**Name of Journal:** *World Journal of Gastrointestinal Surgery*

**Manuscript NO:** 82318

**Manuscript Type:** REVIEW

**Harnessing interventions during the immediate perioperative period to improve the long-term survival of patients following radical gastrectomy**

Liu LB *et al*. Perioperative therapeutic opportunity in GC

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**Author contributions:** Liu LB and Li J are equal coauthors of this article; Li J and Shi S designed the review; Liu LB, Lai JX and Li J reviewed the literature; Liu LB and Lai JX drafted the manuscript; Shi S revised the manuscript; All authors read and approved the final version of the manuscript.

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**Received:** December 15, 2022

**Revised:** January 4, 2023

**Accepted:** March 30, 2023

**Published online:** April 27, 2023

**Abstract**

Although the incidence and mortality of gastric cancer (GC) have been decreasing steadily worldwide, especially in East Asia, the disease burden of this malignancy is still very heavy. Except for tremendous progress in the management of GC by multidisciplinary treatment, surgical excision of the primary tumor is still the cornerstone intervention in the curative-intent treatment of GC. During the relatively short perioperative period, patients undergoing radical gastrectomy will suffer from at least part of the following perioperative events: Surgery, anesthesia, pain, intraoperative blood loss, allogeneic blood transfusion, postoperative complications, and their related anxiety, depression and stress response, which have been shown to affect long-term outcomes. Therefore, in recent years, studies have been carried out to find and test interventions during the perioperative period to improve the long-term survival of patients following radical gastrectomy, which will be the aim of this review.

**Key Words:** Radical gastrectomy; Perioperative events; Gastric cancer; Survival; Metastasis

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**Citation:** Liu LB, Li J, Lai JX, Shi S. Harnessing interventions during the immediate perioperative period to improve the long-term survival of patients following radical gastrectomy. *World J Gastrointest Surg* 2023; 15(4): 520-533

**URL:** https://www.wjgnet.com/1948-9366/full/v15/i4/520.htm

**DOI:** https://dx.doi.org/10.4240/wjgs.v15.i4.520

**Core Tip:** During the relatively short perioperative period, patients undergoing radical gastrectomy will suffer from various perioperative events, which have been shown to affect long-term outcomes. Therefore, in recent years, studies have been carried out to identify and test interventions during the perioperative period to improve the long-term survival of patients following radical gastrectomy. As the majority of these interventions are already safely applied clinically for other indications, are cost-effective and can be administered conveniently, if the desired survival benefits are prospectively confirmed, considerable economic and social improvements can be achieved at little financial cost.

**INTRODUCTION**

Although the incidence and mortality of gastric cancer (GC) have been declining gradually in recent decades, its survival improvement is relatively poorer than that of other common malignancies, such as colorectal and breast cancer[1]. In 2020, there were an estimated 768793 GC-related deaths[2]. Therefore, strategies aiming to decrease the burden of GC are being extensively explored globally. Currently, radical surgical removal of the primary GC is the preferred choice for patients whose disease is still locally resectable[1]. Unfortunately, postoperative development of recurrence and metastasis, the main cause of morbidity and mortality of GC, is inevitable in some patients, especially in those with advanced disease[3]. As no visible evidence of metastasis is the prerequisite for radical gastrectomy, postoperative relapses mainly result from occult cancer cells whose spreading has already occurred or been induced during the perioperative period by the surgery itself or its related events, including anesthesia, pain, intraoperative blood loss, allogeneic blood transfusion, postoperative complications (POCs), and their related anxiety, depression and stress response.

The notion that surgical trauma may enhance the risk of cancer metastasis was already noticed by the ancient Greeks, who cautioned against disturbing cancers[4]. In the 1960s, surgeons found that long-term survival was only moderately improved compared to historical nonoperated controls, and even rapid recurrence and progression were found in patients with cancer following radical resection, indicating the promoting effects of surgery on the spread of cancer cells[4]. However, these negative effects were largely ignored; only when perioperative adjuvant therapies have gained success in survival improvement has interest in this theory reemerged. Typically, perioperative adjuvant therapies for metastasis prevention are administered at least one month before or after surgery for cancer, including GC. The immediate perioperative period is rarely exploited for such interventions, largely owing to concerns over contraindications to surgery[5]. Such a concept has changed in recent years, as the significance of this timeframe in determining long-term oncological outcomes is widely recognized[5]. Therefore, various interventions have been explored during the perioperative timeframe, and some of them show great promise. As the recurrence and metastasis rates are higher and radical gastrectomy is relatively more extensive than surgeries for other malignancies, the immediate perioperative period of radical gastrectomy may be a critical timeframe to improve the prognosis of GC.

In this review, we briefly discuss the mechanisms underpinning the negative effects of radical gastrectomy and its related events and then describe the measures that could be harnessed to mitigate their cancer-promoting effects while improving the long-term survival of patients following radical gastrectomy, with the hope of transforming the perioperative period from a prominent facilitator of cancer recurrence to a window of opportunity for improving oncological outcomes in patients with GC.

**PERIOPERATIVE PHYSIOLOGICAL RESPONSE TO GASTRECTOMY AND ITS RELATED EVENTS**

In clinical settings, extensive surgery for GC always provides no additional survival benefits and even leads to poor survival in some patients[6,7]. Other necessary events during the perioperative period, including anesthesia and analgesia, can shorten the long-term survival of patients with GC if the modality or agents are administered inappropriately[8]. Adverse events during gastrectomy or postoperative recovery, such as intraoperative blood loss, allogeneic blood transfusion, and POCs, were all proven to be independent negative prognostic factors for patients following curative gastrectomy[8]. In addition, concomitant anxiety, depression and stress response can aggravate the cancer-promoting effects of gastrectomy and its related events. The perioperative physiological responses that underpin the cancer-promoting effects of radical surgery and its related events have been extensively studied in surgical oncology and have been excellently reviewed in previous publications[5]. Conclusions relevant to GC have also been discussed in our previous review[8]. Briefly, gastrectomy and its related events activate the sympathetic nervous system (SNS) and inflammatory response, also referred to as the surgical stress response, leading to enhanced growth of residual cancer cells, which may be preexisting micrometastases, incompletely resected fractions of tumor cells or disseminated from the primary tumor during the operation. In addition, following the activation of the surgical stress response, antitumor immunity is suppressed and fails to eliminate these residual cancer cells[5]. Therefore, measures that can alleviate the acute stress response to surgery and liberate the suppression of antitumor immunity are the focus of studies aiming to harness the immediate perioperative period for improving the long-term survival of patients with GC.

**POTENTIAL PERIOPERATIVE INTERVENTIONS**

For a long time, surgeons did everything just for operation and believed that the side effects of surgery must be borne. Therefore, it is not necessary to complicate the perioperative timeframe by additional interventions due to the relatively short time span of tumor evolution- or justified and/or speculated concerns over contraindications to surgery[9]. However, this concept has been challenged by three foundations. First, after curative resection, all visible cancer cells are removed, and the probability of future recurrence or metastasis mainly originates from minimal residual cancer cells, whose metastatic progression can be efficiently arrested or prevented by relatively minimal effort and innocuous therapies. In contrast, when these residual cancer cells evolve into larger and more self-sustaining diseases, this goal becomes more difficult or even impossible. Second, some existing interventions have been demonstrated to be tolerable or circumventable contraindications to surgery and can be administered safely during the perioperative period. Third, a robust biological rationale supports the likely antimetastatic efficacy of various interventions during the perioperative period, including appropriate operation, anesthesia and analgesia selection, approaches to limit stress-inflammatory responses and to preserve or activate anticancer immunity (Figure 1).

***Gastrectomy administered appropriately***

Historically, a series of randomized controlled trials (RCTs) have been conducted to compare the safety and survival benefit between radical gastrectomy with different intensities[10]. The primary principle for surgical management and clinical trial design of curatively resected GC is a balance between resection of tissues with possible cancer cell colonization to achieve long-term survival and acceptable postoperative early death. This high-quality evidence suggests that extensive gastrectomy has few advantages in improving long-term survival compared to less extensive surgery, and some extended resections even lead to increased recurrence or metastasis, indicating that the material benefit conferred by extended surgery may be offset by higher early mortality and increased recurrence resulting from an extensive surgical stress response[6,7]. Therefore, decision-making on surgical approaches for GC should be guided by the latest scientific evidence, and extended gastrectomy without survival benefit should be avoided. Within the modern knowledge of GC management, even in the West, the most appropriate surgery for GC is gastrectomy with D2 lymphadenectomy transabdominally but avoids inevitable paraaortic lymph node dissection, splenectomy, pancreatectomy, and bursectomy[10]. Minimally invasive surgery, such as endoscopic resection and laparoscopic or robotic gastrectomy for early or even advanced GC, is widely adopted as an alternative to traditional open gastrectomy. Although improved long-term survival has not been observed in these studies, less blood loss, fewer POCs, faster recovery, and reduced surgical stress have been found in patients undergoing minimally invasive surgery[11,12]. Therefore, minimally invasive surgery for GC is recommended whenever the conditions of surgeons and patients are feasible. Theoretically, the reduced surgical stress during minimally invasive surgery may translate into a long-term survival benefit when other antimetastatic interventions are coadministered perioperatively.

Nevertheless, both minimally invasive and traditional surgical approaches for GC are highly technically demanding, with a high incidence of intraoperative blood loss and POCs in less experienced hands. Numerous data support the negative effects of intraoperative blood loss, transfusion and POCs on prognosis in patients following radical gastrectomy[8]. In addition, the inferior short-term and long-term outcomes following radical gastrectomy in the East compared with the West were not only determined by more advanced stage and comorbidities but also by the low incidence of GC and uncommon regional specialization. Several other studies have also found that the outcomes of patients with GC were better in experienced and high-volume hospitals[13,14]. Therefore, from the viewpoint of survival benefit, gastrectomy should be centralized in high-throughput centers with the ability to provide standardized perioperative management for GC.

***Choosing the most appropriate anesthesia and analgesia***

Patients diagnosed with locally resectable GC will require anesthesia for endoscopic examination or gastrectomy, and analgesics are commonly prescribed for pain relief following surgery. Currently, several modalities and agents are widely applied for general anesthesia and postoperative analgesia, while total intravenous anesthesia (TIVA), inhalation anesthesia and neuraxial anesthesia are the most studied in cancer surgery. Evidence from studies in GC suggests that these modalities and agents have distinct effects on the stress response, inflammation, anticancer immunity, cancer progression and long-term survival. Inhalational anesthetics, including isoflurane and sevoflurane, have been observed to provide some degree of cytoprotection to organs, which may also support the survival of cancer cells[15]. Although data in GC are lacking, *in vivo* data in other cancer types demonstrate that volatile anesthetics might promote immunosuppression and support tumor cell growth and spreading[16,17]. In contrast, propofol-based TIVA, an alternative to inhalational anesthesia, has appealing anticancer properties and has been extensively studied in GC. Administration of propofol has been shown to inhibit GC cell proliferation, migration and invasion *in vitro* while preserving the cellular immune function of patients undergoing curative gastrectomy[18,19]. Consistent with this evidence, the findings of several retrospective studies or meta-analyses demonstrated that in patients undergoing gastrectomy, long-term survival is better in patients anesthetized with propofol-based TIVA than in those with volatile anesthesia[20]. However, what calls for special attention is that all these studies were retrospectively designed, and selection bias cannot be neglected, providing a limited reference for the choice of anesthesia type during gastrectomy. Currently, a number of prospective trials elucidating the effects of propofol on cancer recurrence and patient survival are ongoing (NCT01975064, NCT03034096, NCT02660411, NCT02840227), and their results may raise the possibility that propofol-based TIVA could become the standard anesthesia approach for cancer surgery, including gastrectomy.

Epidural anesthesia has been widely used jointly with or as an alternative to general anesthesia or for postoperative analgesia in patients undergoing gastrointestinal cancer surgery. Limited clinical evidence suggests the survival benefit of epidural anesthesia for patients with GC[21,22]. The association between epidural anesthesia and decreased cancer recurrence following surgery might reflect the multifaceted effects of this anesthetic and analgesic modality. The addition of epidural anesthesia to general anesthesia significantly decreased the expression of various inflammatory mediators and increased the proportion and activity of antitumor immune cells in patients undergoing GC surgery[23,24]. SNS blockade and inhibition of perioperative lymph flow both contribute to the anticancer effect of epidural anesthesia, although they have not been validated in GC[25,26]. Furthermore, as a widely adopted postoperative analgesic technique, epidural anesthesia significantly decreases the prescription of opioids for postoperative pain relief[22]. Opioids have been found to promote the growth of cancer and inhibit the antimetastatic activity of immune cells, including natural killer (NK) cells, cytotoxic T lymphocytes, dendritic cells and macrophages[27,28]. Nevertheless, the results from two RCTs including lung or unspecified cancer types did not support the positive effect of epidural anesthesia on overall or cancer-specific survival[29,30]. Therefore, determining whether epidural anesthesia provides a recurrence-preventing effect for patients with GC requires further research.

***Inhibiting sympathetic signaling***

Activation of the SNS is one of the responses to surgical stress, resulting in increased levels of circulating catecholamines, including adrenaline and noradrenaline. Consistent with this mechanism, the levels of circulating catecholamines were higher in patients undergoing more extended gastrectomy or with an eventful postoperative recovery[5]. Beta 2-adrenergic receptor (β2-AR), the main type of receptor that mediates the biological functions of catecholamines, was found to be overexpressed in cancer tissues from patients with GC and correlated with metastasis and poor prognosis[31]. The activation of β2-AR by catecholamines leads to increased formation of liver and lung metastases by primary GC cells[31]. Mechanisms underlying the cancer-promoting effect of SNS signaling have also been elucidated in various studies. Activation of β2-AR was shown to induce epithelial to mesenchymal transition and cancer stem cell attributes of GC cells[32,33]. Sympathetic nerves can also help establish premetastatic niches in bone by stimulating host bone marrow stromal cells[34]. On the other hand, SNS activation promotes the establishment of an immune-privileged microenvironment, which is beneficial to tumor growth and metastasis formation[35,36]. Therefore, increasing data reported in recent years indicate that perioperative interventions to attenuate SNS activity are promising for reducing the recurrence risk of GC, which can be achieved by pharmaceutical inhibition (β-AR antagonists) or by epidural anesthesia.

Population-based cohort studies have shown that β-AR antagonists (also known as β-blockers), which are widely used in the clinic for hypertension, can significantly decrease the risk for GC[37]. Therefore, as effective strategies to inhibit sympathetic signaling, which is activated by perioperative events, β-blockers may be used as an effective adjunctive strategy to reduce the risk of cancer recurrence. *In vitro* studies, propranolol, the most commonly used nonselective-adrenergic antagonist, showed the ability to induce apoptosis, repress growth, suppress the expression of matrix metalloproteinases and vascular endothelial growth factor, and inhibit the migration of GC cells[38,39]. In a xenograft mouse model, propranolol decreased the levels of phosphorylated AKT, MEK, and ERK proteins and blocked depression-promoted neuroendocrine phenotypic transformation and lung metastasis of GC[31,40,41]. In patients with GC, treatment with propranolol for one week before surgery significantly inhibited the proliferation of cancer, as measured by Ki-67[40]. In other cancer types, the anticancer ability of propranolol has also been found to be associated with enhanced antitumor immunity[42,43]. However, this effect was not observed in GC animal models or patients[40]. Based on the findings of preclinical studies, several clinical trials have been carried out to assess the effects of perioperative β-blockers on oncological outcomes, but none of them have been designed to focus on survival. Only one small RCT in colorectal cancer collected data on the 3-year recurrence rate and revealed a favorable trend toward reduced cancer recurrence in patients receiving β-blockers[44]. Compared with surgery for breast and colorectal cancer, gastrectomy for GC has a relatively higher intensity. It is reasonable to speculate that the intensity of activation of sympathetic responses in radical gastrectomy will be stronger, and β-blockers do have the ability to decrease the surgical responses and have the probability of decreasing the recurrence in GC patients.

***Anti-inflammatory therapy***

One of the explanations for the lack of effect on long-term outcomes by SNS blockade is that in addition to increased levels of catecholamine, various inflammatory mediators are also abundantly released into circulation during cancer surgery. The levels of inflammatory mediators, including C-reactive protein (CRP), procalcitonin, prostaglandin E2 and plasma cortisol, were elevated significantly after gastrectomy[45,46]. Activation of the inflammatory response plays an important role in residual cancer cell survival and progression through humoral factor release or premetastatic niche establishment[47-49]. A preoperative elevated neutrophil-to-lymphocyte ratio, a systemic inflammation index frequently used in the literature, was shown to have a significant negative effect on survival[50]. Therefore, anti-inflammatory therapy might provide antimetastatic benefits. Selective or nonselective nonsteroidal anti-inflammatory drugs (NSAIDs) for cyclooxygenase (COX) (for example, ibuprofen, celecoxib, and etodolac) are commonly used analgesics following radical gastrectomy, and several studies have shown their anticancer properties. Currently, the most tested anti-inflammatory therapies are selective COX2 inhibitors. In cancer types other than GC, perioperative use of NSAIDs has been shown to attenuate the inflammatory response, enhance the number and function of antimetastatic immune cells and prevent the formation of metastases in mouse models[51,52]. In the clinical setting, studies have reported that perioperative administration of COX2 antagonists decreases the circulating levels of prostaglandins and catecholamines while preserving anticancer immunity by buffering both the elevation of regulatory T cells and the decline in NK cell counts[5]. The anticancer effect of NSAIDs, especially celecoxib, has also been studied extensively in GC, while data are limited to preclinical or GC prevention settings. For example, a population-based intervention trial revealed that celecoxib treatment alone had beneficial effects on the regression of advanced gastric lesions[53]. Selective COX-2 inhibitors were found to suppress GC cell dissemination through apoptosis induction and migration suppression[54,55]. In a mouse model bearing orthotopic xenografts or with carcinomatous peritonitis induced with a highly metastatic human diffuse-type GC cell line, etodolac, a COX-2 inhibitor, significantly decreased tumor lymphangiogenesis and the total weight of metastatic lymph nodes[56]. In the perioperative setting, only one study reported that preoperative treatment with celecoxib significantly promotes necrosis in GC through the induction of apoptosis and the reduction of microvessel density[57].

Data from these preclinical studies and clinical studies on inflammatory biomarker alterations point to an anticancer effect of NSAIDs. Therefore, several clinical trials have also been performed to investigate the long-term outcome impact of NSAIDs on cancer. A retrospective analysis suggested that intraoperative administration of ketorolac decreases the risk of breast cancer relapse compared with other analgesics[58]. A nationwide cohort study involving 15574 patients revealed that the use of NSAIDs can be associated with a reduced risk of early hepatocellular carcinoma recurrence within 2 years after curative liver resection[59]. However, these studies were retrospectively designed and have inherent limitations. Thus, several RCTs aimed at investigating the effect of perioperative administration of NSAIDs on improving cancer-specific survival are currently ongoing. One of these studies was completed and reported that preoperative administration of ketorolac tromethamine does not increase disease-free survival (DFS) or overall survival (OS) in high-risk breast cancer patients[60]. However, because of the relatively low surgical trauma and low recurrence rate in breast cancer, the results cannot be extrapolated to other cancer types. In addition, as mentioned above, activation of the SNS and inflammatory response have redundant roles in promoting the growth and recurrence of cancer, and anticipated survival improvement cannot be achieved by COX2 inhibitors alone. Therefore, the combination of β-blockers and COX2 inhibitors is being investigated in several RCTs. Primary results revealed that perioperative administration of β-blockers and COX2 inhibitors not only improves immune competence and metastatic biomarkers but also shows a favorable trend toward reduced cancer recurrence in treated patients[44,61]. To date, no such RCTs recruiting patients with GC have been reported or registered. As radical gastrectomy activates a relatively more severe surgical stress response, whether pharmacological blockade of these responses will provide a survival benefit deserves further clinical trials.

***Perioperative immune stimulation***

Immunosuppression is a widely recognized phenomenon in patients with cancer, especially during the perioperative period[9]. The total lymphocyte count decreased rapidly from preoperative levels, while the expression of lymphocyte activation gene 3 and programmed cell death 1 on lymphocytes was upregulated, indicating impaired cell-mediated immunity after surgery for GC[62]. Potential contributors to perioperative immunosuppression include the stress-inflammatory response, anxiety, hypothermia, blood loss and transfusion, and the direct and indirect effects of anesthetic and analgesic agents[9]. As immunosurveillance critically determines the fate of minimal residual cancer cells, manipulation of anticancer immunity may provide a promising opportunity to improve the long-term survival of patients following curative surgery[63]. To date, various interventions to activate host anticancer immunity have joined the therapeutic armamentarium for the treatment of many advanced-stage cancers; however, few of them have been tested during the perioperative timeframe, owing to several established and theoretical risks pertinent around the time of surgery[9]. Nevertheless, perioperative immune preservation or stimulation could hold various advantages if the interventions meet the following desired attributes: A rapid immunological response, avoidance of tumor-promoting effects, minimal contraindications to surgery, resilience to perioperative stress and a limited capacity to induce stress responses[9]. Potential strategies have been investigated in various cancers, including GC.

***Cytokines***

Cytokines are essential for an effective anticancer immune response. Treatment with recombinant interleukin-2 (IL-2), a crucial cytokine for various leukocytes, has long been tested in perioperative settings. For example, preoperative treatment with low-dose IL-2 can be safely given in patients with GC and was revealed to prevent postoperative lymphocytopenia and activate peripheral and peritumoral lymphocytes[64]. Furthermore, IL-2 seems to have an impact on the clinical course, reducing the morbidity of surgery and ameliorating OS and DFS[65]. Other cytokines, such as type I interferons and granulocyte colony-stimulating factor, have also been investigated in the perioperative period and were shown to improve anticancer immunity; however, the survival benefits were somewhat heterogeneous[66-68]. Except for these primary promising results, perioperative administration of cytokines has been discontinued in recent years, largely owing to their severe systemic adverse effects, including the induction of fever, weakness and headaches, which cannot be distinguished from signs of infection and might result in surgery being delayed.

***Immunonutrition***

In contrast to other strategies to improve anticancer immunity, data on the beneficial effects of immunonutrients administered perioperatively are accumulating in GC[69]. Immunonutrient interventions are nutrition support that is rich in elements beneficial for the homeostasis of immunity, including ω-3-fatty acids, glutamine, arginine, and nucleotides[70]. Although not all studies found similar clinical effects and some conflicting results have been reported, the influence of immunonutrition on immunological levels, nutrition status and postoperative course was convincing. For example, an early 5-d postoperative enteral immunonutrition supplement significantly improved immune function and the inflammatory response in GC patients following gastrectomy[71]. When immunonutrition was supplied before surgery, the abundance of tumor-infiltrating lymphocytes was upregulated in gastrectomy samples[72]. Several meta-analyses also demonstrated the efficacy of perioperative immunonutrition for improving various immunological indices and decreasing the incidence of POCs[69,73,74]. Although the benefit of improving immune function has been consistently reported, few studies have collected survival data. One study concluded that the 60-d mortality was lower in patients receiving immunomodulating enteral nutrition in the perioperative period, but no improvement in 6-mo and 1-year survival was found[75]. The explanations may be that only a small number of patients were included, and in some studies, prolonged use of immunonutrition increased tumor angiogenesis, which may offset the survival benefit of immunonutrition[72,75]. Therefore, whether perioperative immunonutrition support can provide a survival benefit needs further large prospective studies, and the ingredients and duration of immunonutrition should be determined carefully.

***Toll-like receptor agonists***

Some immunotherapeutic strategies, such as immune checkpoint inhibitors, require repeated administration for weeks or months to induce the desired response, making them inappropriate for perioperative settings. However, other approaches, such as some Toll-like receptor (TLR) agonists, can induce rapid activation of immune responses, which was shown to be effective when administered perioperatively. For example, preoperative treatment with CpG-C oligodeoxynucleotide, a TLR9 agonist, can synergize with propranolol and etodolac to improve cell-mediated immunity and limit metastatic progression in a mouse model[76]. A fully synthetic TLR4 agonist, glucopyranosyl lipid-A, can be safely administered perioperatively and significantly elevates both innate and adaptive immunity, leading to reduced metastatic development[77]. The TLR3 agonist polyinosinic-polycytidylic acid [poly(I:C)] significantly enhanced NK cell activity in preclinical tumor models, healthy human donors and cancer surgery patients[78]. To date, evidence is limited to preclinical models, none of them have been pursued in clinical trials, and the perioperative application potential is not clear in GC. However, as suggested by promising preclinical rationale, research into perioperative immune stimulation is warranted in the future.

***Nutrition support and exercise***

GC patients always present with malnutrition, largely owing to digestion and absorption dysfunction, obstruction attributed to cancer, and anorexia caused by cancer-released cytokines. Numerous studies have reported that malnutrition is associated with poor prognosis, which may be the result of interference with treatment implementation and impaired anticancer immunity[79-81]. Therefore, perioperative nutrition support was proven to improve immune function, weaken the surgical stress response and decrease POCs, thereby theoretically prolonging the survival of GC[82]. However, no survival data are available to date. On the other hand, preclinical studies have suggested that excessive parenteral nutrition support could potentially promote the proliferation of cancer cells[83]. Accordingly, determining the patients who need nutritional support and how to carry out nutritional support optimally are the focus of future studies.

Physical fitness plays an important role in the successful administration of various cancer therapies, including surgery, and thus may determine the long-term survival of patients postoperatively. Low physical performance, partially determined by the loss of muscle mass plus low muscle strength, is associated with POCs and poor prognosis in GC patients[84-86]. Correspondingly, increasing the physical activity of patients through perioperative exercise, always administered simultaneously with nutrition support, decreased the incidence of POCs and enhanced the recovery course following gastrectomy[87]. However, although prolongation of survival has been achieved by exercise in patients with colorectal cancer, whether perioperative exercise programs have clinical benefits with regard to long-term oncological outcomes in GC patients is unclear[88,89]. Mechanisms underlying the protective effect of exercise on cancer mortality are multifarious. For example, exercise was found to decrease the inflammatory marker CRP, indicating an anti-inflammatory effect[90]. Moderate exercise is known to enhance cellular immunity and to decrease the levels of insulin and insulin-like growth factor[91,92]. In addition, activity-induced changes in the body and mental health also support improved tolerance for and the resultant effectiveness of cancer treatment[93]. Therefore, as a simple and convenient intervention that can be safely implemented during the perioperative period, exercise may provide the desired survival improvement in patients following gastrectomy.

***Patient blood management***

Similar to malnutrition, anemia is another most common presentation in GC patients, and the incidence was reported to range from 27% to 44%[94]. Pretreatment anemia predicts increased POCs and decreased long-term survival, including DFS[94]. Therefore, perioperative allogeneic blood transfusion is frequently administered in GC patients. Paradoxically, abundant data suggest that transfusion is intimately associated with cancer recurrence and cancer-related deaths following radical gastrectomy, mainly owing to the inhibition of host immunity and increased risk of POCs[95].

Therefore, in the consideration of long-term survival, the optimization of preoperative anemia treatment is critical for patients with GC. For this purpose, a patient blood management (PBM) strategy was established, which includes different evidence-based interventions, aiming to maintain patients' own blood volume and avoid unnecessary blood transfusion. PBM consists of three parts: Management of anemia through early detection and use of iron preparations to stimulate erythropoiesis; minimization of perioperative blood loss; and optimization of patient-specific physiological tolerance to anemia with a restrictive transfusion strategy[96]. PBM has been successfully applied in orthopedic, cardiac and colorectal surgery, showing the ability to reduce blood transfusion and hospital stay[96]. A Spanish multicenter study applied the PBM strategy to the management of preoperative anemia in GC and reported that PBM can significantly reduce perioperative blood transfusion, especially in patients with preoperative anemia[97]. In addition, reduced postoperative infectious complications, reoperation rate, average hospital stay and mortality were observed in patients under PBM[97]. Although effects on cancer recurrence and long-term survival have not been reported in this study, all short-term benefits mean a possible improved prognosis[97]. PBM requires the completion of laboratory work-up 2-4 wk before surgery, and preoperative iron supplementation at least 7 d before surgery is recommended in patients whose hemoglobin (Hb) is less than 120 g/L and/or ferritin is less than 300 mg/L. For patients with risk factors and/or anemia symptoms, the recommended Hb threshold for blood transfusion is set as 90 g/L; otherwise, transfusion is only recommended for patients with Hb less than 70 g/L[97]. As speculated unfavorable outcomes of delayed surgery for GC, such a long preoperative time spent for PBM may impede its implementation; therefore, only 52% of patients accomplished the preoperative PBM, much higher than that in a previous study on colon cancer, in which only 30% of patients accomplished the preoperative PBM[97,98]. Nevertheless, the implementation of PBM did not significantly prolong the time interval between diagnosis and surgery in the Spanish study[97]. Moreover, there is no evidence that the time required to complete PBM will obviously lead to tumor progression and affect long-term survival. In contrast, the benefits provided by reduced blood transfusion, infectious complications and mortality far exceed the disadvantages of the additional time required for PBM. Therefore, in a German study including some upper gastrointestinal tumors, a preoperative PBM strategy improved the two-year survival rate by 15%, although no subgroup analysis was performed to determine whether the survival benefit remained in GC patients[99]. Therefore, the PBM strategy may balance the contradiction between anemia and transfusion, possibly avoiding the negative effects in both situations while improving the survival of patients following radical gastrectomy, which awaits the results of trials with adequate follow-up.

***Enhanced recovery after surgery***

In the past decade, the effects of numerous perioperative interventions, including nutrition, minimally invasive surgery, nasogastric/nasojejunal decompression, early postoperative diet and mobilization, on immediate postsurgical outcomes have been studied extensively in radical gastrectomy, and these interventions have been integrated into enhanced recovery after surgery (ERAS), an evidence-based, comprehensive, multimodal approach designed to achieve early recovery for patients undergoing radical gastrectomy[100,101]. Published studies on ERAS mainly reported short-term outcomes, with similar POC incidences but reduced postoperative hospitalization and costs[102]. The ERAS approach was also found to improve the postoperative inflammatory response and surgical stress[103,104]. However, no study has yet reported the long-term survival of patients experiencing an enhanced recovery after radical gastrectomy. As the interventions comprising the ERAS approach often overlap with the principles presented herein to limit the deleterious effects of gastrectomy on surgical stress, which may induce the recurrence of GC, it is our recommendation to incorporate them in conjunction with studying oncological outcomes.

**CONCLUSION**

Globally, most of the one million newly diagnosed GC patients require gastrectomy each year. Gastrectomy and its related events, including anesthesia, analgesia, transfusion, POCs and malnutrition, will expose these patients to various stress responses during the immediate perioperative period. Pathophysiological alterations, such as activation of the SNS and inflammatory response and suppression of anticancer immunity, can support the survival and growth of residual cancer cells and promote cancer recurrence. Therefore, exploiting perioperative interventions to reduce the risk of recurrence and metastasis has attracted more attention in recent years. Various approaches, including appropriate operation and anesthesia selection, anti-adrenergic, anti-inflammatory, perioperative immune stimulation, nutrition support, exercise, and PBM, have been widely explored in preclinical or clinical settings, and promising results have been reported. Although data on some approaches are limited or lacking in GC at present, some of them did show the potential to improve the long-term survival of patients with various cancers. However, the majority of evidence was provided by retrospective analysis, and conflicting results have also been observed in clinical trials, perhaps owing to the complex pathophysiological alterations and heterogeneity among patients, leading to the lack of consensus on the optimal approach to perioperative care. In addition, along with the accumulating knowledge of the mechanisms underpinning the invasion-metastasis cascade, the concept of drugging metastasis has attracted more attention[105]. The immediate perioperative period represents a critical timeframe for residual cancer cells to complete the invasion-metastasis cascade, providing an important window for enhancing the efficacy of drugs targeting metastasis. Therefore, a detailed understanding of the changes that occur after surgery in each patient is pivotal for the development of new therapeutic strategies and personalized health care to prevent tumor recurrence. Large-cohort prospective RCTs are required to definitively demonstrate the effects of various perioperative interventions on oncological outcomes after radical gastrectomy. As the majority of these interventions are already safely applied clinically for other indications, are cost-effective and can be administered conveniently, if the desired survival benefits are prospectively confirmed, considerable economic and social improvements can be achieved at little financial cost.

**ACKNOWLEDGEMENTS**

The author thanks the Health Commission of Mianyang City and the Science and Education Department of the Third Hospital of Mianyang for their support. The space limitations of this review have unfortunately meant that we have not been able to separately cite many of the original publications that have contributed substantially to the literature. We sincerely apologize to the authors of these publications.

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**Footnotes**

**Conflict-of-interest statement:** All theauthors report no relevant conflicts of interest for this article.

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**Provenance and peer review:** Invited article; Externally peer reviewed.

**Peer-review model:** Single blind

**Peer-review started:** December 15, 2022

**First decision:** January 3, 2023

**Article in press:** March 30, 2023

**Specialty type:** Gastroenterology and hepatology

**Country/Territory of origin:** China

**Peer-review report’s scientific quality classification**

Grade A (Excellent): A

Grade B (Very good): 0

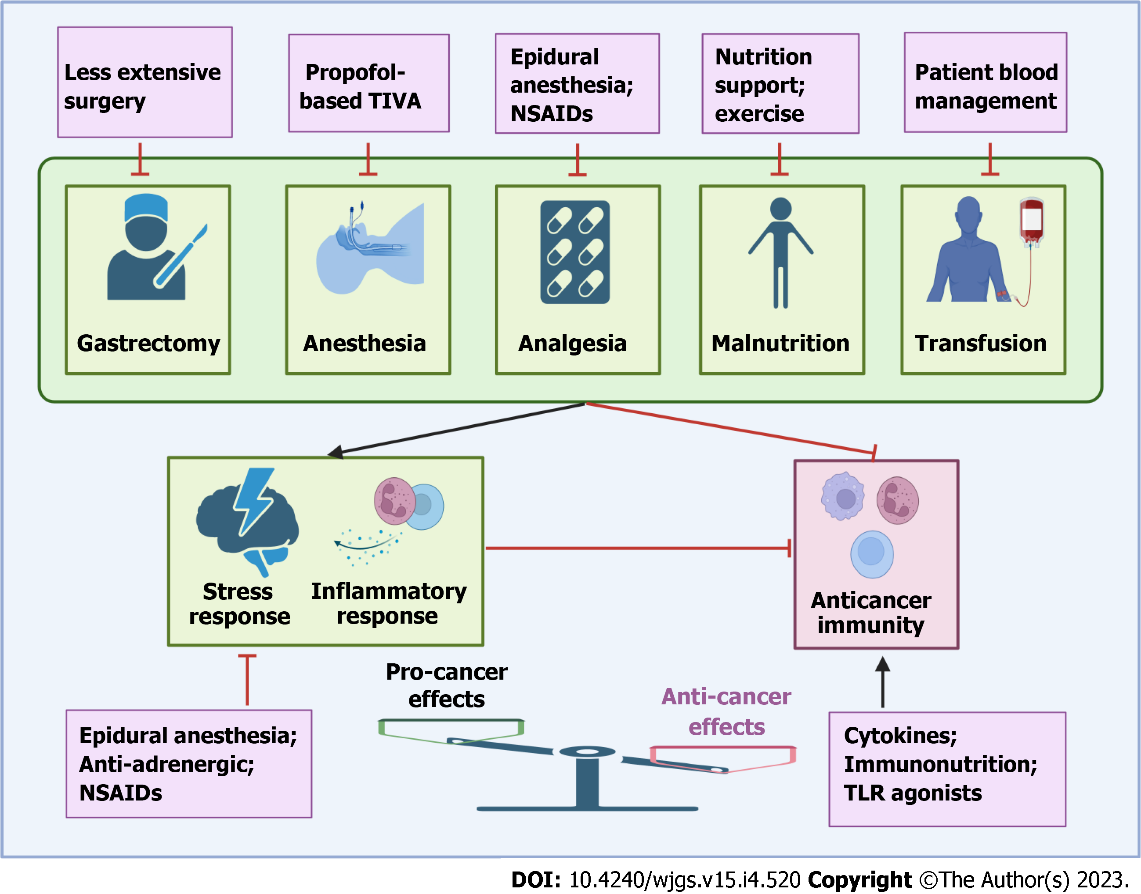
Grade C (Good): C

Grade D (Fair): 0

Grade E (Poor): 0

**P-Reviewer:** Kanaoujiya R, India; Zhi X, China **S-Editor:** Fan JR **L-Editor:** A **P-Editor:** Zhao S

**Figure Legends**



**Figure 1 Potential perioperative interventions for improving the survival of gastric cancer patients following curative resection.** A robust biological rationale supports the likely antimetastatic efficacy of various interventions during the perioperative period, including appropriate operation, anesthesia and analgesia selection, approaches to limit stress-inflammatory responses and preservation or activation of anticancer immunity. The figure in this review was created with BioRender.com. NSAIDs: Nonsteroidal anti-inflammatory drugs; TLR: Toll-like receptor; TIVA: Total intravenous anesthesia.



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