Heterozygote Triplets. A Rare Case of Wilkie’s Syndrome

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Summary
Wilkie’s Syndrome is a rare cause of small bowel obstruction that rarely occurs in a familiar setting. We report a rare variant, related to heterozygote triplets in a case that underwent laparoscopic duodenojejunostomy to bypass the obstructed segment. During the patient’s clinical evolution, we collected and discussed preoperative and post-operative studies. In addition, we performed a review of literature.

Key words. Superior mesenteric artery syndrome, Wilkie’s syndrome, laparoscopic duodenojejunostomy.

Trillizos heterocigotos. Un caso raro de síndrome de Wilkie

Resumen
El Síndrome de Wilkie es una causa poco frecuente de obstrucción del intestino delgado que rara vez tiene una presentación familiar. Reportamos una variante inusual, que involucra trillizos heterocigotos en un caso que se sometió a duodeno yeyunostomía laparoscópica para sortear el segmento obstruido. Los estudios preoperatorios y postoperatorios fueron recolectados y discutidos durante la evolución clínica del paciente. Además, se realizó una revisión de la literatura.

Palabras claves. Síndrome de la arteria mesentérica superior, síndrome de Wilkie, duodenoyeyunostomía laparoscópica.

Introduction
Superior mesenteric artery syndrome (SMAS), or Wilkie’s Syndrome, is one of the rare causes of small bowel obstruction.1 SMAS is a disorder characterized by a vascular compression of the duodenum leading to mechanical obstruction.2 Its prevalence ranges from 0.1 to 0.3%.1, 3 SMAS has been described in a familiar setting,
as well as in twins, suggesting a genetic predisposition to this syndrome.\textsuperscript{3,4} In this report, the genetic consultation ruled out genetic or hereditary association of the SMAS. To our knowledge, this is the first case of SMAS in heterozygote triplets.

**Case report**

A 25 years old woman, with diagnosis of chronic non-progressive encephalopathy, was referred to the E.R. with a history of weight loss associated with postprandial vomiting that intensified in the last 10 days. Nutritional consultation was performed with subjective global assessment score being C and body mass index (BMI) was 12.5 at the time of admission. The patient represents one of two triplets affected by this syndrome. Her sister was diagnosed with SMAS and treated at another institution. The third triplet is being studied at present. Upper endoscopy, esophageal manometry, and laboratory tests showed no abnormalities. The barium swallow test showed megaduodenum with a narrowing in the third portion (Figure 1).

A computed tomography angiography (CTA) was performed, confirming an aorto-mesenteric angle of 17.5° with a distance of 8 mm, both signs consistent with SMAS (Figures 2, 3, 4).

Due to intolerance of enteral feeding, parenteral nutrition was started. After six weeks, conservative treatment failed due to the persistence of vomiting. A surgical approach was indicated, performing a laparoscopic duodenojejunostomy (Figures 5, 6, 7).

After twelve weeks, the patient was discharged with oral and parenteral nutrition.

Six months later, the patient’s BMI was 19, presenting a total weight gain of 16 kg. A new CTA was performed, showing an aorto-mesenteric angle of 44.2° and a distance of 12 mm. (Figures 8, 9, 10, 11, 12).
Figure 5. Duodenojejunostomy. Distented stomach and inframesocolic vision

Figure 6. Duodenojejunostomy. Duodenal dissection and duodenoyeyunal anastomosis

Figure 7. Duodenojejunostomy. Side by side anastomosis
Figure 8. Post-surgical CTA. Aorto-mesenteric angle of $44.2^\circ$.

Figure 9. Post-surgical CTA. Aorto-mesenteric distance of $12$ mm.

Figure 10. Post-surgical CTA 3D. Aorto-mesenteric angle of $44.2^\circ$.

Figure 11. CTA pre-surgical aorto-mesenteric angle of $17.5^\circ$ compared with CTA post-surgical of $44.2^\circ$.

Figure 12. CTA 3D pre-surgical aorto-mesenteric angle of $17.5^\circ$ compared with CTA 3D post-surgical of $44.2^\circ$. 
Discussion

First described by Rokitansky in 1842, the SMAS involves the entrapment and obstruction of the third part of the duodenum between the superior mesenteric artery (SMA) and the aorta. In 1927, Wilkie et al. published the largest study on SMAS based on 75 cases in which the final conclusion was that duodenojejunostomy was the treatment of choice.5

SMAS is characterized by the compression of the third portion of the duodenum due to the narrowing of the space between the SMA and the aorta. This phenomenon is primarily attributed to a loss of the mesenteric fat pad, located in that area.2 Fat loss and secondary compression may be precipitated by progressive weight loss in clinical situations like malignancies, dietary disorders, or trauma. This syndrome may also result from surgical procedures that reduce the aorto-mesentric angle, for example, ileo-anal pouch producing caudal traction of the mesentery or scoliosis correcting surgery.6 SMAS may also be seen in association with psychological symptoms.7

The genetic predisposition to its development has been suggested in a report of its occurrence in identical twins; however, this phenomenon is subject to further research.8 Implicated congenital factors such as a short ligament of Treitz and/or a low implantation of the SMA have also been proposed.3

This disorder is a very rare condition and its incidence is considered to range between 0.2-0.3%.3 However, in patients with scoliosis who undergo corrective spinal surgery, its prevalence increases to 2.4%.9 The SMAS mortality rate is estimated to be 33%.10 A total of 400 cases have been reported in the literature in English with a slight prevalence in young female patients.2, 7 SMAS should be considered when persistent vomiting is present, especially after a history of weight loss.1

Patients usually present a long history of unspecific abdominal symptoms: postprandial epigastric pain, nausea, vomiting, anorexia, weight loss, esophageal reflux, and bloating. The discomfort may be relieved by measures that reduce tension on the small bowel mesentery, such as left lateral decubitus, assuming a ‘knee-chest’ position.5 Dehydration and electrolyte abnormalities due to prolonged vomiting are not uncommon.1

For diagnostic purposes, upper gastrointestinal series, CTA, magnetic resonance, angiography, conventional angiography, ultrasonography, and endoscopy have all been described. The clinical diagnosis can be confirmed by radiologic studies in 95% cases.11

The aorta-SMA angle in patients with SMAS has been reported to range from 6-22° and the aorta-SMA distance to be between 2-8mm, compared with 25-60° and 10-28mm respectively in normal population.2, 7, 9, 12-14 CTA with 3D reconstruction is a useful noninvasive diagnostic tool that may aid in confirming duodenal obstruction due to vascular compression.

A multidisciplinary team should approach these patients. The team should include gastroenterologists, interventional radiologists, dietitians, psychologists and surgeons.6

Although there is no consensus on the optimal therapeutic strategy for SMAS, conservative management is usually the first choice which should be administered for at least six weeks.10 Conservative treatment includes gastric decompression, fluid electrolytes imbalance correction, and nutritional support either through total parenteral or nasojejunal feeding. Conservative treatment also focuses on nutritional support aimed at restoration of retroperitoneal fat and weight gain,1 which may restore the SMA-aortic angle.

When conservative treatment fails, surgery is required.15 Different surgical techniques have been described to treat SMAS. Proposed by Strong, the transection of the ligament of Treitz, and subsequent mobilization of the duodenum, was first prescribed.7 Currently, this technique has been abandoned due to poor results. The gastrojejunalostomy, another treatment option for this condition, also proved to be more susceptible to failure since it does not free the duodenal obstruction. Among surgical techniques, the one that has demonstrated greater effectiveness in solving symptoms has been the duodenojejunosoty.5, 16 After the first successful laparoscopic duodenojejunosotomy by Gersin et al. in 1998,17 the procedure became the most common surgical treatment for this syndrome.15, 18 Benefits of laparoscopic duodenojejunosotomy include those of the minimally invasive approach along with better functional results.7, 19

We can conclude that SMAS is a rare condition that requires a high level of suspicion. Patients should be approached by a multidisciplinary team, which leads to better diagnosis and treatment options. Laparoscopic duodenojejunosotomy is the procedure of choice, when conservative treatment fails.

References


