Neuromuscular function of the human lower oesophageal sphincter in reflux disease and Barrett’s oesophagus

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Abstract

Background—Columnar lined (Barrett’s) oesophagus is often considered a sequel to chronic severe reflux disease. Aberrant lower oesophageal sphincter (LOS) motility associated with Barrett’s oesophagus includes reduced basal LOS pressures. The aim of this study was to characterise neuromuscular function of the LOS in normal (squamous cell carcinoma (SCC) with uninvolved LOS) and reflux affected (Barrett’s) oesophagus in vitro.

Methods—Strips of LOS muscle were prepared at biopsy following oesophagectomy from 16 patients with SCC and seven patients with oesophageal adenocarcinoma and Barrett’s oesophagus associated with a history of reflux disease. LOS smooth muscle responses were recorded in response to electrical field stimulation (EFS), potassium chloride (KCl), DMPP, isoprenaline, capsaicin, bethanechol, and tachykinins.

Results—Basal LOS tone and LOS relaxations in response to isoprenaline, EFS, and DMPP were not significantly altered in the Barrett’s group. After tetrodotoxin pretreatment, responses to KCl and DMPP were significantly reduced in the SCC but not in Barrett’s LOS. Maximal contraction in response to bethanechol was significantly decreased in Barrett’s LOS while substance P and NK-2 receptor mediated contraction was unaltered. Capsaicin, NK-1, and NK-3 receptor agonists exerted negligible effects on LOS tone.

Conclusions—LOS muscle strips from patients with reflux associated Barrett’s oesophagus exhibit a reduction in cholinergic muscle contraction while retaining similar features of basal tone, responses to tachykinins, and inhibitory muscle and neural function. Enteric inhibitory neurons in LOS muscle strips from patients with reflux associated Barrett’s oesophagus display resistance to axonal sodium channel blockade. No evidence for functional NK-1 or NK-3 receptors or capsaicin sensitive axon collateral reflexes was observed in the human LOS.

Keywords: Barrett’s oesophagus; lower oesophageal sphincter; tachykinins; inhibitory motorneurones; tetrodotoxin; smooth muscle

Columnar lined (Barrett’s) oesophagus is often considered a sequel to chronic severe gastro-oesophageal reflux disease, with prevalence rates of 1–10% in patients presenting for endoscopy with symptoms of gastro-oesophageal reflux disease.1 Barrett’s oesophagus is also associated with an increased risk of the development of oesophageal cancer.1 Aberrant upper gastrointestinal motility associated with Barrett’s oesophagus includes reduced basal lower oesophageal sphincter (LOS) pressure, decreased distal oesophageal contraction amplitude, impaired oesophageal clearance, delayed gastric emptying, and increased frequency of transient LOS relaxations,2–5 factors which often predominate in gastro-oesophageal reflux disease. In addition, there are reports of decreased perception of oesophageal acid3 and balloon distension6 in Barrett’s oesophagus. These findings indicate a broad pattern of alterations in both upper gastrointestinal sensation and motility associated with Barrett’s oesophagus. The mechanisms underlying increased frequency of transient LOS relaxations and LOS hypotonia in Barrett’s oesophagus are unknown. Animal surgical models of acute reflux disease, while successful in generating a columnar lined oesophagus,7,8 have yet to assess motility changes to the LOS. While animal models of acute oesophagitis have shown functional deficits in LOS function in vitro,9,10 these are inappropriate models to compare with LOS function of the chronic or severely reflux affected (and columnar lined) oesophagus.

In the present study we have compared LOS smooth muscle and non-adrenergic non-cholinergic (NANC) neural function in LOS biopsies from patients with severe or chronic gastro-oesophageal reflux disease concomitant
with Barrett’s oesophagus, with LOS biopsies from patients with normal squamous epithelium at the gastro-oesophageal junction and no pre-existing oesophagitis or history of reflux disease, to characterise the potential abnormalities in peripheral neuromuscular function of the severely reflux affected columnar lined LOS.

Methods
MUSCLE STRIP PREPARATION
Experiments were performed in tissues from a total of 23 patients referred to the Professorial and Oesophago-gastric Unit of the Royal Adelaide Hospital. All studies were performed in accordance with the guidelines of the human ethics committee of the Royal Adelaide Hospital, Adelaide. Strips of LOS were prepared at biopsy following gastro-oesophagectomy from a maximum of 16 patients with squamous cell carcinoma (SCC; mean age 67 (3) years) and seven patients with Barrett’s oesophagus (suspected adenocarcinoma indication for surgery; mean age 61 (3) years). Only Barrett’s patients with a discernable history of reflux disease were used. Columnar lined mucosa of the distal oesophagus was confirmed at pathology. Immediately following resection and removal of the cardia and oesophagus, sections (approximately 1×1 cm) of tissue were removed from the gastro-oesophageal junction (0.5–1.0 cm above the angle of His, corresponding to sling fibres) and placed in ice cold Krebs solution bubbled with carbogen (95% O2, 5% CO2). Manometric and anatomical evidence suggests that muscle from this region correlates with the functional lower oesophageal sphincter. In all cases only biopsy specimens of the gastro-oesophageal junction that were free of malignancy were used; this was confirmed by tissue pathology at the section margins. After removal of mucosa, submucosa, and longitudinal muscle, tissue was divided under sharp dissection to give four bands of circular muscle and attached myenteric plexus of dimensions 2 mm wide, 5 mm long. These were placed in individual 10 ml water jacketed organ baths containing carbogenated Krebs solution at 37°C of the following composition (mM): NaCl 118, NaHCO3 25, KCl 4.6, MgSO4 1.2, NaH2PO4 1.3, glucose 11, CaCl2 2.5. One end of the tissue was fastened to a support while the other was attached to an isometric force transducer (FPTo3, Grass, Quincy, Massachusetts, USA). On either side of the tissue support a pair of platinum electrodes was used for electrical field stimulation (EFS). Each strip was placed under an initial tension of 20 mN and left to equilibrate for 90 minutes. This corresponded to a degree of tissue stretch of approximately 200% of the initial resting length of the muscle strip, which has been shown to be within the optimal range for mechanical performance of the isolated human LOS. Only muscle strips from the gastro-oesophageal junction developed a stable spontaneous tension at rest and exhibited a typical relaxation profile in response to EFS. These criteria were considered to verify that the origin of the muscle strips was from the LOS.

PHYSIOLOGICAL AND PHARMACOLOGICAL STUDIES
EFS was delivered via rectangular wave pulses from a stimulator (S48, Grass Instruments, Quincy, Massachusetts, USA) with three minute intervals between each stimulus. Responses of LOS muscle strips to EFS (1–20 Hz, 50 V, 1 ms for 5 seconds), KCl (neural depolarization; 20 mM), 1,1-dimethyl-4-phenylpyridinium iodide (DMPP; nicotinic agonist; 10−5 M), isoprenaline (10−5 M), capsaicin (10−6 M), in addition to cumulative increasing concentrations of bethanechol (10−10–10−3 M), substance P, [Sar9, Met (O2)11]-substance P (NK-1 receptor selective agonist), [β-Ala]-neurokinin A 4–10 (NK-2 receptor selective agonist), and [Succinyl-Asp6, MePhe8]-substance P 6–11 (sentkide; NK-3 receptor selective agonist; all 10−10–10−4 M) were measured and recorded onto hard disk using Labview based software (MAD, Charles Malbert). All experiments were performed in the presence of atropine (10−6 M) and guanethidine (3×10−4 M), except where bethanechol was used to stimulate muscle contraction.

DATA ANALYSIS
Relaxation responses of the LOS were measured relative to the response of a supramaximal concentration of isoprenaline (10−3 M) or to baseline tension, while contractions were measured as a percentage of the maximal response to [β-Ala]-neurokinin A 4–10 (10−4 M) or as absolute contraction. Data are expressed as mean (SEM) of n subjects. Non-linear regression of concentration-response data, EMAX and EC50 calculations were performed using Prism 2.0b (Graphpad, San Diego, California, USA). Statistical analysis was performed using a paired Student’s t test for pretreatment studies using tetrodotoxin, an unpaired t test for comparison of individual EMAX and EC50 values between groups, or one way ANOVA for group point comparisons (where EMAX and EC50 values were not appropriate), with group means in this instance compared using Bonferroni’s post hoc test. A p value <0.05 was considered significant.

DRUGS
Substance P, [Sar9, Met (O2)11]-substance P, and sentkide were obtained from Auspep (Melbourne, Australia). [β-Ala]-neurokinin A 4–10, atropine sulphate, 1,1-dimethyl-4-phenylpyridinium iodide (DMPP), isoprenaline hydrochloride, bethanechol chloride, capsaicin, tetrodotoxin, and guanethidine sulphate were obtained from Sigma-Aldrich (Sydney, Australia). All drugs were dissolved in saline, except for capsaicin which was dissolved in saline, ethanol, and Tween 80 (8:1:1 v/v/v).

Results
LOS TONE AND MUSCLE RELAXATION
After placing LOS muscle strips under 20 mN of basal tension they developed additional tone during the equilibration period. Basal tension was reduced in the Barrett’s group (40 (6) mN vs 45 (3) mN in the SCC group) but this was not significant.
neurokinin A 4-10] (M)

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Figure 2 Concentration-response curve representing absolute contraction in response to the NK-2 receptor agonist [β-Ala8]-neurokinin A 4-10 in squamous cell carcinoma (SCC) and Barrett’s lower oesophageal sphincter (LOS) muscle strips. While increased, maximal responsiveness was not significantly altered in Barrett’s group.

LOS responses to electrical field stimulation (EFS) and NANC neural activation

LOS muscle strips responded to EFS (1–20 Hz, 50 V, 1 ms duration for 5 seconds) with a predominant frequency dependent relaxation that was near maximal at 10 Hz (fig 4). While responses to EFS were attenuated in Barrett’s LOS, they were not significantly different (fig 4). Tetrodotoxin significantly inhibited the relaxation in response to EFS at 5 Hz in both Barrett’s (−55.1 (9)% control v −4.7 (2)% with TTX; p<0.05) and SCC (−62.6 (13)% control v −3.33 (0.7)% with TTX; p<0.001) groups relative to maximal relaxation to isoprenaline.

DMPP and KCl administration evoked rapid and powerful relaxations of the human LOS, which were greater than the relaxation elicited by supramaximal concentrations of isoprenaline. However, there were no differences in degree of relaxation between the SCC and Barrett’s groups following either DMPP or

maximal responsiveness when measured relative to the maximum contraction following selective NK-2 receptor activation with [β-Ala8]-neurokinin A 4-10 (Emax Barrett’s 145 (50)% SCC v 145 (50)% Barrett’s, *p<0.05).

[β-Ala8]-Neurokinin A 4-10 administration caused contraction of the human LOS that was dose dependent from 10−6 to 10−4 M and near maximal at 10−5 M (fig 2). [β-Ala8]-Neurokinin A 4-10 displayed similar dose-response curves in LOS muscle strips from the SCC and Barrett’s groups. There were no significant differences in maximal responsiveness when measured in terms of changes in absolute tension.

Substance P mediated contractions were small in comparison (30% of maximal [β-Ala8]-neurokinin A 4-10 evoked contraction) and were not significantly altered in muscle strips from the Barrett’s group (fig 3). [Sar9, Met(O2)11]-Substance P and senktide elicited negligible effects on LOS tone in vitro and these were not significantly different in the Barrett’s group. Capsaicin (10−4 M) failed to elicit a response in LOS muscle strips from either SCC or Barrett’s LOS, as did its vehicle (see Drugs).

LOS Cholinergic, Tachykinergic, and Capsaicin Induced Responses

Bethanechol administration elicited dose dependent contraction that was maximal at 10−4 M. In the Barrett’s group there was a reduction in maximal absolute tension development (fig 1B) and a significant reduction in

Figure 1 Concentration-response curve representing contraction in response to bethanechol in squamous cell carcinoma (SCC) and Barrett’s lower oesophageal sphincter (LOS) muscle strips, expressed in terms of absolute contraction (B) or relative to maximal contraction in response to the NK-2 receptor agonist [β-Ala8]-neurokinin A 4-10 (A). Contraction was suppressed based on both indices and significantly decreased in the Barrett’s group (significant difference in maximal responsiveness: Emax 264.1 (24)% SCC v 145 (50)% Barrett’s, *p<0.05).

LOS muscle relaxation in response to isoprenaline (10 µM) consisted of a prolonged relaxation relative to basal tension (59.7 (4)% Barrett’s, 44.8 (4)% SCC). While the Barrett’s group exhibited greater relaxation following isoprenaline. However, there were no significant differences in the evoked relaxation between groups when measured relative to basal tension.

Supporting Information

Figure 3 Concentration-response curves showing the response to the non-specific neurokinin receptor agonist Substance P in squamous cell carcinoma (SCC) and Barrett’s lower oesophageal sphincter (LOS) muscle strips. Substance P elicited modest contraction in LOS muscle strips of approximately 30% of the maximal contraction to the NK-2 receptor agonist [β-Ala8]-neurokinin A 4-10. There was no difference in tension development in the Barrett’s or SCC group in response to substance P.
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Discussion

In the present study we have investigated the effects of severe gastro-oesophageal reflux disease complicated by Barrett’s oesophagus on the functional activity of the lower oesophageal sphincter (LOS) in vitro, including measurements of smooth muscle and inhibitory motorneuronal activity. While there are many clinical studies showing a range of motility changes in the upper gastrointestinal tract in reflux associated Barrett’s oesophagus,2 none has addressed potential alterations in peripheral neuromuscular function in the LOS. The range of pharmacological markers used in this study reflects characterisation of a range of neurotransmitter functions in the human LOS. Generally there were few differences between SCC and Barrett’s LOS with respect to non-adrenergic non-cholinergic (NANC) inhibitory motorneuronal activation. Muscle strips showed a similar degree of EFS induced relaxation, and maximal nicotinic receptor or [K+] stimulated relaxation (both operating through NANC neurones)1718 was unchanged. A complement of selective neurokinin receptor agonists was used to determine the pharmacological profile of tachykinins in the human LOS. Our findings corroborate those of another study that showed lack of functional NK-1 and NK-3 receptors but a predominant NK-2 receptor mediated contraction in the human LOS.23 The history of reflux, oesophagitis, and/or columnar lined oesophagus did not alter the sensitivity or maximal responsiveness to cholinergic excitation using bethanechol.

Muscular cholinergic responses are believed to occur through muscarinic receptor (M3) stimulation of G protein coupled phospholipase C in the LOS with increased intracellular inositol triphosphate and diacylglycerol leading to increased cytoplasmic calcium and activation of smooth muscle.2021 A similar second messenger pathway is believed to be used by NK-2 receptors in smooth muscle22 but this has not yet been characterised specifically for the LOS. However, NK-2 receptor mediated contractions were not concomitantly reduced (slightly greater, in fact) in the Barrett’s group, suggestive of a selective deficit in cholinergic based muscle contraction rather than a diminished shared second messenger pathway. This occurred in the area of the LOS most reactive to muscarinic activation—the LOS gastric sling fibres that course over the angle of His.1123 We can speculate that LOS hypotonia observed in the Barrett’s patients may be due to a peripheral cholinergic motor deficiency where mean basal LOS pressures are lower than in normal subjects or in patients with oesophagitis alone.2 The extent to which cholinergic mechanisms contribute to basal LOS pressure in humans is reported to be 50–80%2425; this is the extent to which atropine lowers resting LOS pressure when administered to healthy subjects. However, this contribution is also similar in patients with gastro-oesophageal reflux disease where atropine reduced basal LOS pressure by 60%.26 The component of cholinergic tone in basal LOS pressures in

Figure 4 Frequency-response curve representing relaxation in response to electrical field stimulation in squamous cell carcinoma (SCC) and Barrett’s lower oesophageal sphincter (LOS) muscle strips. While attenuated at most frequencies, relaxation was not significantly altered in the Barrett’s group. Relaxation expressed relative to relaxation induced by a supramaximal concentration of isoprenaline (ISO; 10 µM).

Figure 5 Histogram representing lower oesophageal sphincter (LOS) muscle strip relaxation to KCl 20 mM (A) and DMPP 10 µM (B) in the control and Barrett’s group, before and after axonal sodium channel blockade with tetrodotoxin (TTX 1 µM). Relaxation expressed relative to relaxation induced by a supramaximal concentration of isoprenaline (10 µM). Muscle strips from Barrett’s LOS showed similar degrees of relaxation to either DMPP or KCl compared with controls. Tetrodotoxin significantly inhibited relaxation to DMPP in both groups (**p<0.001 controls, *p<0.05 Barrett’s) and inhibited relaxation to KCl in the control (**p<0.05) but not in the Barrett’s group.

KCl administration (fig 5A, B). Following pre-treatment with tetrodotoxin, KCl and DMPP induced relaxations were significantly inhibited in SCC (KCl, p<0.05 (fig 5A) and DMPP, p<0.001 (fig 5B)) but not in Barrett’s LOS.
reflux affected Barrett’s oesophagus has not been determined.

Relaxation in response to isoprenaline in the Barrett’s group was greater than that of the SCC group but this did not achieve significance, indicating that β and adrenoceptors mediated relaxation of the LOS was unaffected by reflux affected columnar lined mucosa in the oesophagus. The human LOS contains both β and β adrenoceptors; activation of either or both causes LOS relaxation by direct activation of LOS muscle. This indicates that the capacity for direct LOS smooth muscle relaxation is unperturbed in reflux associated Barrett’s oesophagus, at least for β adrenergic influences.

The absence of an LOS response to capsaicin administration indicates that either sensory afferent-collateral reflexes are not evident within the sling region of the human LOS or that they are insensitive to capsaicin. The clasp region of the LOS was not studied, so potential activation of capsaicin sensitive pathways has not yet been excluded in this region of the human gastro-oesophageal junction. These pathways, which in the ferret LOS activate inhibitory motorneurones via endogenous substance P release from sensory axon collaterals, are enhanced in experimental oesophagitis. In contrast, exogenous substance P elicited modest excitation of LOS smooth muscle from both SCC and reflux affected Barrett’s LOS, which is likely to be due to NK-2 receptor activation in humans. There was no distinction between responses in either group.

While NANC inhibitory neural activation elicited similar degrees of relaxation in SCC and reflux associated Barrett’s LOS, the SCC LOS displayed greater inhibition of relaxation following maximal NANC neuronal stimulation in the presence of tetrodotoxin. This was evident from inhibitory motorneuronal activation using DMPP or KCl administration. Analysis of individual responses in the reflux associated Barrett’s group revealed a disparity in the efficacy of tetrodotoxin—some showed effective blockade of KCl and DMPP evoked relaxations and some showed little effect (data not shown). Therefore, there may be a subset of reflux affected Barrett’s patients who display resistance to axonal sodium channel blockade in the LOS. Tetrodotoxin resistant sodium channels occur in sensory, small diameter dorsal root ganglion neurones and are upregulated in inflammatory states, but their expression in enteric motorneurones has not been demonstrated and the physiological implications of their expression in Barrett’s oesophagus are not known.

It was not possible to determine if the altered responses of the LOS in Barrett’s patients were due to a process of inflammation in the region or associated with intestinal metaplasia in some way. The extent of active ongoing oesophagitis in the Barrett’s subjects presenting for oesophagectomy was not determined and therefore a correlation between neuromuscular function and inflammatory states in these biopsies could not be undertaken. However, samples of distal oesophageal mucosa from Barrett’s patients show high levels of inflammatory mediators such as leukotrienes, suggestive of prevailing oesophagitis in this group. In conclusion, LOS muscle strips from patients with severe reflux disease associated with Barrett’s oesophagus exhibit selective reductions in cholineric based muscle contraction while retaining similar features of basal tone, tachykininergic contractile, and inhibitory muscle and neural function. However, enteric neurones in LOS muscle strips from a subset of reflux affected patients with Barrett’s oesophagus displayed resistance to sodium channel blockade, which was revealed at more intense measures of neuronal stimulation. NK-2 receptors are the predominant tachykinin receptor in the human LOS; no evidence for capsaicin sensitive axon collateral reflexes or NK-1 or NK-3 receptors was demonstrated. The attenuated cholineric LOS muscle contraction may contribute to deficiencies in basal LOS pressures observed clinically in Barrett’s oesophagus.

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