INTRODUCTION

The oesophago-gastric junction (OGJ) is located between the oesophagus and the stomach. Its dynamic action is dependant upon the intrinsic and extrinsic effects of its own anatomical structure and its position with respect to the surrounding organs in this region of the body. Despite its relationship to diseased states of the oesophagus, no objective measurement technique has been developed to quantify the normal or abnormal functioning of the OGJ. It is generally accepted that sphincters divide the gut into functional segments. They are characterized by a resting tension, generally known as muscle tone which is greater than in the two adjacent segments. It is common nowadays to give the term sphincter a functional meaning rather than its original anatomical meaning[1]. The sphincter at the lower end of the oesophagus, the lower oesophageal sphincter (LOS), forms part of the OGJ structure. At the OGJ the oesophagus opens into the stomach at the cardia. It is important that backflow of stomach secretions into the oesophagus is controlled at the OGJ. The OGJ guards against gastric acid moving up into the oesophagus, opening only transiently to allow passage of the swallowed food into the stomach. Swallowing is a demanding reflex activity[2]. The diaphragmatic hiatus, through which the oesophagus passes just at the OGJ, has a role in this valvular mechanism[3].

Several other structures are important in maintaining a barrier in the OGJ against the back flow of food and the stomach acids into the oesophagus, known as the anti-reflux barrier. The intrinsic muscles of the distal oesophagus and the sling fibres of the proximal stomach make up the internal mechanism of the LOS[4]. The muscles of the diaphragm that connect to the OGJ make up the crural diaphragm and this constitutes the external LOS mechanism. The tissue that connects the distal oesophagus to the crural diaphragm is known as the phreno-oesophageal ligament[5].

The OGJ can be distinguished from the body of the oesophagus by its behaviour pattern characteristic. There is an increase in the tone of the circular muscle in this region. The sphincter relaxes in response to a swallow and this usually precedes the arrival of a contraction wave travelling down the oesophagus. This phase of relaxation is followed by a short-lasting elevation of pressure above resting values[6]. In recent years it has been identified that the sphincter sometimes relaxes even when a swallow does not occur. These relaxations are know as transient lower oesophageal sphincter relaxations (tLOSRs)[7]. Other factors contribute to the role of the OGJ in controlling the flow of solids, liquids and gases between the oesophagus and the stomach. Pressures in the abdominal and thoracic cavities are involved as well as the exact intra-abdominal...
location of the junction. Time and posture are also important factors.

THE ROLE OF THE OGJ IN DISEASE

Disorders of the OGj in general relate to two diseased states. Conditions where the disease may be related to an OGj not closing normally are generally referred to as gastro-oesophageal reflux disease (GORD) and conditions where the OGj fails to relax or open, which is referred to as achalasia.

Gastro-oesophageal reflux disease

GORD symptoms are very common in the general population affecting up to 44% of the US population at least once a month and 20% of the population once a week. It has been shown that patients suffering from GORD have a lower health related quality of life than patients with angina and mild heart failure. There is solid evidence that incompetence or dysfunction of the OGj is a primary determinant of GORD. Three mechanisms at the OGj lead to an increased number of acid reflux events associated with GORD: t:LOSRs, free reflux during periods of low LOS pressure or deglutitive relaxation of the LOS. Much more needs to be learned about the dynamic action of the OGj if we are to fully understand its role in GORD. Current treatment of GORD is with acid suppressing drugs such as proton pump inhibitors but for those refractory to pharmacological treatment surgery is often recommended. The surgery of choice for GORD is the laparoscopic Nissen fundoplication.

Achalasia

Achalasia is an oesophageal disorder of unknown cause characterized by aperistalsis of the oesophageal body and impaired relaxation of the lower oesophageal sphincter. It is a relatively rare disease and in Western countries the annual incidence is in the range 0.5 to 8 cases per 100000 people. It can be difficult to make a diagnosis in some patients because of vague symptoms. Symptoms can emerge slowly and patients adapt. The diagnosis can be confirmed by endoscopy and manometry. Developing a device that can profile the OGj will contribute greatly to understanding the disfunction of this disorder, which often manifests as a permanently constricted or toned sphincter. Surgical performance of an oesophageal myotomy is the treatment of choice for some severe cases of achalasia. Currently the laparoscopic oesophageal Heller-Nissen procedure is commonly used.

BIOMECHANICS OF THE OGJ

Biomechanics are the laws of mechanics applied to biology. There is an increasing recognition of the importance of tissue biomechanical properties for function, e.g. organ geometry, resting conditions and elasticity is determinants of flow in the organ and sensory responses in the neural circuits. When dealing with biological tissues, solid mechanical parameters such as the relation between forces (membrane stress resultant, tension) and deformation (often expressed as a strain) are important to evaluate. The mechanical properties of the OGj are important determinants of its distensibility and competence. In vitro studies have shown potential in measuring biomechanical parameters in the OGj.

Very few methods exist for assessment of mechanical properties in the GI tract in vivo. One method is impedance planimetry from where the cross-sectional area (CSA) and pressure data can be gathered and the wall tension and strain in the organ can be estimated. Other methods rely on the combination of bag distension and manometry with high resolution imaging techniques such as 3D-ultrasound of the gastric antrum or MRI of the rectum. Manometry is of course a simple biomechanical measurement. Its use will be mentioned again later.

METHODS OF EVALUATING OGJ FUNCTION

In 1964 Mann et al. suggested that the zone separating the lower end of the oesophagus from the stomach can be recognized in two different ways, by its distinguishing anatomic features and its unique patterns of behaviour. A huge amount of knowledge about the anatomy of the OGj has been assembled over the last 40 years, but much of our limited understanding of the behaviour has been related to manometric studies using single or multiple pressure recordings or a sleeve placed in the OGj. While manometry provides useful information on motility within the lumens of the digestive tract, it may be more restrictive in what it can tell us about sphincteric regions such as the OGj and their dynamics. Harris and Pope found that sphincters do not necessarily need to squeeze or contract tightly to be competent and therefore resistance to distension by measurement of radial force should be the prime determinant of sphincteric strength. Measurement of luminal pressure, however, is not a direct measure of muscle tone and tension. It is merely a proxy of changes in force generated by the surrounding tissues.

No clinically acceptable quantitative method exists to directly measure OGj mechanical function. Intraluminal pressure recordings is the only objective measurement technique for determining the competence of the LOS, but it is usually the more indirect method of measuring acid reflux in the oesophagus which is the key determinant of failure of the OGj to inhibit flow. Thus, pH-metry rather than manometry is the recommended test for GORD.

Usually in medicine dynamic diagnostic imaging methods can be used to visualize and evaluate anatomical structures similar to the OGj. Currently none of these methods give sufficient dynamic detail to contribute to the routine diagnosis and treatment of diseases of the OGj. Recent developments in material sciences and technology have allowed the OGj’s resistance to distension to be studied using parameters such as distensibility, opening patterns and dimensions and flow rate. It is becoming apparent that these measurements may be useful in determining the success of therapeutic interventions such as surgery. However what is needed is a technique with high spatial and temporal resolution for determination of sphincter geometry and mechanical factors.
**Impedance planimetry**

**Principle of Impedance Planimetry:** Impedance planimetry (IP) is a technique established by Gregersen and coworkers for performing bag distensions in the GI tract (Figure 1). It has been used extensively to provide a single measure of CSA together with bag pressure in animal and human studies using electrical measurements inside the liquid filled bag. Hence the term impedance planimetry. If we fill a polyurethane bag with saline and pass a constant current through it, the CSA is inversely proportional to the voltage across the sensing electrodes. If all other parameters are kept constant then the impedance becomes related to the CSA of the bag between the sensing electrodes.

\[
V = \frac{1}{\text{CSA}}
\]

Many studies have shown the usefulness and reliability of IP in the oesophagus of healthy volunteers and patients. Rather than analyzing distension data based on pressure-volume relationships as is the case with the “Barostat”, IP enables data on the CSA of the bag during distension to be gathered. CSA shows a better relationship to the diametrical changes in the alimentary lumen than volume. However, by far most studies have been done using a single CSA recording inside the bag rather than obtaining a series of CSAs. Due to previous limitations, it has only been used in a few studies in sphincter regions. These limitations related to the difficulties placing the measurement electrodes at the point of most interest and the fact that distending a bag in a high-pressure zone tends to displace the bag during the measurements.

**Multiple measurements:** A recent pilot study in the porcine rectum demonstrates how an IP probe placed in the rectum with 5 sensing electrode pairs with 2mm between pairs and 2cms between sets can detect contractile events at different levels along the longitudinal axis of the gut (Figure 2). This multi-electrode technique demonstrated how a balloon with a conducting liquid and a constant current source set up across two excitation electrodes can measure multiple CSAs by measuring the voltage across multiple electrode pairs within the electric field.

There are a number of potential sources of error in multiple measurement IP systems. Electrodes placed too close together will interfere with each other. The number of wires required to make the electrical connections goes up and therefore the potential for interference, liquid leaking into the probe and bad electrical connections is much.

**Manometry**

Manometry has been used for more than 50 years to monitor motility in the gastrointestinal tract. Since 1977 reliable methods have been available to obtain accurate measurements of force of closure of the lumens of the intestine when contracting. These system often use reliable perfused hydraulic-capillary systems where pressure at a sensor outside the body can be used to accurately measure pressure at a point along a catheter via a low flow rate perfused channel. This system has been used extensively to measure force via a pressure measurement in lumens and in liquid filled balloons. Errors in manometric readings can be caused by air bubbles in the capillary lines, occlusion of the measurement site and the compliance of the recording system but in general if well set up they produce accurate readings. In more recent years catheters with solid state transducers are becoming more popular for pressure reading and these overcome some of the measurement problems.

Using manometry to determine the competence of sphincteric or valvular regions in the lumens of the digestive tract is however dubious. Dent recognized the problem of positioning the measurement point 30 years ago and proposed a solution and others tried to follow suit. But pressure is a measurement of force over area and this may not be a proper measure of the function of a valvular region. As suggested by Pope and Harris we cannot assume that just because a sphincter is closed that it has a contract force or in the terminology of manometry, occlusion of the measurement site and the compliance of the recording system but in general if well set up they produce accurate readings. In more recent years catheters with solid state transducers are becoming more popular for pressure reading and these overcome some of the measurement problems.

**Calculation of wall tension**

**Law of laplace:** For a membraneous cylinder with transmural pressure \( P \) and radius \( r \) the circumferential tension \( T \) can be given by:

\[
Pr = T
\]

Use of Laplace's law is based on the assumption that the geometry of the segment is circular and cylindrical and that the wall thickness is very small.
Measurements using multi-electrode Impedance planimetry in the oesophagogastric junction

We have demonstrated that a multi-electrode probe with a cylindrical bag mounted and filled with saline can be placed in the OGJ and that data on a number of CSAs in this area can be measured. Figure 3 shows results of 3 CSA measurements 8 mm apart measured from distension of a bag in the OGJ of a healthy volunteer. The plot shows how the most distal CSA (D) measurement increases more rapidly than the middle CSA (M) as a constant volume of saline is filled into the bag. For the most proximal CSA (P) the converse is true. This suggests that distally the OGJ is more compliant than proximally. This work also demonstrated for the first time how wall tension in the OGJ can be estimated from simple distension measurements. This preliminary version of the multi-electrode technique demonstrated that a probe can be placed in the OGJ and that data can be successfully collected. However, there were limitations such as the fact that there were only three measurements made. A narrowing region such as a sphincter cannot be profiled very well with three CSA measurements unless the exact position of the sphincter is located and maintained. Another limitation is the outer diameter of the catheter on which the bag is mounted. In this case it was 4.5 mm. If the sphincter is to be distended from as close to the closed position as possible then a smaller catheter needs to be used.

Figure 4 shows how a probe capable of measuring 8 CSAs could be used to profile a sphincter region. It is quite obvious that a larger number of sensing pairs could be more useful in providing a geometric profile. Consequently a more elaborate probe for measurement of up to 8 CSAs in the region of the oesophago-gastric junction was developed. This probe can measure CSAs at 4 mm intervals through the OGJ thus spanning a measurement range of 28 mm.

Figure 5 shows the data from the 8 CSAs during a bag distension at a fixed flow rate of 40 mL/min to a volume of 60 mL. The information in this format can be difficult to decipher. A probe to make these measurements has been validated and tested in a healthy control subject. Since the OGJ is such a dynamic organ it is its functional activity which can be of more interest. Hence it is possible to generate an illustration of the geometric changes in the OGJ.

If we introduce the idea of demonstrating the CSA changes dynamically as reconstructed data we introduce a new concept known as the functional lumen imaging probe (FLIP). The concept suggests that by gathering data on the dynamic changes in the geometry of the OGJ during normal physiological changes we can represent these changes by recreating the geometry in a three-dimensional image. The FLIP technique can be used to monitor geometric changes in the behaviour of the OGJ during bag distension as can be seen in Figure 6. Here the dynamic data is reconstructed demonstrating the effect of the balloon distension on the amount of OGJ opening.
FLIP can quantify changes in compliance of the OGJ after endoscopic suturing at the OGJ in a pig model\textsuperscript{[63]}. Figure 7 indicates how the geometric reconstruction can be used to indicate changes in the profile of the OGJ after endoscopic plication. The pressure is higher, the CSA is smaller and subsequently the wall tension has increased after the plications are inserted. These measurements indicate that the OGJ becomes less compliant after plication. No direct measure of the effects of these therapies on the OGJ has been demonstrated before. Recent studies have expressed concerns about the efficacy and durability of these procedures\textsuperscript{[52-60]}. FLIP may in fact reveal new information on the mechanism of action behind these new reflux therapies. In effect it suggests that endoscopic suturing augments the compliance of the OGJ to make it less compliant. Perhaps patients could now be selected better for these procedures if their compliance can be confirmed to be abnormal in the first place and maybe the success of the procedure could be determined directly at the time of the procedure so for example more stitches can be added if necessary.

Since GORD patients are such a large group perhaps this is the most important area for FLIP to be developed. The probe can distinguish between patients for reflux surgery and normal controls. These are of course the extreme group of patients. Further studies are of course necessary on larger cohorts of patients and controls.

We have recorded with the FLIP the compliance of the OGJ intra-operatively during oesophageal myotomy for achalasia (as yet unpublished findings by the authors). FLIP offers a more sensitive and possibly a better tool than manometry and may be used for evaluating effects of surgery during the procedure as opposed to using manometry more as a blunt instrument before and after as used by Arain et al\textsuperscript{[60]}. FLIP could provide a valuable new tool for the oesophageal surgeon.

**Summary**

Work already carried out shows for the first time that CSA and pressure measurements in the OGJ can be used to measure distensibility by an entirely new technique. It demonstrates how the already established technique of impedance planimetry can be used to measure geometry in vivo in the human oesophagus. It also shows that there is validity of these new measurements, their appropriateness for use in sphincteric regions in the body and their comparable accuracy has been shown with ultrasound. It has been indicated how the measurement could be used practically. Safety and efficacy has been demonstrated. The work also demonstrates how this new measure could be used as a functional imaging technique. Preliminary studies have shown potential applications of the probe. It could prove to be a very valuable tool in endoscopic therapies. It can distinguish between normal and a stretched OGJs in live pigs.

As a resistance to distension test this better describes the sphincteric action of the OGJ than tonic contraction. Even advanced manometry uses such as volume vector analysis, the Dent Sleeve or the Sphinctometer all rely on the principal that there is a force of contraction always present when the sphincter is closed\textsuperscript{[59,60,67-70]}. With FLIP this assumption is not made and it presents more of a challenge test to the sphincter. The technique also represents a more direct technique to evaluate the OGJ using opening patterns with manometry and intraluminal impedance\textsuperscript{[71]}. Some other techniques for measurement of resistance to distension by other investigators are promising and confirm that the principals describe here have great potential for merit. However these other approaches are limited in a number of ways. The technique described by Pandolfini, although it is similar in its measurement design to FLIP, it requires fluoroscopic imaging\textsuperscript{[32,33,35]}. It also depends on the pressure being controlled. Pressure control is difficult to achieve by increasing and reducing volume through a narrow catheter. Using pressure-volume data from the barostat to quantify distensibility does not a good basis in the application of the physical sciences. The bag on a barostat can expand in all directions we cannot be sure that an increase in the volume is reflected by a change in the geometry of the OGJ. This suggest that using this technique is of limited use\textsuperscript{[54]}. Our findings in patients with achalasia confirmed this.

**Perspectives**

FLIP represents the first validated data on a probe capable of measuring OGJ compliance in a simple and time-saving procedure. It represents the opportunity to define the exact role of the OGJ in dysmotility and disease. As a new technique rather than representing a completed work it opens up a new door and perspectives on the junction whose function has been the subject of debate for many years. Future work and improvements in electronics and material sciences will mean that the resolution of the FLIP can be increased dramatically. Design materials have been carefully chosen to minimize signal error. The lower oesophageal sphincter or the oesophagogastriastic junction are often cited as being the prime causes of reflux disease, yet despite this very few research groups are investigating new ways to evaluate this dynamic structure in the gastrointestinal tract\textsuperscript{[32,34]}

Much work has been done in extending manometric measurements enhancing pH measurement techniques to further evaluate the effects of acid in the oesophagus\textsuperscript{[72]} and developing new techniques for measuring bolus transit
both down to the stomach and back up into the oesophagus using intra-luminal impedance\cite{1}. However, none of these techniques have overcome the fundamental question of how the OJG function can be quantified.

FLIP may provide the answer to this question. In a way trialing the probe in the OJG was probably the most challenging and perhaps experiments in another sphincteric region would have been more straightforward where the dynamic activity would be slower. FLIP may have a role in determining biomechanically the root cause of diseases. It can be used for evaluating therapy for example endoscopic and surgical therapy for reflux. It may be possible to use it intra-operatively during Nissen fundoplication, for example where the tension of the wrap is extremely important. It may also be possible to explore other sphincters in the body with FLIP so that the relationship between their biomechanical properties and their relationship to disease can be further defined.

REFERENCES

1. Liebermann-Meffert D. What anatomic structures are undoubtedly responsible for gastroesophageal competence? In: Guil I, Galmiche J, Jamieson G, Scarpignato C, eds. the esophagogastric junction. London: John Libbey Eurotext, 1998: 3-6
2. Goyal RK, Prasad M, Chang HY. Functional anatomy and physiology of swallowing and esophageal motility. In: The Esophagus. Castell DO, Richter JE eds. Philadelphia: Lippincott Williams & Wilkins, 2004: 1-36
3. Boyle JT, Altschuler SM, Nixon TE, Tuchman DN, Pack AI, Cohen S. Role of the diaphragm in the genesis of lower esophageal sphincter incompetence. In: Biomechanics Motion, Flow, Stress and Growth New York: Springer-Verlag, 1990: 499-500
4. Gergersen H, Givorsen IM, Rasmussen LM, Tettrup A. Biomechanical wall properties and collagen content in the partially obstructed opossum esophagus. Gastroenterology 1992; 103: 1547-1551
5. Madsen G, Tettrup A, Andersen KE, Forman A. Mechanical effects of some prostanooids on isolated human esophageal submucosal veins. Pharmacol Toxicol 1992; 71: 330-334
6. Biancani P, Barwick K, Selling J, McCallum R. Effects of acute experimental esophagitis on mechanical properties of the lower esophageal sphincter. Gastroenterology 1984; 87: 8-16
7. Cu A, Murray JA, Conklin JL. Effects of calcitonin gene-related peptide on opossum esophageal smooth muscle. Gastroenterology 1997; 113: 514-520
8. Leichus LS, Thomas RM, Murray JA, Conklin JL. Effects of oxygen radicals and radical scavenging on opossum lower esophageal sphincter. Dig Dis Sci 1995; 40: 1926-1932
9. Gergersen H, Barlow J, Thompson D. Development of a computer-controlled tensiometer for real-time measurements of tension in tubular organs. Neurogastroenterol Motil 1999; 11: 109-118
10. Gergersen H, Gilja OH, Hausken T, Heidmal A, Gao C, Matre K, Ødegård S, Berstad A. Mechanical properties in the human gastric antrum using B-mode ultrasonography and antral distension. Am J Physiol Gastrointest Liver Physiol 2002; 283: C668-C735
11. Frokjaer JB, McMahon BP, Liao DH, Dreves AM, Gergersen H. Modelling of the 3D geometry in the human rectum based on magnetic resonance imaging (MRI). Gastroenterology 2004; 126: A202-A203
12. Harris LD, Pope CE 2nd. “Squeeze” Vs. Resistance: An Evaluation of the Mechanism of Sphincter Competence. J Clin Invest 1984; 43: 2272-2277
13. Gergersen H, Christensen J. Gastrointestinal tone. Neurogastroenterol Motil 2000; 12: 501-508
14. Pandolfini JE, Kahlilas PJ. AGA technical review on the clinical use of esophageal manometry. Gastroenterology 2005; 128: 209-224
15. Pandolfini JE, Shi G, Curry J, Joehl RJ, Brasseur JG, Kahlilas PJ. Esophageal manometry. Lower esophageal sphincter function and distensibility: a factor contributing to sphincter incompetence. Am J Physiol Gastrointest Liver Physiol 2002; 282: G1052-G1058
16. Pandolfini JE, Shi G, Trueworthy B, Kahlilas PJ. Esophageal peristalsis opening during relaxation distinguishes nonhernia reflux patients, hernia patients, and normal subjects. Gastroenterology 2005; 128: 1018-1024
17. Shaker R, Bardan E, Gu C, Massey BT, Sanders T, Kern MK, Hoffman RG, Hogan WJ. Effect of lower esophageal sphincter tone and cranial diaphragm contraction on distensibility of the gastroesophageal junction in humans. Am J Physiol Gastrointest Liver Physiol 2004; 287: G815-G821
18. Pandolfini JE, Curry J, Shi G, Joehl RJ, Brasseur JG, Kahlilas PJ. Restoration of normal distensive characteristics of the esophagogastric junction after fundoplication. Ann Surg 2005; 242: 43-48
19. Gergersen H. Biomechanics of the Gastrointestinal Tract. Lon-
Rao SS, Hayek B, Summers RW. Impedance planimetry: an integrated approach for assessing sensory, active, and passive biomechanical properties of the human esophagus. Am J Gastroenterol 1995; 90: 451-458

Pedersen J, Reddy H, Funch-Jensen P, Arendt-Nielsen L, Gregersen H, Drewes AM. Cold and heat pain assessment of the human esophagus after experimental sensitisation with acid. Pain 2004; 110: 393-399

Andersen IS, Gregersen H, Buntzen S, Djurhuus JC, Laurberg S. New probe for the measurement of dynamic changes in the rectum. Neurogastroenterol Motil 2004; 16: 99-105

Assentoft JE, Gregersen H, O'Brien WD Jr. Determination of biomechanical properties in guinea pig esophagus by means of high frequency ultrasound and impedance planimetry. Dig Dis Sci 2000; 45: 1260-1266

Barlow JD, Gregersen H, Thompson DG. Identification of the biomechanical factors associated with the perception of distension in the human esophagus. Am J Physiol Gastrointest Liver Physiol 2002; 282: G683-G689

Drewes AM, Reddy H, Staahl C, Pedersen J, Funch-Jensen P, Arendt-Nielsen L, Gregersen H. Sensory-motor responses to mechanical stimulation of the esophagus after sensitization with acid. World J Gastroenterol 2005; 11: 4367-4374

Gregersen H, Djurhuus JC. Impedance planimetry: a new approach to biomechanical intestinal wall properties. Dig Dis Sci 1991; 9: 332-340

Gregersen H, Orvar K, Christensen J. Biomechanical properties of duodenal wall and duodenal tone during phase I and phase II of the MMC. Am J Physiol 1992; 263: G795-G801

Gregersen H, Christiansen LL. Pressure-cross-sectional area relations and elasticity in the rabbit esophagus in vivo. Digestion 1996; 57: 174-179

Gregersen H, Vinter-Jensen L, Juhl CO, Dajani EZ. Impedance planimetric characterization of the distal esophagus in the Goettingen minipig. J Biomech 1996; 29: 63-68

Jørgensen CS, Dall FH, Storkholm J, Jensen SL, Gregersen H. Elastic properties of the isolated perfused porcine duodenum. Dig Dis Sci 1991; 9: 401-407

Juhl CO, Vinter-Jensen L, Djurhuus JC, Gregersen H, Dajani EZ. Biomechanical properties of the esophagus damaged by endoscopic sclerotherapy. An impedance planimetric study in minipigs. Scand J Gastroenterol 1994; 29: 867-873

McMahon BP, Frokjaer JB, Drewes AM, Gregersen H. A new measurement of oesophageo-gastric junction competence. A new technique for continuous sphincter pressure monitoring with sphinctometer: in vitro and in vivo studies. Am J Physiol 1999; 277: G57-G68

Petersen JA, Djurhuus C, Koff J, Vinter-Jensen L, Gregersen H. Endoscopic sclerotherapy in porcine esophagus changes luminal cross-sectional area and wall distensibility dose- and time-dependently. Dig Dis Sci 1998; 43: 521-528

Rao SS, Gregersen H, Hayek B, Summers RW, Christensen J. Unexplained chest pain: the hypersensitive, hyperreactive, and poorly compliant esophagus. Ann Intern Med 1996; 124: 950-958

Rao SS, Hayek B, Mudipalli R, Gregersen H. Does esophageal function vary at the striated and smooth muscle segments in functional chest pain? Am J Gastroenterol 2002; 97: 2201-2207

Villadsen GE, Pedersen J, Reddy H, Funch-Jensen P, Arendt-Nielsen L, Gregersen H, Drewes AM. Cold and heat pain assessment of the human esophagus after experimental sensitisation with acid. Pain 2004; 110: 393-399

Villadsen GE, Storkholm J, Zachariae H, Hendel L, Bendtsen F, Gregersen H. Oesophageal pressure-cross-sectional area distributions and secondary peristalsis in relation to subclassification of systemic sclerosis. Neurogastroenterol Motil 2001; 13: 199-210

Patel RS, Rao SS. Biomechanical and sensory parameters of the human esophagus at four levels. Am J Physiol 1998; 275: G187-G191

Sørensen SM, Gregersen H, Sørensen S, Djurhuus JC. Spontaneous anorectal pressure activity. Evidence of internal anal sphincter contractions in response to rectal pressure waves. Scand J Gastroenterol 1989; 24: 115-200

McMahon BP, Frokjaer JB, Liao D, Kunwald P, Drewes AM, Gregersen H. A new technique for evaluating sphincter function in visceral organs: application of the functional lumen imaging probe (FLIP) for the evaluation of the oesophago-gastric junction. Physiol Meas 2005; 26: 823-836

Andorffer RC, Stej J, Dodds WJ, Linehan JH, Hogan WJ. Improved infusion system for intraluminal esophageal manometry. Gastroenterology 1977; 73: 23-27

Dent J. A new technique for continuous sphincter pressure measurement. Gastroenterology 1976; 71: 263-267

Gottle DC, Barham CP, Miller R, Arnold R, Alderson D. The sphinctometer: a new device for measurement of lower oesophageal sphincter function. Br J Surg 1991; 78: 933-935

McMahon BP, Bergstrom M, Frokjaer JB, Hansson L, Drewes AM, Gregersen H, Park PO. Use of FLIP in determining changes in the geometry of the oesophago-gastric junction with endoscopic suturing in a porcine model. Gut 2005; 54(Suppl VII): A104

Schiefke I, Zabel-Langhennig A, Neumann S, Feisthammel J, Moessner J, Caca K. Long term failure of endoscopic gastrostomy (EndoCinch). Gut 2005; 54: 752-758

Arts J, Lerut T, Rutgeerts P, Sifrim D, Janssens J, Tack J. A one-year follow-up study of endoluminal gastrostomy (Endocinch) in GERD patients refractory to proton pump inhibitor therapy. Dig Dis Sci 2005; 50: 351-356

Schiefke I, Soeder H, Zabel-Langhennig A, Teich N, Neumann S, Borte G, Mesnger J, Caca K. Endoluminal gastrostomy: what are the predictors of outcome? Scand J Gastroenterol 2004; 39: 1296-1303

Mahmood Z, McMahon BP, Arfin Q, Byrne PJ, Reynolds JV, Murphy EM, Weir DG. Endocinch therapy for gastro-oesophageal reflux disease: a one year prospective follow up. Gut 2005; 52: 34-39

Arain MA, Peters JH, Tahmankar AP, Portale G, Almobgy G, DeMeester SR, Crookes PF, Hagen JA, Bremner CG, DeMeester TR. Preoperative lower oesophageal sphincter pressure affects outcome of laparoscopic esophageal myotomy for achalasia. J Gastrointest Surg 2004; 8: 328-334

Kahriasis PJ, Lin S, Chen J, Manka M. The effect of hiatus hernia on gastro-oesophageal junction pressure. Gut 1999; 44: 476-482

Strathof JW, Lüchtenborg M, Maslee CA. Comparison of two techniques for lower oesophageal sphincter manometry: sleeve and sphinctometer. Neurogastroenterol Motil 2004; 16: 265-268

Pehlivannov N, Liu J, Arora T, Yamamoto Y, Mittal RK. Lower oesophageal sphincter monitoring with sphinctometer: in vitro and in vivo studies. Am J Physiol 1999; 277: G57-G584

Trudgill NJ, Riley SA. Monitoring the lower oesophageal sphincter: sphinctometer or sleeve? Neurogastroenterol Motil 1999; 11: 173-178

Pandolfino JE, Shi G, Zhang Q, Ghosh S, Bresser JG, Kahri- las PJ. Measuring EGJ opening patterns using high resolution intraluminal impedance. Neurogastroenterol Motil 2005; 17: 200-206

Pandolfino JE, Richter JE, Ours T, Guardino JM, Chapman J, Kahrilas PJ. Ambulatory esophageal pH monitoring using a wireless system. Am J Gastroenterol 2003; 98: 740-749