Brief Report: An Evolutionary Basis to Cancer Growth

Mehdi Fini
University of Colorado Anschutz Medical Campus: University of Colorado - Anschutz Medical Campus

Eric Gaucher
Georgia State University

Brian Boutwell
The University of Mississippi Medical Center

Takahiko Nakagawa
Rakuwakai Health Care System: Iryo Hojin Shadan Rakuwakai

Richard Wright
University of Colorado Anschutz Medical Campus: University of Colorado - Anschutz Medical Campus

Laura Sanchez-Lozada
Instituto de Cardiologia

Peter Andrews
Natural History Museum

Kurt Stenmark
University of Colorado Anschutz Medical Campus: University of Colorado - Anschutz Medical Campus

Richard Johnson (✉ richard.johnson@ucdenver.edu)
University of Colorado Anschutz Medical Campus: University of Colorado - Anschutz Medical Campus
https://orcid.org/0000-0003-3312-8193

Short Report

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Abstract

Background. Recently we have reviewed the evidence that excessive intake of fructose, present in added sugars, may be a unique nutrient that stimulates tumor growth while also causing obesity and metabolic syndrome, thereby providing one of the reasons why obesity is associated with increased risk for various cancers. Prior studies have suggested that one of the reasons fructose increases the risk for cancer may be due in part to its metabolite uric acid, and both a meta-analysis and Mendelian randomization study confirm uric acid is a risk factor. Here we suggest that a mutation in uric acid metabolism may have provided survival advantage to our ancestors but ironically increase our risk for cancer today.

Methods. Ancestral humans lost the gene uricase during the mid-Miocene where it led to higher uric acid levels that facilitated the effects of fructose to stimulate fat accumulation. Here we tested whether a loss of uricase would also facilitate tumor growth. Experiments were performed in mice in which uricase was inactivated by either knocking out the gene or by inhibiting uricase with oxonic acid. We also studied mice transgenic for uricase. These mice were injected with breast cancer cells and followed for 4 weeks.

Results. The inhibition or knockout of uricase was associated with a remarkable increase in tumor growth and metastases. In contrast, transgenic uricase mice showed reduced tumor growth.

Conclusion. A loss of uricase increases the risk for tumor growth. Prior studies have shown that the loss of the mutation facilitated the ability of fructose to increase fat which provided a survival advantage for our ancestors that came close to extinction from starvation in the mid Miocene. Today, however, excessive fructose intake is rampant and increasing our risk not only for obesity and metabolic syndrome, but also cancer. Obesity-associated cancer may be due, in part, to a mutation 15 million years ago that acted as a thrifty gene.

Background

Fructose is a major component of table sugar (sucrose) and high fructose corn syrup (HFCS), which are the two most common added sugars in the western diet. Sugar and HFCS are present in over 70 percent of processed foods and accounts for 15 percent of the calories in the average diet, and in some individuals intake is 25 percent or more. Recent studies suggest that fructose in added sugars is a major risk factor not only for obesity and metabolic syndrome, but also for cancer (1-4). One potential mechanism by which fructose may increase the risk for cancer is by reducing mitochondrial function while stimulating glycolysis, as tumor cells often live in hypoxic microenvironments (3, 5).

A unique aspect of fructose metabolism is that it results in rapid consumption of ATP, associated with adenine nucleotide degradation and the formation of uric acid (6). Our group has found that this pathway is critical for how fructose alters mitochondrial function, as the uric acid causes cellular and mitochondrial oxidative stress that interrupts the Krebs cycle by blockingaconitase while also reducing beta fatty acid oxidation via inhibitory effects on enoyl coA hydratase (7-9). In this regard, hyperuricemia
is also recognized as a risk factor for cancer (10) including by meta-analysis of epidemiological studies (11) as well as Mendelian randomization studies (12).

In this regard, humans have a higher serum uric acid compared to most mammals due to a mutation in the uricase gene. Uricase is an enzyme that degrades uric acid, and most mammals that express uricase maintain serum uric acid levels in the 1 to 3 mg/dl range. However, the great apes and humans lost uricase through a series of mutations that progressively reduced activity of uricase until it was completely silenced around 15 million years ago (13). The consequence was an increase in serum uric acid to the 3 to 4 mg/dl range, which then has subsequently increased with western diets rich in fructose-laden sugars and purines (14). Today nearly 20 million people in the United States have serum uric acid levels greater than 7 mg/dl that place them at increased risk for cancer, obesity, metabolic syndrome and gout (1-4).

We have previously studied the uricase mutation and shown that it likely functioned as a thrifty gene (13, 15-17). Specifically, at the time of the mutation during the mid-Miocene, ancestral apes were living in both Europe and Africa and living primarily on fruits. At that time there was global cooling with a loss of fruit availability during the cooler season in Europe that caused great nutritional stress. Over a period of several million years there was progressive starvation and extinction of apes in Europe. However, based on the fossil record, some apes survived and returned to Africa to become the ancestors of African great apes and humans, while others migrated to southeast Asia to become the ancestor of the orangutan (18, 19). By resurrecting the extinct uricase, we were able to show that its loss in human hepatocytes was associated with enhanced fat accumulation and gluconeogenesis in response to fructose, suggesting the mutation provided survival benefits to ancestral apes (13, 17). However, in today’s society in which excessive intake of fructose and marked hyperuricemia are occurring, the consequence is the emergence of obesity and diabetes (16).

Here we wished to test whether the uricase mutation might also have unwittingly increased our risk for cancer growth and spread.

**Materials And Methods**

**Reagents**

Most reagents, buffers, substrates, uric acid, uricase and oxonic acid were purchased from Sigma Chemical Company (St Louis, MO, USA). Media for cell culture were obtained from Gibco/BRL (Bethesda, MD, USA). Hematoxylin Nuclear Counterstain (H-3401) was purchased from Vector Laboratories (Burlingame, CA, USA) for use in H&E tissue staining.

**Cells and Culture Conditions**

Mouse mammary carcinoma cell line (E0771) were a kind gift from Dr. Mikhail Kolonin, The Brown Foundation Institute of Molecular Medicine, Houston, TX. Mouse mammary carcinoma cell E0771 and
cells were grown in RPMI 1640 containing 2 mM l-glutamine, 2 g/l sodium bicarbonate, pH 7.4, 1X of antibiotic/antimycotic solution, 5 µg/ml insulin, and 10% endotoxin free fetal bovine. Cells were maintained at 37°C in 95% air/5% CO2, fed every 2 days, and split 1:3 when at or near confluence.

**E0771 Syngeneic Breast Cancer Animal Model and Pharmacologic Inhibition of Uricase**

Using pharmacologic approach and to inhibit uricase, control females mice (C57BL/6 JAX) were placed on standard chow (Ctr CHOW) or chow containing oxonic acid (OA-CHOW). After one month of being fed on the special chow, E0771 cells were implanted into the mammary fat pads. After 28 days, mice were sacrificed. Macroscopic tumor volume was measured and lung histology were analyzed for lung metastasis.

**Transgenic Mice**

Homozygous uricase knockout mice (UOX -/-) were backcrossed with C57BL/6 JAX for 10 generations at Denver altitude. Homozygous mice with lack of uricase activity and Heterozygous littermates with normal uricase activity were used for experiments. Uricase overexpressing transgenic mice (ssUOX Tg) were also backcrossed with C57BL/6 JAX for 10 generations at Denver altitude. Transgenic mice with control littermates were used for experiments. Age matched female mice were implanted with E0771 mouse mammary breast cancer cells. Serial tumor volume was measured during a twenty-eight-day growth period at days 7, 14, and 28. Primary tumors were harvested and examined histologically. Lungs, kidneys and heart along with blood serum were recovered and analyzed.

All animal procedures were performed in accordance with the guidelines of the Institutional Animal Care and Use Committee of the University Colorado Denver Anschutz Medical Campus.

**Results**

**Pharmacologic inhibition of uricase stimulates tumor growth, progression and lung metastasis.** C57b/6 mice were placed on standard chow (Ctr CHOW) or chow containing the uricase inhibitor, oxonic acid (OA-CHOW). After 1 month on the different chows, the mice were injected with E0771 breast cancer cells into their mammary fat pads. Four weeks later mice were sacrificed. Double-blind analysis documented significant increase in primary tumor volume (Fig. 1A) and microscopic lung metastasis (Fig. 1B) in the uricase-inhibited mice compared to control mice.

**Genetic manipulation of Uricase modulates tumor growth, progression and lung metastasis.** We also performed similar experiments using homozygous uricase knockout mice (UOX KO) and control littermates. A similar study was performed in which we injected highly aggressive E0771 breast cancer cell lines into the mammary fat pads followed by sacrifice 4 weeks later. Our data show significant augmentation of both primary tumor volume (Fig. 1C) and lung metastases (Fig. 1D) in these mice when compared to their control littermates. Interestingly these were attenuated in transgenic mice overexpressing uricase (UOX Tg). Besides genetic confirmation, we determined the serum level of uric
acid (SUA) as a specific determinant of uricase activity which shows high level of SUA in UOX KO vs UOX Tg transgenic mice (Fig. 1E).

**Discussion**

Here we performed preliminary studies to determine if the loss of uricase could affect cancer growth. Our hypothesis was based on the fact that the uricase mutation has been shown to raise uric acid which not only drives many of the metabolic effects of fructose (1), but also enhances fructose metabolism and production (20, 21). One of the effects of fructose is to stimulate the Warburg effect, by suppressing mitochondrial function while stimulating glycolysis (3). This effect results in less oxygen need for the organism. Recently it was shown that the naked mole rat produces fructose as a mechanism to protect itself from the hypoxia that is present in the underground burrows where it lives (5). However, while the Warburg effect might be of benefit for animals trying to survive hypoxic conditions, it might also be a mechanism for stimulating cancer growth, for this can allow tumor cells to multiply in tissues with minimal blood supply.

Here we tested the hypothesis using two different approaches. First, we inhibited uricase using oxonic acid, and found that this procedure led to a more rapid growth of the breast cancer cells as well as increased metastases. We also showed the same effect when using uricase knockout mice, and a tendency for less tumor growth and metastases in a uricase transgenic mouse. These studies provide tantalizing data that suggest uric acid may have a role in tumor growth.

Supporting this concept has been the observation that hyperuricemia is associated with both obesity and cancer (10), and has been found to increase the risk for cancer incidence and mortality by meta-analysis (11) and Mendelian randomization studies (12). Furthermore, allopurinol, a xanthine oxidase inhibitor that blocks uric acid formation, reduces breast tumor growth and metastases (22) as well as colonic cancer tumorigenesis (23) in murine models.

We would like to suggest that, while the uricase mutation was of benefit by aiding the survival of ancestral hominids, that the mutation is maladaptive in current society in which fructose intake is high. Not only is the mutation increasing our risk for obesity and diabetes, but it may also be increasing our risk for cancer. These findings could explain why cancer is increased in subjects with obesity and metabolic syndrome, for both conditions are associated with excessive fructose intake and hyperuricemia. It also suggests that more basic science and clinical studies should be done on the role of both fructose and uric acid in cancer, and also suggest blocking fructose or uric acid metabolism may be an additional target for cancer therapies.

At a deeper level, this study enforces the concept of how evolutionary pressures can lead to changes that were beneficial in one environment yet be deleterious in another. Specifically, cancer growth and metastases may represent another casualty from the loss of uricase, emphasizing the importance of the ‘thrifty gene’ hypothesis in human biology (15).
Declarations

- Ethics approval and consent to participate. Not applicable.
- Consent for publication. The authors provide consent for publication and assure this manuscript is not under consideration elsewhere.
- Availability of data and materials. The data set in Figure 1 is available upon request to Dr Mehdi A Fini at Mehdi.Fini@cuanshutz.edu.
- Competing interests. RJJ, LGL and ML have equity with Colorado Research Partners LLC that is developing inhibitors of fructose metabolism, and RJJ also has equity with XORTX Therapeutics developing xanthine oxidase inhibitors. Dr Johnson has consulted for Horizon Pharma. All other authors declare no competing interests.
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- Authors' contributions. Dr Mehdi Fini and Dr Richard Johnson wrote the original draft which was reviewed and edited by all authors.
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