Emerging Roles of the Nervous System in Gastrointestinal Cancer Development

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Simple Summary: Nerve–cancer cross-talk has increasingly become a focus of the oncology field, particularly in gastrointestinal (GI) cancers. The indispensable roles of the nervous system in GI tumorigenesis and malignancy have been dissected by epidemiological, experimental animal and mechanistic data. Herein, we review and integrate recent discoveries linking the nervous system to GI cancer initiation and progression, and focus on the molecular mechanisms by which nerves and neural receptor pathways drive GI malignancy.

Abstract: Our understanding of the fascinating connection between nervous system and gastrointestinal (GI) tumorigenesis has expanded greatly in recent years. Recent studies revealed that neurogenesis plays an active part in GI tumor initiation and progression. Tumor-driven neurogenesis, as well as neurite outgrowth of the pre-existing peripheral nervous system (PNS), may fuel GI tumor progression via facilitating cancer cell proliferation, chemoresistance, invasion and immune escape. Neurotransmitters and neuropeptides drive the activation of various oncogenic pathways downstream of neural receptors within cancer cells, underscoring the importance of neural signaling pathways in GI tumor malignancy. In addition, neural infiltration also plays an integral role in tumor microenvironments, and contributes to an environment in favor of tumor angiogenesis, immune evasion and invasion. Blockade of tumor innervation via denervation or pharmacological agents may serve as a promising therapeutic strategy against GI tumors. In this review, we summarize recent findings linking the nervous system to GI tumor progression, set the spotlight on the molecular mechanisms by which neural signaling fuels cancer aggressiveness, and highlight the importance of targeting neural mechanisms in GI tumor therapy.

Keywords: nervous system; gastrointestinal cancer; chronic stress; neurogenesis; neurotransmitter; tumor microenvironment

1. Introduction

Gastrointestinal (GI) cancers, composed of esophageal cancer (EC), gastric cancer (GC), pancreatic cancer (PC), colorectal cancer (CRC), and liver cancer (LC), are some of
the most frequently-diagnosed cancers worldwide. GI tumors are estimated to account for over one quarter of all cancer cases and one third of cancer-associated deaths [1]. The development of these tumors is heavily influenced both by genetic predispositions and microenvironmental factors. While the genetic basis of GI cancer development has been intensively investigated, some microenvironmental grounds remain to be untangled [2].

Studies in recent years uncovered a close relationship between nerves and cancer development, especially in GI tumors [3]. Epidemiological data have provided solid evidence linking psychological factors, such as chronic stress and depression, to increased GI cancer risk in the general population [4]. Nevertheless, mechanistic knowledge of the cross-talk between nerve and GI cancers lags well behind other aspects of tumor biology, such as cancer genetics, immunity and metabolism. This is partially attributable to the complexity of nerve–cancer cross-talk, and the difficulty in establishing reliable experimental models to recapture their interactions [5,6]. For instance, while tumor-associated neurogenesis has been observed since several decades ago, the underlying mechanisms have only been partially elucidated in recent years [3]. During GI tumorigenesis, delicate coordination among different cell types and tumor microenvironment underlies active neurogenesis, whose process is technically intractable using in vitro models [7,8]. Likewise, the roles of nerve fibers in GI cancer malignancy involve various neurotransmitters, neurotrophins and neuropeptides [9]. Their source and impacts on tumor malignancy and microenvironment involve complicated molecular mechanisms, whose elucidation requires vigorous investigations by both basic and clinical researchers. As a consequence, the involvement of the nervous system in GI tumor development remains elusive and needs to be fully understood.

The role of nervous system in tumorigenesis involves a wide range of aspects, at both systemic (neuroendocrine system) and tissue-specific (via tumor-associated nerve fibers) levels [10]. While the role of the neuroendocrine pathway in tumorigenesis has been well-recognized, it is only very recently that tumor-associated neurogenesis has been established as an important driver of GI tumorigenesis [3]. Moreover, mounting data indicated that a variety of neurotransmitter, neurotrophin and neuropeptide receptors are widely expressed in GI tumors. These findings, combined with recent in vitro results showing that nerve fibers fuel solid tumor growth and malignancy, point to a direct involvement of neural infiltration in the development of GI cancers [8]. Neural infiltration and neurotransmitter pathways appear to affect various aspects of GI development and malignancy, such as growth, invasion, chemoresistance, immune evasion and tumor microenvironments (TME). More importantly, these discoveries may lead to novel therapeutic strategies against GI tumors and benefit patients with diminishing responses to current therapies. In this review, we will discuss recent advances in neurological aspects of GI tumorigenesis and present a brief summary of the involvement of nerve fiber outgrowth and neural receptor pathways in GI cancer chemoresistance, malignancy and metastasis. This review will also discuss the potential of targeting neurological pathways as new treatments of GI tumors.

2. Chronic Stress Contributes to GI Tumor Risk

GI tissues are heavily innervated by the enteric nervous system (ENS) [11]. The ENS is the intrinsic nervous system controlling the function of the gastrointestinal tract and other digestive organs [12,13]. Within the GI tract, the ENS consists of thousands of ganglia identified between the longitudinal and the circular muscle layers (myenteric plexuses) and between the muscle and mucosal layers (submucosal plexuses), the neural connections that communicate these ganglia and nerve fibers that innervate effector tissues [14].
nerve fibers of the ENS are distributed extensively within all layers of GI tract, including muscle layers, the submucosa, mucosal crypts and epithelium, to facilitate a wide range of GI functions [15]. While the ENS can function autonomously ex vivo, normal innervation of the digestive system requires integrated communications among the ENS and other nervous systems, particularly the central nervous system (CNS). The bidirectional communication between the ENS and CNS, termed as gut–brain axis (GBA), plays a central role in governing GI function [12,13]. GBA also establishes a link between psychological activity and GI homeostasis, given the fact that mental conditions have solid implications in GI disease progression, and vice versa [16]. Chronic stress conditions, such as depression and anxiety, are established risk factors for various GI disorders, underscoring the importance of psychological factors in GI disease progression [17]. Chronic stress has been linked to the initiation and progression of chronic GI disorders, possibly through aberrant release of neurotransmitters and neuropeptides by the ENS [18].

While the role of psychological factors on GI tumorigenesis remains under debate and inconclusive, some epidemiological data indicated that traumatic events, such as loss of close relatives, may increase the risk of multiple GI tumors, including pancreatic, gastric and colorectal cancers [19,20]. In addition, moderate associations among other stress-related syndromes, such as depression and anxiety, and GI cancer incidence have been revealed [4,21]. These effects of psychological factors were perfectly captured by elegant animal studies showing that enriched physical environments attenuate colon and pancreatic cancer progression [22,23]. Importantly, clinical data and animal studies confirmed that chronic stress may facilitate precancerous lesions in the upper gastrointestinal tract [24,25]. Collectively, these results point to a potential link between chronic stress and increased GI cancer incident, highlighting the importance of psychological factors in the initiation of GI tumors.

The mechanisms by which chronic stress exacerbates GI cancer risk are complex. Studies proposed that the neuroendocrine pathways, including the GBA and the hypothalamus–pituitary–adrenal (HPA) axis, may contribute to chronic stress-augmented GI cancer risk. This effect may involve local immune dysfunction and inflammation via increased levels of stress hormones, such as cortisol, adrenaline and noradrenaline [26,27]. Early studies indicated that psychological factors are associated with increased levels of pro-inflammatory cytokines, such as IL-6, and systemic low-grade inflammation in human populations [28–31]. Animal studies confirmed that chronic stress induces low-grade inflammation and the upregulation of IL-6, TNF-α, and CRP, which may contribute to tumor initiation [32]. In addition to a role in promoting inflammation, recent studies provided other explanations linking chronic stress to GI cancer risk. For instance, it was recently revealed that chronic stress may accelerate the degradation of p53 via the β2-adrenoreceptor (β2-AR)/β-arrestin-1 pathway, leading to genomic instability and the accumulation of DNA damages [33]. Another study showed that glucocorticoids may decrease p53 protein level via SGK1-Mdm2 pathway, which augments ionizing radiation (IR)-induced tumorigenesis [34]. These data surprisingly implicated a mutagenic effect of chronic stress via stress hormone pathways. Together, these findings implicated an involvement of inflammation and genetic damage in chronic stress-induced GI tumor risk (Figure 1).
Figure 1. Schematic diagram showing the molecular mechanisms by which chronic stress increases GI cancer risk. Chronic stress may facilitate \( \beta \)-AR signaling pathways via the ENS and HPA axis, leading to p53 degradation and DNA damage in GI somatic cells and the secretion of pro-inflammatory factors from immune cells. Through these mechanisms, chronic stress results in genetic mutations and systemic low-grade inflammation in GI tissues, causing increased GI cancer risks. GBA, gut–brain axis; ENS, enteric nervous system; HPA, hypothalamus–pituitary–adrenal; \( \beta \)-ARs, \( \beta \)-Adrenergic receptors.

3. GI Tumorigenesis Initiates Active Neurogenesis and Neural Infiltration

While nerve fibers were observed within rectum cancer and other solid tumors nearly a century ago, the role of nerves in tumor progression has long received insufficient attention [35,36]. To date, neurogenesis and nerve fiber outgrowth have been characterized within major GI tumors [37–40]. GI tumors, such as pancreatic cancer, are among those with the highest prevalence of neural infiltration [3]. However, the origin of tumor-associated nerve fibers has been mysterious for a long period of time. Given the facts that the growth of neurons and nerve fibers are mostly developmentally related and that adult neurogenesis only occurs in very restricted brain regions and rarely outside central nervous system, the molecular mechanisms driving neurogenesis and nerve fiber outgrowth in these tumors remain incompletely understood.

Data obtained from gastric and colorectal cancers revealed that neurons may be directly generated by the re-differentiation of cancer stem cells (CSCs) within tumors [41]. Using CSC-derived xenograft models, Ran Lu et al. showed that neural cells with human origin were identified in the ganglia close to and within xenografts. Importantly, blockade of neural differentiation attenuated gastric and colorectal tumor progression in the xenograft models. Another potential source of tumor-resident neurons is the circulating neural progenitors that dissociate from central nervous system [42]. Data suggested that neural progenitors may dissociate from the subventricular zone following the disruption of the blood–brain barrier, circulate in the circulatory system and penetrate into tumor tissues. These landmark discoveries revolutionized our understanding of the diversity
and complexity of neurogenesis within GI tumors, underscoring the importance of active neurogenesis in GI tumorigenesis. It should be noted that neural infiltration is different from perineural invasion (PNI), which is also frequently observed in multiple GI cancers, particularly pancreatic ductal adenocarcinoma (PDAC) and gastric cancer, as PNI essentially represents a chemotactic effect by which tumor cells migrate toward and eventually invade pre-existing nerves [43,44].

It is noteworthy that multiple data have indicated that GI tumors may create a pro-neurogenic microenvironment to facilitate neural infiltration. One of the most obvious mechanisms is the release of a variety of neurotrophins, including nerve growth factor (NGF), brain-derived neurotrophic factor (BDNF) and glial cell-derived neurotrophic factor (GDNF), to facilitate neurogenesis [45,46]. A recent study showed that tuft cells, a type of chemosensory cells resided in intestinal epithelium, as well as nerves, may release acetylcholine to stimulate the production of NGF from gastric epithelial cells, leading to the growth of enteric nerves and accelerated gastric carcinogenesis [47]. Neurogenesis, similarly to angiogenesis, has been proposed to play an indispensable role in GI tumor progression and dissemination [48]. These findings implicated that tumor tissues may fuel neuronal differentiation, as well as neurite outgrowth from pre-existing ENS/PNS, by creating a neurogenic microenvironment [49,50]. Recent studies also indicated a role of immune cells in neural infiltration [51]. It was revealed that macrophages recruited by pancreatic cancer cells may secrete GDNF to promote a cross-talk between nerve fibers and cancer cells [52]. Overall, these findings demonstrate neural infiltration as an integral part of GI tumor development (Figure 2).

Figure 2. The mechanisms underlying neurogenesis during GI cancer development. The release of neurotrophins from various cell types, including tumor-associated macrophages (TAMs), tuft and cancer cells, may lead to neurite outgrowth of ENS/PNS and the differentiation of cancer stem cells (CSCs) and circulating neural stem cells (NSCs) into neurons, to fuel GI tumorigenesis. NE, Norepinephrine; ACh, acetylcholine.

4. Neural Infiltration Fuels GI Cancer Progression

The ENS plays a vital role in tissue homeostasis and metabolism of the gastrointestinal system. Enteric nerve fibers directly control lipid and glucose metabolism, and consequently the development of diabetes and fatty liver diseases [53,54]. However, the roles of nerve
fibers in GI cancer development remain controversial and incompletely understood. Since neural infiltration has been identified across GI tumors, it is conceivable that the nervous system is involved in GI cancer progression. Indeed, nerve fiber infiltration has been linked to worsened prognosis in multiple GI tumors [55,56]. Accordingly, the density of tumoral nerve fibers may serve as indicative biomarkers for clinical outcomes of patients with GI tumors [55,57]. Daniel Albo et al. performed a quantitative analysis of nerve fibers in a colorectal cancer cohort of 236 patients and revealed that patients with high infiltration of PGP9.5-positive nerve fibers showed a 50% reduction in 5 year overall survival and disease-free survival, compared those with no detectable tumor-associated neurogenesis [37]. Furthermore, neural infiltration is also correlated with significantly higher GI tumor recurrence and worsened prognosis in patients with pancreatic cancer [58]. These data implicate a clinical relevance of the nervous system in GI tumors.

Though the nervous system has been linked to the development of all types of GI tumors, the degree of neural infiltration varies greatly among different GI tumors. For instance, nerve fibers have been observed in nearly all pancreatic tumors, 63% of colorectal cancer, 40% of gastric cancer and 38% of esophageal cancers [37,40,44,59]. However, S100-positive nerve fiber was absent within hepatocellular carcinoma (HCC) tumoral specimens, and can be only found in the capsule of a proportion of HCC tissues [60]. Accordingly, nerve fiber density is associated with worsened prognosis in the majority of GI tumors, but has no prognostic value in HCC [40,58,61,62]. These differences are very likely as a result of their histological patterns during the initiation of GI tumors. GI tract is a multi-layer tissue with neuronal bodies and nerve fibers as integral parts of the histological structure. Proximal connections between nerve fibers and various cell types imply an involvement of nerves in the initiation and progression of the tumors arising from GI tracts and pancreas [63,64]. On the other hand, the autonomic nervous system of the liver consists of branches of the splanchnic and vagal nerves. These nerves are mainly distributed around the portal vein, hepatic artery and bile duct, and have no direct connection with a majority of hepatocytes [65]. Nevertheless, neural signaling remains an important driver of hepatic carcinogenesis, despite the fact that nerve fibers have no prognostic merit. For instance, the expression of acetylcholinesterase, a key enzyme responsible for ACh breakdown, is inversely correlated with HCC growth and aggressiveness [66]. Another study showed that catecholamine neurotransmitter degrading enzyme monoamine oxidase A (MAOA) inhibits HCC metastasis via suppressing the adrenergic receptor signalling and the transactivation of EGFR pathway [67].

The mechanisms underpinning nerve-driven GI tumor progression are complex and involve various aspects of tumor aggressiveness. Some fascinating studies have shown that neural infiltration plays multifaceted roles in the malignancy of GI tumors [37,41]. However, because of the complexity of neural infiltration and the difficulty in establishing tumor innervation models in vitro, the precise mechanisms underlying the tumor-promoting roles of nerve fibers in GI cancers remain obscure. Nevertheless, since innervations typically work through the secretion of neurotransmitters and neuropeptides, the discovery of various neural receptors in tumor cells suggests that nerve fibers may directly fuel GI tumor malignancy via neural receptor pathways. Indeed, some delicate models of cancer-nerve cross talk have been established. By establishing an in vitro cross-talk model of murine sciatic nerves and cancer cells, several studies have shown that nerves may facilitate the proliferation, neural invasion and metastasis [68–70]. Notably, several signaling pathways, including GDNF-RET and SLIT2-ROBO, have been implicated in this cross-talk. Moreover, neural signals may also contribute to tumor resistance to chemotherapy. While clinical data supporting an association between nerve fiber density and tumor chemoresistance are lacking, mounting in vitro data have suggested critical involvement of assorted neurotransmitter pathways in the responsiveness to chemotherapeutic agents. Neurological pathways, such as β-adrenergic receptors (β-ARs) and acetylcholine receptors (AChRs), have been shown to promote chemoresistance of GI tumors via downstream oncogenic effectors [71–73]. In addition, neural signaling may promote the epithelial-mesenchymal
transition (EMT) of GI tumors [74–76]. Combined, these findings implied that nerve fibers may facilitate the activation of conventional oncogenic signaling in GI tumors, and contribute to various aspects of tumor malignancies.

5. The Roles of Neurotransmitter, Neurotrophin and Neuropeptide Receptor Pathways in Gastrointestinal Cancer Development

Histological examinations have revealed that nerve fibers have been well characterized within and around GI tumors. Since tumor-associated nerves can be considered as part of peripheral nervous system, tumor innervation, via nerve fibers surrounding and within the tumor, is supposed to work in a mechanism similar to non-tumorous tissues, namely through the secretion of neuroactive chemicals and peptides from nerve terminals. Mostly, these molecules function locally via paracrine routes and act on their receptors that are distributed on the surface of tumor and stromal cells. As stated above, the expression of neural receptors on tumor cells may plays a key role in nerve fiber-primed GI tumor progression. Thus, the distribution and function of tumor-residual neural receptors are of great importance to determine the major pathways involved [77]. Dysregulated expression of assorted neural receptors, including neurotransmitter receptors and neuropeptide receptors, has been reported in GI tumors, and linked to the enhanced malignancy of tumor cells. These studies suggested that a variety of neural receptors are expressed across GI tumors, with many attracting particular attention. Below, we briefly discuss the roles of neural receptor pathways in GI tumorigenesis. In addition, we summarize the current literature indicating the expression, downstream effectors and function of neurotransmitter receptors (Table 1) in GI tumors.

5.1. β-Adrenergic Receptors

β-ARs are critical downstream effectors of neurotransmitters released from sympathetic nerve terminals. As members of G protein-coupled receptors (GPCRs), β-ARs may transmit intracellular signaling via G protein-mediated mechanisms and through adaptor proteins β-arrestin [78]. Studies revealed that all three members of β-ARs (β1-AR, β2-AR and β3-AR) are widely expressed across diverse tumor types [79]. Because of their importance in stress response and the sympathetic nervous system, β-ARs are among the most intensively investigated neurotransmitter receptor pathways in GI tumors [80,81]. β-ARs, particularly β2-AR, have been implicated in various malignant behaviors of GI tumors, including proliferation, chemoresistance and metastasis, using both elegant in vivo and in vitro models. Using a KC (LSL-Kras+/G12D; Pdx1-Cre) pancreatic cancer model, Bernhard Renz et al. [82] revealed that chronic restraint stress (CRS) facilitated β2-AR-dependent PDAC growth, NGF secretion and intratumoral neural density. Blockade of β2-AR and NGF/Trk pathways decreased cancer incidence and extended the survival of KPC (LSL-Kras+/G12D;LSL-Trp53+/R172H;Pdx1-Cre) mice, directly linking chronic stress to PDAC development via a β-AR/neurotrophinloop [82]. The tumor-facilitating effects of β-ARs have also been validated by selective β-AR antagonists, such as atenolol (β1-AR) and ICI 118, 551 (β2-AR) [83,84]. Mechanistic studies revealed that β-ARs have broad impacts on assorted classical oncogenic pathways, such as HIF-1α, AKT and ERK pathways [85,86]. Coinciding with these findings, our study showed that β2-adrenergic receptor (β2-AR) may facilitate the PCBP2-mediated translation of c-Myc to promote the proliferation of pancreatic cancer cells [87]. These data suggested that β-AR pathways are exploited by GI cancers to facilitate downstream oncogenic effectors, highlighting β-AR signaling as integral players of the oncogenic network in GI tumors.

5.2. Acetylcholine Receptors

Acetylcholine receptor (AChR) family proteins are composed of two subtypes, nicotinic (nAChR) and muscarinic (mAChR), both of which have been reportedly involved in GI tumor progression. nAChRs, such as α7nAChR, are of particular concern, partially because of their link to smoking-induced GI tumorigenesis via transmitting nicotine-mediated
As ionotropic receptors, nAChRs may function as ligand-mediated ion channels, but also signal through intracellular signaling transducers [90]. Through these mechanisms, nAChRs may be coupled to various downstream signaling, such as JAK2/STAT3 and NF-kB signaling, leading to aggressive behaviors [91–93]. Different from nAChRs, mAChRs belong to metabotropic GPCRs that primarily transmit signals via G proteins and adenyl cyclase. mAChRs consist of five members, termed m1–5 AChR, all of which are, to some extent, expressed in GI tract [94]. In this regard, m3AChR is a key subtype functionally associated with GI tumorigenesis, probably because of its preferential expression in stem cells [95]. Genetic ablation of m3AChR attenuates colon tumorigenesis, indicating a tumor-promoting function of this receptor [96]. In line with this finding, treatment with m3AChR agonist carbachol activates protein kinase cascades and the proliferation of gastric and colorectal tumor cells [97,98]. Interestingly, studies revealed that mAChR-mediated ERK signaling and cell proliferation may involve the transactivation of the EGFR pathway [99,100]. It should be noted that, while the AChR pathway was frequently associated with enhanced tumor aggressiveness, some studies also revealed tumor-suppressive functions of cholinergic signaling in GI tumors [101–104]. The inconsistency of these studies suggested that cholinergic signaling may function in a tumor context-dependent manner.

5.3. Glutamate Receptors (GluRs)

Serving as a principal neurotransmitter as well as a key metabolite, glutamate plays a unique role in GI tumor progression. Enrichment of glutamate, caused by aberrant metabolism of GI tumors, may trigger the activation of GluRs residing on the tumor cells [105,106]. A variety of GluRs, both ionotropic (iGluR) and metabotropic (mGluR), have been reportedly expressed in GI tumors [107]. Glutamate may promote the invasion of pancreatic cancer via ionotropic AMPA receptor-mediated Kras–MAPK signaling [108]. Multiple members of iGluR and mGluR have been implicated in the progression of colorectal cancer [109–112]. Likewise, NMDA receptor NR1 and NR2A subunits have been reportedly involved in the malignancy of gastric cancer [113,114]. Of great intrigue was Leanne Li et al.’s discovery that NMDAR is highly expressed in invasive fronts of pancreatic neuroendocrine tumorigenesis (PNET), and may drive tumor invasiveness through fluid flow-induced autocrine glutamate signaling circuit [115]. Collectively, these studies demonstrate the crucial roles of GluRs in GI neoplasia and underscore the importance of targeting glutamate metabolism and its receptor pathways as therapeutic strategies for GI tumors.

5.4. Dopamine (DA) and 5-HT (Serotonin) Receptors

Receptors 5-HT and DA are both abundantly produced in GI tract and play vital roles in GI homeostasis [116,117]. Accordingly, it is conceivable that these neurotransmitter pathways play critical roles in GI disease progression, including tumorigenesis. DA receptors (DARs) are composed of five class A GPCRs, and can be divided into D1-type (D1 and D5) and D2-type (D2, D3 and D4) subgroups [118]. Both of D1 and D2 types, to some degree, possess oncogenic properties in GI tumors. For instance, upregulation of DA and DA receptor D1 (DRD1) has been associated with tumor growth and invasion of hepatocellular carcinoma (HCC) [119]. Moreover, upregulated expression of DA receptor D2 (DRD2) is associated with elevated malignant potential of gastric cancer and PDAC, implicating a therapeutic merit of antagonizing DRD2 in these tumors [120,121]. Similar to DARs, serotonin receptor (5-HTR) pathways may also facilitate the progression of various GI tumors. Excessive production of serotonin, as a result of upregulated expression of serotonin biosynthesis rate-limiting enzyme tryptophan hydroxylase 1 (TPH1), contributes to NLRP3 inflammasome activation via serotonin receptor HTR3A and the acceleration of colorectal cancer development [122]. The accumulation of serotonin was observed in a Kras/p53-driven pancreatic cancer model, and linked to Warburg Effect and accelerated growth of pancreatic tumors [123]. An in vitro study also revealed that silencing the expression of 5-HT receptors 5-HT1B and 5-HT1D retarded the proliferation, clonogenicity
and invasion of pancreatic cancer cells, pointing to a direct involvement of these receptor pathways in tumor cell malignancies [124].

Table 1. The expression and roles of neurotransmitter receptors in gastrointestinal cancer.

| Receptors                        | Expression in Cancer/Mechanism | Downstream Effectors/Mechanisms | Effects                                      | Reference |
|----------------------------------|--------------------------------|---------------------------------|----------------------------------------------|-----------|
| **β-adrenergic receptors** (β-ARs) |                                |                                 |                                              |           |
| β1-AR                             | Upregulated in EC              | ERK, COX2                        | proliferation                               | [79,125]  |
|                                  | Upregulated in metastatic GC   |                                 |                                              | [126]     |
|                                  | Expressed in PC                | AKT, ERK, HIF-1α                 |                                              | [127]     |
|                                  | Upregulated in CRC             |                                 |                                              | [79]      |
| β2-AR                             | Upregulated in GC              | STAT3, AP-1, MUC4                | Proliferation, Chemoresistance, metastasis  | [126,128,129] |
|                                  |                                  |                                 |                                              |           |
| β3-AR                             |                                  |                                 |                                              |           |
| Acetylcholine receptors (AChRs)   |                                |                                 |                                              |           |
| α3nAChR                          | Expressed in ESCC              | YAP1                            | Proliferation, migration                      | [135]     |
| α5nAChR                          | Expressed in GC                | AKT                             | Chemoresistance                              | [136]     |
|                                  | Upregulated in EC              | AKT/FOXO1/OTUD3/VEGF            | lymphatic metastasis                         | [137]     |
|                                  | Expressed in GC                | E-cadherin, ZEB-1, fibronectin, AKT, MCL-1, BCL-2 | Migration, chemoresistance | [73,138,139] |
|                                  | Upregulated in PC              | MUC4, 25864419                  | Stemness, metastasis                         | [91,140]  |
|                                  | Expressed in CRC               | NF-kB, Fibronectin, Snail, ZEB1 | Migration                                    | [141–144] |
|                                  | Upregulated in CC              | EMT, ERK                        | Proliferation, viability, migration          | [74,145]  |
|                                  | Upregulated in HCC             | TRAF6/NF-κB                     | Proliferation, chemoresistance               | [72,95]   |
| m1AChR                           | Expressed in HCC               | EMT, PI3K/AKT                   | Invasion                                     | [146]     |
| m3AChR                           | Expressed in GC                | EGFR, AKT, ERK                  | Proliferation, viability                     | [100,147,148] |
|                                  | Upregulated in PC              |                                 |                                              | [149]     |
|                                  | Upregulated in CRC             | Calcium, MMP7, EGFR, p38, ERK, AKT | Proliferation, viability                     | [150–153] |
|                                  | Upregulated in CC              |                                 | Proliferation, metastasis                    | [154]     |
| Glutamate receptors (GluRs)      |                                |                                 |                                              |           |
| AMPA receptor (GluR1–4)          | Downregulated in PC            | Kras-MAPK                       | invasion                                     | [108]     |
|                                  | Downregulated in CRC (GluR4)   |                                  |                                              | [155]     |
|                                  |                                  |                                  |                                              |           |
| NMDA receptors (NR1–3)           | Upregulated in PC              | AKT, ERK, CaMK II, HIF-1α       | Proliferation, migration                     | [156,157] |
|                                  | Expressed in GC                |                                 | proliferation                               | [113]     |
|                                  | Upregulated in CRC (NR2D)      | HIF-1α, AKT, ERK, CaMK II       | migration, angiogenesis                      | [110,158] |
|                                  | Downregulated in GC (GRIK2)    |                                  | Impaired Growth, migration                    | [160]     |
| Kainate receptor                 |                                  |                                  |                                              | [161]     |
|                                  | Upregulated in GC (GRIK3)      |                                  |                                              |           |
| metabotropic glutamate receptors (mGluRs) | Upregulated in PC (mGluR1)    | PEK/AKT/mTOR                    | Viability                                    | [162,163] |
|                                  | Upregulated in CRC (mGluR4)    |                                  | 5-FU resistance, recurrence                  | [109,164] |
|                                  | Expressed in HCC               | Calcium, MAPK                   | Chemoresistance                              | [165]     |
| Dopamine receptors (DRs)         |                                |                                 |                                              |           |
| DRD1                             | Expressed in ESCC              |                                 |                                              | [166]     |
|                                  | Expressed in PC                |                                 | Stemness, growth, migration                  | [167]     |
|                                  | Upregulated in HCC             | cAMP/PI3K/AKT/CREB              | Proliferation, metastasis                    | [119]     |
Table 1. Cont.

| Receptors | Expression in Cancer/Mechanism | Downstream Effectors/Mechanisms | Effects | Reference |
|-----------|--------------------------------|---------------------------------|---------|-----------|
| DRD2      | Upregulated in ESCC            |                              | lymph node metastasis | [166] |
|           | Upregulated in GC              |                              | Proliferation        | [120] |
|           | Upregulated in PC              | Calcium, PKA                  | Proliferation, viability, migration | [121] |
| DRD5      | Upregulated in EC              | mTOR, AKT, Warburg effect     | proliferation       | [168] |
|           | Expressed in GC                | mTOR                           | Impaired growth, autophagy | [169] |
|           | Expressed in CRC               |                                 |                     | [169] |
|           |                                 | CD133, OCT4, and EpCam         | Impaired growth, stemness, migration | [170] |
| 5-HTR     | Expressed in CRC (5-HT1B,      | Calcium/CaMKIIα, NLRP3         | Growth, angiogenesis, viability | [122,171–174] |
|           | 5-HT3A, 5-HT3, and 5-HT4)      | inflammasome, MMP-12           |         |           |
|           | Expressed in PC (5-HT1B,       | PI3K/AKT/mTOR, Warburg         | Growth, invasion    | [123,124] |
|           | 5-HT1D and 5-HT2B)             | effect, uPAR/MMP-2, Integrin/Src/Fak |         |           |
|           | Expressed in CC (5-HT1A,       |                                 | Growth            | [175] |
|           | 5-HT2A, 5-HT2B, 5-HT4 and 5-HT6) |                               |         |           |
|           | Expressed in HCC (5-HT1B and   | AKT, FOXO3a                   | Proliferation     | [176] |
|           | 5-HT2B)                        |                                 |         |           |

EC, oesophageal cancer; ESCC, oesophageal squamous cell carcinoma; GC, gastric cancer; PC, pancreatic cancer; CRC, colorectal cancer; CC, cholangiocarcinoma; HCC, hepatocellular carcinoma.

5.5. Neurotrophin Receptors

While neurotrophin signals appear to promote neurogenesis during GI tumor initiation, neurotrophin receptors, termed tropomyosin-related kinases (TRKs), are also aberrantly upregulated in GI cancer cells [177]. The deregulation of neurotrophin receptors is frequently attributed to epigenetic mechanisms, such as DNA methylation [178–180]. Overexpression of NGF may promote the survival and motility of liver cancer cells via tropomyosin receptor kinase A (TrkA) pathway [181]. A similar effect of NGF-TrkA pathway in other GI tumors, such as colorectal and pancreatic cancers, has also been observed [182,183]. Likewise, BDNF-TrkB signaling fuels the growth and invasion of gastric and pancreatic carcinoma, and may serve as a therapeutic target against these malignancies [184,185]. Interestingly, whereas an earlier study reported a conditional tumor suppressive function of TrkC in colorectal cancer, cell culture and xenograft data indicated that TrkC may also elicit a tumor-promoting effect [180,186]. Similar to their functions in neurogenesis, TRK pathways may activate mitogenic pathways, such as AKT and Ras/MAPK, to drive the malignancy of GI tumors [187]. The expression and effects of neurotrophin receptors in GI tumors were summarized in Table 2.

5.6. Neuropeptide Receptors

Neuropeptides represent another important source of signal molecules bridging nerves and cancer cells. Given the fact that a variety of neuropeptides are abundantly produced in gastrointestinal tract and play key roles in the homeostasis of GI system, it is unsurprising that these neuropeptides are involved in GI tumorigenesis [216,217]. Many neuropeptides are secreted along with neurotransmitters from enteric nerve fibers, and serve as an additional layer of nerve–GI communication [218]. Notably, unlike neurotransmitters that typically play tumor-promoting roles, the roles of neuropeptides in GI tumors are complicated and controversial. An upregulated level of galanin has been linked to enhanced metastasis and chemoresistance of colorectal cancer [219,220]. Intriguingly, studies revealed that galanin may exert a tumor-suppressive effect in gastric and pancreatic carcinogenesis [221,222]. Likewise, methionine enkephalin (MENK) may function as tumor suppressor in gastric cancer development, but possesses oncogenic properties in some other
GI tumors [223,224]. While neuropeptide Y (NPY) plays a critical role in the function of GI system, its role in GI tumorigenesis remains incompletely understood. Early studies suggested that the level of NPY was reduced in the serum of gastric and colorectal cancers [225]. However, recent experiments using genetically modified animals revealed that NPY may promote the proliferation of epithelial cell to facilitate DSS-induced intestinal carcinogenesis [226]. Overall, neuropeptides play complex and diverse roles in GI tumorigenesis. Much of this field remains incompletely understood and needs further investigations.

### Table 2. The expression and roles of neurotrophin receptors in gastrointestinal cancer.

| Receptors | Expression in Cancer/Mechanism | Downstream Effectors | Effects | Reference |
|-----------|-------------------------------|----------------------|---------|-----------|
| p75NTR    | Expressed in ESCC             | Bmi-1                | Self-renewal, proliferation, chemoresistance | [188–190] |
|           | Downregulated in GC/DNA methylation | uPA, MMP-9, NF-κB | Impaired proliferation, invasion and metastasis | [126,191,192] |
|           | Expressed in PC               |                      | Neural invasion, proliferation | [193,194] |
|           | Downregulated in CRC/DNA methylation |                   | Impaired proliferation, invasion and viability | [195] |
|           | Downregulated in HCC          |                      | Impaired proliferation | [196,197] |
| TrkA      | Upregulated in ESCC           |                      | Chemoresistance | [198] |
|           | Expressed in GC               |                      | Chemoresistance | [199] |
|           | Expressed in PC               | PI3K/AKT             | Chemoresistance | [200–202] |
|           | Expressed in CRC              | MAPK/ERK, MMP2, MMP9 | Metastasis | [182] |
|           | Expressed in CC               |                      | | [203] |
|           | Upregulated in HCC/DNA demethylation |              | Proliferation | [178] |
| TrkB      | Expressed in ESCC             |                      | Chemoresistance | [204] |
|           | Expressed in GC               | Nrf2                 | Lymph node metastasis, chemoresistance | [205–207] |
|           | Expressed in PC               |                      | Invasion | [185,208] |
|           | Upregulated in CRC            | ERK                  | Proliferation, invasion, viability | [209–212] |
|           | Upregulated in HCC/DNA demethylation | RhoA, VEGF          | Angiogenesis, proliferation, chemoresistance | [72,178,213] |
| TrkC      | Expressed in GC               |                      | | [199] |
|           | Upregulated in PC             |                      | | [149] |
|           | Downregulated in CRC/DNA methylation |              | Impaired viability | [180,214,215] |
|           | Upregulated in HCC/DNA demethylation |              | Proliferation | [178] |

ESCC, oesophageal squamous cell carcinoma; GC, gastric cancer; PC, pancreatic cancer; CRC, colorectal cancer; CC, cholangiocarcinoma; HCC, hepatocellular carcinoma.

### 6. Neural Infiltration Is an Integral Part of GI Tumor Microenvironment (TME)

Because of the prevalence of nerve fiber infiltration in GI tumors, emerging evidence has indicated that they are an indispensable part of GI tumor microenvironment. Studies revealed that the nervous system is linked to aberrant function of tumor immune cells, fibroblasts and endothelial cells [77,227]. Neurotransmitters and other signal molecules secreted by nerve fibers have profound impacts on tumor environments and various types
of stromal cells. Peripheral and tumor-associated nerves may foster a range of tumor-favoring microenvironments, exacerbating malignant characteristics of GI tumors.

6.1. Angiogenesis

Many neurotransmitter receptors such as β-ARs and ACh receptors (AChRs) have been reportedly expressed in vascular endothelial cells [228,229]. Chronic stress may facilitate VEGF expression and tumor angiogenesis through neurotransmitter signaling pathways [230]. Catecholamines, such as norepinephrine (NE) and epinephrine (E), have been well-recognized to promote tumor angiogenesis, through the secretion of VEGF [231,232]. On the other hand, DA pathway may inhibit angiogenesis in gastric cancer via DA receptor D2 (DRD2) on endothelial cells [233]. This effect of DRD2 may be attributed to enhanced endocytosis of VEGF receptor 2 (VEGFR2) [234].

6.2. Immune Regulation

Intensive studies in the past decades have demonstrated a fundamental role of the nervous system in immune regulation [235]. While the nervous system may fine-tune systemic inflammation and immune responses via direct innervations of lymphoid organs, nerve fibers within GI tumors play a vital role in the tumor immune microenvironment (TIME) via a paracrine mechanism. Neurotransmitters and neuropeptides secreted from local nerve terminals have strong influences on TIME through direct intervention on assorted immune cells. It has been well-documented that the receptors of various neurotransmitters, including catecholamines, GLU, 5-HT and ACh, are broadly expressed in diverse immune cells, including those infiltrating tumor tissues. For instance, functional β-ARs and α7nAChR have been reportedly expressed in tumor-associated macrophages (TAMs) in GI tumors [236,237]. Likewise, neural receptor pathways, including glutamate receptors and β-ARs, play indispensable roles in regulating tumor-eradicating activity of CD8+ cytotoxic T lymphocytes (CTLs) [238,239]. Aside from TAMs and CTLs, chronic stress has also been reported to regulate other immune cells, such as Treg and CD4 cells, via β-adrenergic receptor pathway in a pancreatic cancer model [240]. In agreement with these findings, studies based on an orthotopic PDAC model revealed that nerve fibers reprogram TIME and suppress intratumoral T-cell response to favor tumor progression [241]. Overall, in line with a tumor-promoting speculation of neural infiltration, most of these studies suggested that neural pathways may facilitate TIME in favor of tumor progression.

6.3. Chemotactic Effects

Studies in the past decades have indicated that neurotransmitters may serve as chemoattractants to guide cancer cell invasion towards nerve fibers. Perineural invasion has been observed in multiple GI tumors, which essentially represents a chemotactic effect of nerve fibers to tumor cells [242–244]. Using both in vivo and in vitro models, studies revealed a critical involvement of assorted neural pathways, such as substance P (SP)/NK-1R, NGF and β-adrenergic signals in pancreatic cancer PNI [193,245,246]. It is revealed that nerve fiber and neurotransmitters may increase the mobility of GI cancer cells in terms of inducing cytoskeleton remodeling and epithelial–mesenchymal transition [146,247]. In this regard, neural receptors may activate some key signals involved in cytoskeleton remodeling and cell migration, such as Kras–MAPK, MMPs and STAT3, and potentially drive directed migration of tumor cells along with neurotransmitter gradients [108,245].

7. Intervention of Neural Infiltration as Measurements of Cancer Treatment

Because of the critical role of neural infiltration in fueling GI tumor malignancy, it is unsurprising that intervention of nerve fibers may serve as a strategy to retard GI tumor progression. At present, multiple strategies have been proposed to block nerve–cancer cross-talks, among which denervation and pharmacological blockade of neural receptor pathways are some most well-characterized approaches.
Studies in the past decades have indicated promising effects of denervation in reducing cancer incidence and progression. Chun-Mei Zhao et al. reported that both surgical and pharmacological denervation reduced gastric tumor incidence, stemness and chemoresistance [248]. Similarly, denervation of the pancreas reduces cancer initiation and invasion in a Kras-driven PDAC model [249]. Data obtained from colorectal cancer also support the notion that denervation may be beneficial to the prevention of tumor incidence and progression [250,251]. However, it remains unclear whether denervation may prevent tumor-associated neurogenesis, or simply block the influence of the pre-existing peripheral nervous system on tumorigenesis.

Despite the fact that denervation exhibits considerable merit in preventing tumor progression, the potential side-effects may hinder its translation into clinical practice. On the other hand, pharmacological inhibition of neural receptors appears to be a reliable intervention that may be applied into clinical use [252,253]. Coinciding with the presumed tumor-facilitating role of β-AR signaling, epidemiological studies revealed that administration of β-blockers, such as propranolol, may reduce GI cancer risk [254–256]. Data obtained from in vitro and in vivo experiments also indicated that pharmacological inhibition of β-ARs retard GI cancer growth, invasion and chemoresistance [87,257]. These findings suggested a promising future of β-blockers in GI tumor prevention and therapy.

Other than β-blockers, nAChR inhibitors, such as α-bungarotoxin and mecamylamine, are candidate therapeutic agents for GI cancers [91,258]. While pre-clinical data have indicated apparent effect of these chemicals in blocking GI cancer malignancy, the in vivo efficacy and potential side-effects need to be addressed before they are considered for clinical use. In vitro studies also discovered anti-tumor effects of several other neurotransmitter pathway inhibitors in GI tumors. For instance, 5-HT receptor inhibitor vortioxetine may inhibit the viability, proliferation and invasion of gastric cancer cells [259]. Likewise, treatment with 5-HT2A inhibitor ketanserin and 5-HT3 ondansetron exacerbated ionizing radiation-induced cell death of colorectal cancer cells [260]. Recent studies also showed that administration with DAR antagonist pimozide reduced the growth and lymph node metastasis of colorectal cancer xenografts [261]. It should be noted that a significant proportion of these pharmacological agents have been under clinical use for a long period and can be quickly repurposed to cancer therapy. Cumulatively, these studies suggest that pharmacological blockade of neurotransmitter signaling pathways may serve as a valuable therapeutic strategy against GI cancer.

8. Conclusions
Mounting evidence has suggested that neural infiltration and neural signaling pathways play crucial roles in the development of GI tumors. Tumor-associated nerve fibers may fuel various aspects of tumor progression, including growth, metastasis, chemoresistance, angiogenesis and immune suppression. Neural receptor pathways, particularly those critically involved in the sympathetic and parasympathetic nervous systems such as β-AR and AChR pathways, are key players linking innervation and tumor progression. Pharmacological inhibitors of neurotransmitter pathways, such as β-blockers, have shown promising potential in the prevention and treatment of GI tumors. Herein, we reviewed recent progress in dissecting the role of tumor innervation in GI cancer progression. Given the fact that most of GI tissues are heavily innervated by enteric nerve fibers, it is conceivable that neural mechanisms play particularly important roles in GI tumor development. With the fast growth of new studies, the fascinating story of nerve and GI cancer partnership will continue.
Author Contributions: C.W. and X.Z. conceived and wrote the manuscript. B.H. and X.Y. drew the figures and tables. All authors revised the manuscript. All authors have read and agreed to the published version of the manuscript.

Funding: This study was supported by the National Natural Science Foundation of China (no. 81972279 and 31171038), and Natural Science Foundation of the Jiangsu Higher Education (21KJB320016).

Conflicts of Interest: The authors declare no conflict of interest.

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