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Commentary

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Epoxyeicosatrienoic acids: a double-edged sword in cardiovascular diseases and cancer

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High dietary fat intake is reported to be associated with several human diseases, including diabetes and heart disease. Moreover, epidemiologic and experimental observations support the hypothesis that high dietary fat intake is also a risk factor for cancers. However, the mechanisms underlying the link between high dietary fat intake and cancer progression are poorly understood. One factor thought to be involved is arachidonic acid (AA), a major component of animal fats that is primarily found in red meats, egg yolks, and organ meats. The bioactive lipids derived from AA play critical roles in cancer progression (1).

Eicosanoid synthesis pathways

AA is a polyunsaturated omega-6 fatty acid that constitutes the phospholipid domain of most cell membranes. AA is liberated by cytoplasmic phospholipase A₂ (cPLA₂) (Figure 1). Free AA can be metabolized to eicosanoids through three major pathways: the prostaglandin-endoperoxide synthase/cyclooxygenase (PTGS/COX) pathway, the lipooxygenase (LOX) pathway, and the cytochrome P450 (CYP) pathway. Prostanoids are the eicosanoids generated by the PTGS/COX pathway, while the LOX pathway generates leukotrienes and hydroxyeicosatetraenoic acids (HETEs). The CYP enzymes that convert AA into eicosanoids include CYP epoxygenase and CYP o-hydroxylase enzymes. CYP epoxygenases, such as members of the CYP2C and CYP2J families, metabolize AA to four biologically active epoxyeicosatrienoic acids (EETs) (5,6-EET, 8,9-EET, 11,12-EET, and 14,15-EET). CYP ω-hydroxylases, such as members of the CYP4A and CYP4F families, convert AA to HETEs. Among the members of the CYP2C and CYP2J families, CYP2C9 are the predominant epoxygenase isoforms that convert AA into EETs. All EETs are then further metabolized to less active dihydroxyeicosatrienoic acids (DHETs) by soluble epoxide hydrolase (sEH).

EET signaling in cardiovascular and kidney diseases

EETs are produced primarily by endothelial cells, although they are also produced by other cell types, such as astrocytes and cardiomyocytes. Because EETs induce vasodilation and exert antiinflammatory effects in blood vessels in an autocrine manner (2), they can lower blood pressure, protect the myocardium and brain from ischemia, attenuate hypertension-induced renal damage, and reduce cigarette smoke–induced lung inflammation (3). Increasing evidence reveals that EETs govern these various biological functions by inducing endothelial cell proliferation, survival, and tube formation and stimulating renal epithelial cell proliferation and survival through multiple signaling pathways (4). Hence, great efforts have been made to develop drugs targeting these pathways. For example, 11,12-EET has been shown to improve coronary artery endothelial function when it is added to transplant preservation solutions (5). Moreover, an sEH inhibitor (AR9281) is currently under evaluation in phase II clinical trials as a treatment for patients with hypertension and type 2 diabetes on the basis of evidence that sEH inhibitors have beneficial effects in animal models of hypertension and cardiovasual diseases (3, 6). However, emerging evidence shows that EETs can promote cancer progression by directly promoting cancer cell proliferation, survival, migration, and invasion. In this issue of the JCI,
Panigrahy et al. report breakthroughs in our understanding of how EET signaling in the tumor microenvironment contributes to tumor growth and metastasis (7). These results raise concerns about using EET analogs and agonists as well as sEH inhibitors to treat cardiovascular diseases.

EET signaling and cancer

Although little is known about the role of EET signaling in cancer progression, emerging evidence indicates that CYP epoxygenases and the metabolites they generate are involved in tumor biology. CYP2J2 expression is elevated in human malignant solid tumors in esophageal, liver, breast, lung, and colorectal organs, and high levels of EETs have been detected in urine and blood samples obtained from patients with these cancers (8, 9). Interestingly, CYP2C9 is specifically expressed in the tumor-associated vasculature of human renal carcinomas (10). Moreover, hypoxia induces the expression of CYP2C8 and CYP2C9 (11). These results indicate that CYP epoxygenases may be involved in tumor-associated angiogenesis.

In vitro studies have shown that both overexpression of CYP2J2 in cancer cells and treatment of cancer cells with EETs stimulate cell proliferation, survival, migration, and invasion (8, 12). Further, in mouse xenograft models, overexpression of CYP2J2 in breast cancer cells promotes lung metastasis (12), whereas treatment with a selective inhibitor of CYP2J2 suppresses breast tumor growth and lung metastasis (13). Moreover, disruption of the Cyp2c44 gene of the xenograft recipient inhibits tumor growth and tumor-associated angiogenesis (10). Yet another study showed that preventive treatment of mice with CYP2J2 peptide inhibited tumor growth by activating host antitumor immunity at an initial stage of an implanted murine bladder tumor, whereas continuous treatment of mice with this peptide accelerated tumor growth by suppressing host antitumor...
immunity at an advanced stage (14). These findings indicate that CYP epoxygenases and EETs may play an important role in the tumor microenvironment.

Although EETs are well known to stimulate angiogenesis by promoting endothelial cell proliferation, survival, migration, and tube formation, surprisingly little research has directly addressed the question of how modification of EET signaling in endothelial cells affects neoplastic growth and metastasis. In this issue of the JCI, Panigrahy and colleagues present the first direct evidence showing that elevation of EET levels in endothelial cells leads to the promotion of tumor-associated angiogenesis and metastasis (7). They found that treatment of endothelial cells with 14,15-EET promoted primary tumor growth and metastasis. By contrast, reduction of EET levels by overexpression of either CYP2C8 or CYP2J2 in endothelial cells and by cell type-nonspecific deletion of sEH also stimulated tumor growth and metastasis. By contrast, elevation of EET levels by overexpression of either CYP2C8 or CYP2J2 in endothelial cells and by cell type-nonspecific deletion of sEH also stimulated tumor growth and metastasis. These data generated by Panigrahy and colleagues significantly extend our understanding of how EET signaling in the tumor microenvironment contributes to tumor growth and metastasis.

The data from human specimens and the in vitro and in vivo studies discussed here support the hypothesis that EETs may promote cancer progression by directly inducing cancer cell proliferation, survival, migration, and invasion and/or by changing the tumor microenvironment by influencing angiogenesis and immunosuppression in an autocrine and/or paracrine manner. This hypothesis has been tested in preclinical studies in which inhibitors of epoxygenase and EET antagonists were evaluated for their ability to inhibit tumor formation and growth (Figure 1). For example, treatment of glioblastoma-bearing rats with CYP epoxyenase inhibitors was found to attenuate tumor growth and tumor-associated angiogenesis (15). Similarly, the work reported in this issue of the JCI by Panigrahy et al. (7) showed for the first time that an EET antagonist could inhibit tumor growth and metastasis as well as prolong survival in several animal models. These in vivo data are consistent with a previous study in which an EET antagonist inhibited EET-induced prostate carcinoma cell migration and invasion in vitro (16). Collectively, the results discussed here not only raise concerns about developing sEH inhibitors as well as EET analogs and agonists for human use to treat cardiovascular diseases, but also support the rationale for developing EET antagonists and inhibitors of CYP epoxyenase enzymes as antitumor agents (Figure 1).

EET downstream signaling pathways in cancer

Although no EET receptor(s) have yet been clearly identified, EETs have been shown to bind to GPCRs (17, 18) and to facilitate binding activity of the PPAR/RXR heterodimer to a peroxisome proliferator response element (19, 20). Moreover, 14,15-EET induces EGFR transactivation in cancer cells in vitro (21). Indeed, EETs induce cancer cell proliferation via the EGFR/Pi3K/Akt and EGFR/MAPK pathways and promote cancer cell survival through multiple pathways, including the TNF-α pathway and antioxidant enzyme-mediated pathways (8, 22). Moreover, pro-metastatic MMPs may mediate the effects of EETs on metastasis (12). The report by Panigrahy et al. (7) reveals that a VEGF signaling pathway is affected by EETs in endothelial cells. Furthermore, they found that VEGF signaling was required for EET-induced tumor-associated angiogenesis, which accelerated tumor growth and metastasis. However, it remains unclear whether EETs promote cancer progression by binding to cell-surface receptors and/or intracellular receptors such as nuclear receptors, with subsequent enhancement of cell proliferation, promotion of angiogenesis, inhibition of apoptosis, and stimulation of invasion/motility. Identification of specific EET receptors will be critical not only to further understanding of the molecular, cellular, and biological mechanisms underlying the involvement of EETs in malignant diseases, but also to enable the development of EET receptor-specific antagonists as antitumor agents.

Summary

CYP epoxygenases and the metabolites they generate, EETs, clearly have cardiovascular protective effects. However, the findings by Panigrahy et al. in this issue of the JCI (7) and other published results (8–16) indicate that EETs also promote tumor growth and metastasis in some contexts. This warrants further investigation before sEH inhibitors as well as EETs and their analogs and agonists can be considered as therapies for cardiovascular disease. Clarifying this issue is of critical importance in order to avoid harmful effects in patients who may be considered for treatment with this class of drugs.

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11. Michaelis UR, Finslhalter B, Barbosa-Sicard E, Falcik JR, Fleming I, Busse R. Cytochrome P450 epoxygenoaromatics 2J2 and 2J9 are implicated in hyposia-induced endothelial cell migration and angiogenesis.
The number of people who suffer from obesity and one or more of its adverse complications is rapidly increasing. It is becoming clear that diet, exercise, and other lifestyle modifications are insufficient strategies to combat this growing problem. Greater understanding of the mechanisms controlling our desire to feed and our ability to balance energy intake with energy expenditure are key to the development of pharmacological approaches for treating obesity. Although great strides have been made in our understanding of how the hypothalamus regulates feeding and energy balance, much less is known about how obesity affects the structure of the hypothalamus. The authors of two papers in this issue of the JCI have addressed this issue by examining the effects of obesity on neurons and glia in the hypothalamus. These studies reveal that obesity may be in part due to hypothalamic injury, which leads to inflammation and reduced neurogenesis. These findings support the notion that obesity is a disease that affects multiple organs, including the brain, and that disruption of normal brain function leads to abnormal regulation of peripheral metabolism.

Central regulation of peripheral metabolism
The role of the hypothalamus in the regulation of feeding and energy balance was first highlighted by lesion studies in rodents (1, 2). Although these classic studies proposing the existence of “feeding” and “satiety” centers in the hypothalamus lacked anatomic precision and were overly simplistic in their interpretation, the importance of the hypothalamus in the regulation of feeding and energy balance was subsequently highlighted by the discovery that hormone and peptide regulators of feeding and metabolism act on the hypothalamus as well as the brainstem and other areas of the brain (3). The adipose tissue–derived hormone leptin exerts its inhibitory effects on food intake by modulating the function of neurons in the arcuate nucleus (an aggregation of neurons in the medial basal region of the hypothalamus). Specifically, leptin suppresses neurons that release neuropeptide Y (NPY) and agouti-related peptide — neuropeptides that normally increase appetite — while stimulating neurons that release proopiomelanocortin (POMC) — the precursor of several neuropeptides, including some that suppress appetite (3). Leptin, ghrelin, and other hormones that control energy metabolism also affect synaptic plasticity — the structural connections of neurons in the hypothalamus and other areas of the brain (4–7).

A hallmark of obesity is the ability of adipose tissue to expand and undergo significant remodeling in order to fulfill its role as the major energy-storing organ. This involves coordinated responses among various cell types, including adipocyte precursors, blood vessels, and immune cells (8). Inflammation in obese adipose tissue is related to adipocyte death, accumulation of macrophages and other immune cells, and metabolic dysfunction (8). Recent studies have shown that the presence of cytokines and inflammatory molecules in the adipose tissue can lead to the activation of proinflammatory signaling pathways in the hypothalamus, which in turn can alter the expression of neuropeptides that regulate feeding and energy balance (9–11).

Conflict of interest: The authors have declared that no conflict of interest exists.

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