub>2</sub> in all animals and plants. New research is showing that CA malfunction due to elevated CO<sub>2</sub> causes a range of health problems including cancer and diabetes (Aspatwar et al 2021). Perhaps the most serious of these is tissue calcification which can result in kidney and cardiovascular failure. Furthermore, the incidence and prevalence of human kidney calcification (i.e. stones) is increasing globally (Romero et al. 2010; Turney et al. 2011; Kittanamongkolchai et al. 2018) and it is possible that rising indoor CO<sub>2</sub> levels (boosted by increasing ambient CO<sub>2</sub>) is the contributing cause.

Perhaps the most informative study about CA behaviour is that of Rodriguez-Navarro et al (2019). CA catalyses the formation of the reactive precursors (i.e., HCO<sub>3</sub><sup>-</sup> and CO<sub>3</sub><sup>2-</sup> ions) required for mineralization and accelerates the precipitation of metastable amorphous calcium carbonates. Ca<sup>+</sup> and CO<sub>3</sub><sup>2-</sup> ions promote the partial unfolding and oligomerization of CA, resulting in fibril- and sheet-like supramolecular assemblies that template nanostructured calcium carbonate crystallization. The enzyme then loses its CO<sub>2</sub> hydration catalytic activity and elicits a mechanism for arresting calcium carbonate mineralization. Such a negative feedback mechanism would first help jump-start CaCO<sub>3</sub> mineralization and subsequently contribute to the arrest of this process, offering a simple mechanism for organisms to control CaCO<sub>3</sub> biomineralization. Otherwise, if CA endlessly remained catalytically active once secreted, it would be difficult for an organism to stop the biomineralization process (Rodriguez-Navarro et al 2019). Despite the significant reduction observed, some enzyme activity remains even after precipitation of calcite. In a future where increasing atmospheric CO<sub>2</sub> will potentially result in excess CO<sub>2</sub> in the body, the resulting excess CA will be converted into calcium carbonate factories creating an ongoing calcification problem.

What level of permanent CO<sub>2</sub> will cause significant calcification and other protein malfunction effects? It has been suggested that blood pH would be reduced to dangerous levels, if there were no physiological compensation, at CO<sub>2</sub> levels as low as about 430 ppm (Robertson 2006) implying that

ongoing compensation would occur at this level. Ambient conditions may already be dangerously close to CO<sub>2</sub> levels that cause human tissue calcification, particularly when considering the additive effect of increased ambient levels on indoor CO<sub>2</sub> concentrations. In the final paper of the US Navy CO<sub>2</sub> research program in the 1960's and 1970's, Schaefer (1982) indicated that this issue had "become the concern of the Department of Energy and other US government agencies" although it appears to have been largely forgotten since. If allowed to persist, problems such as kidney and artery calcification could lead to cardiovascular failure. In the extreme case, lifespans could become shorter than the time required to reach reproductive age. Calcification of kidneys and arteries can be fatal through renal and cardiovascular failure. This could threaten the viability of human and animal species without interventions such as the creation of artificial living environments.

Cognitive decline due to CO<sub>2</sub>, evidenced by definitive studies (Satish et al 2012; Allen et al 2016; Allen et al 2018) of indoor environments, would logically produce lower intelligence in humans which is now being measured around the world (Bratsberg and Rogeberg 2018). It is feasible that rising outdoor CO<sub>2</sub> levels are the cause of the measured decline in human intelligence (Bierwirth 2018). It is possible that such effects occur without recognition in daily life (Satish et al. 2012). The modest reductions in multiple aspects of decision making, seen as low as 950 ppm (Allen et al. 2016), may not be critical to individuals, but at a societal level or for employers an exposure that reduces performance even slightly could be economically significant. The observation of brainwave changes at elevated levels of CO<sub>2</sub> (around 1000 ppm and greater) (Lee et. al. 2022) is significant being physiological evidence of cognitive impairment. This is concerning since it suggests our brains may be increasingly affected by rising environmental CO<sub>2</sub> levels.

The impacts on students including sickness, reduced attendance and reduced learning abilities should also be a concern for society. Moreover, the relatively high levels of CO<sub>2</sub> in vehicles associated with declining concentration and fatigue has serious implications for the safety of drivers and their passengers. This is an issue that does not appear to have been raised in research on driver fatigue illustrating the general lack of awareness about CO<sub>2</sub> effects.

Carbon dioxide is known to cause anxiety and panic attacks in humans (Battaglia 2017). CO<sub>2</sub> sensitivity is one of the most basic and general alarm/avoidance systems within the realm of biology. Once it permeates the blood-brain barrier, CO<sub>2</sub> causes temporary acidification of extracellular brain fluids resulting in enhanced arousal, and subsequent anxiety (Battaglia 2017). While panic and anxiety attacks occur at high levels of CO<sub>2</sub>, the distribution of liability to CO<sub>2</sub> sensitivity is continuous and normally distributed in humans and animals. This means that there are potentially small anxiety effects even at the current elevated levels of atmospheric CO<sub>2</sub>. Even a small permanent increase in global human anxiety could have a serious impact on societies – mental disturbance, crime, conflicts, etc.

The human species is already impaired in indoor environments and this is likely to get worse as rising outdoor levels of CO<sub>2</sub> contribute to increased indoor concentrations (Azuma et al. 2018). It is not only humans that are at risk. It has been demonstrated that animals have varying degrees of susceptibility to carbon dioxide (Schaefer et al. 1971). The impacts of elevated CO<sub>2</sub> are even greater for water breathing animals than air breathing animals. In general, land animals have much higher blood CO<sub>2</sub> than aquatic animals and can compensate for hypercapnia by increasing ventilation. In



aquatic animals, compensation by increased ventilation is rare and a small increase in ambient CO<sub>2</sub> causes hypercapnic acidosis (Portner et al. 2004; Knoll et al. 1996; McNeil and Sasse 2016). Studies have shown that hypercapnia in fish produces substantial neurological, behavioural and physiological effects (Ishimatsu et al. 2005; Heuer and Grosell 2014) for even short-term exposures at a CO<sub>2</sub> concentration predicted to be persistent in the ocean before the year 2100; this level corresponding with an atmospheric concentration of 650 ppm CO<sub>2</sub> (McNeil and Sasse 2016).

Most of the problems associated with elevated indoor CO<sub>2</sub> levels greater than about 800 ppm, can be alleviated by spending time in fresh air. The indoor environments can be restored to acceptable CO<sub>2</sub> levels with effective ventilation although this is often not being achieved. The available resource of fresh air may be the underlying misguided reason why there is a lack of concern for pollution and its effects. Significantly this resource may not be available in the future as rising atmospheric CO<sub>2</sub> associated with climate change could exceed the 800 ppm level in the current century (Smith and Woodward, 2014). At that stage, there would be no outdoor escape from the described symptoms. Under such a condition of permanent exposure, there could be health impacts at levels less than 800 ppm.

## **10. Conclusions**

There is now a significant body of research work that demonstrate (1) current health impacts of rising CO<sub>2</sub> levels and (2) more serious CO<sub>2</sub> health impacts for humans at some time in the near future.

Current impacts of elevated and increasing ambient CO<sub>2</sub> in indoor environments include inflammation, respiratory diseases, headaches, fatigue, increased heart rate, increased blood pressure, and other symptoms at levels above about 800 ppm. This finding together with the impairment of cognitive abilities at CO<sub>2</sub> levels just above ambient (between 600 and 950 ppm), along with observed associated brainwave patterns, is significant in that it has implications at a societal level for human function particularly for jobs with critical responsibility (e.g., surgery, air-traffic controllers, drivers etc.) together with the impact on learning, human development and economies. It also appears likely that there will be growing global levels of human anxiety from breathing higher levels of CO<sub>2</sub>. Physiological CO<sub>2</sub> effects will be increased and more permanent in a future with elevated outdoor ambient CO<sub>2</sub> concentrations. Ongoing impacts may include the exacerbation by CO<sub>2</sub> of cellular oxidative stress and protein malfunctions resulting in an increase in cancers, neurological diseases, viruses and many other conditions. Studies of health effects at higher levels of CO<sub>2</sub> at around 2,000-5,000 ppm demonstrate the impact of the inability of the body to fully compensate for increased CO<sub>2</sub> and acidity in the blood. These effects include human tissue calcification and bone degradation; the former, associated with the overexpression of the enzyme carbonic anhydrase that is responsible for processing CO<sub>2</sub>, might represent the greatest existential threat for many animals. Given there is a lack of studies in humans at lower but elevated CO<sub>2</sub> levels, demonstrated effects in animals and symptoms experienced by humans indicate that longer-term mechanisms compensating for increased blood CO<sub>2</sub> are active when breathing at around 800-1000 ppm CO<sub>2</sub>. Long-term exposure to these and potentially lower levels would likely cause severe illness. These levels are predicted for the ambient atmosphere by the end of the century in a “business as

usual” world. At some stage in the near future humans could be experiencing persistent physiological effects resulting in serious health problems.

The risk from rising CO<sub>2</sub> levels for human and animal population health in the near-future is extremely high with the health or survival of species under threat. Communication and global awareness of this issue alongside climate change would further strengthen the need to drastically reduce CO<sub>2</sub> emissions. New research on the health effects of long-term exposure to realistic future atmospheric CO<sub>2</sub> levels is urgently needed to quantify this risk.

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