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Cross-disease communication between cancer and heart failure provides a rational approach to prevention and treatment of both diseases

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Accumulating clinical data have demonstrated a clear positive association between cancer and cardiac disorders, particularly chronic heart failure (CHF). These two diseases can be mutual drivers of each other, and hence frequently co-occur in patients. The immune system is the core mechanism that eliminates transformed cells from our bodies. However, immune cells often play distinct or even conflicting roles in cancer and CHF. Moreover, CHF alters the properties of immune cells, particularly those of regulatory T cells. Our previous study showed that the oxidative phosphorylation capacity of peripheral blood mononuclear cells is impaired in CHF, leading to the increased production of reactive oxygen species. Therefore, the co-occurrence of cancer and CHF becomes a serious problem, affecting the treatment of both diseases, and consequently negatively affecting patient survival rates. To date, few methods have been identified that effectively treat both diseases at the same time. Mitochondria activity may change in immune cells during their activation and exhaustion, and in CHF. Mitochondria activity is also largely affected in myocardia in CHF. We here focus on the mitochondrial abnormalities of immune cells in cancer and CHF, and discuss possible ways to treat cancer and CHF at the same time by targeting mitochondrial abnormalities. Many cancer cells are inevitably produced daily in our bodies, mostly owing to enzymatic nucleotide errors of DNA replication and repair. Therefore, the possibility of ways to prevent cancer by preventing the onset of heart failure will also be discussed.

KEYWORDS
immune-checkpoint inhibition, mitochondrial oxidative phosphorylation, disease prevention, myokine, PBMC, exercise, diet, reactive oxygen species
Introduction

Cancer and cardiac disorders, including chronic heart failure (CHF), represent two major causes of morbidity and mortality in developed countries (1, 2). Epidemiological studies have shown that the risk of developing cancer in patients with CHF is approx. four times greater than in those without CHF (3–6). Conversely, cancer patients can be at increased risk of cardiac disease due to deterioration of their lifestyle behaviors (e.g., inactivity and an unbalanced diet) (7), and also due to treatment toxicity, as many anticancer drugs are known to cause cardiotoxic side effects (8–11). Therefore, cancer and cardiovascular diseases can be mutual disease drivers, and hence co-occur frequently in patients (Figure 1). Moreover, immune cells, particularly regulatory T (Treg) cells, play distinct or even conflicting roles in cancer and CHF (12, 13).

Hence, the co-occurrence of cancer and cardiovascular disease is a serious problem, affecting the treatment of both diseases, and consequently negatively affecting survival rates (14, 15). To date, however, treatments exist only for each disease. Mitochondria are central to ATP production by oxidative phosphorylation (OXPHOS) and to metabolism. To address above problems, we here focus on the mitochondrial abnormalities of immune cells during CHF and cancer, and discuss possible methods to treat cancer and CHF at the same time by targeting these mitochondrial abnormalities; and, moreover, discuss possible ways to prevent cancer by preventing the onset of CHF.

Immune system mediates the crosstalk between cancer and CHF

T-cell dysfunction, particularly of tumor-infiltrating lymphocytes (TILs), is highly detrimental to antitumor immunity and immunotherapy (16). Recently, Koelwyn et al. reported that the adjusted relative risk of death from breast cancer is increased by approx. 60% in the presence of a cardiovascular event (17). They also demonstrated by using mouse models that myocardial infarction (MI), which leads to HF, accelerates breast cancer development (17). Molecularly, it was shown that MI epigenetically reprogrammed Ly6C(high) monocytes, which are macrophage precursors in the bone marrow reservoir, to an immunosuppressive state, and increased their circulation and infiltration into tumors, whereas their depletion abrogated tumor growth (17). Moreover, tumors of MI mice had fewer T lymphocytes than control mice, in which T(reg) cells are predominant. These changes that occur in MI mice may be beneficial to the heart, but they all promote tumor growth and survival (17). Therefore, certain populations of immune cells clearly play a central role in cross-disease communication between cancer and CHF.

CHF affects mitochondrial OXPHOS of immune cells

Mitochondrial OXPHOS plays a central role in lymphocyte activity (18). Mitochondria are also fundamental to the development and fate determination of peripheral lymphocytes (19, 20). Suppressed glycolysis and OXPHOS were shown to be early drivers of CD8+ T-cell exhaustion (21). Moreover, TILs are constantly exposed to tumor antigens, and may also experience metabolic stress, which is thought to occur frequently in the tumor microenvironment. A recent report demonstrated that continuous antigen stimulation together with hypoxia impairs the mitochondrial functions of T cells, and hence promotes terminal T-cell exhaustion (22).

Our group has found that mitochondrial respiratory capacity of peripheral blood mononuclear cells (PBMCs), which are predominantly lymphocytes, declines with the progression of CHF, with class III (i.e., moderate to severe CHF) patients by New York Heart Association (NYHA) criteria having 10-24% lower mitochondrial respiratory capacity than NYHA class I/II (i.e., mild CHF) patients, in which mitochondrial ROS production of PBMCs was increased by 13-24% in patients with NYHA class III compared to those with NYHA class I/II (24). Such changes were observed even in the early stages of HF, and were closely associated with the severity of CHF. We moreover found that the capacity of complex II, but not complex I, of the mitochondrial OXPHOS of PBMCs was specifically decreased in CHF (24). It has been reported in monkeys that there is a close association among the mitochondrial OXPHOS activities of circulating monocytes, cardiac cells, and skeletal muscle cells (25). Therefore, ROS levels in PBMCs can be a marker indicating the onset and the severity of HF. As PBMCs mostly consist of unprimed lymphocytes, it awaits to be clarified whether activated lymphocytes are also affected in CHF patients.

Activating mitochondria of immune cells improves tumor immune therapies

Mitochondrial function of CD8+ T cells in lung cancer patients can be a marker for determining the efficacy of anti-PD-1 immune checkpoint inhibition therapy (26). Scharping et al. have shown that restoration of mitochondrial activity and T-cell function by reversing the loss of PGC-1α in tumor-specific T cells resulted in increased antitumor immune responses (23). Yu et al. demonstrated that administering nicotinamide riboside (NR), a
precursor of nicotinamide adenine dinucleotide, may be able to
restore mitochondrial activity, prevent T-cell exhaustion, and
sustain the antitumor responses of T cells in tumor-bearing
mice (27). NR supplementation was moreover found to facilitate
antitumor immune activity, when used in conjunction with the
anti-PD-1 antibody (27). Vardhana et al. demonstrated that N-
acetylcysteine (NAC), which is known to increase glutathione
synthesis and neutralize ROS, reverses the metabolic defects of
exhausted T cells, and promotes their antitumor immune activity,
to act synergistically with anti-PD-L1 immunotherapy in
lymphoma and melanoma (28). Therefore, activating immune
cell mitochondria may improve the efficacy of immune checkpoint
inhibition-based tumor immunotherapy. However, it should be
noted that the administration of molecules such as NR or NAC
may also activate cancer cells to more malignant states, and it is
hence unclear whether they will be effective in the treatment of
patients. It is also well documented that the reinvigoration of T
cells, once they are deeply exhausted, might be very difficult (29).
Another way to improve the efficacy of cancer immunotherapy
would be to enhance the new production of T cells, and diversify
the T-cell receptor repertoire, as has been demonstrated with
radiation (30), but this might also be difficult in patients with CHF
because of their poor health condition.

Future perspectives

When cancer and CHF coexist, the treatment of either disease
alone is inadequate. Normalization of mitochondrial activity and
the function of immune cells, which are frequently impaired in
CHF, is a rational strategy to improve cancer therapeutics. For
example, identification of a molecular basis for the downregulation
of mitochondrial respiratory capacity in the PBMCs of CHF
patients, which we have shown previously (24), and if such a
mechanism occurs specifically in PBMCs but not in tumor cells,
improving mitochondrial respiratory capacity in PBMCs may be
promising for the treatment of cancer in patients who also have
CHF. Such a strategy targeting immune cells’ mitochondria may
also enhance tumor growth suppression in cancer treatment by
immune checkpoint inhibitors, although cardiac assessment with a
careful follow-up is necessary because immune checkpoint
inhibitors are known to have a cardiotoxicity with low incidence
rate (<1%) with single use of them (31). Furthermore, activation of
immune cells is beneficial for chemotherapy (32), and thus,
immuncell-targeted treatment strategy may help
chemotherapy improve outcomes of cancer patients with or
without CHF, although robust clinical evidence is still lacking.

Lifestyle habits, such as a proper diet and daily exercise are
important preventive measures of cancer and CHF. Regarding
the molecular bases, skeletal muscles secrete various myokines,
which have positive effects on mitochondria in different organs
and tissues, and may also promote immunity (33–35). Proper
exercise by patients can also suppress tumor growth and
promote anti-tumor immunity, and may improve the
therapeutic effects of immune-checkpoint inhibitors, whereas
the types of myokines and immune cells therein involved have
been shown to differ depending on the types of cancer (36–38).
On the other hand, muscle dysfunction occurs not only in CHF

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**FIGURE 1**

Cancer and HF mutually promote each other. An unhealthy lifestyle contributes to the development of cancer and CHF, and they are mutual
disease drivers. Anticancer drugs and regulatory T cells appear to have conflicting roles in cancer and HF. Mitochondrial function and reactive
oxygen species (ROS) production in immune cells are potential therapeutic targets in both diseases.
References


