Carbon dioxide deficiency in the human body can be fatal under certain conditions. However, there is still much to be learned about the ways in which hypocapnia (low levels of carbon dioxide) can affect both mental and physical health via altered intracellular pH (during short or longer periods). Dr András Sikter, although now retired, continues to work alongside the Municipal Clinic of Szentendre, Hungary, to explore this further. One aspect of his work investigates the relationship between hypocapnia and the body’s need to meet increased energy demands.

The primary function of the human respiratory system is to breathe in oxygen and exhale carbon dioxide. Oxygen passes from inhaled air in the lungs into the circulatory system. In contrast, carbon dioxide passes from the blood into the cells and alveoli in the lungs and is then exhaled. The goal is to have enough oxygen and maintain normal levels of carbon dioxide in the body, the latter is just as important as the former.

Special receptors called chemoreceptors are primarily able to sense CO2 (carbon dioxide) and pH, and only secondly oxygen levels. When these parameters change, the counter-regulation implements changes which aim to restore the normal levels – sometimes it conflicts with unsolvable contradictions, if all three cannot be met.

Hypocapnia (a reduced carbon dioxide level in the blood) is usually caused by deep or rapid breathing (hyperventilation). It is often associated with high arousal states including anxiety and panic disorder; anxiety can be the cause or a consequence of the hypocapnia, possibly both.

Hypocapnia can alkalise both the blood serum and the inside of the body’s cells. This can have rather significant effects on other molecules in the blood and cells, such as calcium ions which are essential for nerve and muscle excitability. Consequently, the symptoms of low levels of carbon dioxide may include tingling limbs, disrupted heart rate, muscle cramps, seizures and many more.

Opposed to this, hypercapnia is the occurrence of increased levels of carbon dioxide in the blood, often secondary to increased breathing rate, lung disease or a state of reduced consciousness.

Dr András Sikter works alongside the Municipal Clinic of Szentendre, Hungary, to explore how changes in hydrogen ion concentration (pH) due to hypocapnia and hypercapnia impact upon cell metabolism and how they can induce diseases of civilisation. In this work, he has explored those vicious circles in which hypocapnia on its own can accelerate metabolism through intracellular alkalinosis and ultimately lead to fatal outcomes. If his hypothesis is true, the therapeutic strategy of emergency medicine should be modified because hypocapnia often leads to death in critically ill patients for example, due to congestive heart failure, lung diseases, high altitude sickness, stroke, brain trauma, meningitis, sepsis, COVID-19 infection, etc.

CHANGES IN ACIDITY LEVELS
Any change in carbon dioxide levels, and therefore respiration, will affect the pH of cells. It takes hours or days for the compensatory mechanisms to kick in, by which time full functional damage may have already occurred in the case of hypocapnic emergency. The compensatory processes are also not completely effective. In the numerous counter-regulation mechanisms appear to contradict each other or create additional error-synergy to try to restore intracellular pH; this is the case in chronic ventilatory disorders.

Intracellular pH has a number of very significant impacts on the rates of metabolic reactions and energy demand. Due to this, if it differs from the normal levels, the regulation of intracellular H+ concentration (pH) has priority above other ions, often at the expense of them.

Intracellular pH regulation

Generally, alkalinosis (increased pH) causes cell metabolism to increase, whereas acidosis (decreased pH) slows it down. As a result, energy demand also increases or decreases. A small increase in pH results in a huge increase in cell metabolism and excitability of neurons, muscles cells and cardiac cells. Cell excitability is defined as cells being activated by, and reacting to, a stimulus such as an electrical signal.

The regulation of intracellular H+ concentration (pH) has priority above other ions, often at the expense of them.

It is often induced by changes in the cell membrane potential, i.e. the differences in the concentrations of ions on opposite sides of the cell membrane. During steady-state, the ratio of these ions remains unchanged.

ENERGY DEMAND AND PRODUCTION
Adenosine triphosphate (ATP) is the energy currency of a cell and plays a key role in metabolism. Hypocapnia increases cells’ oxygen and ATP energy utilisation. Young, healthy tissues are able to adjust their ATP production to meet increased demands. However, the cell membranes can become more permeable to other molecules, such as calcium ions, which can lead to excitability problems and symptoms of functional disorders.

Dr Sikter explains that in contrast, seriously damaged tissues are unable to meet this demand; for example, the heart tissue may be impaired due to another illness/disease. In these cases, the attempt to meet energy demands, and the subsequent failure to provide an adequate amount of ATP, can lead to a series of further complications such as fluid accumulation in the lungs. Dr Sikter highlights that low phosphate levels in the blood can also arise from hyperventilation, resulting in further ATP deficiency and exacerbating the whole issue.

PERMISSIVE HYPERCAPNIA
Critically ill hypocapnic patients suffering from ARDS or COVID-19 infection tend to maintain mechanical ventilation are often placed in the state of ‘permissive hypercapnia’ by physicians. A controlled high carbon dioxide level has a therapeutic effect on infected lungs and is not life-threatening if sufficient oxygen is available at the same time. Because the developing acidosis has the ability to slow down metabolic and enzymatic processes, this can potentially be disadvantageous in the short term. The ‘permissive hypercapnia’ theory and practice, formed by experience of
the exhaustion of the heart's energy markers for poor prognosis as it reflects during congestive heart function is a harder to match the work of a normal heart. This correlation is also seen when those who are seriously damaged or for young, healthy tissues but not for mechanisms kick in. This is achievable demand alone before compensatory each cell is left to meet increased energy to provide energy supplies can be an important dividing line between organic and functional diseases. THE VICIOUS CYCLES OF MENTAL AND SOMATIC ILLNESSES The link between mental and physical health is well known. For example, people will increase or decrease their breathing rate in line with emotions and arousal, thus inducing hypo- or hypocapnia. Dr Sikter identified that vicious cycles will develop involving somatic and functional changes; they almost inevitably develop in the case of hypocapnic emergency. However, there is a narrow path (such as ‘permissive hypocapnia’) to recovery. The occurring acidosis is able to break the vicious circle and turn it into a virtuous cycle.

Dr Sikter concludes that ‘the ability to provide energy supplies can be an important dividing line between organic and functional diseases’. In contrast, acute hypocapnia can further endanger critically ill or organs due to the increased energy demands which they are unable to meet. Hypocapnia has an immediate impact on cell metabolism, whereas counter-regulation takes longer to respond. This means that each cell is left to meet increased energy demand alone before compensatory mechanisms kick in. This is achievable for young, healthy tissues but not for those who are seriously damaged or impaired (‘the weakest link’).

This correlation is also seen when heart function is considered; heart failure increases oxygen demand as the damaged tissues have to work harder to match the work of a normal heart. The development of hypocapnia during congestive heart function is a marker for poor prognosis as it reflects the exhaustion of the heart's energy reserves and hypocapnia demands a further increase in ATP. It was found by others that low-dose carbon dioxide treatment could theoretically reduce the risk of hypocapnia generated by congestive heart failure. However, this could also go some way to explain the link between hypocapnia, anxiety and long-term disease; hypocapnia may kill or damage the injured brain cells further, suggesting an explanation for the progression of underlying diseases during delirium.

Hyperventilation hypocapnia in individuals with panic disorder can result in cardiac arrhythmias and, while panic disorder does not cause coronary artery disease, it can double the chance of death if the two disorders occur at the same time. Dr Sikter's earlier work has also explored the relationship between physical and mental health. He suggests that protracted or repeated episodes of major depression could lead to actual physical damage to the brain and other critically ill organs. That is why Major Depression is more of an organic rather than a functional disorder – according to Dr Sikter's theory, chronic (respiratory or metabolic) acidosis promotes its development. He concludes that ‘the ability to provide energy supplies can be an important dividing line between organic and functional diseases.’ To expand on this, organic diseases involve an observable and measurable disease process, such as inflammation, whereas functional diseases have no definite somatic direction to explain disease occurrence.

Although generally thought not to be dangerous, Dr Sikter discusses that hypocapnia can have fatal results for those who are critically ill, and the deciding factor appears to be whether the cells are able to mobilise sufficient ATP to meet increased energy demands. He believes that administering physiological doses of carbon dioxide levels. In the long run, the compensation promotes only the survival of respiratory disorders, which can be important stages in the development of the diseases of civilization. We have to work out a widely applicable strategy to restore CO2 levels to normal. As we demonstrated, even hyperventilation alone can be fatal in critically ill patients (permissive hypocapnia) can also be used only in the short term). All cells have to fight alone against the intracellular alkalosis caused by hypocapnia, the critical ATP deficiency will develop in the Paths of Least Resistance.

Should hypo- and hypercapnia be given more consideration in the pathophysiology of human diseases?

We must not tolerate the persistence of higher or lower carbon dioxide levels. In the long run, the compensation promotes only the survival of respiratory disorders, which can be important stages in the development of the diseases of civilization. We have to work out a widely applicable strategy to restore CO2 levels to normal. As we demonstrated, even hyperventilation alone can be fatal in critically ill patients (permissive hypocapnia) can also be used only in the short term). All cells have to fight alone against the intracellular alkalosis caused by hypocapnia, the critical ATP deficiency will develop in the Paths of Least Resistance.

References