A review of Kurozu, amber rice vinegar made in pottery jars

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ABSTRACT:
Brewed rice vinegar, Kurozu, is a traditional Japanese vinegar with a dark amber color. Kurozu is produced in a regional area of Japan using traditional techniques and made inside handcrafted pottery jars. Kurozu is used as both a seasoning and a healthcare supplement. In vitro and in vivo investigations of ingredients in Kurozu have been carried out. Studies of the functional aspects of Kurozu began in the 1980s, and the health promoting and disease preventing effects of Kurozu have since been elucidated. It was reported that Kurozu improved the symptoms of hypertension, allergies, hypercholesterolemia, enhanced carbohydrate metabolism, and inhibited tumor growth. Kurozu-Moromi is an insoluble product created from the fermentation of Kurozu. Kurozu-Moromi also shows valuable properties, including improvement in dyslipidemia, prevention of hyperglycemia, antitumor effect, and antiallergic activity.

Keywords: Amber color, Brewed vinegar, Functionality, Moromi, Pottery jars

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INTRODUCTION: Vinegar has been used as a seasoning for several thousand years. In ancient times, there was a record of Hippocrates using vinegar as medication. Traditional vinegar is produced from raw materials containing sugars or starches, such as grape, apple, malt, and rice. The production of vinegar typically involves a first fermentation where simple sugars in the raw material are converted to alcohol by yeasts. The resultant alcohol is further oxidized into acetic acid by acetic acid bacteria (AAB), which are present everywhere in the environment [1].

Kurozu, which contains rice, spring water, and koji, is made using pottery jars. Koji is a culture of a certain mold that grows on rice and is used when it produces fermentation products, such as rice wine. Kurozu changes to an amber color with aging (Figure 1). The change in color was the origin of black vinegar, and it was named “Kurozu” in 1975 by Sakamoto Kurozu, Inc. “Kuro” indicates black color, while “zu” indicates vinegar in Japanese. It is believed that sugars react with amino acids by aminocarbonylation, resulting in the product’s amber color.

Kurozu has been produced in the Fukuyama region of Kagoshima prefecture for about 200 years. The Fukuyama region is suitable for brewing vinegar because of its warm climate and ready supply of spring water [2, 3]. Kurozu is handcrafted using traditional techniques and craftsmanship.

In Japan, there are many regional brand products that have obtained strong reputations because of their unique production methods and natural characteristics. The Ministry of Agriculture, Forestry and Fisheries (MAFF) of Japan certifies Kurozu as a specific agricultural product.

![Figure 1. Color change of Kurozu with aging. Left, 1 year old; Middle, 2 years old; Right, over 3 years old.](image)

The Geographical Indication (GI) Act protects the names of specific agricultural, forestry, and fishery products and foodstuffs and was implemented by MAFF in accordance with a treaty with the World Trade Organization, which defines intellectual property rights of specific agricultural products.

BREWING: Kurozu is produced in outdoor environments using Japanese pottery jars (typical size: trunk diameter 40 cm, caliber 14 cm, height 62 cm, volume 54 L; Figure 2, 3). Koji-containing steamed rice, steamed rice, spring water, and Koji are layered in the jar. Koji is sprinkled on the surface of the water at a constant thickness (Figure 3). Kurozu is brewed twice a year during spring and autumn. Koji is widely used in Japanese fermented foods, including sake (alcoholic beverage), soy sauce, and rice vinegar. Most Japanese fermented foods contain koji (rice fermented with the non-pathogenic fungus *Aspergillus oryzae* or *A. luchuensis*) as the saccharifying agent of the starch contained in crops. Koji initiates a dynamic microbial succession, resulting in the saccharification of starch and subsequent alcoholic fermentation [4-7]. Kurozu does not require the addition of yeast or acetobacter to the pottery jars or thermoregulation of the brewing process (Figure 4).
The rice ferments for approximately 3–6 months. Kurozu develops a mellow taste and flavor during the aging process. The typical duration of the aging process ranges from 6 months to 3 years. During the aging process, the color of the vinegar turns from a light color to dark amber. The most common brewed vinegar is made through two phases of saccharification, alcohol fermentation and acetic acid fermentation by AAB. A distinctive feature of Kurozu is that acetic acid fermentation does not need the addition of AAB. In the initial stages of brewing, the starch of rice is saccharified by Aspergillus oryzae in Koji, and alcohol fermentation proceeds with Koji containing Saccharomyces under anaerobic conditions. Koji sprinkled on the water ferments the initial fermentation products, resulting in the formation of a lid-like layer that prevents the invasion of outside microbes. In the subsequent stage, the surface lid of Koji precipitates and the fermented products enter aerobic conditions. When alcoholic fermentation is saturated, the alcohol produced is converted into acetic acid by AAB in the jar. The insoluble residue (Kurozu-Moromi: KM, Figure 4) precipitates with aging. After fermentation the jars are washed with water and dried outdoors. The jars are never sterilized and are reused directly [2].

**COMPONENT:** The Japanese Agricultural Standard (JAS) system defines Kurozu as a brewed rice vinegar. Brewed vinegars are made with cereal, fruits, vegetables, or saccharides. Kurozu has to be made with more than 180 grams of rice per 1 L and has to be amber colored through fermentation and aging. The JAS system approves brewed cereal vinegar containing organic acids, including acetic acid at a concentration of 1.5–8.0%. The GI Act approves the standard quality of Kurozu in more detail. Kurozu is a vinegar that is colored amber or blackish brown as a result of fermentation and aging. Kurozu includes amino acids (approximately 5 g/L) and organic acids (0.2–0.3%
organic acids except acetic acid) derived from unpolished rice. In addition, Kurozu has to conform to the following quality standards:

1. All nitrogen sources comprise more than 0.12% 
2. The concentration ratio of amino nitrogen is more than 50% in all nitrogen sources 
3. The concentration of direct reducing sugar is less than 0.30%, as evaluated using the Soy sauce test. The test is used as the analysis method for ingredients in soy sauce, which is a fermentation product of soybean.

Kurozu was previously shown to contain organic acids, amino acids, and inorganic cations. Kurozu is rich in alanine, leucine, lysine, valine, and glycine. Kurozu contains other organic acids such as lactic acid and pyrogallic acid along with acetic acid. Furthermore, major inorganic cations such as sodium, potassium, and magnesium are found in Kurozu. The compositions are greatly different between products. Koizumi Y et al. reported that 11 kinds of Kurozu contained 0.01–0.48% saccharide, 724–4,593 mg/L organic acids without acetic acid, and 569–3,626 mg/L free amino acids. It was estimated that differences in the composition were caused by differences in the raw materials and preparation methods. Ingredients such as amino acids, organic acids, and minerals differ in their density depending on the products and may influence differences in functionality [8]. Kurozu contains peptides, such as Ile-Tyr-Pro, Phe-Phe, Gln-Leu-Pro and Asn-Pro [9].

Over the past decades, the presence of D-amino acids has been shown in bacteria, plants, animals, and fermented foods [10]. Large amounts of D-Ala, D-Asp, and D-Glu have been observed in Kurozu [11]. Shimoji Y et al. reported that Kurozu contained polyphenolic compounds with anti-oxidative activities, such as dihydroferulic and dihydrosinapic acids [12]. Yamagishi K et al. reported that Kurozu contained histamine, which improves blood fluidity [13]. Recently, Kurozu was shown to contain a lipopolysaccharide compound produced by AAB [14 - 16].

KM is a product created from fermentation and comprises the insoluble residue, which may be taken in small amounts from the jar. Components of KM have been investigated. Kamata S et al. reported that KM contained saturated fatty acids (palmitate and stearate), unsaturated fatty acids (oleinate and linoleate), and their esters [17]. The odorant compounds in Kurozu were then analyzed. Kurozu contained ketones (e.g. 2-butanon, 2,3-butanedione, 3-hydroxy-2-butanone, 2,3-pentanedione), aldehydes (e.g. benzaldehyde, 2-methyl-butanal, 3-methyl-butanal, acetaldehyde), alcohols (e.g. ethanol, 2-ethyl-1-hexanol), and amines (e.g. 2,3-dimethylpyrazine, trimethylpyrazine) [personal communications, provided by Dr. H Kanouchi]. These odorant compounds are well known as ingredients in brewed sake made from rice [18, 19].

**FUNCTIONALITY:** Functional studies of Kurozu have dated back to the findings of empirical health benefits. Studies on the functions of Kurozu began in the 1980s, and both *in vitro* and *in vivo* investigations of ingredients in Kurozu have been carried out [3]. Recently, clinical studies have also been performed.

**Metabolic effects:** Health-related properties of Kurozu were initially reported on dyslipidemia. Tanizawa H et al. reported that Kurozu decreased serum cholesterol levels in both normal and high cholesterol diet-fed mice. Kurozu ingestion also inhibited increases in lipid peroxide levels induced by adriamycin injection in mice myocardium [20]. Oominami and Okuda also reported that supplementation of Kurozu significantly decreased the levels of triglycerides, total cholesterol, and free fatty acids in plasma following corn oil ingestion. Constituents of Kurozu inhibited lipid synthesis stimulation with insulin in rat adipocytes [21]. Concentrated Kurozu administration significantly
reduced the average size of adipocytes, but the numbers of adipocytes were increased and fatty acid excretion was significantly increased [22].

In a case study, treatment with an extract of Kurozu showed significant decreases in total cholesterol in patients with cardiovascular disease [23]. A randomized control study showed that combined intake of Kurozu and garlic may have the potential to improve serum levels of total cholesterol and low-density lipoprotein cholesterol in a subject population comprised of prehypercholesterolemic and mild-to-moderate hypercholesterolemic subjects [24]. Hamadate N et al. reported that concentrated Kurozu supplementation significantly decreased body weight, and may decrease body mass index in healthy adults with obesity [25]. Hamadate N et al. also reported that supplementation of Kurozu decreased hip circumference. Kurozu also increased fat-derived energy consumption while suppressing carbohydrate-derived energy consumption in obese but otherwise healthy subjects [26]. Abe A et al. reported that Kurozu decreased visceral fat accumulation in obese healthy subjects [27]. Oral administration of concentrated Kurozu decreased adipocyte size via inhibition of dietary fat absorption and reduction of PPARγ and mRNA levels of fatty acid binding protein 2 in adipocytes in rats [22]. It was reported that acetic acid improved obesity and glucose tolerance. Exogenously administered acetic acid may have effects on lipid metabolism via alteration of glucose transporter 4 and myoglobin gene expression, and it may also function against obesity and obesity-linked type 2 diabetes [28, 29]. Publications about the function of Kurozu evaluated concentrates without acetic acid. It is thought that both the concentration of ingredients in Kurozu and acetic acid contribute to the lipid metabolism improvement.

Recently, Shibayama Y et al. reported that Kurozu inhibited the incidence of hepatic steatosis with a high-fat diet and induced the expression levels of hepatic Sirt1, Pgc-1α, Lpin1, and Igfbp1 in mice [30]. Shibayama Y et al. also reported that concentrated Kurozu supplementation reduced the onset of hepatic hyperplasia and increased hepatic microRNA-34a, -149-3p, and -181a-5p [31]. An effect on glucose metabolism was also reported by Nagano M et al., where the intake of KM and Kurozu concentrated liquid prevented increases in blood glucose level in mice [32].

**Antioxidants effects:** Kurozu contains polyphenolic compounds, and antioxidant effects of Kurozu have been reported. Kurozu scavenged 1,1-diphenyl-2-picrylhydrazyl radicals and decreased antioxidant effect on copper-mediated human low-density lipoprotein oxidation in vitro [33]. In an animal study, an extract of Kurozu inhibited 12-O-tetradecanoylphorbol-13-acetate-induced edema formation and myeloperoxidase activity in female ICR mouse skin [34]. Kurozu increased the serum biological antioxidant potential level gradually, and after 30 days it was significantly higher compared with the pre-study level. The serum level of diacron-reactive oxygen metabolites and blood filtration time decreased in 10 healthy female subjects [35]. Kurozu improved inflammatory bowel diseases in mice; Fukuyama N et al. reported that concentrated Kurozu supplementation inhibited inflammation and oxidative stress induced by dextran sulfate sodium treatment in ulcerative colitis [36]. Recently, a notable observation of antioxidant action was reported. Shin M et al. reported that the size of the intracerebral hemorrhage-induced nerve injury lesion and apoptotic neuronal cell death in the striatum was significantly suppressed by Kurozu intake. It was thought that the anti-apoptotic effect occurred with decreasing expression of c-Fos and caspase-3. It was assumed that the decreasing effect of the expression suppressed the oxidative stress and increased the antioxidant level [37].
Hematologic and vascular effects: Kurozu influences cardiovascular function and blood fluidity. Fujino T et al. reported that Kurozu ingestion improved red cell filterability in patients with cardiovascular disease [23]. Kurozu also influences function of immune cells. Constituents of Kurozu improved the activity of natural killer (NK) cells, and upregulated interferon-γ, tumor necrosis factor-α, and interleukin-12 in tumor-bearing mice [3]. Kurozu significantly decreased the whole blood filtration time in male long-distance runners [39]. NK cell activity was significantly decreased by training in a Kurozu non-consuming group, but not in a Kurozu-consuming group. It was estimated that Kurozu prevents the degradation of NK cell activity induced by intense training, suggesting its contribution to the physical condition of bicycle racers [40]. Significant differences in blood pressure were observed between a concentrated Kurozu intake group and a non-intake group, both comprised of healthy adult subjects [41]. It was reported that peptides in Kurozu lowered blood pressure [9, 42].

Anti-neoplastic effects: Kurozu inhibited tumor growth in vitro and in vivo. Nanda K et al. reported that an extract of Kurozu inhibited the proliferation of cancer cell lines in a dose-dependent manner, and apoptosis was caused by G0/G1 arrest through p21 induction [43]. Kurozu inhibited cancer development in vivo. Extract of Kurozu inhibited the incidence and multiplicity of colon adenocarcinoma in rats [44] and tumor growth in tumor-bearing mice [3, 45]. It was estimated that inhibition effect of tumor growth occurred through upregulation of tumor necrosis factor (TNF)-α [3] and downregulation of matrix metalloproteinase (MMP)-2 [45]. Shizuma T et al. also reported that Kurozu supplementation showed anti-colitis activity, and in 800–4,000 Da fractions, Kurozu had anti-oxidative or anti-nitration effects [46]. KM had similar effects. Fukuyama N et al. reported that supplementation of KM reduced tumor volume and inhibited MMP activation in tumor-bearing mice [47]. Shizuma T et al. also reported that KM decreased the size of hepatocellular carcinomas and levels of activated matrix MMP-2 and -9 in rats. KM also prolonged survival in mice with diethylnitrosamine-induced hepatocellular carcinomas [48].

Other effects: Kurozu shows protection effects against hepatic damage. Kurozu inhibits decreases in the cell number of primary cultured cells from rat livers [49]. Concentrated Kurozu decreases IgE levels in the blood in allergy model mice by regulating the tyrosine kinase signaling pathway in the spleen [50]. It was reported that Kurozu contributed to cognitive function. Concentrated Kurozu and KM improved memory impairment in the Morris water maze test, which used senescence-accelerated mouse prone 8 (SAMP8) mice [51]. Ingestion of Kurozu upregulated the expression levels of heat shock protein (HSP) in the brain in SAMP8 mice. It is known that HSP repairs and prevents protein degradation. It was assumed that upregulated HSP prevented the recognition functions of SAMP8 mice [51]. Kurozu and KM suppress inflammatory effects in mice. Ohkura et al. reported that supplementation with Kurozu inhibited lipopolysaccharide (LPS) stimulated TNF-α production in adipocytes and macrophages. KM suppressed pathways of both LPS-stimulated TNF-α production in adipocytes and macrophages and TNF-α induced plasminogen activator inhibitor 1 (PAI-1) production. Kurozu and KM decreased thrombotic tendencies by decreasing PAI-1 production in inflammatory states and have potential as antithrombotic foodstuffs [52]. The representative functions of Kurozu and KM are indicated in Table 1.
Table 1. Summary of the representative functions of Kurozu and KM

<table>
<thead>
<tr>
<th>Effect</th>
<th>Species</th>
<th>Ref. No.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Metabolism</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Decreased serum cholesterol</td>
<td>Mouse</td>
<td>20</td>
</tr>
<tr>
<td>Decreased levels of triglyceride, total cholesterol and free fatty acid</td>
<td>Rat</td>
<td>21</td>
</tr>
<tr>
<td>Inhibited lipid synthesis</td>
<td>Rat</td>
<td>22</td>
</tr>
<tr>
<td>Increased fatty acid excretion</td>
<td>Mouse</td>
<td>22</td>
</tr>
<tr>
<td>Decreased total cholesterol in serum</td>
<td>Human</td>
<td>23</td>
</tr>
<tr>
<td>Decreased total cholesterol and LDL cholesterol</td>
<td>Human</td>
<td>24</td>
</tr>
<tr>
<td>Decreased body weight and BMI</td>
<td>Human</td>
<td>25</td>
</tr>
<tr>
<td>Decreased hip circumference</td>
<td>Human</td>
<td>26</td>
</tr>
<tr>
<td>Decreased visceral fat accumulation</td>
<td>Human</td>
<td>26</td>
</tr>
<tr>
<td>Inhibited incidence of hepatic steatosis</td>
<td>Mouse</td>
<td>30</td>
</tr>
<tr>
<td>Prevented the increase in blood glucose level</td>
<td>Mouse</td>
<td>32</td>
</tr>
<tr>
<td><strong>Antioxidation</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Scavenged radicals</td>
<td>In vitro</td>
<td>33</td>
</tr>
<tr>
<td>Decrease LDL oxidation</td>
<td>In vitro</td>
<td>33</td>
</tr>
<tr>
<td>Inhibited increase of lipid peroxide levels</td>
<td>Mouse</td>
<td>20</td>
</tr>
<tr>
<td>Inhibited myeloperoxidase activity</td>
<td>Mouse</td>
<td>34</td>
</tr>
<tr>
<td>Increased serum biological antioxidant potential level</td>
<td>Human</td>
<td>33</td>
</tr>
<tr>
<td><strong>Hematologic and vascular</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Decreased blood filtration time</td>
<td>Human</td>
<td>23, 40</td>
</tr>
<tr>
<td>Decreased NK cell activity</td>
<td>Human</td>
<td>40</td>
</tr>
<tr>
<td>Decreased blood pressure</td>
<td>Human</td>
<td>41</td>
</tr>
<tr>
<td><strong>Anti-neoplastic</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Inhibited the proliferation of cancer cell lines</td>
<td>In vitro</td>
<td>43</td>
</tr>
<tr>
<td>Inhibited growth of colon cancer</td>
<td>Rat</td>
<td>44</td>
</tr>
<tr>
<td>Inhibited growth of colon cancer</td>
<td>Mouse</td>
<td>45, 47</td>
</tr>
<tr>
<td>Decreased the size of hepatocellular carcinomas</td>
<td>Rat</td>
<td>48</td>
</tr>
<tr>
<td><strong>Others</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Prevented hepatic cell damage</td>
<td>Rat</td>
<td>49</td>
</tr>
<tr>
<td>Improved allergy</td>
<td>Mouse</td>
<td>50</td>
</tr>
<tr>
<td>Improved memory function</td>
<td>Mouse</td>
<td>51</td>
</tr>
<tr>
<td>Upregulated expression of HSP in the brain</td>
<td>Mouse</td>
<td>51</td>
</tr>
</tbody>
</table>

**Safety concerns**: The safety of Kurozu is controversial. A case record reported that a Kurozu supplement induced hepatic dysfunction, but interruption of the Kurozu supplement led to recovery of hepatic function.
Another case series described that Kurozu might induce intoxication dermatosis, bleeding tendency in eyeball, and gastroenteritis [54]. The properties of food–drug interactions were reported by Iwao K et al., wherein Kurozu reduced the permeation of glibenclamide in vitro, indicating that Kurozu might modulate the absorption of glibenclamide in vivo [55].

Oguma T et al. reported that concomitant administration of Kurozu increased the absorption of itraconazole in achlorhydria patients by oral proton pump inhibition [56]. It is well known that drug interactions are induced by the alteration of metabolic enzymes and drug transporters. It was reported that Kurozu did not alter the expression levels of principal drug metabolism enzymes and transporters in rats [57].

CONCLUSION

The functionalities of Kurozu and KM have been revealed by in vitro and in vivo study. A number of studies have demonstrated that Kurozu and KM improved the symptoms of hypertension, allergies, and hypercholesterolemia, enhanced carbohydrate metabolism, and inhibited tumor growth. However, the functional properties of Kurozu and KM in humans have not yet been explored in detail. Further investigation is needed to determine their functionality in humans.

List of abbreviations: Acetic acid bacteria, AAB; Heat shock protein, HSP; Kurozu-Moromi, KM; Lipopolysaccharide, LPS; Low-density lipoprotein, LDL; Matrix metalloproteinase, MMP; Natural killer, NK; Plasminogen activator inhibitor 1, PAI-1; Senescence-accelerated mouse prone 8, SAMP8; Tumor necrosis factor, TNF.

Competing interests: AF and MN are employees of Sakamoto Kurozu, Inc.

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