Pancreatic and neuroendocrine free papers

**OC-072 USE OF A NOVEL SELF-EXPANDING METAL STENT TO ALLOW FOR ENDOSCOPIC DRAINAGE AND NECROSECTOMY OF PANCREATIC FLUID COLLECTIONS**

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**Introduction** Post-inflammatory peri-pancreatic fluid collections are frequent sequelae of severe acute pancreatitis. Collections are at risk of suppurative infection complicated by pancreatic necrosis. Over the last decade there has been an increasing emphasis on minimally invasive drainage procedures, including EUS-guided cyst-gastrostomy, and these approaches seem to be associated with lower morbidity and mortality. Access to the necrosis cavity has however been severely limited by having to maintain the tract with small diameter plastic stents. Recently, a novel flanged fully covered self-expanding metal stent (FCSEMS; NAGI stent, Taewoong Medical, Korea) has been developed to allow for better drainage of infected necrosis and easier endoscopic access into the cavity.

**Setting** A non-randomised prospective multicentre phase II study to determine the safety and efficacy of FCSEMS endoscopic cyst-gastrostomy in the management of complex/infected pancreatic fluid collections.

**Methods** Patients were included if they had evidence of a pancreatic fluid collection which was deemed to be amenable for EUS-guided drainage after discussion at a HPB multidisciplinary meeting. Patients selected for EUS-guided drainage had cross sectional imaging (MR or CT) performed within 2 weeks of the procedure and then an EUS assessment was made of the necrotic component. The collection was punctured using a cystotome and the FCSEMS inserted over a guidewire with fluoroscopic control. Repeat procedures were performed as necessary.

**Results** A total of 11 patients (8 male, 3 female) were included in the study. Median age was 57.3 years. The aetiology of the collection was gallstones in 6 patients, idiopathic in 3, ischaemic in 1 and drug-induced in 1. Ten patients had evidence of at least 30% necrosis within the collection. Mean diameter of the collection was 15 cm and EUS-guided puncture was initially performed in all patients. The tract was dilated with a balloon in 6 patients. Stent insertion was either with a 20 mm (7 patients) or 30 mm (4 patients) length FCSEMS. Ten patients underwent endoscopic necrosectomy, with a median of 3 procedures (range 1–10). Significant reduction in the size of collection was achieved in all patients. Adverse events included stent migration in 3 (2 spontaneously and 1 during necrosectomy). Two patients died of complications of severe acute pancreatitis.

**Conclusion** FCSEMS insertion is feasible and safe for drainage of pancreatic fluid collections. It allows repeated through the stent necrosectomy procedures and appears to be a major advance in the management of infected pancreatic necrosis.

**Disclosure of Interest** None Declared.

**OC-073 USE OF BOTULINUM TOxin TO PREDICT MANOMETRY RESULTS IN TYPE III SPHINCTER OF ODDI DYSFUNCTION; A RETROSPECTIVE SINGLE CENTRE REVIEW**

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**Introduction** Management of type III sphincter of Oddi dysfunction (SOD) remains controversial. A recent large multi centre study of manometry and sphincterotomy in type III SOD (EPISO-D) found that sphincterotomy was no more effective than sham treatment. Botulinum toxin (botox) injection to the papilla has been shown to be safe and lead to improvement in symptoms, it may also predict response to sphincterotomy.

This study reviewed use of botox in patients with type III within a single tertiary centre to guide decision making.

**Methods** The endoscopy unit database was searched for cases between January 2008 to August 2013 who received botox for SOD. Records were reviewed to identify those who had type 3 SOD as per Rome 3 criteria. Response to botox was graded as no response, partial (reduction but not resolution of pain) or complete response. Complications, manometry and sphincterotomy results were recorded.

**Results** 63 patients had botox injection for SOD 46 were classified as type III and formed the study group. All received 100IU of botox. Following the procedure 3 of 46 patients required overnight observation for abdominal pain, there were no cases of pancreatitis. 14 patients had no response to botox, 7 partial response, 24 a complete response, 1 did not attend follow up.

Of those that had a complete response 14 patients proceeded to manometry; reasons not to proceed included failure to attend follow up (2) patient declined treatment (1) other co-morbidities (1). One underwent a second botox procedure with no relief in symptoms and was not offered further treatment, 5 patients (21%) were not offered manometry. Of those that proceeded to manometry, 10 (71%) had elevated pressures 8 biliary, 2 pancreatic. All proceeded to sphincterotomy with good response in 9 (64%) of Botox responders, 3 of the 7 patients who partially responded proceeded to manometry; 1 of these had elevated biliary pressures. 3 patients (18%) had an episode of pancreatitis following manometry.

**Conclusion** Response to botox appears to show moderate correlation with abnormal manometry findings and response to sphincterotomy. ERCP and manometry was associated with a significant risk of pancreatitis. Randomised sham controlled studies are required to ascertain whether a response to botox can accurately select patients who benefit from a sustained response to sphincterotomy.
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