Glycogen Synthase Kinase-3beta regulates Snail and beta-catenin during gastrin-induced migration of gastric cancer cells

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Abstract

Background: The gastrointestinal peptide hormone gastrin is known to regulate various cellular processes including proliferation, migration and metastasis in gastrointestinal (GI) cells. The studies described here were undertaken to elucidate in detail the signaling pathways mediating the migratory responses of amidated gastrin (G17) and to understand the involvement of the serine/threonine kinase Glycogen Synthase Kinase-3 beta (GSK3β) in this.

Results: Our results indicate that incubation of gastric cancer cells overexpressing CCK2 receptor (AGSE cells) with G17 results in a dose and time dependent increase of GSK3βSer9 phosphorylation, indicative of an inhibition of the kinase. Pretreatment with a pharmacological inhibitor of PI3Kinase pathway (Wortmannin) was unable to antagonize G17-induced GSK3βSer9 phosphorylation, suggesting that this might involve PI3Kinase-independent pathways. Treatment with G17 was also associated with increased Snail expression, and β-catenin nuclear translocation, both of which are GSK3β downstream targets. Pretreatment with a pharmacological inhibitor of GSK3β (AR-A014418) augmented Snail expression and β-catenin nuclear translocation in the absence of G17, whereas overexpression of a phosphorylation deficient mutant of GSK3β (S9A) abrogated Snail promoter induction. These suggested that G17 modulates Snail and β-catenin pathways via inhibiting GSK3β. In addition, overexpression of GSK3β wild type (WT) or S9A mutant inhibited G17-induced migration and MMP7 promoter induction. G17 studies designed following small interference RNA (siRNA)-mediated knockdown of Snail and β-catenin expression indicated a significant reduction of G-17-induced migration and MMP7 promoter induction following combined knockdown of both proteins.

Conclusion: Our studies indicate that inhibition of GSK3β is necessary to activate G17-induced migratory pathways in gastric cancer cells. Inhibition of GSK3β leads to an induction of Snail expression and β-catenin nuclear translocation, both of which participate to promote G17-induced migration.

Background

Gastric cancer is the second leading cause of cancer-related deaths worldwide [1], and are often characterized as highly aggressive and unresponsive to therapy [2]. The major risk factor contributing to this disease include Helicobacter pylori (H. pylori) infection, diet as well as genetic background [3,4]. Interestingly, studies during the past two decades have also demonstrated that the gastrointestinal (GI) peptide hormone gastrin might contribute to the pathobiology of gastric cancers. In addition to regulating gastric acid secretion, mature gastrin (G-17) and its unprocessed intermediate forms progastrin and glycine extended gastrin (Gly-G) can regulate growth in a variety of cancer cells [5,6]. Results from transgenic mice show that mice overexpressing the amidated form of gastrin have increased proliferation of gastric mucosa [7], which can synergize with Helicobacter infection leading to the development of invasive gastric cancer [8]. Prolonged hypergastrinemia increases the relative risk of developing colon cancer [9] and might promote adenoma to carcinoma progression [10]. Recent studies have confirmed gastrin to be an essential cofactor for carcinogenesis of gastric corpus [11]. In addition, significantly high levels of plasma gastrin has been reported in patients with gastric cancer, with high expression of gastrin and...
its receptor (CCK2R) in gastric cancer cell lines [12]. All these studies indicate an important role of gastrin-and its receptor system in mediating gastric cancer. G17 as well as invasion [31-33] which are prerequisites for in vivo metastasis. Our earlier studies in gastric cancer cells have demonstrated that G17-induced migration involves an activation of the Mixed-Lineage-Kinase 3/ JNK1 signaling axis [31]. Due to a close connection of GSK3β in regulating cell migration, we designed these studies to understand its role in G17-induced migration. We show that incubation with G17 increases GSK3βSer9 phosphorylation in a transient manner, which was also associated with a corresponding increase in the expression and promoter activation of Snail and an increase in the nuclear translocation of β-catenin. Inhibition of GSK3β via a pharmacological inhibitor resulted in increased Snail expression and β-catenin nuclear translocation in the absence of G17 and overexpression of a phosphorylation deficient mutant of GSK3β (S9A) antagonized G17-mediated induction of Snail promoter. Similarly, ectopic overexpression of Wild type (WT) or S9A mutant of GSK3β antagonized G17-induced migration and MMP7 promoter induction. Our studies also indicate that, combined knockdown of Snail and β-catenin by small interference RNA (siRNA) significantly attenuated G17-induced migration and MMP7 transcription. These studies indicate that G17 modulates Snail and β-catenin pathways via inhibiting GSK3β, both of which in turn participate to mediate G17-induced migration.

Results

Effect of Gastrin (G17) on GSK3βSer9 phosphorylation

In order to determine the role of G17 on GSK3β pathway, Western Blot analysis was performed with G17-treated gastric cancer cell lines overexpressing the CCK2 receptor (AGSE) [34]. These indicated a time (Fig 1A, pGSK3βSer9 panel) and dose-dependent (Fig 1B) increase in GSK3β Ser9 phosphorylation, which was maximal after 1 hour of G17 treatment. Pretreatment with an antagonist of the CCK2R (YM 022), inhibited G17-induced GSK3βSer9 phosphorylation (Fig 1C), indicating that this is mediated via CCK2R pathway. Since Ser 9 phosphorylation is an inhibitory phosphorylation site of GSK3β, these studies indicated a G17-induced inhibition of GSK3β pathway, possibly via activation of an upstream kinase (for example AKT). In fact, Western analysis also showed an increase in AKTser473 phosphorylation (Figs 1A, B pAKTser473 panel) corresponding to the time of GSK3βSer9 phosphorylation, indicating a simultaneous activation of AKT. However, pretreatment of the cells with PI3Kinase inhibitor (Wortmannin) was unable to antagonize G17-induced GSK3βSer9 phosphorylation (Fig 1D, compare lanes 2 and 4, pGSK3βSer9 panel), although it completely antagonized AKT phosphorylation (pAKTser473 panel). In addition, treatment of another gastric cancer cell line (MKN45) with G17 showed an increase in GSK3βSer9 phosphorylation without any increase in AKT phosphorylation (Fig 1E). These studies indicated that G17-induced increase in GSK3βSer9 phosphorylation might involve a PI3 Kinase independent pathway.
Effect of G17 on Snail expression

To understand the consequences of G17-mediated inhibition of GSK3β, G17 studies were performed to determine changes in the expression of GSK3β downstream target Snail. Incubation of AGSE cells with G17 resulted in an increase in Snail protein expression in a time (Fig 2A) and dose-dependent (Fig 2B) manner, which was also associated with an increase in Snail transcription (Fig 2D). In addition, G17 induction of Snail expression was mediated via CCK2R, since pretreatment with YM 022 abolished G17-induced Snail expression (Fig 2C).

G17 induces Snail expression and β-catenin nuclear translocation via inhibiting GSK3β

In order to determine whether G17 increased Snail expression via inhibiting GSK3β, G17 studies were performed following pretreatment of the cells with a pharmacological inhibitor of GSK3β (AR-A014418) [35]. These studies showed an induction of Snail expression following pretreatment with two different concentrations of AR-A014418 (AR) in the absence of G17 (Fig 3A, compare lanes 1, 3 and 5). Pretreatment with 5 μM of AR pro-
have demonstrated that G17 treatment increases β-catenin transcription in the absence of G17 (lanes 1 & 3), which was equal to the G17-treated levels (lanes 3 & 4). The same extracts were also blotted with GAPDH (cytoplasmic protein) and Lamin A/C (nuclear protein) to show the purity of the nuclear preparation. To understand any crosstalk between MLK3/JNK1 axis [31] and GSK3β axis, Snail and β-catenin studies were performed following pretreatment with the pharmacological inhibitor of JNK (SP600125). These studies indicated a complete inhibition of JNK downstream c-Jun phosphorylation with SP600125 (Fig 3D, lower panel, compare lanes 2 and 6, pc-Jun panel). SP600125 however, was unable to inhibit G17-mediated induction of Snail expression (compare lanes 2 and 6, Snail panel) or β-catenin nuclear translocation (compare lanes 2 and 6, β-catenin panel). These suggested that G17-mediated activation of MLK3/JNK1 and inhibition of GSK3β might be parallel pathways operating independent of each other.

G17-induced migration involves GSK3β inhibition

To understand whether G17-induced inhibition of GSK3β was critical to induce migration, wound-healing assays were carried out following overexpression of either wild-type or mutant forms of GSK3β. As shown in Fig 4A, G17-induced migration results in wound closure in the cells overexpressing an empty vector or GSK3β-K/A mutant (8 hr, Empty vector and GSK3β-K/A panels). Ectopic overexpression of GSK3β-WT or GSK3β-S9A on the contrary, significantly inhibited G17-induced migration (compare GSK3β-WT, S9A and Empty vector panels). The average gap of migration in these cells were also measured and plotted as graphs, which indicated a complete wound closure at 8 hrs of G17 treatment with Empty vector (Fig 4B, lane 2) and GSK3β-K/A (lane 6), and an inhibition of migration with GSK3β-WT (lane 4) and GSK3β-S9A (lane 8). Western Blot analysis of these cell extracts is shown in Fig 4C, which indicates the expression of the various ectopic GSK3β forms. In these samples, overexpression of GSK3β-WT and S9A resulted in a decrease in the expression of endogenous β-catenin (β-catenin panel, lanes 3-6), suggesting that these ectopic proteins retain GSK3β activity.

G17-induced migration involves Snail and β-catenin pathways

Since GSK3β inhibition in these cells was enough to induce Snail expression and β-catenin nuclear translocation (Figs 3A, D), and inhibition of GSK3β was necessary for G17-induced migration (Fig 4A), it was conceivable that Snail and β-catenin are involved in G17-induced migration. To address this possibility, wound-healing
assays were performed following knockdown of endogenous Snail and β-catenin protein expression (alone or in combination) utilizing corresponding siRNAs. Transfections of Snail or β-catenin siRNA lead to a significant decrease in the expression of the respective proteins as shown in Fig 5C. Wound-healing assays performed under these conditions indicated a complete closure of the wound following G17 stimulation in the presence of control siRNA (Fig 5A, control siRNA panel), which was inhibited significantly in the presence of Snail or β-catenin siRNA alone (lanes 4, 6) and significant inhibition with the combination (lane 8).

G17-mediated induction of MMP7-promoter activation involves GSK3β, Snail and β-catenin

To elucidate further the mechanism by which G17/GSK3β axis mediated migration, studies were also designed with the MMP7 promoter which was shown to be induced by G17 in our earlier studies [31]. Treatment with G17 resulted in an increase in MMP7 promoter activity when transfected with an empty vector (Fig 6A, compare lanes 1, 2), which was inhibited significantly in the presence of GSK3β-S9A (compare lanes 2, 4). Overexpression of GSK3β-K/A on the other hand, increased MMP7-promoter activity in the absence of G17 (lane 5), and produced synergistic effects when combined with G17 (lane 6). These suggested the involvement of GSK3β in mediating G17-induced MMP7 promoter induction. Since Snail and β-catenin are the two downstream targets of GSK3β mediating G17-induced migration (Fig 5A), it is likely that they are also involved in inducing MMP7 pro-
moter activity. Luciferase assays indicated a partial inhibition of G17-induced MMP7 promoter activity following knockdown of β-catenin expression alone (Fig 6B, compare lanes 2 and 4), which was significantly inhibited following knockdown of Snail alone (lane 6) or with combined knockdown of Snail and β-catenin (lane 8).

Discussion

The GI peptide hormone gastrin (G17 and its unprocessed forms) can regulate various cellular processes involved in cancer [37,38]. The studies described here were designed to elucidate in depth the mechanism by which G17 induces migration in gastric cancer cells, which in our earlier studies have shown to involve an acti-

vation of the MLK3 and JNK1 signaling axis [31]. In this axis G17-induced activation of JNK1 leads to phosphorylation and activation of c-Jun, leading to induction of MMP7 transcription and increased migration. The current studies were performed to determine whether the serine/threonine kinase GSK3β plays any role in this, due to its close connection in regulating cellular migration. A crosstalk of MLK3 pathway with GSK3β has also been reported earlier [39]. GSK3β has been shown to regulate migration both in a positive and negative manner. For example inactivation of GSK3β can increase migration in fibroblasts [16], and induce EMT in nontumorigenic breast epithelial cells [18]. In other studies, GSK3β was shown to promote cancer cell migration by cooperating with h-prune [40], or with small GTPase Rac [41]. To obtain a mechanistic insight towards the role of GSK3β in G17-induced migration, overexpression studies were performed with either wild-type or mutant forms of the kinase. As shown in Fig 4A, ectopic overexpression of GSK3β-WT as well as S9A mutant significantly attenuated G17-induced migration. MMP7 is known to mediate migration of gastric cancer cells [42], the transcription of which was induced by G17 [31]. Studies described here also revealed an inhibition of G17-induced MMP7 promoter activity following overexpression of GSK3β-S9A (Fig 6A), which was increased following overexpression of GSK3β-K/A in the absence of G17.

In many cells, GSK3β is constitutively active, which can be inactivated by various signaling mechanisms including Wnt signaling pathway [26,43] and PI3K/AKT pathway [44]. Although the detailed mechanism how Wnt pathway inactivates GSK3β is still unclear, PI3K/AKT inhibits GSK3β via increasing its Ser 9 phosphorylation [45]. In our studies, treatment with G17 also produced an increase in GSK3βSer9 phosphorylation (Figs 1A, 1B), suggesting an inactivation of the kinase during G17/CCK2R activation. This was associated with a corresponding increase in AKTSer473 phosphorylation, indicating the possibility that G17 might induce GSK3βSer9 phosphorylation and downstream cellular responses via PI3K/AKT activation. However, pretreatment with Wortmannin (pharmacological inhibitor of PI3Kinase), was unable to antagonize G17-induced GSK3βSer9 phosphorylation (Fig 1D, pGSK3βSer9 panel), despite a complete inhibition of AKTSer473 phosphorylation (pAKT Ser473 panel). In addition, treatment of another gastric cancer cell line (MKN45) with G17 produced an increase in GSK3βSer9 phosphorylation without any effect on AKT phosphorylation (Fig 1E). These results suggested that G17-induced increase of GSK3βSer9 phosphorylation was mediated via PI3K/AKT independent pathway. AKT-independent phosphorylation of GSK3β has been reported earlier [29,46], including those mediated by members of the PKC
pathway [47]. It will thus be important to determine the contribution of any of these signaling pathways in mediating G17-induced GSK3βSer9 phosphorylation.

The detailed mechanism by which GSK3β regulates migration is still unknown and might involve specific downstream targets. Since Snail and β-catenin are both downstream targets of GSK3β [22,26], which are also involved in mediating EMT, migration and proliferative responses, the next set of studies were specifically focused on understanding the role of these molecules on G17-induced events. Snail has been shown to mediate inflammation-linked migration in cancer cells [48] and promote EMT, a phenomenon that is a prerequisite for cellular migration, invasion and normal development process [19,49]. Our studies with G17 indicated a transient increase in Snail protein expression as well as transcription (Figs 2A, 2D) corresponding to the time of increased GSK3βSer9 phosphorylation. These studies also show that the increase in Snail expression was mediated via inhibition of GSK3β pathway, since pretreatment with a pharmacological inhibitor of GSK3β (AR-A014418) induced Snail expression and transcription in the absence of G17 (Figs 3A, B). In addition, ectopic overexpression of GSK3β-S9A inhibited G17-induced Snail transcription (Fig 3C). The other GSK3β target β-catenin is considered to be a major oncoprotein and a mediator of conventional Wnt/β-catenin pathway [26]. In normal cells, constitutively active GSK3β negatively regulates Wnt/β-catenin signaling via phosphorylation-induced degradation of β-catenin, thus limiting β-catenin expression and activation of Wnt/β-catenin signaling. Interestingly, in our studies, despite an induction of GSK3βSer9 phosphorylation (indicating inactivation), we observed no increase in β-catenin total protein expression at any time point following G17 treatment as reported earlier [36]. However, Western analysis with nuclear extracts showed a distinct increase in β-catenin nuclear translocation with G17, which was also induced following pretreatment with AR in the absence of G17 upto similar levels (Fig 3D, upper panel). This indicated a link between GSK3β inactivation and β-catenin nuclear translocation. The results from these studies and those of others thus suggest that GSK3β activation can regulate β-catenin signaling at two distinct levels: (i) it inhibits β-catenin expression via activating the conventional degradation pathway (ii) it inhibits β-catenin nuclear translocation via a yet unknown mechanism. The former event seems to be lacking in these cells with G17 stimulation, since G17 does not lead to an increase in β-catenin total protein expression [36]. The second event is present in the G17 pathway, since inhibition of GSK3β by AR increases β-catenin nuclear translocation. The mechanism how GSK3β inhibits β-catenin nuclear translocation is still unclear, and might involve a similar phosphorylation dependent mechanism as was reported in the nuclear export of cyclin D1, another GSK3β downstream target [50]. Recent studies by another group have demonstrated that p21-activated kinase 1 (PAK1) is also involved in regulating various steps of β-catenin signaling and migration following G17 stimulation [51]. It is thus tempting to speculate that a crosstalk between GSK3β and PAK1 might be mediating this process. Since JNK pathway can regulate β-catenin nuclear translocation [52] and G17 can activate JNK [31], G17 studies were also performed following pretreatment with an inhibitor of JNK pathway (SP600125). SP600125, however, was unable to show any increase in β-catenin nuclear translocation or Snail induction (Fig 3D) despite a complete antagonism of c-Jun phosphorylation, suggesting that these two proteins are specifically regulated by GSK3β axis.

To understand whether Snail and β-catenin were specific downstream targets of GSK3β to mediate G17-induced migration, G17 studies were performed following siRNA-mediated knockdown of endogenous Snail or β-catenin expression. Knockdown of either protein alone produced a partial inhibition of G17-induced migration (Fig 5A) as well as MMP7 promoter induction (Fig 6B). However, combined knockdown of both proteins significantly antagonized these events. Although a crosstalk between β-catenin and Snail has been reported earlier [53], we were unable to detect any endogenous interaction between these two proteins (data not shown). Further studies are needed to elucidate how Snail and β-catenin coordinate with each other to mediate G17 effects. Interestingly, our earlier studies have indicated the involvement of MLK3/JNK1/c-Jun axis in regulating MMP7 promoter activation and migration following G17 stimulation [31]. Taken together it seems that G17 stimulation operates via two independent signaling axes which ultimately leads to migration: one involves activation of MLK3/JNK1 axis which operates via activation of c-Jun, and the second one involves inhibition of GSK3β axis leading to an induction of Snail expression and β-catenin nuclear translocation. Both of these axes converge to induce MMP7 transcription and migration (Fig 7), and thus represent two potential targets for future drug development.

**Conclusions**

The present study demonstrates that G17-induced migration and MMP7 promoter induction in gastric cancer cells involve an inhibition of GSK3β pathway. G17-induced inhibition of GSK3β leads to an increase in Snail expression and β-catenin nuclear translocation, both of which collectively mediate the migratory response. Thus β-catenin and Snail serve as two downstream targets of GSK3β in G17-induced migration pathway. However, this G17/GSK3β/Snail-β-catenin axis seems to be indepen-
dent of a previously identified G17/MLK3/JNK1 axis [31] and both operate parallel to each other to induce migration. Targeting of both of these axes might be beneficial to antagonize G17-induced migratory effects in gastric cancer.

**Methods**

**Reagents**

DMEM, LipofectAMINE 2000 and β-Galactosidase assay kit were purchased from Invitrogen (Carlsbad, CA); amidoated gastrin (G17) from Bachem (King of Prussia, PA) and the luciferase assay kit from Promega (Madison, WI). The antibodies utilized were obtained from the following sources: GSK3β, pGSK3βSer9, AKT, pAKTSer473, pc-Jun, c-Jun, Snail and Lamin A/C (Cell Signaling Technology, Danvers, MA), GAPDH (Ambion, Austin, TX), β-catenin (BD Biosciences, San Jose, CA), HA.11 (Covance, Berkeley, CA). The CCK2R antagonist YM 022 was from Tocris Bioscience (Ellisville, MO), PI3 Kinase inhibitor Wortmannin and GSK3β inhibitor AR-A014418 from EMD Biosciences (Gibbstown, NJ) and JNK inhibitor SP600125 from Alexis Biochemicals, Axxora (San Diego, CA). The AGSE cells were obtained from Dr. Timothy C. Wang (Columbia University Medical Center, New York, NY) as described earlier [34,36]. The MMP7-luciferase promoter construct was obtained from Dr. Howard Crawford (Stony Brook University, Stony Brook, NY) [54] and the Snail-luciferase promoter was obtained from Dr. Antonio Garcia de Herreros (Universitat Pompeu Fabra, Barcelona, Spain) [18]. The HA-tagged GSK3β expression vectors (WT, K/A and S9A) were from Dr. James R Woodgett (University of Toronto, Toronto, Canada) [55].

**Cell Culture**

The AGSE cells used in these studies were derived from AGS cells stably overexpressing CCK2R as described previously [34]. These cells were maintained in DMEM supplemented with 10% FBS and 100 IU/ml penicillin as reported [31,36]. Wherever indicated, confluent populations of cells were treated with 100 nM G17 in serum deficient media, and subjected to Western Blot analysis, luciferase or migration assays. In the studies with various inhibitors, cells were pretreated with the specific inhibitors followed by G17 treatment.

**Luciferase assays**

Subconfluent populations of cells were transiently transfected with Snail-luciferase [18] or MMP7-luciferase reporter constructs [54] along with β-Galactosidase vector (to correct for transfection efficiency) using lipofectAMINE 2000 as per manufacturer’s instructions. To determine the effect of overexpression of GSK3β (WT or mutants) or knockdown of Snail and β-catenin, the corresponding overexpression vectors or siRNAs respectively were co-transfected along with the luciferase reporters. Each transfection was performed in triplicate and each experiment was repeated at least twice. Following 48 hours of recovery in the growth medium, the transfected cells were treated with either vehicle or G17 for 24 hours. Luciferase and β-Galactosidase (β-Gal) assays were performed as described [31] using a luminometer (Berthold Technologies, Centro XS3 LB 960) and a plate reader (Power Wave XS, Biotek) respectively. The results obtained were calculated as the ratio of relative light units (RLU) to β-Gal values (RLU/β-Gal) and expressed as % increase compared to controls.

**Western Blot analysis**

Whole cell extracts were prepared from cells treated with G17 by RIPA extraction buffer, and equal amounts of total protein were subjected to Western Blot analysis utilizing procedures described previously [31,36]. Nuclear protein extraction was performed following protocols as described [56]. Briefly, cells were incubated with a lysis buffer (containing 1% Triton X-100, 50 mM Hepes pH 7.6, 150 mM NaCl, 100 mM NaF, 50 mM Na-pyrophosphate, 4 mM EDTA and 10 mM Na2VO4 supplemented with protease inhibitors) and rotated at 4°C for 30 min-

![Figure 7 Model representing the signaling pathway of G17-induced migration. Stimulation of G17/CCK2R pathway leads to activation of two separate signaling axes: (i) an activation of MLK3/JNK1 axis, which via activation of its downstream transcription factor c-Jun induces MMP7 transcription leading to increased migration; (ii) an induction of GSK3βSer9 phosphorylation (inhibition of axis) via a PI3Kinase-independent (and yet unknown) mechanism. This leads to an increase in Snail protein expression and β-catenin nuclear translocation, combination of which lead to increased MMP7 transcription and cell migration.](image-url)
utes. This was followed by centrifugation at 13,000 rpm for 15 minutes to pellet nuclei, washing the nuclear pellet once with wash buffer (containing lysis buffer + 25% glycerol) and centrifugation again. To obtain nuclear extract, the nuclear pellet was lysed in nuclear lysis buffer (wash buffer + 300 mM NaCl), sonicated and incubated on ice for 30 minutes followed by centrifugation. Western Blot analysis was performed with equal amounts of nuclear protein with the indicated antibodies. To determine the purity of the nuclear preparation, they were blotted with antibodies against GAPDH (cytoplasmic protein) and Lamin A/C (nuclear protein).

Small interference RNA (siRNA)
The β-catenin siRNA [57] was synthesized from Dharmacon (Lafayette, CO) and the Snail siRNA was from Invitrogen (Carlsbad, CA) [58]. The control-siRNA was from Ambion (Austin, TX). siRNA transfection was performed using lipofectAMINE 2000 as per manufacturer’s instructions following protocols described earlier [57]. G17 treatment was performed after 48 hours of siRNA transfection, and the cells were then subjected to Western Blot, luciferase or migration assays.

Wound healing assay
These were performed following procedures described earlier [31]. Confluent populations of cells were wounded linearly using a small pipette tip, washed with PBS and treated with the various agents in serum deficient media for various periods of time. For overexpression experiments, cells were transfected with the corresponding overexpression vectors or siRNAs, wounded 48 hrs following transfection, and allowed to recover overnight before adding G17. The migratory cells were visualized and photographed using inverted phase-contrast microscopy (Axiovert 200 inverted microscope, Zeiss), interfaced with a camera (AxioCam) and the image analyzer software (Axiovision, Zeiss). To estimate relative migration, the unclosed distances at 3 points in each scratch were measured using the Axiovision software, their average calculated and plotted as “Average Gap”.

Abbreviations
APC: Adenomatous Polyposis Coli; B-Gal: β-Galactosidase; CCK2R: CCK2 receptor; EV: Empty Vector; EMT: epithelial-to-mesenchymal transition; G-17: ammated gastrin; G-Gly: glycine-extended gastrin; GI: Gastrointestinal; GSK3β: Glycogen Synthase Kinase 3beta; HRE: hormone responsive element; HSF1: heat shock factor 1; HSF: heat shock factor; JR: JNK response; MIP: matrix metalloproteinase; MLK3: Mixed lineage kinase-3; MMPs: matrix metalloproteinases; PCNA: proliferating cell nuclear antigen; RLU: relative light units; siRNA: small interference RNA; TCF/LEF: T cell factor/lymphoid enhancer factor; WT: Wild Type;

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
PM performed Western blot analysis, Luciferase analysis, overexpression and migration experiments. SS performed some of the gastrin-related migration and Western experiments in various gastrointestinal cells, obtained photographs of migration and contributed to the editing of the manuscript. AR helped with the analysis of the GSK3β overexpression experiments and provided intellectual input in this collaborative study. BR contributed to the overall study design, interpretation of the results during all phases and drafted/editd the final manuscript. All authors read and approved the final manuscript.

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