CASE REPORT

ANESTHETIC MANAGEMENT OF A PATIENT WITH SITUS INVERSUS POSTED FOR LAPAROSCOPIC CHOLECYSTECTOMY
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ABSTRACT: Situs inversus totalis is a congenital visceral malrotation anomaly that results from disturbances in establishment of left-right asymmetry and it is characterized by total transposition of thoracic and abdominal viscera, and the predicted incidence is one in 10,000 among the general population. In a patient with situs inversus totalis, not just the diagnosis of any acute abdomen pathology is difficult but equally challenging is the anesthetic management during the respective surgical procedure. We are reporting a patient who had situs inversus totalis and was operated for laparoscopic cholecystectomy under general anesthesia with I-gel, a new supraglottic airway device as an airway conduit. Though the problems related to such patients are mainly of surgical feasibility, an anesthesiologist must be aware of the associated problems of both, situs inversus and the laparoscopy. The present case report lays an emphasis on the potential difficulties during anesthetic management and its various implications. To the best of our knowledge, we report the first case in India of a successful laparoscopic cholecystectomy in a patient with situs inversus totalis with use of I-gel.

KEYWORDS: Cholecystectomy, Dextrocardia, I-gel, Situs inversus.

INTRODUCTION: The first case of situs inversus in humans was reported by Fabricius in 1600.¹ This condition of abnormal visceral rotation was known in animals since the days of Aristotle.² Situs inversus with dextrocardia is also termed situs inversus totalis because the thoracic as well as the abdominal viscera, is a mirror image of the normal anatomy. The incidence of situs inversus totalis has been thought to be 1:5000 to 1:20000.²,³

During the embryological development, a 270-degree clockwise rotation instead of normal 270 degree anti-clockwise of the developing thoraco-abdominal organs results in mirror image positioning of the abdominal and thoracic viscera.⁴ This defect may present difficulties in the diagnosis and management of abdominal pathology due to mirror-image anatomy. The association of situs inversus totalis with syndromes such as Kartagener's syndrome, cardiac anomalies, spleen malformations and other such clinical entities makes the clinical scenario extremely challenging for the concerned anesthesiologist.⁵

We are reporting a case of situs inversus totalis who underwent laparoscopic cholecystectomy, under general anesthesia with I-gel as an airway conduit. The anesthetic considerations and implications associated with such anatomical abnormalities are discussed.

CASE REPORT: A 25 year old male, weighing 35 kg, presented with pain in abdomen for the last 3 days. He was mentally retarded and had tremors since childhood. There was no history suggestive of repeated respiratory tract infections like bronchiectasis and pan sinusitis. Ultrasonography abdomen for the evaluation of pain revealed situs inversus and X-ray chest revealed dextrocardia with fundic
gas shadow on right side [Figure 1]. Computed tomography (CT) thorax and abdomen revealed situs inversus totalis. Echocardiograph revealed normal cardiac parameters with ejection fraction of 62% and dextrocardia. On examination, he had bilateral hand tremors. Airway examination revealed Mallampatti class II. Hemodynamic parameters were within normal limits. He was scheduled to undergo laparoscopic cholecystectomy.

In the preanaesthetic room intravenous access was secured. Inj Pantoprazole 20 mg and inj. Ondansetron 6 mg were given intravenously. In the operation room, ECG (leads, mirror image of normal), NIBP and pulse oximeter were connected. Premedicated with inj. Midazolam 1 mg and inj. Fentanyl 80 mcg.

Patient was induced with inj. Propofol 100 mg and 2% Sevoflurane. Neuromuscular blockade was achieved with inj. Vecuronium 6 mg. Cuff less supraglottic airway device I-gel was inserted, placement confirmed with bilateral air entry on auscultation and ETCO₂ wave form. I-gel was fixed and controlled ventilation was instituted with ventilator settings of tidal volume-12 ml/kg, Respiratory rate-12/min and I: E = 1:2. Ryle’s tube No. 12 was passed through I-gel [Figure 2].

During laparoscopic surgery orientation and ergonomics were altered, surgeon stood on right side and video monitor was shifted to left side of patient. Total operating time was 95 min.

Anesthesia was maintained with O₂: N₂O=60:40, isoflurane 0.6-1% and intermittent doses of inj. Vecuronium. SpO₂ was ranging between 98-99%. ETCO₂ was ranging between 28-36 mmHg. At the end of surgery, when the patient had respiratory attempts, residual neuromuscular blockade was reversed with Inj. Neostigmine 2.5 mg. and inj. Glycopyrolate 0.4 mg. After Ryle’s tube and oral suctioning, I-gel was removed and patient was shifted to recovery room. Perioperative period was uneventful and patient was discharged from hospital on fifth post-operative day.

**DISCUSSION:** Situs inversus totalis is a rare congenital disorder occurring in 0.01% of the population. This entity is considered to have a genetic predisposition that is autosomal recessive with the defect being localized on the long arm of chromosome 14.

It is characterized by the transposition of the major thoracic organs and all the visceral organs of the abdomen to the side opposite to normal position in the body. The liver and gall bladder are located on the left, while the stomach and the spleen are on the right. The normal development requires a 270 degree counterclockwise rotation that yields the normal anatomy. In situs inversus totalis, the 270 degree rotation is in the clockwise direction.

Early embryonic process of determination of normal body situs is complex and probably controlled by several genes. The exact etiology is unclear; however, it is thought to be due to a single autosomal recessive gene of incomplete penetration. The male to female ratio is 1:1 and there is no racial predilection.

In acute abdomen, it is important to be aware of the presence of situs inversus to ensure the correct diagnosis and treatment. Acute appendicitis causes left lower quadrant pain, whereas cholecystitis causes left upper quadrant pain in these patients. CT scanning is the preferred diagnostic modality as it shows the anatomical details.

In this case the patient presented with right upper quadrant pain only and had no definite left upper quadrant pain. It has been noted in 30% of previous reported cases of acute cholecystitis in patients with situs inversus that the pain was felt in the epigastrium alone and in 10% the pain was
localized to the right upper quadrant. The proposed explanation for this is that the central nervous system may not share in the general transposition.

Patients with situs inversus who are scheduled for laparoscopic cholecystectomy should be assessed pre-operatively for any potentially serious cardiac or respiratory abnormalities.

While there is no evidence to suggest that there is an increased risk of bile duct injuries in patients with situs inversus, the orientation and ergonomic challenges may result in an increased operative time. The procedure becomes difficult because of the fact that the anatomy is mirror image of that of routinely seen and most surgeons are right handed.

A case of prolonged paralysis after administration of succinylcholine has been reported earlier in a patient with situs inversus totalis. However we avoided succinylcholine since there was no anticipation of difficult airway. Use of I-gel, which is a cuff less supraglottic airway device with a gastric channel, is well accepted during laparoscopic surgeries.

I-gel has advantages like easier insertion, minimal risk of tissue compression, stability after insertion and smooth awakening. It produces lower hemodynamic instability during placement and avoids stimulating the infraglottic structures. High leak pressure and low peak pressure ensure safe ventilation and decreased the risk of gastric insufflations, even in obese patients (body mass index >35 kg/m²) who were ventilated with 22 cm H₂O peak pressure.

An integrated gastric channel facilitates venting of gas from the stomach and allows for the passing of a nasogastric tube to empty the stomach contents. The positive pressure should be kept below 40 cm of H₂O during intermittent positive pressure ventilation and in our case it did not cross 26 cm of H₂O. Though mucociliary dysfunction leading to increased secretions may be associated with situs inversus, there were no signs and symptoms suggestive of increased mucoserous secretions (history of repeated sinusitis, bronchiectasis) in our case. Hence use of a supraglottic device is justified and there is less stimulation of infraglottic structures which itself can lead to increased secretions and bronchospasm.

In situs inversus, special considerations are:

a. ECG electrodes and defibrillator pads are placed reverse.

b. The association of situs inversus with other syndromes and diseases such as Kartagener's syndrome, mucociliary dysfunction, airway anomalies, etc., which may predispose the patient to numerous varieties of airway difficulties and pulmonary infections that can have considerable implications during induction of anesthesia and intubation.

c. The syndrome is associated with numerous cardiac anomalies such as atrial septal defects, ventricular septal defects, transposition of great vessels, absent coronary sinus, double-outlet right ventricle, total pulmonary anomalous venous defect and pulmonary valve stenosis either singly or in combination.

d. In case of inversion of great vessels preference, should be given to left internal jugular vein for cannulation (to avoid thoracic duct and to ensure direct access to right atrium).

e. Main stem intubation occurs often to the left.

f. The spinal deformities like split cord, spina bifida, meningomyelocele, scoliosis, etc. have been described in the literature, and one has to evaluate the patient carefully if any surgery is planned under neuraxial anesthesia.
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g. In thoracic surgery the anatomy of the bronchi should be considered before selecting a double-lumen tube. Insertion of a double-lumen tube will pose a multitude of challenges, and the successful intubation and separation of lungs cannot be accomplished without the aid of fibreoptic bronchoscope. The transposition of the thoracic viscera also alters the various anatomical landmarks, and one has to be well acquainted with ultrasound-guided procedures if in case a need arises for invasive central venous cannulation and brachial plexus blockade.

h. Uterine displacement in a parturient is logically to the right in these patients.15

i. Situs inversus in Kartagener’s syndrome is invariably associated with mucociliary dysfunction.

Primary ciliary dyskinesia is present in 25% of the patients with situs inversus totalis with an increased probability of developing respiratory complications. Therefore, moist and filtered mixture of gases should be administered during mechanical ventilation. The role of bronchodilators, chest physiotherapy, postural drainage, antibiotics and incentive spirometry cannot be underestimated and is mandatory in optimizing the pulmonary status before any surgical procedure.

CONCLUSION: To conclude with meticulous planning and care, patients with Situs inversus can be successfully managed. A Supraglottic device is a good option if situs inversus is not associated with Kartagener’s syndrome.

REFERENCES:


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Figure 1: X-ray chest showing dextrocardia with fundic gas shadow on right side

Figure 2: I-gel with Ryle’s tube in situ
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