Multifactorial Distress, the Warburg Effect, and Respiratory and pH Imbalance in Cancer Development

Gabi Drochioiu

Biochemistry Group, Faculty of Chemistry, Alexandru Ioan Cuza University, 11 Carol I, 700506 Iasi, Romania; gabidr@uaic.ro; Tel.: +40-722280328

Abstract: Oncogenes are thought to play an important role in aberrant regulation of growth factors, which is believed to be an initiation event of carcinogenesis. However, recent genetic and pharmacological studies have shown that the Warburg effect (WE) is needed for tumour growth. It refers to extensively studied aerobic glycolysis over the past decade, although its impact on cancer remains unclear. Meanwhile, a large body of evidence has indicated that oxidative stress (OS) is connected with the occurrence and progression of various forms of cancer. Psychosocial factors (PSF), such as chronic depression, sadness, stressful life experiences, stress-prone personality, and emotional distress or poor quality of life affect the immune system and contribute to cancer outcomes. Here, we examine the relationship between WE, OS, PSF, metal ions, other carcinogens, and the development of different cancers from the viewpoint of physiological and biochemical mechanisms.

Keywords: Warburg effect; oxidative stress; psychosocial factors; pH imbalance; distress; cancer

1. Introduction

Metabolic changes in cancer are no longer seen as an indirect response to signals of cell proliferation and survival. Rather, impaired metabolism status is the basic hallmark of cancer [1]. The hypothesis that oncogenic transformation alters cellular metabolism to sustain high rates of growth and division has been extensively explored [2]. Recent genetic and pharmacological investigations have shown that the Warburg effect (WE) is also required for cancerous growth [3,4]. During cancer progression, oxygen respiration always decreases, fermentation takes place, and highly differentiated cells switch to anaerobic fermentation, having lost all their previous physiological functions and only retaining the now useless property of proliferating and multiplying [5,6]. Cancer metastasis and therapeutic resistance are usually studied as separate areas using different strategies. However, metastatic progression and therapeutic resistance signalling are mediated by common mechanisms, such as the involvement of integrins and other contextual receptors, cell–cell communication, stress responses, and metabolic reprogramming [7]. During proliferation and metastasis, malignant cells adapt to oxidative stress by increasing NADPH in a variety of ways, including by activating AMPK, PPP, and reductive glutamine, as well as folate metabolism [8]. Indeed, reactive oxygen species (ROS) influence the progression of cancer, either by initiating or stimulating tumorigenesis and supporting the transformation and proliferation of cancer cells, or by causing their death [9–11]. While cancer cells have increased levels of ROS, and increased ROS concentrations are associated with various carcinogenic processes, some drugs destroy cancer tumours via toxic levels of ROS [12]. Psychosocial factors are stressors that have a negative impact on cancer patients, but their effects vary depending on the type of psychosocial factor, cancer location, and cancer outcome [13]. Thus, stressful life experiences have been shown to be associated with lower cancer survival and higher mortality, but not higher incidence.

Heavy metal-induced oxidative stress can promote various cancers and diseases by ROS-based mechanisms [14]. Heavy metals stimulate tumour progression and reduce...
tumour sensitivity to treatment, while tumour tissue shows a different level of DNA methylation [15].

Biochemically, the less-differentiated cell structure of the cancer tissue somewhat resembles that of foetal tissue [16–18]. Consequently, the importance of studying the embryo to understand the evolution of the tumour and contribute to the development of effective therapeutic strategies was highlighted [19]. In addition, high concentrations of alphafetoprotein are normally found in foetal blood but are almost undetectable in adult blood. Therefore, this protein has attracted increasing interest because of its connection with carcinogenic events [20]. It is precisely this feature that should be investigated, as it may hide the underlying mechanisms of cancerogenesis.

Hypoxia is the condition in which tissues are exposed to oxygen deficiency and is an essential phenomenon influencing cellular health. The effect of hypoxia on human cells can be either positive or negative depending on the severity, duration and context [21]. Multi-cellular organisms have developed both systemic and cellular responses to hypoxia [21]. The generation of adenosine triphosphate (ATP) in mitochondria is particularly sensitive to changes in oxygen tension. For that reason, the hypoxic state is an aggravating factor commonly seen in cancer, multiple sclerosis, heart disease, kidney disease, liver disease, etc. [22]. Hypoxia can play an important role in the regeneration of damaged tissues, in particular by acting on tissue-specific stem cells. However, its role can be a drawback when it involves neoplastic stem cells.

Here, we first analyse the function of WE, highlight its significance and discuss its shortcomings. Our analysis, carried out from the viewpoint of physiological and biochemical mechanisms, mainly focuses on the relationship between WE, OS, non-specific stress (NSS), hypoxia, heavy metals, and other carcinogenic factors involved in the occurrence of different cancers. Since the normal cells of the body meet their energy needs by breathing oxygen while cancer cells do this mainly through fermentation, we have introduced the term respiratory imbalance, which refers to an impairment of respiration. Further, glycolysis generates lactic acid, which alters the pH of body fluids. We have therefore added the notion of pH imbalance. All findings discussed here suggest that lifestyle, food, distress, carcinogens, ROS, and heavy metals can be environmental factors involved in cancer aetiology and progression. We are also pursuing common physiological mechanisms capable of shedding light on the carcinogenic effect of so many carcinogens involved in so many different cancers and their link to the Warburg effect.

2. Oxidative Stress

Molecular stress is considered to be involved in cancer initiation and progression [23,24]. Oxidative stress from endogenous and exogenous sources leads to mutations and epigenetic deregulations, which contribute to the development of neoplastic diseases [25]. Among the first category of stressors are peroxisomes and enzymes, such as NADPH oxidase [26], xanthine oxidase [27], dihydrolipoamide dehydrogenase [28], etc., most of which are found inside the mitochondria. Other stressors, such as alcohol, nicotine, exercise, or UV radiation, are responsible for an increase in the intracellular level of several reactive species, such as reactive oxygen species (ROS), reactive nitrogen species (RNS), and reactive sulphur-based species (RSS) [25]. ROS appear during mitochondrial aerobic metabolism, being a reaction of human and animal cells to bacterial invasions, presence of xenobiotics, on-going distress, or X-ray exposure [29]. The relative excess of ROS, when compared to antioxidants, has been linked to multiple pathologies, such as neurodegenerative disease, cardiovascular disease, diabetes mellitus, etc. [30]. The cancer cell is also known to show aberrant redox natural balance. While ROS are pro-tumorigenic, a high level of ROS can be cytotoxic [31]. Excessive production of ROS is associated with many types of diseases, such as chronic inflammation [32,33] and a variety of cancers [34–37]. Thus, the proliferation of malignant cells is associated with high ROS production. Notably, these cells are adapted to grow under conditions where this oxidative stress shifts the redox homeostasis away from a reduced status; tumour cells achieve this balance by increasing their antioxidant potential to
optimize ROS-driven proliferation [38,39]. Since there is a link between the redox potential of a cell and its tolerance to high levels of ROS, biochemistry of reduced glutathione (GSH), thioredoxins (TXN), and NADPH becomes important. Normal metabolic processes typically generate ROS and reactive nitrogen species (RNS) that are potentially harmful in high concentrations; these species are intracellular signalling molecules [40]. However, cells possess an array of antioxidant systems to ensure that ROS and RNS signalling mechanisms are preserved and oxidative injury is avoided. The increase in the level of ROS and RNS in relation to antioxidant activity of the cell means that oxidative stress (OS) is counteracted by the normal cell with the help of GSH and TXN. Their action is supported by NADPH, which keeps them both in a reduced state. Oxidation of polyunsaturated fatty acids by ROS results in lipid peroxidation, while the peroxidised compounds and their breakdown products may act as signalling molecules to stimulate inflammation and apoptosis [38,41].

The main resources of ROSs include: electron flow to $O_2$ in mitochondria and the reaction between coenzyme Q10 (CoQ) found in the semiquinone form and the molecular oxygen at complex III of the respiratory chain [42]. In addition, NADPH oxidases reduce $O_2$ to superoxide, $O_2^•−$ [43]. The formation of highly reactive hydroxyl radicals (HO$^•$) by Fenton’s chemistry from $H_2O_2$ molecules usually implies heavy metal ions, such as copper, iron, or manganese. One of the main RNS species is the vasodilator NO$^•$, which is liberated by nitric oxide synthase using L-arginine. The reaction between NO$^•$ and $O_2^•−$ produces ONOO$^−$ [44].

Antioxidant defence prevents the accumulation of ROS and RNS through several scavenger molecules, such as GSH, melatonin, α-lipoic acid, bilirubin, melanin, or uric acid. Vitamin E, vitamin C, β-carotene, and plant polyphenols are also antioxidants [40]. GSH is subject to homeostatic regulation and is often increased in some forms of cancer [45]. On the other hand, the cytoplasmic copper/zinc superoxide dismutase (SOD1), mitochondrial manganese superoxide dismutase (SOD2), and extracellular superoxide dismutase (SOD3) catalyse the conversion of $O_2^•−$ to $H_2O_2$ and $O_2$ [46]. Since SOD1 and SOD2 protect against spontaneous malignization and are defined as tumour suppressors, they are up-regulated during oncogenesis [47]. Next, catalase (CAT) is involved in the decomposition of $H_2O_2$ into $H_2O$ and $O_2$ [48].

Cancer cells have to deal with OS at the onset, during matrix cleavage, during entry into circulation, and when the disease relapses following treatment [8]. Tumour cells are able to adapt by various means to ensure that ROS activity is limited to a dynamic threshold that allows them to proliferate while avoiding cell mortality [49,50].

Oxidative stress is connected to the progression of the most common form of liver cancer [51]. Nevertheless, the mechanisms are still unclear. Typically, OS happens when the body detects any danger signal, either from an internal or external source [47]. Reactive oxygen species (ROS) are permanently generated in peroxisomes, mitochondria, cytosol, and apoplast. The imbalance between ROS generation and detoxification leads to oxidative stress, and the accumulation of ROS is harmful to cells. In addition, ROS function as signalling molecules and activate signal transduction processes in response to various stresses [52]. Uncontrolled overproduction of ROS, resulting from an imbalance between ROS production and removal, leads to vascular disease [53]. It then induces oxidative damage to the DNA and abnormal protein synthesis, putting the body in a condition of susceptibility to developing various diseases, including cancer. Many factors are involved in liver carcinogenesis, including hepatitis B virus and hepatitis C virus infection, alcohol abuse, and non-alcoholic fatty liver disease. Elucidation of the influence of OS in cancer aetiology is important for the prevention and treatment of various cancers. Treatment with OS antioxidant drugs can control OS lesions in vitro [54]. However, in the case of liver cancer, chronic viral infections can induce inflammation and necrosis of liver cells [55]. DNA damage caused by ROS leads to the accumulation of cancer-related genetic mutations. Chronic inflammation is one of the causes of human cancer [56,57]. Oxidative stress and accumulation of DNA damage play an important role in virus-induced cancer [58].
addition, miRNA dysfunction in inflammatory reactions is believed to be the central event in the occurrence of some cancers [54].

High levels of ROS have been detected in almost all cancers, where they promote many aspects of tumour development and progression. However, tumour cells also express increased levels of antioxidant proteins to detoxify from ROS, suggesting that a delicate balance of intracellular ROS levels is required for cancer cell function [59,60].

There is an increased risk of cancers among obese patients; some forms of fatness explain cancer risk in obese patients, while oxidative stress may play a role in obesity-related cancers [61]. The enzyme involved in triacylglyceride synthesis and lipid droplet formation, diacylglycerol O-acetyltransferase 1 (DGAT1) is commonly up-regulated in melanoma, allowing these cells to support excess fatty acids [62]. DGAT1 inhibitors induce OS in melanoma cells, which adapts by increasing cell defences against ROS. Inhibition of both DGAT1 and superoxide dismutase 1 profoundly inhibit tumour proliferation due to exaggerated OS.

### 3. The Non-Specific Stress

Several studies suggest that stress may influence oncogenic development, and data from subhuman experiments have shown that aggressive offenses can enhance or suppress oncogenicity [63,64]. Thus, psychological factors may affect the risk and progression of tumour proliferation [65]. Increased tumour progression is evident following acute exposure to severe stress, and the impact of aggressive stimuli varies according to previous distress history and social living conditions [66]. In addition, prolonged depression can be associated with an increased risk of cancer [67]. Moreover, providing psychosocial support may help reduce depression, angst, and hurt, and can prolong the survival period with cancer. The relationship between sadness and cancer evolution assumes dysregulation of the hypothalamic-pituitary-adrenal alignment, particularly daytime changes in melatonin and cortisol. In general, depression affects the immune system, which can affect cancer control [68]. People with psychiatric disorders are no more susceptible than the general population to developing cancer, but they are more inclined to die from this disease [69]. All these findings suggest that there may be a close relationship between other forms of stress, mainly related to human personality.

Stress induces or worsens cardiovascular diseases, non-alcoholic fatty liver disease, depression, neurodegenerative disease, and cancer through peripheral inflammation as well as neuroinflammation [70]. Stress endangers central microglia and astrocytes, blood vessels, and immune system. It has been suggested that inflammation may be the common pathway for stress-related diseases, which may act as a contributing factor to disease progression or may occur very early during the disease development [70].

### 4. The Warburg Effect

A century ago, Otto Warburg and his colleagues observed that growing ascites cells converted most of their glucose to lactate, even under O₂-rich surroundings [71]. He thought that such altered metabolism was specific to cancer cells, and that it arises from mitochondrial deficiencies that inhibit their capability to efficiently oxidize glucose to CO₂. He concluded that the existence of dysfunctional mitochondria is one of the causes of cancer [72]. However, injured mitochondria have been shown not to affect aerobic glycolysis in most tumour cells; mitochondria in cancer cells are not damaged, but simply dysfunctional [73]. The metabolism of the cancer cell is altered and involves augmented glucose uptake and glucose fermentation to lactate [74]. This condition is known as the Warburg effect or aerobic glycolysis and is observed even in the presence of fully functional mitochondria, or even in the presence of oxygen [71,75]. Since respiration can maintain tumour viability, it was thought that these cells can be killed by depriving tumour cells of energy, so both glucose and oxygen should be removed [76]. However, Herbert Crabtree reported the heterogeneity of glycolysis in various tumours. Therefore, variable intensity
of respiration in tumours was discovered [77]. Crabtree established that there is also variability in fermentation, probably due to environmental or genetic influences.

Nevertheless, Warburg proposed later that the origin of aerobic glycolysis is dysfunctional mitochondria [78]. Yet, Efraim Racker showed that tumours have respiratory capability. He advanced his own hypothesis about the Warburg effect by studying intracellular pH imbalances that disrupt the ATPase activity [79]. It has also been observed that aerobic glycolysis can be controlled by growth factor signalling. However, the identification of genes with a potential role in oncosogenesis led to the conclusion that aberrant growth factor regulation could be the initial event in tumorigenesis [80,81]. Nonetheless, WE is necessary for tumour growth [3,4]. Therefore, targeting both aerobic glycolysis and mitochondrial metabolism might be required in cancer therapy [82–84]. Nevertheless, the functions of the Warburg effect have remained controversial for a long time.

The Warburg effect confers direct tumour cell signalling functions [85–88]. Thus, a direct contributory role of glycolytic metabolism in stimulating carcinogenesis through this signal transduction affecting other cellular processes is suggested. It has been thought that aerobic glycolysis may offer some advantage as it provides a favourable tumour microenvironment for cancer cell multiplication [1,89,90]. Under certain conditions, the Warburg effect could be the choice of an energy metabolism based on high glucose consumption.

Most tumour mitochondria are functional and are therefore able to perform oxidative phosphorylation. Nevertheless, mitochondrial metabolism in proliferating cells seems to be directed to macromolecular syntheses. Warburg and his colleagues did not consider such a possibility [1]. However, some authors consider that the Warburg effect is an initial event in carcinogenesis, being a direct result of an oncogenic mutation, which occurs before abnormal cell multiplication, and also in benign and early-stage lesions [91,92].

The Warburg effect was extensively investigated from multiple points of view [93]. Thus, metabolic alteration was understood as a necessity for rapid multiplication. It instantly generates energy in the form of ATP molecules, supports the biosynthesis of macromolecules and maintains the redox state of cells. Processes such as pH modification of tumour microenvironment, the stabilization of hypoxia-inducible factor (HIF), some mutation of tumour suppressor genes, and dysfunctions of mitochondria have been discussed. In addition, selective targeting by miRNA, altered glutamine metabolism and post-translational modifications were also investigated. These authors considered that a holistic understanding is needed to discover novel metabolism-based therapeutic strategies to hinder the Warburg effect and cancer advancement. Other authors found that the Warburg effect stimulates cancer metastasis and changes the tumour microenvironment; it may play a role in promoting angiogenesis, formation of cancer-associated fibroblasts, immune suppression, and drug resistance [94]. High uptake of glucose by cancer cells reduces considerably its accessibility in the tumour microenvironment, which results in a low-glucose extracellular environment and disturbs the activity of immune cells [74]. Moreover, tumour cells release high amounts of lactate, which induces an increase in the acidity of the microenvironment. Lactate can be utilized by some non-tumour cells in the liver to produce glucose with high energy consumption [95]. An acidic tumour microenvironment stimulates local invasion, and then metastasis and diminishes the anti-tumour action of immune cells [96,97].

In cancer cells, respiratory function decreases, and an increase in glycolysis proportional to the increase in the growth rate is observed [98]. However, decreased cellular respiration is not obligatory for an increased rate of cell proliferation.

Myc and HIF-1 activate the Warburg effect in reaction to growth factors and hypoxia. It is an important metabolic and energetic process that meets the requirements for fast gene replication [99]. Paradoxically, cancer appears to be a normal physiological phenomenon that follows precise rules, but it is also a degeneration, a dysregulation caused by a multitude of factors: lifestyle, diet, carcinogens, etc.
5. The Aerobic Glycolysis

Glycolysis is a primitive metabolic pathway that is essential for rapid multiplication of cancer cells, tissue regeneration, but also for growth of bacteria and viruses [99]. Aerobic glycolysis, which occurs not only in cancer cells, can be defined as an exaggerated increase in glucose consumption compared to oxygen supply, even when oxygen levels and delivery in the blood are sufficient to meet demand [100]. Therefore, this type of glycolysis is uneconomical and ATP generation is very low compared to the ATP produced by respiration [101,102]. Nevertheless, the amount of ATP synthesized in a given period of time is similar in both forms of glucose metabolism [103]. Therefore, the reason why the cancer cell uses aerobic glycolysis should be investigated and a suitable explanation should be found for this inherent difference in kinetics. A very simple explanation would be that there is a precise ratio between the concentration of ADP and ATP. Thus, decreasing the concentration of ADP will cause a phosphorylation reaction of ADP in order to keep the ratio of adenosine-di and triphosphate as constant as possible. It has been hypothesized that cells with higher glucose consumption, albeit with lower efficiency in ATP production, may have an advantage when competing for common and restricted energy resources [104,105]. On the contrary, we think differently: low oxygen concentration can lead to glycolysis [106]. Thus, it has been shown that when the cellular environment is altered to greatly increase ATP requirements, aerobic glycolysis increases rapidly, and oxidative phosphorylation remains constant [107]. In such cases, the aerobic glycolysis is considered an adaptive process to sustain the conditions for the biosynthesis of macromolecules and other compounds required by the uncontrolled multiplication. Therefore, increased glucose uptake is a carbon source for syntheses required to sustain tumoral growth [72,108–110]. The aerobic glycolysis is also required to support the rapid generation of ATP required to sustain chemical synthesis. However, the ATP requirement for cell growth and division is much lower than required, and ATP demand may never reach threshold values during cancer cell growth [111]. Similar mechanisms are also observed in other cell types linked to a rapid demand for ATP are also present in tumour cells. Thus, fast ATP synthesis based on creatine kinases in muscle is manifest in most tumour cells [74]. Compounds resulting in glycolysis are required for nucleotide, lipid, and protein synthesis [112–115]. Proliferating cells have a greater need for NADPH or NADH [116]. Increased synthesis of the reducing equivalents implies a higher utilization of glucose, which is then employed in the biosynthesis of lipids, amino acids and other biomolecules [1]. It was considered that the role of aerobic glycolysis is to regenerate NAD$^+$ in the reaction of NADH+H$^+$ with pyruvate, which produces lactate, the final product of aerobic glycolysis [117,118]. Lactate is not just a by-product of glycolysis, but has an important role in tumour metabolism, as identified by the Warburg effect studies [119]. Lactate plays a major role in cancer cell proliferation, but is also involved in inflammation, neural excitation, and many other biological processes.

NADH is generated in the reaction catalysed by glyceraldehyde phosphate dehydrogenase and is oxidized to NAD$^+$, thus keeping glycolysis active. Glycolysis enables 3-phosphoglycerate to convert to serine for the production of NADPH and nucleotides [120]. NADPH homeostasis is regulated by several metabolic enzymes that undergo adaptive changes in cancer cells [121]. It is thought that modulating NADPH homeostasis in cancerous cells could be an effective strategy to eliminate them.

Aerobic glycolysis maintains a fertile environment that supports rapid biosynthesis to sustain multiplication and proliferation [122]. Cancer cells use aerobic glycolysis for energy metabolism, and a method to deprive malignant cells of glucose would prevent these cells from surviving and induce apoptosis in several types of cancer, which could be the basis of a potential treatment [123]. Cancer cells use glycolysis as an energy source although oxygen is present, this changed metabolism may provide a selective benefit for survival and growth, consistent with the Warburg effect. In addition, several molecules, such as NADPH, HIF, PKM2, and others, are important for the reproduction of cancer cells in the abnormal hypoxic medium.
It was also suggested that aerobic glycolysis is a pathway to support biosynthesis [124,125]. Although ATP production is inefficient, it can come at the cost of maintaining anabolic pathways, such as those involved in nucleotide and lipid metabolism. There may also be a limited number of mitochondria; thus, the necessary energy and biomass beyond mitochondrial capacity must be produced from aerobic glycolysis [126–128]. Therefore, aerobic glycolysis is considered to support biomass production when ATP production is limited. There is an apparent correlation between aerobic glycolysis and cell proliferation. The demand for NADPH is higher than the ATP requirement for biosynthesis. However, in aerobic glycolysis, the carbon atoms are not sequestered but are liberated extracellularly as lactate [2,111]. Acidosis can be beneficial for cancer cells; protons, H+, secreted by tumour cells may be liberated into the environment and modify the tumoral–stroma interface, permitting increased invasion [129]. Tumour-derived lactate has also been shown to contribute to tissue-associated M2 macrophage (TAM) polarization [130].

Glucose availability seems to be a result of intensive competition between resulted tumours and tumour-infiltrating lymphocytes (TILs) [131–133]. Intense aerobic glycolysis limits the availability of glucose to tumour-infiltrating lymphocytes, which need abundant glucose for their physiological functions [134–136]. Consequently, evidence is sought that inhibition of aerobic glycolysis in the tumour would allow increased glucose supply to TILs, thereby stimulating their function to eradicate tumour cells [137,138]. All these observations may suggest that malignant cells are in contact with cells in the immune system to sustain pro-tumour immunity [139–143].

Lactic acid plays a key role as it is capable of translocating through cell membranes, contributing to the cell-pH state, as well as influencing the complex immune response due to acidosis of the tumour microenvironment [99]. Even working brain tissues partly oxidize glucose and produce some lactic acid [100,144–147]. Therefore, aerobic glycolysis occurs normally when cells are stressed. Aerobic glycolysis occurs in astrocytes, where the Crabtree effect coincides with the Warburg effect. At the same time, neurons use both glucose and lactate, and there is a balance between glycolysis and respiration. This leads to the activation of Warburg and Crabtree effects in brain tissue, resulting in a high degree of aerobic glycolysis, indicating stimulation of astrocytes to generate neuronal ATP.

6. Copper and Cancer

Copper (Cu) is involved in numerous cellular processes, which include mitochondrial respiration, anti-oxidative defences, redox signalling, autophagy, kinase signalling, and regulation of protein quality [148]. Specific abnormalities of copper metabolism appear to have clinical potential as prognostic and predictive biomarkers [149]. Cu2+ ions are also capable of binding to growth factors, cell signalling proteins, or even structural proteins [150]. These ions can regulate the activity of several proteins; thus, many signalling metabolic reactions are dependent on copper. Mitochondria also play an essential role in copper homeostasis, which is important for mitochondrial physiology [151]. Cu is a component of cytochrome c oxidase, which is present in the respiratory chain in mitochondria [152]. Therefore, Cu is involved in energy production via oxidative phosphorylation [153]. In addition, copper is present in cell lysosomes, and some metallo-reductases maintain it in the Cu(I) form because lysosomes are an oxidative environment [154,155]. Such reductases are mainly located in the intracellular vesicles [156,157]. They are involved in the regulation of cell proliferation and apoptosis.

Glutathione (GSH) and metallothioneins (MTs), which are cysteine-rich cytoplasmic proteins, are greatly engaged in intracellular storage of excess copper [158]. GSH is implicated both in numerous mechanisms of metabolism and in the transfer and removal of metal ions, including copper ions, as Cu(I)–GSH complexes [159,160]. These complexes are thought to be related to the exchangeable pool of cytosolic copper [161,162].

Changes in Cu levels or in the Cu: Zn ratios have been observed in several forms of cancer [163,164]. Nevertheless, the Cu: Zn ratio changes with aging, inflammation, nutritional status, and OS. Increased copper concentration is accompanied by diminished
levels of zinc in bladder cancer [165] and other forms of cancer [166–169]. However, certain authors testified that there are decreased copper levels in colorectal and breast cancers [170,171].

Increased levels of copper have been reported in tumour areas [172,173]. Copper is involved in proliferation and angiogenesis, two phenomena seen in tumorigenesis and cancer development. In addition, specific copper accumulation was reported in cancer cells themselves [174]. Thus, high levels of Cu were reported within the tumoral cells of breast cancer [175]. It is likely that copper ions induce the formation of secondary tumours by activating some enzymes implicated in cell multiplication [176]. Moreover, increases in serum copper in cancer pathology were sometimes correlated with cancer stage. In addition, in the case of patients who are resistant to chemotherapy, increased levels of serum copper have been measured [176]. Data on different types of cancer on this subject are contradictory. We suggest that a link between copper level and pH may exist. The isotopic $^{63}$Cu/$^{65}$Cu ratio in the serum of tumour patients also seems to be altered [74], with increasing levels of the lighter isotope. These changes could be due to increased glycolysis and lactate formation. Furthermore, the Cu isotopic ratio could be used as an early diagnostic biomarker for cancer [176].

Some copper-related proteins, such as ATP7B and Ctr1, have been found to increase in breast cancer [177]. Dysregulation of several proteins involved in copper metabolism has an influence on cell migration and metastasis formation. Thus, Atox1 protein is elevated in several malign tissues [178,179]. It may also promote inflammatory neovascularization by acting putatively as a transcription factor and as a copper chaperone [180].

Cu-dependent LOX metalloenzymes play a significant function in tumour metastasis [181]. Thus, cancer cells produce LOX protein to promote collagen cross-linking and fibronectin biosynthesis. However, the pathway by which copper ions are delivered to copper-dependent LOX metalloenzymes is still unclear. ATP7A/B protein is used to limit copper toxicity and up-regulates cancerogenic enzymes, such as LOX and LOX-like proteins [182,183].

7. Cancer and Lifestyle

Only 5–10% of cancers are thought to be caused by inherited genetic defects. Numerous cancers are not inherited and are caused by various agents (environmental factors, physical factors, and hormones) [184]. Environmental factors encompass lifestyle (nutrition and overweight, over-smoking, over-drinking, stress, physical inactivity); physical factors (environmental pollutants, virus, bacteria and parasitic infections, ionizing and non-ionizing radiation); as well as socio-economic and attitudinal factors. Therefore, most cancers have multiple possible concurring causes.

A healthy lifestyle includes a healthy diet, weight control, physical exercise, reducing alcohol drinking, and smoking avoidance [185]. About 25–30% of total cancer deaths are caused by tobacco, 30–35% are diet-related, approximately 15–20% are caused by infections, and the rest are attributable to other agents, such as radiation, stress, environmental pollutants, etc. In spite of medical progress, cancer incidence is expected to increase substantially in the near future [186]. It is also thought that all these carcinogens associated with lifestyle factors and all chemopreventive agents are connected with the long-term inflammation. Chronic inflammation is strongly associated with the tumorigenic trajectory, as evidenced by multiple findings [187]. Carcinogens activate while chemopreventive agents suppress NF-κB activation, which is a mediator of inflammation.

All living beings are constantly under stress, which is the nonspecific response of the body to any demand made upon it [188]. Unlike eustress or adaptive stress, distress affects immune responses, generally by exerting a suppressive effect. The stress-induced increases in tumour size are most probably a consequence of immunosuppression [189,190]. Some other authors showed that psychological stress is weakly associated with increased mortality from colon cancer [191]. However, chronic stress is associated with neuroendocrine abnormalities that can up-regulate inflammation and down-regulate protective immunity.
Thus, the affected immune cells may not effectively control cancer cells and act as stromal cells, communicating with the tumour microenvironment and circulating cancer cells to promote tumour growth mechanisms, invasiveness, extravasation into the circulation and metastasis [192].

The response to physical and social stress involves a complex reaction at the cellular and molecular level [193]. NF-κB plays a key role in the cellular response to stress. Thus, stress up-regulates some genes, such as transcriptional genes that control cell growth, chromatin structure, cell cycle activation, and enzymes involved in nucleic acid and protein biosynthesis. Under stress, cell cycle inhibitors, the NF-κB inhibitor, apoptosis-related genes, antiproliferative cytokines, and Apo J are down-regulated. NF-κB is activated in response to many inflammatory factors such as carcinogens, chemotherapeutic agents, cytokines, hormones, mitogens, viral products, eukaryotic parasites, endotoxin, fatty acids, metals, radiation, hypoxia, and psychological, ROS, and chemical stresses [194]. Drugs that prevent cancer or inflammation have been proven to suppress NF-κB up-regulation. Curcumin and other polyphenols inhibit NF-κB, p53 pathways and potentiate Nrf2 activation [195].

Protein p53, which is a universal sensor of genotoxic stress, coordinates the cellular response to various genotoxic stimuli, determining cell death or survival [196]. ROS also appear to be involved in p53 signalling, being effective activators of p53 function. Some chemotherapeutic agents activate p53 due to their involvement in ROS production. However, ROS, generated following p53 activation, play a role in mediating apoptosis [196].

The role of transcription factors nuclear factor erythroid 2–related factor 2 (Nrf2) and nuclear factor-κB (NF-κB) related to OS was also investigated [197]. Thus, in response to OS, the transcription factor Nrf2 up-regulates the expression of antioxidants and detoxifying enzymes involved in antioxidant protection, being considered as the master regulator of redox homeostasis. The activation of the transcription factor NF-κB leads to the production of proinflammatory cytokines and chemokines, prostaglandins, free radicals such as NO and superoxide anions, and ultimately leads to chronic inflammation [198–201].

A healthy lifestyle has been associated with a substantial reduction in the overall risk of developing liver cancer [202]. Lifestyle improvements to combat cancer have long been recommended; however, there has been a renewed appreciation of their importance and relevance given the growing number of cancer survivors seeking alternative options for prevention and secondary cure [203]. Tumour survivors often face drug toxicity, also being at risk of cancer recurrence, a second primary cancer and high cause of mortality [204–207]. Most cancer survivors live with higher risks of complications and relapses, lower quality of life and reduced life expectation. There is an immediate need to improve cancer survivorship by improving lifestyle beyond clinical interventions. The association between a sedentary lifestyle and worsened survival after cancer was also noticed [208,209]. Physical exercises may be positively correlated with the control of tumour biology through specific effects on intrinsic tumour factors, such as Warburg-type high glycolytic metabolism [210]. Tumour metabolism can be selectively influenced by single exercise as well as by regularly applied exercise, depending on the intensity, duration, frequency, and mode of exercise. High intensity anaerobic exercise has been shown to inhibit glycolysis, and some animal studies have shown that the effects on tumour growth may be stronger compared to moderate intensity aerobic exercise. Of course, early detection and treatment can result in growing prevalence of survivors of cancer.

Stress-prone personalities, unfavourable coping styles and negative emotional responses, and poor quality of life were related to higher cancer incidence, poorer cancer survival, and higher cancer mortality [13]. Site-specific analyses indicate that PSF are associated with a higher incidence of lung cancer and poorer survival in patients with breast, lung, head and neck, hepatobiliary, and lymphoid or hematopoietic cancers. These analyses suggest that stress-related PSF have an adverse effect on cancer incidence and survival, although there is evidence of publication bias and results should be interpreted with caution. Some clinical studies have shown that psychological and/or pharmacological inhibition of excessive adrenergic and/or inflammatory stress signalling, especially
Stresses 2023, 3

in conjunction with cancer treatments, would improve prognosis [211]. There are some critical phases of cancer progression that are more sensitive to stress. Therefore, there is a need to focus on more vulnerable populations using individualised pharmacological and psychosocial approaches [212]. Addressing psychosocial stressors also raises the issue of distinguishing between human and laboratory animal cancers [211].

8. Respiration and pH Imbalance

In numerous pathologies, including cancer, an impaired respiration can be observed, along with a change in pH, due to a multitude of stress factors. In order to suggestively explain the imbalance between respiration and glycolysis, which leads to pH alteration, or more precisely, the imbalance of both respiration and pH of body fluids, the so-called respiratory and pH imbalance (RpHI) has been introduced [106]. There are striking similarities between the metabolic profiles of cancer cells and those of rapidly multiplying normal cells, such as aerobic glycolysis and increased biosynthesis [2]. The role of aerobic glycolysis in malignant growth should be elucidated, including whether there is metabolic reprogramming that may be related to chronically sustained proliferation [213]. However, altered metabolism is a hallmark of cancer, and metabolic reprogramming in cancer cells is seen in the main pathways of central carbon metabolism [214]. Different cancers are characterised by an intra-tumour hypoxia resulting from deregulated cell proliferation [215]. Tumour hypoxia is associated with poor prognosis and resistance to therapy [216]. Physiological responses triggered by hypoxia can impact all critical aspects of cancer progression, including immortalization, transformation, differentiation, genetic instability, angiogenesis, metabolic adaptation, autocrine growth factor signalling, invasion, metastasis, and resistance to treatment. Hypoxia-inducible factors (HIF) are key oxygen sensing factors that mediate the response to low oxygen pressure [217]. These transcription factors regulate cellular adaptation to hypoxia and protect cells by reacting acutely and inducing the production of endogenous metabolites and proteins to promptly regulate metabolic pathways. Therefore, hypoxia itself could be the trigger for the induction of aerobic glycolysis without any mitochondrial damage [218,219]. HIF1α activates via Activin/nodal signalling, and its increased expression redirects ATP production from oxidative phosphorylation to glycolysis. In addition, it has been reported that HIF1α-dependent expression of BNIP3 (a member of the apoptotic Bcl-2 proteins) promotes mitophagy to control ROS production and ROS-induced cell death [220].

Hypoxia, which is closely related to glycolysis, is common during carcinogenesis; it is associated with functional and structural modifications in proliferating cells [221]. Hypoxia also underlies the energetics processes of various activities in brain-like alertness, sensory processing, cognition, and physiological conditions. Its specific functions performed in cells are still less understood [100]. Aerobic glycolysis is characterized by excessive glucose utilization relative to oxygen consumption, even when oxygen levels and availability are adequate. Propranolol blocks aerobic glycolysis, including adrenal release of epinephrine, brain signalling through the vagus nerve, and an enhanced liberation of norepinephrine in the locus coeruleus. Sugar utilization is stimulated by norepinephrine and not oxygen consumption.

Glycolysis stoichiometry does not allow both biomass production and lactate generation, and NAD+ regeneration by lactate alone is not possible. It is therefore hard to see how the Warburg effect can directly stimulate biosynthesis. In general, cells allocate half of their genes to synthesize proteins engaged in glycolytic processes [208,222]. However, the cellular biosynthesis programs require lower amounts of protein. Therefore, the cost to produce proteins for aerobic glycolysis may be higher than the cost of producing proteins needed for biosynthesis. There is evidence that mitochondrial functions run concurrently with the Warburg effect, and thus, during aerobic glycolysis, mitochondrial activity is not impeded. It is thus unclear whether the Warburg effect functions to facilitate the various biosynthetic pathways. However, increased consumption of glucose to liberate lactate lowers the pH in the microenvironment [96].
In principle, RpHI can be regarded as a physiological reaction against any stress agents, and if the stressors are strong, an oxygen crisis within the body occurs, the busiest cells divide faster and faster, producing first preneoplastic cells and, in time, malignant tumours. The tumours acquired in this way, as well as their fermentation products, may disturb the normal functions of most cells and tissues in the body [223].

Since there are many forms of cancer depending on the organ or organs affected, the prognosis of the disease, and its stages, there are many forms of RpHI. Thus, a carcinogen can cause irritation of a tissue, followed by chronic inflammation and malignancy, while slow debilitation can lead to degenerative disease and, ultimately, to cancer. One can also speak of an increased concentration of glucose in cells and body fluids which can ferment in the presence of insufficient amounts of oxygen in the tissue although the patient’s breathing may appear normal. Thus, epinephrine (adrenaline) secretion during stress may explain the increased flow of glucose into certain active tissues, and higher concentrations in relation to oxygen intake may lead to fermentation and lactic acid formation. Therefore, a full description of RpHI requires further work.

Thus, under hypoxic conditions, overstressed cells, which receive less oxygen than necessary, undergo anaerobic fermentation to produce adenosine triphosphate (ATP). This process is associated with excessive multiplication and, ultimately, tumour growth. Indeed, severe hypoxia due to profoundly low arterial O₂ content (hypoxemia) results in hypercapnic and metabolic acidosis, developed together with extensive lactate generation, with pH decreasing to under 6.8 [224]. Because hypoxia is dependent on the magnitude and duration of action of the causative factors, human and animal organisms can only compensate for hypoxia if the causative agents stop acting for a long time. Normally, various stress agents, such as physical and chemical stressors, viruses, other infectious agents, hormones in excess, but also, long-lasting anxiety, emotions, conflict states, etc., are able to affect the body as a whole [225–230].

Confronted with the external environment, the living organisms have several coordinate physiological processes to keep their internal states of equilibrium [231]. The living bodies react against aggression using metabolic energy obtained by the oxidation of organic substances, including organic acids in the mitochondria [232]. Under normal conditions, when the supply of oxygen is sufficient, cells can carry out aerobic respiration [233,234]. Hence, when busy, cells produce the energy they need from the food stores, including the organic acids they possess. During the Krebs cycle, carbon dioxide, of a weaker acid type, is released, normally outward [235,236]. Since stronger organic acids, such as succinic, malic, 2-ketoglutaric, and oxalylacetic acids, are replaced by carbon dioxide; the cell milieu tends to be more alkaline, i.e., just blood and urine [237]. The tendency of blood alkalization entails retention of carbon dioxide so that the pH of blood should alter as little as possible. The retention of carbon dioxide in blood does not allow the oxygen to shift at a normal rate in lungs [238,239]. As a consequence, oxygen pressure in blood and tissues decreases although it stays normal overall. However, oxygen may reach quite a low level without affecting cell breathing in any drastic way. Unfortunately, the busiest cells in the body need more oxygen. As oxygen partial pressure decreases lower, these cells receive less oxygen than they need, and fermentative processes develop alongside with breathing.

In fact, hypoxia is the natural environment in which DNA auto-replication and transcription take place in vivo in all eukaryotes [240–243]. Nuclear division unfolds anaerobically using the energy produced by glycolysis [244–246]. Consequently, the cells forced to manifest themselves in rather anaerobic conditions will divide more intensely [247]. Aerobic breathing, which provides cells with a great amount of energy, creates the necessary conditions for the existence of fine structures of the cells, and the specific functions run unimpaired [248]. Lack of oxygen, even partial, causes rupture of these structures, leading to the gradual disappearance of specific functions of cells as well as contact inhibition. At the same time, lack of oxygen entails cell-division to a greater extent than necessary for the tissue in question. It follows the first stage of the RpHI, which affects cell respiration and division. Nevertheless, the organism possesses buffer systems, lung
ventilation, and kidney mechanisms to control the concentration of hydrogen ions within the cellular milieu (Figure 1) [2,249]. Alveolar ventilation is responsible for carbon dioxide elimination [250]. Mild acidosis occurs primarily when alveolar airflow is reduced or if CO₂ generation is elevated. However, the organism has several compensatory systems to minimize a decrease in pH. For example, non-oxygenated haemoglobin easily buffers the blood environment to prevent significant pH alterations. Normally, carbon dioxide, a stimulant of respiration, induces an increase in minute ventilation to normalize the pH by eliminating increased quantities of CO₂. Unfortunately, this effect is mitigated when CO₂ concentrations remain elevated for more than a few hours. The kidneys are also capable of controlling both the blood pH and some other blood parameters. However, this process is slow and lasts for several hours or days. In fact, renal compensation begins in 6–12 h, but maximal compensation occurs in 3–5 days. The kidneys enhance the expulsion of protons, predominantly as ammonia. If the stress agents act continuously, the blood will become slightly more alkaline than usual, and the blood oxygen concentration will be lower than normal. Getting less oxygen than they need will lead to anaerobic fermentation in the overstrained cells. The overstrained cells also cause a lower content of NADH+H⁺ and NADPH+H⁺ a higher content of NAD⁺ and NADP⁺. Therefore, a decrease of the oxidation reduction potential will occur as well. The quantity of sulphhydryl groups in the blood and tissues also decreases. A marked decrease in succinic dehydrogenase and slight increase in cytochrome oxidase levels could be found, suggesting the alteration of the Krebs cycle. For this reason, a cell with excessive fermentation will not reach an upper energetic state if neighbouring cells and blood do not interfere with its metabolism. The second stage of the RpHI is reached when lactic acid is produced due to hypoxic conditions. In this case, the CO₂ concentration may decrease; however, part of the produced carbon dioxide is not removed because the cell content may remain slightly alkaline in spite of the lactate production. Again, the kidneys should control the hydrogen concentration in blood, releasing acidic species, such as ammonium ions or phosphates, into urine [251]. The process is complicated by the existence of lactic acid in blood which decreases blood pH, while intracellular pH of overstrained cells is increased. However, if the stress agents act continuously, the neighbour cells involved in curing or rebalancing the overworked cells will also work hard while being overstrained and deprived of oxygen. There follows a third stage of the RpHI in which a real state of illness (infections, viruses) occurs. A very special balance between the two types of cells (attacked cells and neighbour ones) is established. The blood pH value is a little altered. If the stress agent is very strong, it can be lethal to the organism, and such a case does not reach the cancer stage. It is the case of microorganism- or virus-induced diseases, which, if untreated, can have a bad prognosis. On the contrary, a long-standing action by the stress agents may cause a slow shift in this imbalance even if the action is mild, resulting in a stepwise decrease of blood pH value. Long-standing infectious pathologies can thus lead to an RpHI, which creates the conditions for either the transformation of normal cells into preneoplastic cells, followed by the preneoplastic to neoplastic pathway, or the multiplication of malignant cells. It is well-known that only bodies with a significant amount of morbidity may become cancerous. Most cells become glycolytic, while fewer remain normal but overstressed. This pathway leads to the fourth stage of the RpHI, which is that of tumour formation.

Respiration and glycolysis are two independent biochemical processes that may occur simultaneously in the living cell (Figure 2). Given an oxygen concentration of 10 per cent or more in the surrounding atmosphere, respiration occurs in the living cell. Given a concentration below 3% of oxygen, fermentation (glycolysis) will occur. Both processes occur in the range from 3% to 10% oxygen (Figure 2). Growth and lifespan of human diploid cell strains at oxygen levels below 20% is increased, and an enhancement of around 25% in the lifetime of both cell types has been achieved by long-term cultivation under 10% oxygen [252]. It is well known that there is extremely low O₂ content in growing tumours [253].
Indeed, if mitochondrial respiration in tumour cells were down-regulated, the accumulation of substrates from the Krebs cycle could also serve as a signal to stimulate glycolysis [254].

Whenever a cell within the metabolic system (muscle, liver, kidney, lung, etc.) uses oxygen at a faster rate than can be provided by the circulatory system, the cell begins to function anaerobically, reducing the pyruvate to lactate instead of oxidizing it further, as would happen if oxygen supplies were adequate [255–257]. Moreover, excess glucose can be glycolytically converted to lactic acid if the glucose: oxygen ratio increases. Lactate thus accumulates in that cell, diffuses through the bloodstream, and eventually reaches the liver, where it is re-oxidised to pyruvate and converted to glucose via the gluconeogenic pathway [258].

In fact, prokaryotic cells produce energy for their needs by glycolysis in the absence of oxygen. Glucose is thus metabolized into lactate or ethyl alcohol, depending on the cell type. In the cytoplasm of eukaryotic cells, including human cells, glycolysis takes place with the release of pyruvic acid or lactic acid [259,260]. These acids enter the mitochondria where they are degraded to CO₂ and H₂O, with the formation of reduced forms of NADH+H⁺.
and FADH₂. In the respiratory chain, the hydroxyl-rich compounds are oxidized and the chemical energy they contain is liberated and stored as ATP molecules. If there is insufficient oxygen, then the respiratory chain is blocked and NADH+H⁺ and FADH₂ are no longer oxidized. Under these conditions, no new quantities of reduced NADH+H⁺ and FADH₂ are formed; lactic and pyruvic acids accumulate in cells or are excreted extracellularly. This is a very simple mechanism for switching from respiration to glycolysis \[258,261–263\].

Figure 3 better suggests the molecular and biochemical pathways linking respiration and glycolytic processes to oxygen supply and hypoxia. The human cell is made up of the nucleus, which plays a role in the division and transmission of genetic information, the cell membrane, through which oxygen, glucose and other nutrients flow, and the cell organelles. Among the latter are mitochondria, which have an essential role in energy production in the form of adenosine triphosphate (ATP) molecules, oxidative degradation of fatty acids, metabolism of pyruvic acid and acetyl-Coenzyme A from fatty acids with the formation of NADH+H⁺ and FADH₂ molecules. These reduced forms of nicotinamide dinucleotide (NAD⁺) and flavin adenine dinucleotide (FAD) compounds are involved in cellular energetics and multiple biochemical syntheses and constitute the fuel from which ATP is formed in the so-called electron transport chain (ETC). Glycolysis takes place in the cell cytoplasm and not in the mitochondria.

Glucose is degraded via the glycolytic pathway to pyruvate, which is either used as acetyl-CoA in mitochondria or released into the cytoplasm as lactate (Figure 3). Most of biochemical reactions are reversible. Several enzymes catalyse the conversion of glucose to pyruvic acid (PYR) with the release of two molecules of ATP. Pyruvic acid then reacts with Coenzyme A to form acetyl-CoA, which is broken down in the Krebs cycle with the release of CO₂, while its hydrogen atoms reduce NAD⁺ and FAD to NADH+H⁺ and FADH₂. These energy-rich molecules are used in the electron transport chain to form ATP.

Figure 3. Schematic presentation of the biochemical process of spontaneous transition from ATP formation by respiration (OXPHOS) to ATP production by glycolysis, depending on oxygen availability. Here: G6P-glucose-6-phosphate, F6P-fructose-6-phosphate, FBP-fructose-1,6-bisphosphate, GAP-glyceraldehyde-3-phosphate, DHAP-dihydroxyacetone phosphate, BPG-1,3 bisphosphoglycerate, GAP-glyceraldehyde-3-phosphate, DHAP-dihydroxyacetone phosphate, BPG-1,3 bisphosphoglycerate, 3PG-3-phosphoglycerate, 2PG-2-phosphoglycerate, PEP-phosphoenolpyruvate, PYR—pyruvate, Lac—lactate, HK—hexokinase, PGI—phosphoglucoisomerase, PFK—phosphofructokinase, ALD—aldolase, TPI—triosephosphoisomerase, GAPDH—glyceraldehyde-phosphate dehydrogenase, PGK—phosphoglycerate kinase, PGM—phosphoglycerate mutase, ENO—enolase, PK—pyruvate kinase, LDH—lactate dehydrogenase, O₂—molecular oxygen, ADP—adenosine diphosphate, ATP—adenosine 5’-triphosphate, OXPHOS—oxidative phosphorylation, PDH—pyruvate dehydrogenase.
of CO₂, while its hydrogen atoms reduce NAD⁺ and FAD to NADH+H⁺ and FADH₂. These energy-rich molecules are used in the electron transport chain to form ATP in the presence of molecular oxygen. However, if oxygen is insufficient, the reduced NADH+H⁺ and FADH₂ species cannot be used in ETC to generate the energy-rich molecules of ATP. The Krebs cycle is thus blocked, and acetyl-CoA is no longer needed. As a result, the pyruvic acid formed in the glycolytic process is no longer required in the mitochondria and is reduced to lactic acid by the existing NADH+H⁺. In this way, glycolysis occurs in the presence of insufficient oxygen levels to provide the required ATP, but glucose consumption is greatly increased for the same ATP concentrations required by the cells. Furthermore, the increase in NADH+H⁺, produced in the Krebs cycle due to hypoxia, inactivates phosphatase and tensin homolog (PTEN), which is encoded by the PTEN gene [264]. PTEN is involved in the regulation of the cell cycle, preventing cells from growing and dividing too rapidly. Thus, a close link between the Warburg effect and metabolic alterations in cancer cells has been found; it may gain a survival advantage and withstand therapeutic agents. The microenvironment of solid tumours is characterised by hypoxia, high lactate levels, extra-cellular acidosis, and depletion of glucose and glutamine [220]. Nevertheless, hypoxia might be responsible for the autophagy induction in tumour cells via HIF1α. NRF2 promotes HIFx activation, the metabolic switch, and colony formation [265]. ROS-induced NRF2 activates HIFx and drives the metabolic switch toward glycolytic energy production. However, further research is needed as such phenomena may be secondary to all physiological reactions.

However, if the glucose level is high enough, fermentative processes may occur due to the presence of glycolytic enzymes. Therefore, the so-called RpHI appears to be more complex and may comprise several biochemical pathways, which are interdependent. Therefore, this review suggests that aerobic glycolysis and malignant transformation could be controlled, at least in their early stages.

### 9. Discussion

Stress factors influence the body neurochemically, hormonally, and immunologically, and these factors have an impact on the carcinogenic process, suggesting a relationship between them and stress-induced changes in tumour growth [266]. Social stress influences tumour growth [267]. Reactive oxygen species (ROS) are also stressors that play important roles in a variety of normal biochemical functions and abnormal pathological processes [268]. Thus, ROS can induce cellular aging and cell death [269]. Conversely, an increase in ROS is related to an abnormal growth of cancer cells and indicates a disturbance of redox homeostasis, either due to an increase in ROS output or a decrease in ROS scavenging activity [270]. When ROS increases to a certain critical threshold that is incompatible with cell survival, ROS can cause a cytotoxic action, leading to cancer cell death and reduced cancer proliferation. Nevertheless, under intrinsic oxidative stress, most cancer cells adapt well to such stress and develop enhanced endogenous antioxidant ability.

Thus, low levels of ROS can increase the ability of cells to cope with stress, while an increase in ROS can cause damage to normal cells that can be killed or transformed into cancer cells. Furthermore, an exaggerated increase in ROS concentration can cause apoptosis of normal cells, but not cancer cells. Only very high amounts of ROS work as anti-tumorigenic agents [271]. Adequate levels of ROS are essential as excess ROS damages cellular membranes and nucleic acids [272]. Inadequate levels of ROS disrupt signalling mechanisms, which are useful for cell growth-like inactivating phosphatases and tensin homologues as well as tyrosine phosphatases. The Warburg effect can alter the redox potential of mitochondria, leading to ROS formation [273].

Hypoxia can induce enzymatic breakdown of cellular constituents into simple subunits, a phenomenon capable of sustaining glycolysis to maintain cellular ATP production [220].

It has been hypothesized that the primary reason for cachexia is elevated acidity of body tissues, which leads to increased and non-specific proteolysis of cell proteins. Hence, moderate hypoxia may be tightly linked to lactic acid formation throughout the body, not just around the cancer cells [274]. Indeed, hypoxia promotes acidosis by shifting from
oxidative phosphorylation to glycolytic metabolism [275]. Inhibition of mitochondrial respiration induces increased NADH+H⁺ concentration, which can subsequently inactivate PTEN (phosphatase and tensin homologue) through a redox modification mechanism [264]. Cachexia can be a progressive body wasting disorder marked by loss of adipose tissue and skeletal muscle tissue in cancer, infection, acquired immunodeficiency status, and heart congestion [276,277]. Sarcopenia judged by skeletal muscle mass volume is a prognostic marker in some cancer patients [278,279]. It is related to ageing, but can also be caused by poor nutritional status, and inflammatory, endocrine, and malignant diseases [280]. The relationship between cancer and sarcopenia is well-recognized. Numerous inflammatory agents that facilitate tumour progression are also associated with cancer cachexia, pain, weakness, and poor survival.

Inflammation is a critical component of tumour progression [281]. The role of inflammation in the pathogenesis of various diseases has also been examined [282]. Inflammatory responses may occur acutely following traumatic tissue injury or infection, or may be induced chronically by malignant cells, degenerative alterations or tissue ischemia due to oxygen deprivation [283]. Many cancers arise from areas of infection, chronic irritation and inflammation. Inflammatory cells clearly participate in the neoplastic process, promoting proliferation, survival, and migration. Furthermore, tumour cells take up innate immune system signalling molecules, such as selectins, chemokines, and their receptors for invasion, migration, and metastasis. Therefore, an anti-inflammatory therapeautic approach can be considered in cancer.

DNA damage mediated by chronic inflammation increases cytokine expression or ROS release contributes to type 2 diabetes, heart disease, various cancers, and stroke [282]. The release of proinflammatory cytokines, such as TNF-α and IL-1, modulates innate immune cells that release inflammatory mediators, chemokines, interferons, recruited neutrophils, and adhesion molecules. Figure 4 shows a suggestive scheme illustrating the relationship between stressors, chronic inflammation, the Warburg effect, and various medical conditions. TNF-α stimulates COX-2 expression and nitric oxide synthesis by activating NF-kB [284,285]. COX-2 catalyses the synthesis of inflammatory prostaglandins (PGs) from arachidonic acid, which in turn causes chronic inflammation [286].

Figure 4. Suggestive presentation of the relationship between stressors, chronic inflammation, the Warburg effect and various medical conditions such as arthritis, obesity, myocardial infarction, stroke, cancers and others. Stressors include heavy metals, psychosocial stressors, carcinogenic chemicals, asbestos, viruses, bacteria, etc.
Disturbed copper homeostasis is seen in many types of cancer. It may be related to increases or decreases in protein status. Moreover, copper consumption by growing cancer cells increases. Both proteins involved in copper metabolism and copper-containing proteins are exposed to multiple dysregulations, which results in higher carcinogenicity. Oxidative stress, aerobic glycolysis, hypoxia, etc., create conditions for increased acidity around cancer cells and body fluids and increased intracellular pH. Copper ions react with phosphate ions with the formation of insoluble copper phosphate, resulting in decreased copper concentration in some biological fluids. Excess copper, relative to the amount of phosphate available, can react with lactic acid produced by glycolysis. The result would be increased copper ion concentration in other media. In addition, as the pH of a solution increases, copper ions increasingly bind to proteins and peptides to form complexes. This may explain the measurements of variable copper concentrations in patients with malignant tumours.

However, a correct understanding of biochemical and physiological processes that manifest in a cancer pathology is only possible if our conception of living organisms is greatly improved. Eugen Macovschi advanced a so-called biostructural theory on cancerogenesis [287,288]. In addition, older hypotheses and theories, all based on the molecular outlook on living systems, should be replaced by others that are more adequate to understanding living phenomena [289]. For example, carcinogens in the environment, acting on living tissues, cause partial, sometimes reversible, breakdown of the biostructure described by Eugen Macovschi [290]. They cause cellular hypoxia which induces alteration of the state of the biostructure, and thus, of living matter. The whole body is affected, and malignant cells are in fact normal cells that receive less oxygen than necessary and undergo glycolysis, dividing more than necessary. Therefore, carcinogens that come into contact with living tissue must be removed at all costs. These carcinogens cannot be destroyed or removed by the body alone despite its exhausting efforts. The main cause of cancer would thus be a so-called respiration and pH imbalance, and not gene mutations, as the molecular theory claims.

Current molecular medicine relies on physico-chemical laws to investigate biological phenomena associated with cancer, which are thought to occur only at the molecular level of living organisms. Only chemical reactions take place there [223]. However, a structural-phenomenological outlook seems to be more appropriate to illustrate the observed aspects of living organisms and the relationships between the biological levels and soul (psychostructural level), mind (noesistructural level) or between mind and consciousness. In other words, cancer can be conceived as a phenomenon which occurs on both biological (biostructure) and physico-chemical structures, while cancer aetiology seems to be related to specific breakdown of the biostructure of the whole organism. The cancer cell biostructure is found under an altered, abnormal state and the investigation of this state is essential to understand carcinogenesis. In addition, a mathematical hypothesis of networks of multidimensional hierarchic evolution with various ranks was advanced, and the self-organization of living was analysed in the frame of Macovschi’s biostructural conception [291,292]. The network’s complexity varies horizontally, within the same level, and vertically, from the lower to the upper level.

The Warburg effect appears to be a normal physiological process due to the low oxygen supply relative to the needs of rapidly multiplying cells. Nerve cells and circulatory system cells are exceptions to this rule. Their overload usually leads to their destruction as opposed to excessive multiplication. Malignant cells adapt to the partially anaerobic environment, undergo protein degradation processes, and are unable to behave normally in the presence of oxygen. The phenomenon of excessive multiplication is similar to wound healing. The cells involved are too overworked to cope with the stress of the external environment, receive too little oxygen in relation to their increased needs, multiply excessively, and then the body eliminates the unnecessary cells. In the case of cancer cells, the body is no longer able to destroy cells that have multiplied in excess of the needs of the tissue in question.
Pathways leading to increased glycolysis can also cause inhibition of mitochondrial activity [293]. HIF-1 is thought to stimulate essential glycolysis pathways, but also regulates genes that control angiogenesis, cell survival, and invasion. However, high levels of HIF-1 are observed in some tumours, even in the presence of oxygen. Therefore, not only hypoxia, but also other factors (e.g., hormones and growth factors) could induce stabilisation of HIF-1 expression [294].

The cell’s response to hypoxia is also controlled by HIF-1, which activates expression of specific genes involved in angiogenesis, glucose uptake, glycolysis, growth factor signalling, apoptosis, invasion, and metastasis [295]. Hypoxia can induce enzymatic breakdown of cellular constituents into simple subunits, a phenomenon capable of sustaining glycolysis to maintain cellular ATP production [220]. Thus, HIF-1 not only stimulates glucose influx and utilization in tumour cells, but also stabilizes mitochondria through various mechanisms. Stimulation of mitochondrial activity would cause cellular energy metabolism to return to the phenotype characteristic of non-malignant cells and would also promote ROS production by mitochondria, leading to apoptotic cell death of tumour cells [218,296]. We believe, however, that hypoxia causes the breakdown of the triplet states of the biologically active molecules in ECT responsible for the electromagnetic transfer of energy from NADH and FADH$_2$ to ATP [297]. This hypothesis is also supported by the fact that hypoxia greatly reduces the number of mitochondria in the body’s cells [298,299].

The anti-cancer effect of many conventional treatments, such as ionising radiation, etoposide, and arsenic trioxide, is based on the stimulation of ROS production [218,300]. According to the General Adaptation Syndrome (GAS) described by Hans Selye, there are different stages of stress, namely (i) alarm, (ii) resistance, and (iii) exhaustion [189]. The psychological factors can also play a significant role in the stress process. Prolonged stressors can cause psychosomatic disorders, depending on their intensity and duration. Under the action of stressors, at any stage, a person can die. However, we believe that there may be a fourth, cancerous stage in which the body fluids become more acidic due to fermentative processes.

Studies during the past decade suggest that the WE is more closely related to alterations in signalling pathways that govern glucose uptake than to mitochondrial defects [93]. Although glycolysis is indeed greatly increased in cancer cells, mitochondrial respiration continues to function normally at rates proportional to oxygen uptake. There is, instead, an up-regulation of glycolysis, not a switch from OXPHOS to glycolysis [301].

10. Concluding Remarks

A substantial body of research links stressors, including psychosocial factors, to the Warburg effect and increased cancer incidence. The literature findings reviewed here support the idea of a close relationship between oxidative stress, or other forms of stress, hypoxia, aerobic glycolysis, and carcinogenesis. Currently, there are all the prerequisites for a correct understanding of the aetiology of the various forms of cancer and their development and treatment. However, given the multitude of cancer-causing factors as well as different forms of cancer, it is difficult to design a unified theory of oncogenesis. In addition, theories based solely on chemical reactions appear to be unable to provide a satisfactory explanation for the relationship between stress and disease. A nature-of-life approach is possible, which takes into account the whole biological phenomenology of disease and not just its molecular aspects. In this brief review, we have sought general information, leaving aside some particular aspects of several forms of cancer, although these too would have better completed the picture of this disease. From the few findings reviewed here, a picture emerges in which the importance of physiological and biochemical aspects is highlighted. The role of hypoxia and the way in which it occurs and manifests itself has thus been highlighted in greater depth, although it is more likely to be a matter of oxygen insufficiency of overstimulated cells. It was highlighted that respiration and glycolysis are two biochemical processes that can occur simultaneously in living cells, and aerobic glycolysis also takes place under normal physiological conditions. The Warburg effect can
thus be considered a normal physiological process due to the low oxygen supply in relation to the needs of the overworked cells. The recently advanced physiological mechanism may better explain the Warburg effect than the older theories on cancer occurrence. Starting from molecular biology and medicine, supramolecular theories could be developed, followed by improved biostructural and structural-phenomenological concepts of the disease state to fully understand and design revolutionary therapies for different forms of cancer.

**Funding:** The present research has not received any external support.

**Data Availability Statement:** Not applicable.

**Acknowledgments:** The authors greatly appreciate the constructive suggestions of the reviewers, which have very much contributed to the improvement of this review. In addition, GD greatly addresses the Editorial Team who greatly improved the style and English language of this work.

**Conflicts of Interest:** I declare that there is no conflict of interest associated with this review.

**References**


98. Pinto, M.M.; Paumard, P.; Bouchez, C.; Ransac, S.; Duvezin-Caubet, S.; Mazat, J.P.; Rigoulet, M.; Devin, A. The Warburg effect and mitochondrial oxidative phosphorylation: Friends or foes. BBA-Bioenergetics 2023, 1864, 148931. [CrossRef]


107. Epstein, T.; Xu, L.; Gillies, R.J.; Gatenby, R.A. Separation of metabolic supply and demand: Aerobic glycolysis as a normal physiological response to fluctuating energetic demands in the membrane. Cancer Metab. 2014, 2, 7. [CrossRef]


149. Lelièvre, P.; Sancey, L.; Coll, J.L.; Deniaud, A.; Busser, B. The multifaceted roles of copper in cancer: A trace metal element with dysregulated metabolism, but also a target or a bullet for therapy. *Cancers* 2020, 12, 3994. [CrossRef]


269. Williams & Wilkins: Baltimore, MD, USA; Philadelphia, PA, USA, 2007; pp. 15–24.


274. Drochioiu, G. Chronic metabolic acidosis may be the cause of cachexia: Body fluid pH correction may be an effective therapy. Med. Hypotheses 2008, 70, 1167–1173. [CrossRef]


**Disclaimer/Publisher's Note:** The statements, opinions and data contained in all publications are solely those of the individual author(s) and contributor(s) and not of MDPI and/or the editor(s). MDPI and/or the editor(s) disclaim responsibility for any injury to people or property resulting from any ideas, methods, instructions or products referred to in the content.