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**Cellular physiological approach for treatment of gastric cancer**

Shiozaki A *s*. Cellular physiological approach for cancer therapy

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**Abstract**

Recent studies show that ion channels/transporters play important roles in fundamental cellular functions that would be involved in the cancer process. We review the evidence for their expression and functioning in human gastric cancer (GC), and evaluate the potential of cellular physiological approach in clinical management. Various types of ion channels, such as voltage-gated K+ channels, intracellular Cl- channels and transient receptor potential channels have been found to express in GC cells and tissues, and to control cell cycles. With regard to water channels, aquaporin 3 and 5 play an important role in the progression of GC. Regulators of intracellular pH, such as anion exchanger, sodium-hydrogen exchanger, vacuolar H+-ATPases and carbonic anhydrases also involved in tumorgenesis of GC. Their pharmacological manipulation and gene silencing affect cellular behaviours, suggesting their potentiality as therapeutic targets for GC. Our studies indicate the intracellular Cl- concentration could act as a mediator of cellular signaling and control cell cycle progression in GC cells. Further, we demonstrate the cytocidal effects of hypotonic shock on GC cells, and indicated that the blockade of Cl- channels/transporters enhances these effects by inhibiting regulatory volume decrease. A deeper understanding of molecular mechanisms may lead to the discovery of these cellular physiological approaches as a novel therapeutic strategy for GC.

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**Key words:** Gastric cancer; Ion channels; Water channels; Intracellular pH; Intracellular chloride; Osmolality

**Core tip:** This article aims to systematically review the current knowledge on expression and functioning of ion transporters in gastric cancer (GC). Various types of ion channels, water channels and regulators of intracellular pH have been found to express in GC, and to control tumorgenesis. Our studies indicate the intracellular Cl- concentration could control cell cycle progression in GC cells. Further, we demonstrate the cytocidal effects of hypotonic shock, and indicate that regulation of ion transport enhances these effects. A deeper understanding of molecular mechanisms may lead to the discovery of these cellular physiological approaches as a novel therapeutic strategy for GC.

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**Introduction**

Gastric cancer (GC) represents the second most common cause of cancer-related deaths in the world[1]. Recently, the prognosis of GC is improved with advances in surgical techniques, adjuvant therapy, chemoradiotherapy and molecular targeted therapy[2]. However, long-term outcomes of patients with GC remain dismal, especially for the advanced disease. An improvement in the treatment of recurrent or metastatic GC depends on understanding of the molecular mechanisms regulating the tumorgenesis and the progression of the disease.

Over the past few decades, many reports have revealed that ion channels and water transporters play important roles in fundamental cellular functions. Particularly, their physiological roles in cell proliferation have been considered since cell volume changes, which require the participation of ion and water movement across the cell membrane, are indispensable in cell cycle progression. Recently, the roles of ion and water channels/transporters have been studied in cancer cells[3-7] and various types of transporters have been found in cancers of digestive organs.

This article aims to systematically review the current knowledge on expression and functioning of ion and water channels/transporters in GC cells and tissues. The ultimate objective is to evaluate the potential of cellular physiological approach, such as regulation of ion channels, water channels, intracellular pH, intracellular ion concentration and osmolality, in clinical management of GC.

**Regulation of ion channels**

Recent studies have demonstrated that several subtypes of K+ channels are expressed in human GC cells, and are associated with cell proliferation. Altered expression of several voltage-gated K+ channels (Kv) has been observed in GC. Lan *et al*[8] have demonstrated that Kv1.5 protein is frequently detected in GC tissues and down-regulation of the expression of Kv1.5 in SGC7901 cells inhibits the proliferation and tumorigenicity. Further, Han *et al* have shown that up-regulation of Kv1.5 increases the K+ current density and sensitivity of SGC7901 cells to multiple chemotherapeutic drugs, adriamycin or 5-fluorouracil[9]. Expression of Kv4.1 has been found in the human GC cell lines, and its down-regulation inhibited cell proliferation via the blockage of G1-S transition[10]. Eag1 (Kv10.1) was aberrantly expressed in GC tissues and associated with cancer lymph node metastasis and stage[11]. Human ether-a-go-go-related gene (HERG) encodes one of the components of delayed rectifier K+ currents. In GC, HERG channel has revealed cancer-limited expression and its blocker diminishes the G1-S transition[12,13]. Ding *et al*[14] have shown that the survival rates for the hERG1-positive expression group are significantly lower than the negative group, and hERG1 expression is found to be an independent prognostic factor. Further, HERG expression has been reported to be essential for cisplatin to induce apoptosis in human GC[15]. Disruption of K+ channel protein, voltage-gated K+ channel subfamily E member 2 (KCNE2), has been shown as a possible risk factor for gastric neoplasia[16]. Kuwahara *et al*[17] have analyzed the expression of KCNE2 in surgically excised tissue from human GC associated with gastritis cystica profunda and confirmed that reduced KCNE2 expression correlates with disease formation. It has been proposed that atrial natriuretic peptide modulates the proliferation of human GC cells *via* voltage-gated K+ channel, KQT-like subfamily, member 1 (KCNQ1)[18]. Inwardly rectifying K+ channels (Kir) has been also implicated in GC. Lee *et al*[19] have demonstrated that knockdown of Kir2.2 suppresses tumorigenesis by inducing reactive oxygen species-mediated cellular senescence.

There is evidence also for Cl- channels involvement in GC. Overexpression of channels concerns chloride intracellular channel 1 (CLIC1) is shown to be a potential prognostic marker for GC[20]. Elevated CLIC1 expression is strongly correlated with lymph node metastasis, lymphatic invasion, perineural invasion and pathological staging, suggesting that it is a potential prognostic marker[20]. Zheng *et al*[21] have shown that PA28β regulates cell invasion of GC by modulating the expression of CLIC1 .On the other hand, Ma *et al*[22] have shown that high CLIC1 expression inhibits proliferation and enhance apoptosis, migration and invasion of GC cells.

The transient receptor potential (TRP) superfamily consists of a highly diverse group of ion channels that are mostly permeable to monovalent and divalent cations. TRP channels may be divided into seven subfamilies, including the classical (TRPC), the vanilloid receptor related (TRPV) and the melastatin related (TRPM) channels. In GC, Cai *et al*[23] have shown that Ca2+ elevation regulated by TRPC6 channels is essential for G2/M phase transition and suppresses growth in human GC cells. TRPV6 has been implicated in capsaicin-induced apoptosis in GC cells[24]. Further, several reports have shown important roles of TRPM7 in apoptosis and cell viability of GC cells[25-28].

There is significant evidence for involvement of these ion channels in GC cell proliferation and disease progression. Hence, their clinical potential would be worth investigating further.

**Regulation of water channels**

Aquaporins (AQPs) are transmembrane proteins that facilitate transport of water and, in some cases, small solutes across membranes; charged species are not permeated. Thus, AQPs are important for cell volume regulation and electrolyte balance under both physiological and pathophysiological conditions. To date, 13 AQP subtypes and their pathophysiologic roles have been characterized in humans. In GC, several reports indicated the role of AQP3 in signal pathway. Huang *et al*[29] have shown that AQP3 plays a critical role in human epidermal growth factor (hEGF) -induced cancer cell migration and proliferation and that hEGF induced AQP3 expression via ERK signal transduction pathways. Wang *et al*[30] have demonstrated that c-Met regulates the expression of AQP3 via the ERK signalling pathway in GC. Xu *et al*[31] have shown that AQP3 positively regulates matrix metalloproteinases (MMPs) proteins expression through PI3K/AKT signal pathway in human GC cells. Recently, the microRNA-mediated gene repression mechanism involved in AQP3's role has been investigated in GC[32]. AQP5 also plays an important role in the tumorigenesis, progression and differentiation of human GC cells[33,34]. Shen *et al*[35] have reported expression profiles of multiple AQPs in human GC and its clinical significance. AQP3 and AQP5 are detected remarkably stronger in the carcinoma tissues than that in normal mucosa by immunofluorescence. They have shown that both AQP3 and AQP5 expression are associated with lymph node metastasis and lymphovascular invasion in patients.

These results indicate that AQPs play important roles in the tumorigenesis and progression of human GC and suggest that especially, AQP3 and 5 can become potential therapeutic targets against GC.

**Regulation of intracellular pH**

Anion exchanger (AE) proteins facilitate the electroneutral exchange of Cl- for HCO3- across the plasma membrane of mammalian cells and thus contribute to regulation of intracellular pH. The AE family is now comprised of three members, AE1, AE2 and AE3. AE1 is frequently expressed in GC, where it fails to traffic to the plasma membrane, but interacts with the tumor suppressor p16 in the cytoplasm. Down-regulation of AE1 in gastric cancer SGC7901 cells is shown to inhibit cell growth and clinical analyses have indicated that AE1 expression is associated with a low survival rate of GC patients. Suppression of AE1 induces cell death in human GC cells[36-39]. Expression of AE2 in human GC has been also investigated, and AE2 is associated with gastric carcinogenesis and the achlorhydria[40]. Wang *et al*[41] have shown that early growth response protein 1 is critical for gastrin-dependent up-regulation of AE2 in GC cells.

The sodium-hydrogen exchanger (NHE) mediates a coupled counter-transport of one H+ ion in exchange for one Na+ ion. The basic role is to maintain intracellular pH, but NHE proteins are also important for regulation of cell volume and growth. Liu *et al*[42] have shown that NHE1 antisense gene significantly suppresses cell growth and induced cell apoptosis in SGC7901 cells. Nagata *et al*[43] have shown that rapid and extensive decrease of intracellular pH caused by NHE1 inhibitor leads cells to apoptotic and cytotoxic events in the MKN45 and MKN74 cells.

Vacuolar H+-ATPases (V-H+-ATPases), as the specific proton pump of the cell, play an important role in maintaining intracellular pH. Proton pump inhibitors (PPI), mainly treating acid-related diseases, could inhibit the expression of V-H+-ATPases. Chen *et al*[44] have shown that PPI decreases the intracellular pH of SGC7901 cells, by inhibiting V-H+-ATPases, and enhanced the cytotoxic effects of antitumor drugs.

The carbonic anhydrases (CAs) are a family of zinc metalloenzymes that have an important role in cellular pH regulation through reversible hydration of carbon dioxide to carbonic acid. To date, 16 isozymes have been identified, which differ in tissue distribution, subcellular localization, and catalytic activity. Expression of CA IX has been found at the invasion front of gastric cancers[45]. Kato *et al*[46] have shown that the CA IX expression level is significantly high in cases of type 4 GC and diffuse type GC, and significantly correlates with the invasion depth, lymph node metastasis. The prognosis for CA IX -positive patients is significantly poorer than that of CA IX -negative patients.

These results suggest that pH regulators, such as AEs, NHEs, V-H+-ATPases and CAs are potentially key therapeutic targets and the silencing of their expression could provide a new therapeutic approach for treating GC.

**Regulation of intracellular ion concentration**

Several reports indicating the important roles of Cl- channels/ transporters on cell proliferation suggest that the intracellular chloride concentration ([Cl-]i) regulated by them would be one of critical messengers. We have investigated roles of the [Cl-]i in cell cycle progression of human GC cells[47]. We have found that furosemide, a blocker of Na+/K+/2Cl- cotransporter (NKCC), diminished cell growth by delaying the G1-S phase progression in GC cells with high expression and activity of NKCC[48]. NKCC is one of the important transporters controlling the [Cl-]i *via* uptake of Cl- into the intracellular space and, therefore, furosemide decreases the [Cl-]i[49]. Cl- channels also contribute to the regulation of [Cl-]i which is related to cell volume. When cell shrinkage occurs isosmotically, [Cl-]i decreases because the major membrane-permeable anion is Cl-[50]. Furthermore, we have found that the decrease of the [Cl-]i inhibits cell growth of GC cells and that this inhibition of cell growth is due to cell cycle arrest at the G0/G1 phase caused by diminution of CDK2 and phosphorylated Rb[51]. The decrease of the [Cl-]i significantly increased expressions of p21 mRNA and protein[51]. In addition, we revealed that the [Cl-]i affects the cell proliferation *via* activation of MAPKs through up-regulation of p21 in GC cells[52]. Similar phenomena are also observed in GC cells with low [Cl-]i caused by inhibition of NHE[53]. These findings suggest that the [Cl-]i regulates important cellular functions in GC cells, leading to the development of novel therapeutic strategies.

**Regulation of osmolality**

Several previous studies have indicated the cytocidal effects of hypotonic stress on cancer cells. Lin *et al*[54] have reported that peritoneal lavage with distilled water improves the survival rate in patients with spontaneously ruptured hepatocellular carcinoma. Huguet *et al*[55] have discussed the optimal method for peritoneal lavage with distilled water during colorectal cancer surgery. In GC, Mercill *et al*[56] have reported that exposure to distilled water reduces the number of surviving gastric cells. Tsujitani *et al*[57] have shown that hypotonic intraperitoneal cisplatin treatment with distilled water at the time of a gastric resection is well tolerated for patents with GC. Recently, we have analyzed the changes in the cellular morphology and volume of GC cells subjected to hypotonic stress using several unique methods and apparatus, such as a differential interference contrast microscope connected to a highspeed digital video camera and a high-resolution flow cytometer[58]. Video recordings by high-speed digital camera have demonstrated that hypotonic shock with distilled water induces cell swelling followed by cell rupture. Measurements of cell volume changes using a high-resolution flow cytometer indicate that severe hypotonicity with distilled water increases broken fragments of GC cells within 5 min. In addition, we treated the GC cells with 5-nitro-2-3-phenylpropylamino)-benzoic acid (NPPB), a Cl- channel blocker, to enhance the cytocidal effects of the lavage by increasing their cell volume during the hypotonic stress via the inhibition of regulatory volume decrease (RVD)[59,60]. RVD occurs after hypotonicity-caused cell swelling. RVD is caused by activation of ion channels and transporters, which cause effluxes of K+, Cl-, and H2O, leading to cell shrinkage. NPPB is the broad spectrum Cl- channel blocker which is fat-soluble and inhibits both Cl- channels in cell membrane and CLIC. In the MKN45 and Kato-III cells, treatment with NPPB increases cell volume by inhibiting RVD and enhanced the cytocidal effects of the hypotonic solution (Figure 1). We have found similar phenomena in esophageal[61] and pancreatic cancer cells[62]. AQPs also contribute to RVD[63]. On the other hand, NKCC plays some roles in regulatory volume increase (RVI)[64].

These findings demonstrate the cytocidal effects of hypotonic shock on GC cells, and suggest that the regulation of ion transport enhances these effects. A deeper understanding of ion transport mechanisms in gastric cancer cells during hypotonic shock could lead us to the development of novel therapeutic strategies.

**Conclusion**

This review shows a variety of ion channels, AQPs and pH regulators are expressed in human GC cells and tissues. Their expression relates to the pathological character of the GC tissues. Pharmacological manipulation and gene silencing affect their activities and fundamental cellular functions that would be involved in the GC process. Overall, we can suggest that ion, water channels and pH regulators are functional biomarkers and therapeutic targets in GC. A deeper understanding of molecular mechanisms may lead us to the discovery of these cellular physiological approaches, such as regulation of ion channels, water channels, intracellular pH, intracellular ion concentration and osmolality as a novel therapeutic strategy for GC.

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**Figure 1Blockade of Cl- movement (channel) enhances the cytocidal effect of hypotonic solution *via* the inhibition of regulatory volume decrease in cancer cells.** Rvd: regulatory volume decrease.