Potential significance of the regucalcin gene in human carcinoma

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I am pleased to prepare inaugural editorial for a new Journal “Integrative Cancer Science and Therapeutics (ICST)”. ICST is an open access, international peer reviewed journal that enlightens the cancer research community by providing an insight on breakthrough discoveries that cover integrative fields of basic and clinical cancer research. Cell proliferation is mediated through various intracellular signaling transductions that are stimulated by many hormone and cytokines. Enhanced cell proliferation may lead to carcinogenesis. However, mechanism of carcinogenesis is complexity and its therapy is not established. Cancer is a pathological condition, where assemblage of cells displays uncontrolled growth, invasion and metastasis. Cancer medicine is based on diagnosis, therapy and post therapy procedure. Cancer therapeutics involves various pathological, pharmacological, medical and clinical approaches with several implications. ICST will contribute to development of cancer sciences and therapy.

Regucalcin is a novel suppressor protein in cell signaling discovered by the author of this editorial, Yamaguchi, in 1978 [1,2]. Regucalcin is demonstrated to play a multifunctional role in cell regulation in various types of cell and tissues [3,4]. Regucalcin is predominantly expressed in liver and kidney tissues. Interestingly, overexpression of the regucalcin gene was found to suppress liver cell proliferation and carcinogenesis in animal models [3-7]. Moreover, there is growing evidence that the regucalcin gene expression is uniquely suppressed in various human carcinoma tissues using analysis with multiple gene expression profiles and proteomics. Suppression of the regucalcin gene may lead to development of carcinogenesis. This editorial focuses a potential significance of regucalcin in the gene therapy for human hepatocarcinoma.

Hepatocellular carcinoma (HCC), the most common primary liver cancer, is one of the most prevalent malignant diseases worldwide, and the third most common causes of cancer-related death [8-10]. Globally, there are approximately 750,000 new cases of HCC reported per year. The incidence of HCC is increasingly in the United States and other developed countries. Moreover, features of HCC are also the cause of liver cirrhosis that includes viral infections (hepatitis B and C) and alcohol consumption; further risk factors include tobacco smoking, exposure to aflatoxin B1 and vinyl chloride, diabetes, and genetic disorders, such as hemochromatosis and alpha-1 antitrypsin deficiency [11-15].

Hepatocarcinogenesis is a multistep process initiated by external stimuli that lead to genetic changes in hepatocytes or stem cells, resulting in proliferation, apoptosis, dysplasia and neoplasia. The majority of HCC cases are also related to chronic viral infections. Hepatitis B virus (HBV) DNA integrates into the host genome, inducing chromosome instability and insertion mutations that may activate various oncoproteins, such as cyclin A [16-19]. Viral proteins, in particular X protein (HBx), act as transactivators to upregulate several oncoproteins (such as c-myc and c-jun) and transcriptional factors (such as nuclear factor-κB) [20-22]. Additionally, HBx activates promoters of genes encoding interleukin-8 (IL-8), tumor necrosis factor (TNF), transforming growth factor (TGF)-β and epidermal growth factor receptor (EGFR) [23]. HBx can also stimulate several signal transduction pathways, including the JAK/STAT, RAS/RAF/MAPK, and Wnt/β-catenin pathways [23,24]. The contributions of hepatitis C virus (HCV) to hepatocarcinogenesis are mediated through viral proteins, including core, NS3 and NS5A proteins. HCV core protein can promote apoptosis or cell proliferation through interaction with p53 or upregulation of Wnt-1 at the transcriptional level [25-27].

The prognosis of advanced HCC remains poor in spite of the development of novel therapeutic strategies [28]. Traditional therapies are not effective for HCC and are too toxic for patients with cirrhosis. Transarterial chemoembolization and radioembolization are the main treatments for intermediate- stage HCC at the present time. Improved knowledge of the oncogenic processes and signaling pathways, which regulate tumor cell proliferation, differentiation, angiogenesis, invasion and metastasis, has led to the identification of several potential therapeutic targets that have driven the development of molecularly targeted therapies [28]. An ideal cancer target meets the following criteria: the target is relatively specific for cancer cells (not expressed or expressed at very low levels in normal cells but over expressed in cancer cells) [28]. Meanwhile, over expression of the target is associated with malignant biological phenotypes and/or poor prognosis; the target plays an essential role in cancer initiation and progression, and inhibition of expression or activity of the target induces growth suppression and/or...
apoptosis in cancer cells. The target is "drugable" as an enzyme (e.g., a kinase) or cell surface molecule (e.g., a membrane-bound receptor) that can be easily screened for small-molecule inhibitors or targeted by a specific antibody [28,29]. The only systemic therapy available for advanced HCC is based on the multikinase inhibitor sorafenib [29], which is the most effective therapeutic tool for advanced nonresectable HCC. The survival of patients with advanced HCC treated with sorafenib depends on the absence of liver dysfunction and on the status of the patient [30]. In the past few years, the use of sorafenib in combination with transarterial chemoembolization has improved survival rates in patients with advanced HCC. Recently, new perspectives in cancer treatment have appeared with the advent of microRNAs, a novel class of noncoding small RNAs [31].

Regucalcin may play a pivotal role in the suppression of hepatocarcinogenesis [5-7]. Regucalcin plays a role as a suppressor protein in various cell signal transductions [3,4]. The regucalcin gene is located on the X chromosome in consisting of seven exons and six introns [reviewed in Ref. 32]. Regucalcin and its gene (rgn) are identified in over 15 species consisting of regucalcin family, and the gene species are highly conserved in vertebrate species [32]. The regucalcin gene expression is regulated through various transcription factors (including AP-1, NF1-A1, RGPR-p117, β-catenin, SP1 and others), which are identified as the enhancer and suppressor, and this expression is regulated with hormonal stimulation and physiological state [32,33]. Regucalcin plays a pivotal role as a suppressor protein in various signal transductions to maintain cell homeostasis for stimuli, and it plays a multifunctional role in cell regulation through maintaining of intracellular Ca^{2+} homeostasis and suppressing of signal transduction in various cell types [reviewed in Refs. 3,4]. Interestingly, the cytoplasmic regucalcin was translocated into the nucleus that is mediated through protein kinase C-dependent signaling [31,34]. Nuclear regucalcin suppressed Ca^{2+}-dependent and -independent protein kinases and protein phosphatases, Ca^{2+}-activated endonuclease, and DNA and RNA synthesis in the nucleus [reviewed in Refs. 33].

Overexpression of regucalcin was found to play a role as a suppressor protein in cell proliferation that is mediated through various signaling stimulations in the cloned normal rat kidney proximal tubular epithelial NRK52E cells and the cloned rat hepatoma H4-II-E cells [31,34,35]. Regucalcin caused G1 and G2/M phase cell cycle arrest in these cells [31,36]. The anti-cell proliferation effect of regucalcin was not dependent on apoptosis; regucalcin suppresses apoptosis induced through multisignaling pathway [reviewed in Ref. 37]. Molecular mechanisms by which regucalcin suppress the promotion of cell proliferation was elucidated. Regucalcin directly inhibited the activities of various Ca^{2+}/calmodulin-dependent enzymes, protein kinases and protein phosphatases in the cytoplasm and nuclei [3,4,33]. Nuclear regucalcin was found to inhibit nuclear DNA and RNA synthesis and suppress the gene expression of c-myc, Ha-ras, and c-erb, a tumor-stimulator gene, and stimulate the gene expression of p53 and Rb, a tumor-suppressor gene [33,38]. Moreover, regucalcin was demonstrated to inhibit protein synthesis due to inhibiting aminoacyl-tRNA synthetase and stimulate protein degradation due to activating cysteiny1 protease [reviewed in Refs. 3,4]. Thus, suppressive effects of regucalcin on cell proliferation are mediated through molecules with multi targets in liver cells [3,4,33,37].

The gene expression of regucalcin was found to suppress in hepatocarcinogenesis. Liver regucalcin gene expression was suppressed at earlier periods of carcinogenesis in rats treated with diethylnitrosamine and then 2-acetylaminofluorene combined with partial hepatectomy, which induces an increase in proliferating cells [6]. The suppression of regucalcin protein expression was identified in proteomic analysis that was differentially expressed in the livers of rats fed 5% ethanol for 1 and 3 months [7]. Liver regucalcin mRNA expression was suppressed by disorder of liver metabolism induced by administration of carbon tetrachloride [39], galactosamine [40] and phenobarbital [41] in rats. In addition, liver regucalcin level was reduced with the conditions of diabetes and ethanol ingestion [42] that lead to cirrhosis and HCC. The suppression of regucalcin gene expression may lead to the development of HCC. Noticeably, the regucalcin gene and its protein levels was found to specifically suppress in human HCC using analysis with multiple gene expression profiles and proteomics [43-47]. Suppressed regucalcin gene expression may lead to development of hepatocarcinogenesis.

Regucalcin may be a key molecule as a suppressor in cell proliferation and carcinogenesis in various types of cells and tissues. Overexpression of the regucalcin gene in cancer cells may reveal preventive and therapeutic effects on the development of carcinogenesis. Development of the regucalcin gene deliver system will be expected as a novel gene therapy in clinical aspects for cancer treatment.

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References


