Plasma chitinase 3-like 1 is persistently elevated during first month after minimally invasive colorectal cancer resection

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AIM: To assess blood chitinase 3-like 1 (CHI3L1) levels for 2 mo after minimally invasive colorectal resection (MICR) for colorectal cancer (CRC).

METHODS: CRC patients in an Institutional Review Board approved data/plasma bank who underwent elective MICR for whom preoperative (PreOp), early postoperative (PostOp), and 1 or more late PostOp samples [postoperative day (POD) 7-27] available were included. Plasma CHI3L1 levels (ng/mL) were determined in
duplicate by enzyme linked immunosorbent assay.

RESULTS: PreOp and PostOp plasma sample were available for 80 MICR cancer patients for the study. The median PreOp CHi3L1 level was 56.8 CI: 41.9-78.6 ng/mL (n = 80). Significantly elevated (P < 0.001) median plasma levels (ng/mL) over PreOp levels were detected on POD1 (667.7 CI: 495.7, 771.7; n = 79), POD 3 (132.6 CI: 95.5, 173.7; n = 76), POD7-13 (96.4 CI: 67.7, 136.9; n = 62), POD14-20 (101.4 CI: 80.7, 287.4; n = 22), and POD 21-27 (98.1 CI: 66.8, 137.4; n = 20, P = 0.001). No significant difference in plasma levels were noted on POD27-41.

CONCLUSION: Plasma CHi3L1 levels were significantly elevated for one month after MICR. Persistently elevated plasma CHi3L1 may support the growth of residual tumor and metastasis.

Key words: Colorectal cancer; Recurrence; Minimally invasive colorectal resection; Chitinase 3-like 1; Metastasis

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Core tip: Colorectal cancer (CRC) resection surgery is well known to be associated with short lived immunosuppression and transient plasma protein changes. We have documented that a second set of blood protein alterations that last for 3 to 5 wk after CRC; interestingly, all of these proteins play a role in angiogenesis. This group of pro-angiogenic proteins includes vascular endothelial growth factor, placental growth factor, angiopoietin-2, monocyte chemo-attractant protein-1 and matrix metalloproteinase-2. Our published data further confirms that pro-angiogenic plasma compositional changes that may support the establishment of metastases and the growth of already present tumor deposits during the early post-surgical period.

Shantha Kumara HMC, Gaita D, Miyagaki H, Yan X, Hearth SAC, Njoh L, Cekic V, Whelan RL. Plasma chitinase 3-like 1 is persistently elevated during first month after minimally invasive colorectal cancer resection. World J Gastrointest Oncol 2016; 8(8): 607-614 Available from: URL: http://www.wjgnet.com/1948-5204/full/v8/8/607.htm DOI: http://dx.doi.org/10.4251/wjgo.v8.i8.607

INTRODUCTION

Colorectal cancer (CRC) is the second most diagnosed cancer in the United States, with an expected 142820 cases and 50830 deaths in 2013[1]. Surgical resection is the primary treatment for the 80% of CRC patients who present without metastatic disease. While this procedure is considered curative for these patients, more than 40% who present with stage II or III disease will develop a recurrence[1,2].

It has been hypothesized that surgical resection of tumors may paradoxically contribute to the development of cancer recurrences. Murine studies have shown those laparotomy and bowel resections are associated with increased tumor growth and establishment vs results in anesthesia control mice[3-5]. In humans, numerous case reports have noted increased tumor growth soon after surgery in patients with residual cancer[6-9]. A number of mechanisms, including surgery-related immunosuppression and the removal of primary tumor generated anti-angiogenic factors, have been proposed to account for accelerated tumor growth after surgery[10,11]. Of interest, the last decade has seen the emergence of another possible mechanism, namely surgery-related proangiogenic plasma compositional changes that may support the establishment of metastases and the growth of already present tumor deposits during the early post-surgical period.

Angiogenesis or neo-vascularization is necessary for tumor growth greater than 1-2 mm[3,12,13]. During a nascent tumor’s initial avascular dormant stage, cells obtain nutrients by passive diffusion only. The activation of what has been called the “angiogenic switch” leads to the development of new vessels that infiltrate and extend beyond the tumor mass which permits growth and, later, metastasis. Similarly, further growth of established metastases requires new vessel formation.

It has been shown that colorectal resection is associated with continual elevations (2-5 wk) in the blood levels of a number of angiogenesis promoting proteins including vascular endothelial growth factor (VEGF), angiopoietin-2 (Ang-2), placental growth factor (PIGF), soluble vascular cell adhesion molecule-1 (sVCAM-1), monocyte chemotactic protein-1 (MCP-1), and matrix metalloproteinase-3 (MMP-3)[14-18]. Furthermore, human plasma from the second and third weeks after surgery has been shown to promote endothelial cell (EC) proliferation, invasion, and migration, which are key steps in angiogenesis[14-19]. It is possible, therefore, that the postoperative (PostOp) plasma composition may encourage the growth of residual tumor metastases after “curative” colorectal resection of a primary tumor. In an effort to better understand the impact of surgery the search goes on for other proteins with proangiogenic effects whose blood levels may be increased after surgery.

Chitinase 3-like 1 (CHi3L1), also named as YKL-40 and human cartilage glycoprotein-39 (HCgp-39), is a member of the glycosyl hydrolase 18 protein family[20]. The substrate for this family of proteins is chitin, a polymer of N-acetyl-glucosamine. Importantly, chitins are not found in mammals yet these proteins are produced by a number of mammalian cell types including EC’s neutrophils, macrophages, and vascular smooth muscle cells. CHi3L1 does not have any known enzymatic activity in mammals, however, it is believed that this protein binds to endogenous carbohydrates such as hyaluronic
For CRC blood levels of CHi3L1 might be increased. This following minimally invasive colorectal resection (MICR) remodeling and angiogenesis we hypothesized that surgery are unknown. Because CHi3L1 involved in tissue the impact of surgical resection on blood levels after of CHi3L1 are elevated prior to surgery in CRC patients, to be strongly associated with increased microvascular levels of CHi3L1 and a poor prognosis a correlation has been demonstrated between blood ovarian, thyroid, and malignant melanoma and a wide variety of cancers including colorectal, breast, prostate, lung, thyroid, endometrial, pancreatic, hepatocellular, ovarian, gastric cancer, and malignant melanoma.

Additionally, for the majority of these malignancies, a correlation has been demonstrated between blood levels of CHi3L1 and a poor prognosis. Also, increased expression of CHi3L1 in CRC has been shown to be strongly associated with increased microvascular density. Although it has been shown that serum levels of CHi3L1 are elevated prior to surgery in CRC patients, the impact of surgical resection on blood levels after surgery are unknown. Because CHi3L1 involved in tissue remodeling and angiogenesis we hypothesized that following minimally invasive colorectal resection (MICR) for CRC blood levels of CHi3L1 might be increased. This study was carried out to investigate plasma levels of CHI3L1 during the first 4 to 6 wk following MICR for CRC.

**MATERIALS AND METHODS**

**Patient selection for study**

Patient populations who underwent elective MICR were selected for this study from an Institutional Review Board (IRB) approved multi center plasma and data bank that was organized by Columbia University and that included the following institutions: New York Presbyterian Hospital, Columbia University; the Ferguson Clinic, Grand Rapids, Michigan and Mount Sinai West Hospital Center, New York. The broadly stated purpose of this effort is to study the physiologic, immunologic, and oncologic ramifications of open and minimally invasive surgical methods. Prospective data including demographic, operative, and short term recovery statistics were collected for all patients. Patients who were immunosuppressed or transfused peroperatively were excluded. Patients undergoing urgent or emergent surgery were, likewise, excluded.

**Blood sampling and processing**

To be eligible for entry into this study preoperative (PreOp) and, at least, several PostOp plasma samples had to be available for CRC patients who underwent MICR. Of note, blood samples after postoperative day (POD) 7 were obtained at follow up office appointments but were not scheduled on a specific POD. Late post-operative samples were not available on the same day from all patients so the late samples were bundled into 7 d (or longer) time periods (POD 7-13, POD 14-20, POD 21-27, and POD 28-41). Samples were collected in tubes containing heparin. The samples were processed within 6 h of collection, and the plasma fraction stored in aliquots at -80 °C until the assay was performed.

**Plasma CHI3L1 analysis**

Plasma CHI3L1 levels were determined in duplicate using commercially available enzyme linked immunosorbent assay (ELISA) (R and D Systems, Minneapolis). CHI3L1 concentrations are reported as nanograms per milliliter (ng/mL).

**Statistical analysis**

For continuous variables, data are expressed as mean ± SD. Frequencies and percentages were determined for categorical variables. In regards to the CRC Pre vs PostOp CHI3L1 comparisons, the results are reported as the median and 95%-CIs and the Wilcoxon signed rank test was used to analyze the data. Correlation between PostOp CHI3L1 plasma levels and incision size and length of surgery was evaluated by the Spearman’s rank correlation coefficient (r). All data analysis was performed utilizing SPSS version 15.0 (SPSS, Inc., Chicago, IL). Because the sample size varies for the POD 7-13, POD 14-20, POD 21-27 and POD 28-41 time points a separate Pre-Op bar is included for each time point in the Figure 1.
RESULTS

A total of 80 MICR patients with CRC (42 male/38 female, age 65.66 ± 12.83 years) were included in the study. The majority of patients underwent right colectomy, lower anterior resection/anterior resection, or sigmoidectomy (Table 1). Laparoscopic-assisted methods were used in 59% while hand-assisted minimally invasive methods were utilized in 41% of the patients. There were 12 conversions (15%) to open methods (defined as final incision > 7 cm in laparoscopic-assisted cases and incision > 11 cm in hand-assisted cases). The mean surgical incision length was 7.8 ± 3.6 cm for the entire population, the mean operative time was 308.3 ± 124.2 min. The mean length of stay was 6.6 ± 4.3 d. Seven complications (8.8%) were noted including small bowel obstruction (3), wound infection (1), hernia (1), hyperkalemia (1), and hematoma (1). There were no perioperative deaths. The cancer pathological stage breakdown of the study population was as follows: Stage I, 20 (25.0%); stage II, 25 (31.3%); stage III, 32 (40%); stage IV, 3 (3.7%).

The median PreOp CHI3L1 level was 56.8 CI: 41.9, 78.6 ng/mL (n = 80). Significantly elevated median plasma levels were noted on POD 1 (667.7 CI: 495.7, 771.7 ng/mL, n = 79, P < 0.001), POD 3 (132.6 CI: 95.5, 173.7 ng/mL, n = 76, P < 0.001), POD 7-13 (96.4 CI: 67.7, 136.9 ng/mL, n = 62, P < 0.001), POD 14-20 (101.4 CI: 80.7, 287.4 ng/mL, n = 22, P < 0.001), and POD 21-27 (98.1 CI: 66.8, 137.4 ng/mL, n = 20, P = 0.001) when compared to PreOp levels. The percent increase from median baseline at each time point was 1068% at POD 1, 157% at POD 3, 88% at POD 7-13, 50% at POD 14-20, and 64.0% at POD 21-27. No significant difference found at the POD 28-41 time point.

To determine whether there was a statistical correlation between incision length and post surgery plasma CHI3L1 levels, the laparoscopic-assisted group (mean incision size 5.8 cm) and the hand-assisted group (mean incision, 10.7 ± 3.5 cm) were compared. While the mean incision size for the hand assisted group was higher, there was no statistically significant difference in PostOp plasma CHI3L1 levels between the groups. Furthermore, at 5 of the 6 PostOp time points there was no correlation between CHI3L1 levels and incision length. There was a statistically significant correlation between incision length and CHI3L1 levels at the POD 14-20 time point. There was no significant correlation between cancer stage and PreOp CHI3L1 plasma level.

DISCUSSION

We have demonstrated in several studies that elective surgery for CRC is associated with persistent plasma protein changes that render the blood proangiogenic for 2-3 wk as judged by in vitro invasion, migration and EC proliferation analysis[14-18]. This is in contradiction to the vast majority of serum proteins that have been studied perioperatively whose levels are transiently elevated after major surgery (alterations resolving in < 1 wk such as IL-2, IL-6, CRP, etc). CHI3L1, which has been shown to promote EC proliferation and tube formation among its other effects, can now be added to the list of persistently elevated proangiogenic proteins.

In this study, mean plasma CHI3L1 levels of 80 CRC patients rose to a peak of 1068% over the PreOp baseline value on POD1, and remained significantly elevated at 50%-157% over baseline for four weeks after MICR until returning to baseline at the POD 28-34 time point. Of note, the percent changes in CHI3L1 levels over baseline were amongst the highest observed when compared to the results of the 9 other proteins previously shown to have long duration plasma increases post MICR.

Because CHI3L1 levels have been shown to be increased preoperatively in cancer patients when compared to tumor free patients, a fall in blood levels would be anticipated after the primary tumor is resected. Why then do plasma levels of CHI3L1 rise during the first month after surgery? Most likely there are several mechanisms. As regards the initial elevation during the first 3-4 d after MICR, although unproven, it is possible that macrophages and polymorphonuclear leukocytes (PMNs), which play a vital role in the acute inflammatory response that follows surgery, generate the added CHI3L1. This hypothesis is based on the following observations. Both PMNs and macrophages have been shown to generate CHI3L1 in the setting of chronic inflammation (e.g., IBD and rheumatoid arthritis) and blood levels of CHI3L1, known to be increased in these patients, have been shown to directly correlate with disease severity.

As regards the persistent plasma elevation noted during weeks 2 to 4, the authors believe, also admittedly without direct evidence, that the healing surgical wounds are the source of the added CHI3L1. Since CHI3L1 has

Table 1 Demographic and clinical characteristics of the study population

| Surgical method, & (%)| Laparoscopic-assisted | Hand-assisted/hybrid laparoscopic |
|------------------------|-----------------------|----------------------------------|
| Type of resection, & (%)| Subtotal/total 4 (5.0) | LAR/AR 26 (32.5) |
|                      | Right 24 (30.0)       | APR 2 (2.5)                     |
|                      | Transverse 5 (6.2)    | Sigmoid/rectosigmoid 13 (16.3)  |
|                      | Left 6 (7.5)          | LAR 26 (32.5)                  |
|                      |                      |                                |
| Sex, n (%)           | Female 38 (47.5)      |                                |
|                      | Male 42 (52.5)        |                                |
| Age, years (mean ± SD)| 65.66 ± 12.83        |                                |
| Length of stay, d (mean ± SD) | 6.68 ± 4.30   |                                |
| Incision length, cm (mean ± SD) | 7.78 ± 3.61 |                                |
| Operative time, min (mean ± SD) | 308.3 ± 124.2 |                                |

(Please note: The table is not fully visible in the provided text, but it contains demographic and clinical characteristics of the study population.)
been shown to play key role in tissue remodeling as well as angiogenesis it is certainly possible that it is involved with the wound healing process. The hypothesis is that wound levels are very high such that the CHI3L1 spills over into the blood stream raising plasma levels. Support for this concept can be found in several human studies that measured VEGF levels in wound fluid and the blood of surgery patients.

In a Dutch study of MICR patients, on POD 4, wound VEGF fluid levels were found to be 7 times higher than serum concentrations which were elevated over PreOp baseline levels. The same research team, in a study of mastectomy patients noted that on POD 4 wound VEGF levels were between 23 and 32 times greater than serum levels. There is also strong preliminary unpublished data of the authors suggesting that wound levels of ANG2, and MMP2 are elevated and many times higher than plasma concentrations after MICR (plasma levels also elevated 3-4 wk). Further, the fact that the proteins shown to have markedly elevated wound levels play roles in neovascularization (like CHI3L1) indirectly suggests that the origin of the CHI3L1 during most of the first PostOp month is the wound. As mentioned above, the authors are currently conducting a study that is simultaneously assessing wound and plasma protein levels which will soon shed more light on this topic.

What are the potential clinical ramifications, if any, of the month long elevation in CHI3L1 levels? There is some research evidence that CHI3L1 may play a direct role in chronic inflammation related epithelial cancer development via up-regulation of β catenin. As mentioned earlier, in addition to promoting angiogenesis in general there is strong in vivo evidence that CHI3L1 promotes tumor angiogenesis in particular. Macrophages are known to promote cancer progression by producing a number of growth and proangiogenic factors. Tumor associated macrophages lack antigen presenting abilities and have been found to cluster in avascular areas of breast carcinomas. Kawada et al revealed that the CHI3L1 increased the secretion of inflammatory chemokines, IL-8 and monocyte chemoattractant protein-1 (MCP-1), from SW480 cells via mitogen-activated protein kinase (MAPK) pathway in vitro. Furthermore, CHI3L1 expressing colon cancer cells significantly increased macrophage recruitment in xenograft mice, as well as tumor growth and angiogenesis. Persistently elevated levels of plasma CHI3L1 levels together with IL8 and MCP-1 may collectively enhance the blood angiogenic property after MICR.

The proangiogenic effects of CHI3L1 should be considered in the context of the other 9 proteins (all with proangiogenic effects) whose levels have also been shown to be persistently elevated after MICR. There is a concern that these plasma changes might promote tumor angiogenesis early after resection of the primary cancer in patients with established distant micrometastases. Further, these compositional changes may encourage the establishment of new metastases by circulating viable tumor cells present in the blood stream after surgery. Is there any evidence of increased tumor growth after surgery in humans?

The medical literature contains numerous case reports of rapid growth of residual metastases after major surgery. In particular, rapid growth of pre-existing metastatic cancer in the liver after resection of the primary colorectal tumor has been noted by several investigators. Of note, in one investigation the vascular density of metastases has been noted to be significantly increased over the pre-resection baseline several months after primary resection; this suggests that tumor angiogenesis is stimulated after surgery. Finally, there is strong experimental data showing that surgical trauma is associated with accelerated tumor growth.

The data from this study and others mentioned above suggest that the first month after surgery may be a precarious time for cancer patients who harbor residual tumor since the plasma composition is proangiogenic for up to a month. The fear is that these conditions will foster blood vessel formation in the tumor and, in so doing, will promote tumor growth. Because conventional adjuvant or palliative PostOp chemotherapy is usually started 4 to 8 wk post operatively the patient is left to their own during this time period. It is logical to administer some type of anti-cancer treatment in this unused time window. The challenge is to find effective anti-cancer agents that do not interfere with the process of wound healing. Presently, a phase I clinical trial is underway that is assessing perioperative treatment with polyphenon E (a green tea extract) and a milk thistle plant component called siliphos. Both agents have been shown to inhibit tumor growth while not inhibiting wound healing which makes them safe for the early PostOp period.

A weakness of this study was the relatively small number of late plasma samples collected. Most of the late samples were obtained during follow-up visits scheduled by the patients; unfortunately, many patients refused late PostOp blood draws. Thus, there were fewer late samples than at the PreOp, POD 1, and POD 3 time points. This made it necessary to bundle the late samples into 7 to 13 d blocks. Also, the timing of the late samples varied considerably. A larger study will allow a more meaningful correlation of PreOp CHI3L1 levels and cancer stage. Similarly, a follow up study with more uniform and comprehensive PostOp blood sampling should allow a better assessment of PostOp levels, in general, as well as the impact of surgical method, if any, on post surgery plasma levels.

The results of this study demonstrate that surgical stress, in particular MICR, is associated with notably increased plasma levels of CHI3L1 that persist for 1 mo after surgery. The source of this elevation is unclear. The clinical implications of these findings, if any, are uncertain. In theory, among other possible effects, this transient plasma alteration may promote tumor angiogenesis in patients with residual cancer after surgical resection of the primary tumor. The fact that plasma levels of 9 other
proteins with proven proangiogenic effects have been shown to be elevated for 2 to 4 wk after MICR lends support to this hypothesis. Further study of perioperative plasma CHI3L1 levels and of the clinical ramifications of the surgery-related long duration proangiogenic plasma compositional changes appear to be warranted.

**COMMENTS**

**Backgrounds**

Major abdominal surgery is well known to be associated with a brief period of immunosuppression and short lived plasma protein changes. Recently, another group of blood protein alterations that lasts at least 3 to 5 wk after colorectal cancer (CRC) resection have been noted. All of these proteins play a role in angiogenesis. This set of proangiogenic proteins includes vascular endothelial growth factor (VEGF), placental growth factor (PIGF), angiopoietin-2, soluble vascular adhesion molecule-1 (sVCAM-1), monocyte chemo-attractant protein-1 (MCP-1) and matrix metalloproteinase-3. Proangiogenic chitinase 3-like 1 (CHI3L1) protein promotes in vitro human endothelial cell (EC) migration and tube formation, however, the impact of minimally invasive colorectal resection (MICR) for CRC on plasma levels of CHI3L1 is unknown.

**Research frontiers**

CHI3L1 is a member of the Chitinase family of proteins and has chemotactic and proangiogenic properties similar to those of MCP-1 and IL8 that are mediated, in part, by MCP-1. Colon, breast, and hepatocellular carcinomas have been shown to express higher levels of CHI3L1. It has been shown that CHI3L1 may utilize its chitin binding ability to communicate with other signal transduction pathways to modulate inflammation, apoptosis, tissue remodeling, cell growth and angiogenesis. CHI3L1 has been shown to promote in vitro cancer cell proliferation, macrophage recruitment, human EC migration and tube formation, and contributes to wound healing. The authors analyzed preoperative (PreOp) and post-MICR CHI3L1 levels in CRC patients. Significantly elevated blood levels of CHI3L1 may promote tumor angiogenesis and therefore, the growth of residual tumor during the first month after MICR.

**Innovations and breakthroughs**

Persistently elevated plasma levels of the proangiogenic proteins including VEGF, Ang-2, PIGF, sVCAM-1 and MMP3 have been noted for 3-5 wk after CRC resection. Plasma from the 2nd and 3rd weeks after CRC resection has been shown to stimulate EC proliferation and migration in vitro when compared to EC culture results with PreOp plasma. This study reports that plasma CHI3L1 levels are significantly elevated over PreOp levels for a month after MICR for CRC. These persistent plasma compositional changes may promote tumor angiogenesis and growth in patients with residual cancer deposits in the 1st postoperative (PostOp) month.

**Applications**

This study results further support that persistent plasma compositional changes may promote tumor angiogenesis and growth in patients with residual cancer deposits in the first PostOp month. Thus, it is logical to give anti-cancer therapy perioperatively, for safe use in this period, in addition to having anti-cancer effects, candidate agents must not impair wound healing.

**Terminology**

The significantly increased blood proangiogenic protein levels during the early PostOp period after MICR and open colorectal resection may be associated with the short lived acute inflammatory response that occurs after surgery and resolves in the first week. In contrast, the later and persistent elevation noted during weeks 2-4 after surgery may be related to wound healing. Persistently elevated plasma CHI3L1 levels shown in this study, together with the similarly increased levels of the other proangiogenic proteins such as VEGF, ANG2, PIGF, sVCAM-1/MCP-1 and MMP3 may collectively promote tumor angiogenesis and therefore, the growth of residual tumor during the first month after MICR.

**Peer-review**

The paper is written in a good language, the logic is clear and the subject and results are discussed graphically and meaningfully.

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