Site and mechanism of pain perception with oesophageal balloon distension and intravenous edrophonium in patients with oesophageal chest pain

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Abstract
Ten healthy volunteers and 13 patients with oesophageal motility disorders whose primary presenting complaint was chest pain were studied by distending an intraoesophageal balloon in 1 ml steps to the point of a sensation of discomfort. The net balloon pressure (intra-balloon pressure when inflated within the oesophagus minus the pressure recorded at the same volume outside the patient) was measured at each volume increment and the distension volume at the perception of discomfort was noted. The measurements were repeated after intravenous injection of edrophonium (80 μg/kg) and again after 1-2 mg intravenous atropine. Oesophageal wall compliance was similar in patients and controls, and the two groups showed a similar effect of decreased compliance with edrophonium and increased compliance after atropine. There were no significant differences between patients and controls of distending volume at perception of discomfort. Edrophonium, however, resulted in a significant reduction in distension threshold for pain (p<0.03) in patients. A similar though non-significant trend was seen in controls. In both controls and patients, distension volume for pain production after atropine was significantly (p<0.01) higher than after edrophonium. From these results and other published data, we suggest that the pain receptor for noxious stretch and after edrophonium challenge is likely to be an 'in series' mechanoreceptor located in oesophageal longitudinal muscle.

The mechanism of production of pain with edrophonium is unclear: patients who develop their typical symptoms with the drug have a significantly greater increase in peristaltic duration (but not amplitude) than those who do not. A large overlap exists, however, so that some patients with large increases in peristaltic duration do not have symptoms whereas others develop symptoms with little manometric change. Furthermore, provoked chest pain is frequently continuous over several minutes while oesophageal pressure waves are only observed intermittently. A possible reason for this failure to correlate pain and manometric findings is that pain does not directly result from abnormal motility but rather from heightened sensory perception. This could result in otherwise painless 'physiological' stimuli being perceived as painful. It is possible and likely that the manometric changes observed in some patients are modulated by the sensory receptors, probably by an axon reflex influence on the effector neurones of the myenteric plexus. Balloons have a long history of application to the study of oesophageal pain. The use of oesophageal balloons has recently been revived after a long lapse, perhaps resulting from the observation that patients with documented ischaemic heart disease could not differentiate between the pain of their exertional angina and that induced by oesophageal balloon distension (despite the lack of electrocardiographic changes during balloon induced pain). These reservations persist, particularly now that microvascular angina has emerged as an alternative explanation for chest pain in patients with chest pain but anatomically normal coronary arteries. Nevertheless, oesophageal balloon distension can reproduce pain in 40 to 60% of patients with non-cardiac chest pain suspected to be of oesophageal origin. There is some evidence that these individuals have increased oesophageal sensitivity to stretch, while others have pointed out that such patients show abnormalities of the oesophageal peristaltic reflex during balloon induced pain.

The purpose of this investigation was to study the effects of edrophonium and atropine on the perception of discomfort and oesophageal wall compliance during oesophageal balloon distension in an attempt to throw some light on the mechanisms and site of action of edrophonium.
and balloon distension as provocative manoeuvres.

Methods

SUBJECTS
Ten healthy volunteers were recruited (seven men and three women, mean age 28 (SD 4.8) years). Three had experienced occasional heartburn less than once a month; otherwise no volunteer had any oesophageal or gastrointestinal symptoms. Thirteen patients agreed to participate (five men and three women, mean age 48 (SD 8) years. All had been found to have oesophageal motility disorders during previous oesophageal manometry and all had had a normal endoscopic examination of the oesophagus. The manometric diagnoses were: diffuse oesophageal spasm (six), high amplitude and/or prolonged duration peristalsis (six patients—three had also experienced their typical symptoms with provocation tests during manometry) and vigorous achalasia (one). This latter patient had previously undergone forceful pneumatic dilatation of the cardia and had been shown radiologically not to have a dilated oesophagus. All patients had originally presented with chest pain, 10 also complaining of dysphagia, a further two of odynophagia and two of heartburn (distinct from their chest pain) in addition. At the time of the study, 12 of the 13 patients were still symptomatic.

An Arndorfer ESM3 polynvinyl multilumen oesophageal manometry catheter connected to a low compliance constant water perfusion pump (Arndorfer Medical Specialities Inc, Greendale, Wisconsin, USA) was adapted by tying a rubber balloon over one of the recording ports so that it was placed 10 cm above the distal recording port. The balloon was fashioned from a latex rubber finger cot cut to 3 cm in length. The ends were secured to the polyvinyl tube by winding and tying a silk suture tightly around either end. The loose ends were trimmed and a silicone glue seal applied to make the balloon airtight. The balloon channel was filled with air and connected to an external pressure transducer and chart recorder (Elcomatic, Neilston, Glasgow) using a three way tap. This enabled inflation of the balloon with air in steps of 1 ml, and measurement of intra-balloon pressure at each step.

Inflations were carried out on the bench in air before each patient study, and the balloon was checked in water for leaks on each occasion. The same balloon was used for all experiments in patients and controls, as preliminary trials indicated that different sized balloons or even balloons of the same size constructed out of similar latex rubber finger cots had different compliances. This problem has been noted previously. For the balloon used in the studies, the relationship between balloon diameter (measured with a micrometer) and volume during benchtop inflations, although not entirely linear, showed no evidence of flattening over the range of volumes used during intraballoonal inflations (Fig 1). Although no in vivo measurements were made, two studies have indicated that during inflation in the oesophagus, similar rubber balloons are only slightly deformed. For patient studies, the deflated balloon and manometry catheter assembly was passed through the nose and positioned so that the balloon lay 9 cm above the lower oesophageal sphincter. After inflation in 1 ml steps, the balloon was inflated in 0.4 ml steps in such a fashion that the patient did not know when air was being added to the balloon, until a continuous sensation of discomfort was felt. A short period was allowed to elapse after each step of the inflation for the balloon pressure tracing to stabilise, but the balloon was not deflated between steps. Once a stable pressure trace was observed, the minimum intraballoon pressure at each volume step was measured with reference to the basal pressure reading when the balloon had been deflated within the oesophagus.

Dry swallows were sensed by a hand on the patient’s throat so that each could be marked on the pressure tracing chart. The patient was instructed to stop the period of balloon inflation to report the point at which any abnormal chest or back sensation occurred, and also the point at which this became definitely uncomfortable. No attempt was made to prompt the patients who were required to volunteer the information requested. The point at which definite discomfort was reported was recorded on the chart. In the case of patients with proven motility disorders, each was asked whether the discomfort resembled their spontaneous symptoms. The balloon was deflated once this point had been reached and patients were asked to describe the location and character of the symptom that had been provoked.

In each case, the same procedure was repeated after intravenous injection of edrophonium (80 µg/kg). In four subjects, an end point of discomfort was not achieved after edrophonium, as the balloon had to be deflated at the patient’s request because of intolerable nausea or nasal discomfort from aboral balloon traction. Twenty five to 30 minutes after edrophonium, atropine 1-2 mg was administered intravenously and the balloon inflation procedure repeated. In some instances, more than one inflation procedure

Figure 1: Volume—diameter relationship for the balloon used in all the studies. The diameter of the balloon at its widest point was measured using a micrometer. The individual data points are the mean of 2 separate benchtop inflations; the mean difference between the 2 measurements at each volume increment was 0.4 mm.
was carried out during each stage of the procedure to assess the reproducibility of the results. The time in minutes taken for each inflation (mean (SD)) was, for control subjects and patients respectively, 3.1 (1.2) and 3.1 (1.1) for the basal period, 3.0 (0.9) and 2.7 (1.4) after edrophonium and 3.1 (1.4) and 3.3 (1.7) after atropine. In particular, in no subject did the inflation period exceed four minutes 40 seconds after edrophonium, an important consideration as this drug has a short duration of action.

During each recording, three channels (one at 5 cm and one at 10 cm above the balloon and one in the lower oesophageal sphincter) were perfused with water from the Arndorfer hydraulic capillary infusion pump. A continuous record of oesophageal motor activity at these points was obtained. Oesophageal motor activity proximal to the balloon was assessed qualitatively, but no attempt was made to measure the amplitude or duration of pressure waves, as all swallows were 'dry'. Because the recording channel in the lower oesophageal sphincter tended to slip into the stomach (presumably because of oesophageal shortening known to occur with distension of the oesophageal body), readings from this channel were not analysed.

After each experiment, the observed intra-balloon pressure at each volume step during benchtop inflation was subtracted from the corresponding value during intraoesophageal distension to obtain a more accurate reflection of the contribution to the measured balloon pressure resulting from the oesophageal wall. The pressure measurements obtained are termed 'net balloon pressure'. According to Laplace’s law, tension (T) in the wall of a cylinder is given by the relationship T=π×r, where p=intraluminal pressure and r=the radius of the cylinder. As oesophageal diameter was not measured, however, the tension could not be calculated, so the pressure measurements are only an indirect reflection of oesophageal mural tension.

**STATISTICAL ANALYSIS**

Statistical tests used included analysis of variance (either repeated measures or factorial, depending on whether paired or unpaired samples were being compared). Where significant overall differences were found, Scheffe’s test was used as a multiple range test to locate the differences, with an adjustment of the p value for each comparison to reduce the chance of a type I error. Pearson’s product moment correlation coefficient was calculated as a measure of reproducibility. A p value of <0.05 was taken to indicate a significant result.

**Results**

**SITE AND NATURE OF DISCOMFORT**

Discomfort was experienced by control subjects retrosternally in six, with radiation to the back (two), abdominal right upper quadrant (two) and throat (one). Two experienced throat discomfort only, one back pain alone and one pain in the back and right upper quadrant. The pain was described as burning by one subject, and by the others as a ‘pressure’ or ‘ache’.

Among the 13 patients, eight experienced retrosternal discomfort only. In three others, the discomfort was felt in the back as well as retrosternally and in the final two, the throat was the only site of discomfort. Two patients described the pain as heartburn and five (38%) felt the pain during the initial inflation period (without administered drugs) was the same as their usual spontaneous symptom. Of these five, three were cases of high amplitude and/or prolonged duration peristalsis and two had diffuse oesophageal spasm.

Several patients and control subjects commented that the discomfort was initially felt at one site but, on subsequent balloon inflations, was also experienced at another site. A minority of patients and controls noted that the symptoms worsened transiently coincident with peristaltic waves reaching the site of balloon distension.

**OESOPHAGEAL MOTOR ACTIVITY DURING BALLOON INFLATION**

This was only analysed qualitatively, as wet swallows were impossible with the oesophagus occluded by a balloon. Dry swallows were permitted, but are known to give variable results in terms of patterns of pressure change and amplitude or duration of pressure waves.
In the majority (60 – 80% – the proportion varying during different phases of the study) of controls, all pressure waves were peristaltic during the balloon inflation, with a few (20–30%) having spontaneous non-propagated waves occurring more than once a minute during balloon inflation. In only one did the non-peristaltic activity occur only during discomfort.

Not surprisingly, in less than half (46%) of the patients was pressure activity only peristaltic, as seven were known to have either diffuse oesophageal spasm or vigorous achalasia. The proportion with peristaltic activity but no abnormal pressure waves increased following edrophonium to 69%. Although a high proportion (31–46%) of patients exhibited frequent spontaneous non-peristaltic waves, in only a small number (8–23%) did this abnormality coincide with the development of symptoms. Even fewer controls exhibited this feature.

Non-swallow related peristaltic waves proximal to the distending balloon were only observed in 40% of patients and controls, being reduced or abolished by atropine. The infrequency of these waves may be attributable to the fact that patients were allowed to swallow at will, resulting in inhibition of this activity. Atropine diminished peristaltic amplitude in all patients and controls, causing apertistalsis in 23% and 40% respectively. A steady rise in baseline proximal intraoesophageal pressure to between 6 and 12 mm Hg above basal pressure was observed in 10% of controls (but only in one of the 13 patients) during balloon inflation, being more frequently seen (40% of controls) after atropine. This occurred independently of the development of symptoms, and represents a phenomenon already reported by Enzmann and colleagues.4

Finally, multipeaked peristaltic waves were observed in 31% of patients (but not controls) after edrophonium, always independent of symptoms.

RELATIONSHIP OF BALLOON VOLUME AND PRESSURE TO DISCOMFORT

The balloon distension volumes and pressures at which discomfort was first perceived by controls and patients is shown in Figures 2 and 3. For patients and controls considered together, there was a significant overall difference in volumes at first perception of pain by ANOVA (f=2.51; p<0.02), but not for pressure (f=1.09; p>0.3). Using Scheffe’s multiple range test, mean inflation volume at perception of discomfort was significantly lower for patients after edrophonium than after either atropine (p<0.01) or initial basal inflation period before drugs were administered (p<0.03; Fig 2). A significant reduction was found in controls in balloon volumes to produce discomfort after edrophonium compared with balloon distension after atropine (p<0.01; Fig 2); although there was a trend for a reduction in distension threshold with edrophonium compared with basal inflation, this was not statistically significant in the controls.

No significant differences in threshold balloon volumes at perception of discomfort were apparent between controls and patients with chest pain. There was a non-significant trend for lower intraballooon pressures at the perception of discomfort in controls and patients after atropine compared to basal and post edrophonium periods (Fig 3).

The reproducibility of the measurement of balloon volume and pressure at first perception of discomfort was assessed by comparing duplicate balloon inflations which were carried out on a total of 45 occasions. Reproducible results were obtained for volume measurement at the first perception of discomfort (r=0.89; p<0.0001) but less so for pressure (r=0.61; p<0.001).

EFFECTS OF EDROPHONIUM AND ATROPINE ON OESOPHAGEAL COMPLIANCE

Net intraoesophageal balloon pressure values are shown in Figures 4 to 6 over the range 1 to 10 ml of air introduced into the balloon. Results for balloon volumes above 10 ml are not given, because the number of control and patient volunteers became too few for meaningful comparison.

An overall comparison was performed between six sets of values (three for patients, three for control subjects) at each balloon volume in the range 1 to 10 ml using one way analysis of variance. Significant overall differences were found at 1 and 2 ml (p<0.05). No other significant differences overall were observed. There did not appear to be any consistent reason for the significant differences observed at 1 and 2 ml. Because significant differences could not be detected at other volumes, it seems reasonable to conclude that there were no differences between the patient and control groups with regard to oesophageal compliance resulting from administration of the drugs used. For clarity, the pressure volume relationship is displayed in Figure 4 for the basal inflation period only; there is no consistent difference in oesophageal compliance between the two groups.

Further comparisons were made within groups – that is, for the patients and controls separately, using analysis of variance (repeated measures) with Scheffe’s multiple range test to locate significant effects. Although consistent results were not obtained at low volumes (1 and 2 ml) probably because the balloon was not sufficiently distended for the pressure to reflect oesophageal wall tension, and at volumes exceed-
results in a sustained contraction at the level of the balloon\(^1\) producing a strong aboral traction force. This is transiently inhibited by swallowing but reinforced by the arrival of the primary peristaltic wave at the balloon.\(^2\) This contraction, which continues as long as distension is maintained, results in oesophageal shortening below the balloon, and has been shown in the opossum to be the result of the contraction of muscularis mucosae and the longitudinal muscle of the muscularis propria.\(^3\) The temporal characteristics of the ‘oesophageal propulsive force’ observed in man by Winship and Zboralske\(^4\) and the ‘duration response’ observed by Christensen and Lund\(^5\) make it likely that these represent the same phenomenon. This response is abolished by anticholinergic drugs.

In the present study, significant differences were found in oesophageal compliance after edrophonium and atropine. From the foregoing discussion, a likely cause for changes in oesophageal wall compliance measured during balloon distension after administration of drugs affecting cholinergic pathways is contraction of oesophageal muscularis mucosae and longitudinal muscle.\(^6\) No differences were found between patients with motility disorders and controls, however, with respect to oesophageal compliance. These findings agree with previous observations in the monkey\(^7\) and in man.\(^8\)\(^9\)

The administration of edrophonium to patients resulted in a significantly lower balloon volume, but not pressure, at pain perception compared with both basal and postatropine balloon distension. A similar significant effect or trend was observed among controls.

These findings may be interpreted as indicating that because administration of a cholinergic blocker reduced oesophageal wall compliance, a critical intraoesophageal pressure (for pain perception) was reached at a lower distension volume, with the converse applying after administration of an anticholinergic agent. This is in keeping with the general observation that changes in gut smooth muscle tension could modulate changes in pain threshold to distension.\(^7\) We were not able to show a clear difference between patients and controls with respect to this effect.

Oesophageal ischaemia consequent on distension has been suggested as a mechanism for pain production.\(^10\) If so, cholinergic stimulation could enhance oesophageal wall ischaemia, perhaps simply by allowing a critical oesophageal wall tension inhibiting blood flow to be reached at an earlier stage of distension. Our findings would be compatible with this hypothesis, but other preliminary studies have failed to find evidence of oesophageal ischaemia either in patients during episodes of chest pain\(^11\) or in a rabbit model during swallowing before or after intravenous edrophonium.\(^12\)

Can any conclusions be reached from these observations regarding the mode of action of oesophageal balloon distension and intravenous edrophonium when used as provocative agents in patients with non-cardiac chest pain? While our results apply only to patients who have been found to have motility disorders during oesophageal manometry (and so may not necessarily be
extrapolated to all patients with non-cardiac chest pain), we believe that some conclusions may be drawn.

With regard to balloon distension, motility changes proximal to the balloon did not show any consistent relationship with pain perception. Other work suggests that a proportion of such individuals developing their symptoms during balloon distension have abnormalities distal to the balloon.11 Whether such abnormal distal activity is the cause of pain or (as seems more likely) a marker of an abnormal oesophagus is at present not clear.

We could not show abnormally low thresholds to stretch in our patients, but our sample size was small and highly selected. A reduced threshold has been reported by Richter and colleagues in patients with chest pain,12 but they studied larger numbers of patients who differed from ours in that although all had non-cardiac chest pain, only a minority had manometric abnormalities. In addition, there are methodological differences in the techniques of balloon inflation, although we do not believe these are likely to have influenced the results. In agreement with the findings of this group, we did not observe any difference in oesophageal compliance between our patients and healthy controls. Furthermore, we did not find any differences between patients and controls in oesophageal wall compliance resulting from cholinergic or anticholinergic drug administration.

Our results are consistent with the hypothesis that stretch sensitive mechanoreceptors located in longitudinally oriented muscle are involved in the perception of pain induced by balloon distension. It has recently been shown that mechanoreceptors with the electrophysiological characteristics of nociceptors appear to be located 'in series' with oesophageal longitudinal muscle but not circular muscle.16 Mechanoreceptors in circular muscle layers by contrast do not appear to have the characteristics of nociceptors,17 responding maximally only to physiological changes in tension that result from peristaltic activity.

The observation that edrophonium resulted in a lowering of the pain threshold to distension and also in decreased oesophageal wall compliance (which we have argued represents an effect on longitudinal muscle) prompts the speculation that pain developing with this drug is sensed by the same receptors as those responding to balloon distension. If so, it would explain why there appears not to be a close correlation between the manometric changes which occur after the drug and the appearance of symptoms. Longitudinal muscle contraction is not directly recorded by conventional manometry, although peristaltic duration might be expected to be prolonged by longitudinal muscle contraction and is the parameter best correlating with a positive edrophonium pain response.17 Why edrophonium should cause pain only in individuals with clinical symptoms is unclear, unless the neural pathways responsible for pain perception are already 'sensitised' in these individuals. Perhaps the nociceptors themselves are sensitised in these patients by quite a different mechanism – for instance in response to refluxed gastric acid. There is cer-

1 Richter JE, Hackshaw BT, Wu WC, Castell DO. Edro-


25 Winsans CA. Alteration of lower esophageal sphincter charac-


