Alterations of the gastric stump and not resected stomach mucosa after truncal vagotomy in rats

Paulo Antônio Rodrigues\textsuperscript{1}, Luiz Eduardo Naresse\textsuperscript{2}, José Guilherme Minossi\textsuperscript{3}, Claudia Nishida Hasimoto\textsuperscript{4}

\textsuperscript{1}PhD, Assistant Professor, Departament of Surgery & Orthopedics, Faculty of Medicine, Sao Paulo State University (UNESP), Botucatu-SP, Brazil. Conception, design, intellectual and scientific content of the study; acquisition and interpretation of data; technical procedures; manuscript writing.
\textsuperscript{2}PhD, Associate Professor, Departament of Surgery & Orthopedics, Faculty of Medicine, UNESP, Botucatu-SP, Brazil. Interpretation of data, technical procedures.
\textsuperscript{3}PhD, Assistant Professor, Departament of Surgery & Orthopedics, Faculty of Medicine, UNESP, Botucatu-SP, Brazil. Interpretation of data, technical procedures.
\textsuperscript{4}Fellow Master degree, Postgraduate Program in General Basis of Surgery, Faculty of Medicine, UNESP, Botucatu-SP, Brazil. Acquisition and interpretation of data.

ABSTRACT

PURPOSE: To evaluate morphological changes of the gastric stump and not resected stomach mucosa after the completion of truncal vagotomy.

METHODS: Eighty male Wistar rats were divided into four groups: CT, TV, RY and RYTV. In CT group, abdominal viscera were manipulated and the abdominal cavity was closed, in TV vagal trunks were isolated and sectioned, in RY a partial Roux-en-Y gastrectomy was performed and in RYTV the vagal trunks were sectioned and a partial Roux-en-Y gastrectomy was performed. At the 54th week after surgery, the rats were euthanized. The findings were submitted to histological analyses.

RESULTS: None macroscopic or histological alterations in groups TV and CT was observed. Specimens from RY and RYTV groups did not show alterations in the gastric stump mucosa. At the jejunal side of the gastroenterostomy we found shallow ulcerative lesions always single, well-defined and with variable diameter 3 to 6 mm, six times in the RY group and none in the RYTV group (RY>RYTV, p=0.008). Neoplastic or preneoplastic lesions were not diagnosed in all groups.

CONCLUSION: Truncal vagotomy is a safe and non-carcinogenic method in not resected and partially resected stomach.

Key words: Vagotomy, Truncal. Adenocarcinoma. Stomach. Rats.
Introduction

Truncal vagotomy is a surgical technique still used nowadays in elective and emergency surgical procedures. For patients presenting with complicated peptic ulcer disease such as perforation and/or hemorrhage not controlled by endoscopic and clinical treatment, truncal vagotomy associated with a drainage procedure is a fast and effective treatment. This technique can also be used in elective situations as a complement to reducing acid in distal gastrectomies or when the surgeon chooses the reconstruction of alimentary tract through a Roux-en-Y partial gastrectomy. In Roux-en-Y procedure there is a risk of postoperative development of anastomotic ulcers at jejunal side of gastroenterostomy, caused by diversion of biliopancreatic secretions from the gastroenterostomy and therefore damaging the buffering of the chlordropeptic secretion produced by the stomach remnant. Moreover, truncal vagotomy has been identified as a risk factor for the development of gastric cancer in epidemiological and experimental studies. The mucosa denervation would lead to a decrease in the gastric mucosa cell function, cytoprotection, and an increase in regeneration of immature cells.

Methods

The study was developed after approval of the Ethics Committee of Experimental Research of UNESP (Protocol 612) and follows the Council for International Organization of Medical Sciences (CIOMS) ethical code for animal experimentation.

Eighty male Wistar rats weighing approximately 150 grams provided by the Central Animal Facility of the Faculty of Medicine of Botucatu were used. Animals, five in number per unit, were housed in plastic boxes wrapped in room with steam exhaust air, artificial light cycles of 12 hours and room temperature maintained at 24°C. The animals were offered water ad libitum and standard rat chow. After the end of the adaptation period, they were divided in four experimental groups of 20 rats in each group, as described:

Control (CT): underwent to laparotomy and manipulation of the abdominal viscera.

Truncal vagotomy (TV): underwent to section of the anterior and posterior vagal trunks.


Roux-en-Y + Truncal Vagotomy (RYTV): underwent to the same surgical procedure of group RY, adding bilateral truncal vagotomy.

In all surgical procedures, animals received anesthesia by intraperitoneal injection of sodium pentobarbital at a dose of 30 mg / kg body weight. Access to the peritoneal cavity was obtained by laparotomy with 3 cm long. In CT group, abdominal viscera were manipulated and the abdominal cavity was closed. In TV group, vagal trunks were dissected, isolated and sectioned. In RY group distal half of the glandular stomach was resected, the duodenal stump was buried. The gastrointestinal transit was reconstituted by a Roux-en-Y gastroenterostomy between the proximal cut edge of afferent loop and the efferent limb 10 cm below the gastroenterostomy. RYTV group animals were subjected to a combination of the procedures performed in RY and TV groups. All sutures were done with a 6.0 polypropylene (Ethicon). The layout of surgical procedures is illustrated in Figure 1.

![FIGURE 1 - Schematic representation of the surgical procedures.](image-url)

After the surgical procedure, the animals had access to water only on the first 24h postoperatively, freeing up access to feed after this period. The control of body weight was done weekly in the morning and deaths recorded. At the 54th week after surgery, the rats were euthanized. They were initially anesthetized with the same methodology used for surgical procedures. Through sternotomy and laparotomy intracavitary organs were checked, the stomach were removed, photographed and fixed in 10% formalin for 48h. After fixation, each specimen was sectioned through successive longitudinal sections including the pyloric and oxyntic mucosa and separately squamous epithelium.

At the gastroenterostomy, the sections were perpendicular to it one. We used conventional methods for the preparation of histological slides stained by the hematoxylin/eosin. Histopathological examination was performed using an optical microscope. For the statistical analysis was adjusted a generalized linear model assuming a binomial distribution with LOGIT link. Differences were considered significant when the p<0.05.
Results

The effective number of animals in each group was CT = 20, TV = 20, RY = 13 and RYTV = 12. Postoperative complications, malnutrition and intra-abdominal abscesses were the causes of 15 deaths recorded during the experiment. The examination of the abdominal cavity revealed no significant changes.

The examination of the surgical specimens of CT and TV groups showed no significant gross lesions in both glandular and squamous portion of stomach.

The macroscopic appearance of specimens of the RY and RYTV groups did not show alterations in the squamous portion and in the gastric stump mucosa. The general appearance of specimens from RYTV is illustrated in Figure 2A. At the jejunal side of the gastroenterostomy we found shallow ulcerative lesions always single, well-defined, variable diameter 3 to 6 mm, six times in the RY group and none in the RYTV group (RY>RYTV, p = 0.008), (Figure 2B).

Histological analyses of ulcerated lesions showed that the bottom of the lesions consisted of fibrin crust, variable inflammatory infiltrate and proliferation of collagen. There was no cellular atypia to suggest malignancy (Figure 3). None other significant histological alteration was observed in all groups.

FIGURE 2 - Macroscopic aspect of the specimens from RYTV at the 54th week (A). Ulcerated lesion located in the jejunal side of gastroenterostomy (black arrows) in specimen from RY at the 54th week (B). Sq = Squamous portion; G1 = Glandular portion; J = Jejunum.

FIGURE 3 - Ulcerated lesion in the jejunal side of gastroenterostomy (black arrows) showing proliferation of collagen and mild inflammatory process, (H&E x40).

Discussion

The only significative difference observed among the groups was the occurrence of ulcerative lesions in the jejunal side of the gastroenterostomy, observed in six specimens of the RY group. These lesions may have been caused by the contact between chlorydropeptic secretions, produced in the gastric stump, with the jejunal mucosa. Roux-en-Y procedure causes a decrease in the release of hormones such as secretin and cholecystokinin that, in addition to stimulating the release of pancreatic secretion and gallbladder contraction, results in inhibition of gastric secretion11,12. Furthermore, in this type of surgery, the chlorydropeptic secretion produced in the gastric stump is not buffered by the neutralizing effect of biliopancreatic secretions, which transits the gastroenterostomy, as occurs in the Billroth II gastrectomies. The truncal vagotomy was proved effective in preventing the development of gastroenterostomy ulcers in animals that underwent Roux-en-Y partial gastrectomy. Previous studies also showed the same results13,14. A study compared the frequency of anastomotic ulcers in rat’s subjected to partial Roux-en-Y gastrectomy with or without truncal vagotomy and observed that the occurrence of anastomotic ulcers was significantly higher in the group without vagotomy (92.8% versus 14.3 %) after a ten months follow up15. Another
study in pigs submitted to a Roux-en-Y gastrectomy, with or without truncal vagotomy, obtained a 50% incidence of ulcers in the non-vagotomized and only 20% in the underwent to a truncal vagotomy \(^4\). In the present study, it was not observed the development of benign or malignant proliferative lesions in intact stomach or gastric stump mucosa. However, we cannot fail to mention that truncal vagotomy has the potential to facilitate the development of such lesions. Truncal vagotomy causes several changes in gastric physiology since it decreases the ability of the stomach to secrete hydrochloric acid causing an increase in intragastric pH and consequently the formation of carcinogenic N-nitroso compounds \(^5\). TV may also change the permeability of the mucosa of the stomach due to decreased in secretion of neutral mucins and lose of regulation of submucosal blood flow, facilitating the contact of the N-nitroso compounds with the proliferative compartment of gastric epithelium \(^6\).

Some epidemiological studies suggest that vagotomy can increase the risk of gastric cancer development. In one of them, the authors found an increase risk of gastric cancer development 1.6 times higher than expected comparing patients underwent to a vagotomy for treatment of peptic ulcer and controls \(^4,5\). In another study, which examined cancer incidence in 1,992 patients undergoing to a gastric surgery found that, particularly, the vagotomized had a significantly increased risk of gastric cancer and questions the safety of prolonged acid suppression through use of inhibitors of the proton pump \(^7\). Some experimental studies also suggest that vagotomy can raise the risk of developing gastric cancer: in one study the author refers to be possible to obtaining gastric cancer in rats subjected to a truncal vagotomy and drainage procedure without using carcinogens \(^8\); in another study was reported that vagotomy significantly increases the incidence and number of gastric adenocarcinoma obtained from rats treated with MNNG \(^8\); and finally one study reports that vagotomy associated with duodenogastric reflex in rats treated with MNNG increases the number of tumors obtained when compared to duodenogastric reflex only, and believe that denervation of the gastric mucosa not only leads to decreased of gastric mucosa cell function and cytoprotection but it also increases the regeneration of immature cells \(^8,18\).

**Conclusion**

Truncal vagotomy is a safe and effective method in not resected and partially resected stomach, since it prevents the development of anastomotic ulcers and does not causes the development of lesions considered preneoplastic or neoplastic.

---

**References**

Acknowledgement

To Professor César Tadeu Spadella by a critical review of the study, formatting and proofreading the manuscript in English.

Correspondence:
Paulo Antônio Rodrigues
Alameda das Grevilias, 300
18607-360 Botucatu – SP Brasil
Tel.: (55 14) 3880-1435
paulor@fmb.unesp.br

Received: Oct 24, 2013
Review: Dec 20, 2013
Accepted: Jan 21, 2014
Conflict of interest: none
Financial source: none

1Research performed at Laboratory of Surgical Technique and Experimental Surgery, Faculty of Medicine, Sao Paulo State University (UNESP), Botucatu-SP, Brazil.