Motor responses of the oesophagus to intraluminal distension in normal subjects and patients with oesophageal clearance disorders

G P N KENDALL, D G THOMPSON, S J DAY, AND N GARVIE

From the Department of Medical Research, St Mark's Hospital, London and Departments of Gastroenterology, Biometry and Nuclear Medicine, The London Hospital, London

SUMMARY Oesophageal motor responses to intraluminal distension were studied manometrically in 16 healthy volunteers and in nine patients with disordered swallowing, who had prolonged oesophageal clearance without structural abnormality. In the normal subjects distension was associated with an increased number of secondary contractions above the balloon, decrease of all contractile activity below the balloon and was accompanied by an aborally propulsive force which occurred independently of the perception of discomfort. Cholinergic blockade abolished the proximal distension induced contractile response, but did not affect primary peristalsis. Despite normal sensory thresholds, proximal excitatory responses to distension were absent in six and distal inhibition was absent in seven patients. These results show that the normal human oesophagus responds to distension with a proximal enhancement of propulsive motor activity, mediated through a cholinergic pathway. This may be defective in some patients with disordered oesophageal transit. Investigation of the motor responses to intraluminal distension may thus be a useful adjunct to standard manometry for studying patients with suspected oesophageal clearance dysfunction and might allow identification of disordered enteric nervous control.

Oesophageal transport occurs through two peristaltic mechanisms.1 Primary, or swallow initiated, peristalsis which is centrally mediated originates in the pharynx and progresses aborally to the stomach. If food remains in the oesophagus distending the lumen then local neural reflexes induce secondary, or non-swallow initiated, peristaltic activity to clear it.1 Supine radionuclide oesophageal transit has been advocated as a sensitive screening method for detecting oesophageal motility disorders.2,3 In some cases, however, no structural or manometric cause for delayed transit can be found and some people consider these cases to be false positive results.4 Radionuclide transit studies,2,4 measure the time taken for an ingested isotopically labelled bolus to traverse the oesophagus and provide information on the efficiency of oesophageal clearance. The usual procedure involves one swallow at the start of the test so that speed of clearance is a function of the efficiency of one primary peristaltic contraction and any subsequent secondary activity. Standard oesophageal manometry5 is useful for defining abnormalities of primary peristalsis, but secondary peristalsis, in contrast, is not usually demonstrated and transit of intraluminal contents is not evaluated. Secondary peristalsis has been studied experimentally and can be induced by intraluminal balloon distension,6,7 but it has not previously been investigated in patients with swallowing difficulties.

We hypothesised that patients with delayed oesophageal clearance but apparently normal manometry might have abnormal secondary peristaltic activity. We therefore performed the series of studies described below with the aim of defining the normal responses of the human oesophagus to localised distension and comparing this information with that obtained from patients with unexplained oesophageal clearance disorders.

Address for correspondence: Dr D G Thompson, Department of Medicine, Hope Hospital, Salford M6 8HD.

Received for publication 11 July 1986.
Motor responses of the oesophagus to intraluminal distension in normal subjects

Table

<table>
<thead>
<tr>
<th>Case no</th>
<th>Age</th>
<th>Sex</th>
<th>History</th>
<th>Barium swallow report</th>
<th>SROT</th>
<th>Standard manometry</th>
<th>Distension proximal excitation</th>
<th>Response distal inhibition</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>41</td>
<td>M</td>
<td>17 yr effortless regurgitation of food.</td>
<td>Normal</td>
<td>43 sec</td>
<td>Normal peristalsis, LOS 10 mmHg. Normal relaxation</td>
<td>Absent</td>
<td>Absent</td>
</tr>
<tr>
<td>2</td>
<td>43</td>
<td>F</td>
<td>3 yr food sticking on swallowing, regurgitation, retrosternal pain with food</td>
<td>Normal</td>
<td>&gt;60 sec</td>
<td>Normal peristalsis, LOS 6 mmHg. Normal relaxation</td>
<td>Absent</td>
<td>Absent</td>
</tr>
<tr>
<td>3</td>
<td>67</td>
<td>F</td>
<td>3 yr food sticking on swallowing</td>
<td>Tapered lower oesophagus Hold up at lower oesophagus</td>
<td>22 sec</td>
<td>Normal peristalsis, LOS 12-5 mmHg. No relaxation</td>
<td>Absent</td>
<td>Absent</td>
</tr>
<tr>
<td>4</td>
<td>55</td>
<td>F</td>
<td>4 yr food sticking on swallowing, retrosternal pain with food</td>
<td>Normal</td>
<td>&gt;60 sec</td>
<td>Low amplitude peristalsis, LOS 12:5 mmHg. No relaxation</td>
<td>Absent</td>
<td>Absent</td>
</tr>
<tr>
<td>5</td>
<td>39</td>
<td>M</td>
<td>1 yr retrosternal pain and dysphagia with food</td>
<td>Hold up at lower oesophagus</td>
<td>23 sec</td>
<td>Multipieked peristalsis, LOS 12:5 mmHg. No relaxation</td>
<td>Present</td>
<td>Present</td>
</tr>
<tr>
<td>6</td>
<td>79</td>
<td>F</td>
<td>1 yr choking and retrosternal pain with food</td>
<td>Poorly contractile oesophagus, hold up at lower oesophagus</td>
<td>20 sec</td>
<td>Low amplitude peristalsis, LOS 27.5 mmHg. Normal relaxation</td>
<td>Absent</td>
<td>Absent</td>
</tr>
<tr>
<td>7</td>
<td>29</td>
<td>F</td>
<td>10 yr retrosternal pain on eating</td>
<td>—</td>
<td>25 sec</td>
<td>Normal peristalsis, LOS 12-5 mmHg. Normal relaxation</td>
<td>Present</td>
<td>Absent</td>
</tr>
<tr>
<td>8</td>
<td>38</td>
<td>M</td>
<td>2 yr retrosternal pain with food</td>
<td>Tertiary peristaltic waves</td>
<td>17 sec</td>
<td>Multipieked peristalsis. No LOS demonstrated</td>
<td>Absent</td>
<td>Absent</td>
</tr>
<tr>
<td>9</td>
<td>41</td>
<td>M</td>
<td>1 yr chest pain, regurgitation with food</td>
<td>Tertiary peristaltic waves</td>
<td>16 sec</td>
<td>Multipieked peristalsis. No LOS demonstrated</td>
<td>Present</td>
<td>Present</td>
</tr>
</tbody>
</table>

SROT = supine radionuclide oesophageal transit; LOS = lower oesophageal sphincter.

Methods

Subjects

The study protocols were approved by the London Hospital Ethics Committee. Sixteen healthy adult volunteers (age 19–37), without history of gastrointestinal disorder, and nine patients were studied. The patients comprised all those referred to the gastroenterology unit for oesophageal manometry, over a period of 12 months, with a history of swallowing difficulty that included regurgitation, dysphagia, odynophagia and aspiration (Table). Patients with a dilated oesophagus, endoscopic evidence of oesophagitis or a structural disorder of the oesophagus were excluded from the study and none had obvious evidence of achalasia. All participants gave their informed consent before manometric examination.

Oesophageal Transit Studies

Supine oesophageal transit was measured according to a standard protocol. A 5 ml bolus of technium 99m sulphur colloid (150 μCi, 5-55 MBq) was placed in the mouth and the patient was asked to swallow once. Gamma emissions were counted, over the thorax, for the next 60 seconds using a wide field of view gamma camera (Siemens 37 ZLC) and the data were analysed by computer. From previous validation studies, transit was considered to be abnormally prolonged if the time taken for the level of gamma emission over the thorax to fall below 10% of its original activity was greater than 10 seconds.

Oesophageal Manometry

Intraluminal oesophageal pressure activity was measured using a standard multilumen perfused tube system. The tube was constructed from 12 radiolucent polyvinyl chloride capillary tubes (internal diameter (ID) 0.63 mm, external diameter (ED) 1.4 mm) bonded around a central radio-opaque PVC tube (ID 1.5 mm, ED 2.5 mm). A balloon, constructed from a 5 cm length of condom rubber, was attached 15 cm from the distal end and was inflated with water through another PVC tube (ID 1 mm, ED 1.6 mm) (Fig. 1). Each channel was perfused with distilled water, at 0.4 ml per minute, through a pneumatic system which gave a pressure response of 100 mmHg/sec/m on occlusion of the most distal port. Intraluminal pressure changes were detected, proximal and distal to the balloon, using strain gauge transducers (Gaeltec 8th, Skye, Scotland) attached to the proximal end of each perfused lumen, and were recorded on a multi-channel chart recorder (Watanabe Linear Corder mark VII, Tokyo, Japan) at a paper speed of 100 mm/min.

Swallowing was detected via the most proximal channel of the assembly, which was sited in the lower oropharynx, thus enabling primary and secondary motor activity to be distinguished.
lay just below the diaphragm, identified by positive inspiratory pressure changes. This manoeuvre ensured that the balloon was consistently positioned 10–15 cm above the diaphragm in all studies. Thirteen of the normal subjects and all the patients were studied over three consecutive five minute periods. Control periods of recording, without balloon distension, were done before and after a test period during which the balloon was slowly distended with water until retrosternal discomfort was just occurred. Persistence of balloon inflation throughout the study period was confirmed by complete recovery of the water instilled into the balloon, at the end of the procedure. Throughout the study the tube was held in position by the observer, at the angle of the mouth, to prevent tube displacement and to detect the presence of any traction forces applied to the tube.

Pilot studies indicated that balloon distension caused an increase in the volume of salivation and swallowing frequency. As swallowing activity may inhibit the generation of spontaneous oesophageal contractions, all individuals were asked to restrict the frequency of swallowing during both the test and control periods and to help control swallowing frequency saliva was continuously aspirated from the mouth during the test period.

Six of the 13 subjects underwent a repeat study after an intravenous injection of the cholinergic antagonist (Hyoscine Butylbromide, 20 mg). This was administered at the beginning of balloon distension.

Three additional subjects were studied during a period of subthreshold oesophageal distension. This was achieved by inflating the balloon until retrosternal discomfort was just felt, and then removing half the volume instilled into the balloon for the study period.

DATA ANALYSIS
Standard oesophageal manometry
The oesophageal motility recordings were examined by inspection and measurements of the amplitude and duration of intraluminal pressure changes were made manually.

Distension studies
The number of contractions in each channel were counted during both the control and test periods and divided into those related to swallowing (primary) and those unrelated to swallowing (secondary). Comparison of the number of primary and secondary contractions was then made between the two control periods and between the pre-inflation period and the test period. Initial inspection of these data indicated they were positively skewed, a square root transfor-
mation was therefore performed, before calculating means and standard deviations, in order to render their distribution more normal. The mean (and confidence limits) results quoted in the text are the retransformed values calculated from the square rooted data. To assess the effect of distension on the propagation of swallowing activity, the ratio of primary contractions below and above the balloon was calculated using the channels 5 cm distal and 5 cm proximal to the balloon during both the control and test periods. These data appeared normally distributed and thus required no transformation. In all cases statistical comparison was performed using a paired *t* test to determine the probability that the observed differences could have occurred by chance.

**Results**

**Normal subjects**
The volunteers all had a normal standard manometric examination.

**Control period**
During the control period (Fig. 2) normal primary peristaltic activity was seen together with a few spontaneously occurring secondary contractions. There was no significant difference in the frequency of either primary or secondary contractile activity during the predistension and postdistension periods (Fig. 3).

**Balloon distension**
Balloon distension induced retrosternal discomfort described in similar terms by all 13 subjects (median volume of water instilled 13 ml, range 6–30 ml; balloon diameter 0.8–3.8 cm), and was accompanied by an alteration in the pattern of oesophageal motor activity. This alteration always preceded the perception of the retrosternal discomfort. A marked increase in the number of secondary contractions occurred above the balloon (Figs. 2, 3), the mean (confidence limits) number changing from 2.0 (0.9–3.7) secondary contractions/5 minutes during the predistension period to 18.1 (12.1–25.1) during the distension period (*p*<0.001). Primary peristaltic activity started and propagated normally above the balloon during both the test and control periods (Fig. 3). During distension, however, the number of primary contractions traversing the balloon was markedly reduced, the ratio of contractions either side of the balloon (distal/proximal) changing from 1±0.18 (mean±confidence limits) for the predistension period, to 0.38±0.18 for the distension period, (*p*<0.001). These changes in motor activity were associated with the development of a variable but persistent aborally directed force on the tube, which was detected manually by the observer and had to be resisted by traction to prevent tube displacement.

On balloon deflation the pattern of motor activity rapidly returned to normal; no specific pattern of activity was related to deflation.

---

*Fig. 2 Oesophageal contractile activity during the predistension (left) and the distension (right) periods in a normal subject. During the predistension period normal primary peristalsis is seen together with one secondary contraction. In the distension period there is an increase in secondary contractile activity above the balloon while the propagation of contractions across and below the balloon is reduced.*
observed. On these occasions, however, primary peristalsis traversed the balloon and migrated through the distal oesophagus suggesting that distal inhibition required a greater distension threshold than proximal stimulation.

**Cholinergic blockade studies**
Administration of hyoscine butylbromide (20 mg) during balloon distension did not alter the sensation of retrosternal discomfort and had no effect on either the initiation or propagation of the primary peristalsis proximal to the balloon. In contrast the distension induced increase in secondary contractile activity was virtually abolished, within one minute of drug injection; the mean number of contractions changing from 22.1 (11.3–36.5) per five minutes for the distension period without cholinergic blockade to 2.9 (0.2–8.8) during cholinergic blockage (p<0.01). The persistant aborally directed force on the tube was also abolished.

**PATIENT STUDIES**
Details of the symptoms and investigations carried out on the patients are shown in the Table. All suffered from chronic difficulty in swallowing and had delayed isotope transit, but normal oesophageal endoscopy. Barium swallow study suggested a failure of lower oesophageal sphincter relaxation in four patients (case numbers 3–6) although oesophageal diameter appeared normal. On manometry three of the patients (case numbers 3–5) showed abnormal lower oesophageal sphincter responses, with absent relaxation on swallowing, suggesting early achalasia. Abnormalities of primary peristalsis were present in five people; three (case numbers 5, 8, and 9) had multiphasic waves consistent with a diagnosis of diffuse oesophageal spasm,11 the others (case numbers 4 and 6) had low amplitude peristalsis. In the three remaining cases (case numbers 1, 2, and 7) no abnormality was detected on either standard manometry or other investigations.

Balloon inflation caused retrosternal discomfort in all the patients at similar volumes to those used in the normal subjects (Median volume of water instilled 15 ml, range 4–30 ml), indicating similar distension thresholds. As with the normal subjects neither the frequency of swallowing nor its propagation proximal to the balloon was altered during distension. In contrast with the normal subjects, however, six of the nine patients (case numbers 1–4, 6, and 8) failed to show an increase in proximal secondary contractile activity (Figs 4, 5, 6), the mean number of secondary peristaltic contractions proximal to the balloon being 0.2 (0–1) for the predistension period and 0.4 (0–1.6) for the distension period (p>0.05). Furthermore all primary peristaltic contractions propagated across

---

**Subthreshold distension studies**
In the three individuals in whom the balloon was distended without retrosternal discomfort (volume of balloon inflation 3.5, and 7 ml), the proximal stimulation of secondary contractile activity was still
Motor responses of the oesophagus to intraluminal distension in normal subjects

Control

<table>
<thead>
<tr>
<th>Pharyngeal</th>
<th>Distension</th>
</tr>
</thead>
<tbody>
<tr>
<td>10 cm</td>
<td></td>
</tr>
<tr>
<td>15 cm</td>
<td></td>
</tr>
</tbody>
</table>

Balloon

| 20 cm      |            |
| 25 cm      |            |
| 27.5 cm    |            |
| 30 cm      |            |

Fig. 4 Effect of oesophageal distension in a patient (no 1) with normal standard manometry. During the control period normal peristalsis is seen. In contrast with Figure 2 there is neither a proximal increase in secondary activity nor a distal inhibition of primary peristalsis during distension.

directed motor inhibition.

Discussion

Our results show that balloon distension of the normal human oesophagus induces a specific motor response, characterised by increased secondary motor activity above, and reduced motor activity below, the distension together with a strong aborally directed propulsive force. The term secondary contractile activity, rather than secondary peristalsis, has been used to describe this response because the activity observed by us differs from that previously described as secondary peristalsis. Experimentally induced secondary peristalsis usually refers to distally propagating motor activity induced by deflation rather than that seen during distension. This may in time prove to be an unnecessary distinction but has been made to avoid possible confusion until the exact nature of this motor activity and its relationship to previous studies has been better defined.

Other workers have studied the motor responses of the human oesophagus to distension, but the results have been variable possibly because of differ-

Fig. 5 Effect of oesophageal distension in a patient (no 6) with oesophageal hypomotility is shown. Poor primary peristalsis is seen during both control and distension periods, secondary contractile activity did not increase during distension.
ences in the experimental protocols. Some workers have observed proximal motor excitation together with distal inhibition but others recognised no consistent pattern of motor activity even though an aboral force was noted. These findings contrast with our normal studies where proximal excitation was consistently seen. Such differences result from the type of distension used. We used a sensory end point to define adequate distension while others have used predetermined volumes, which may have produced more variable distension.

Motor responses to oesophageal distension have also been studied, both in vivo and in vitro, in animals. Similar motor responses have been recorded under both experimental conditions indicating that the mechanisms for producing such motor activity are intrinsically mediated, though these responses can be modified through the vagus nerve. Three motor responses of the opossum oesophagus in response to distension have been described. On balloon inflation there was a myogenically mediated response, the 'on response', during distension a cholinergically mediated contraction of the proximal longitudinal muscle, the 'duration response' and on balloon deflation a distally propagating contractile wave starting just above the level of the distension, the 'off response'. The 'off response' appears to be mediated via non-adrenergic non-cholinergic nerves and is manometrically similar to secondary peristalsis. It is likely that the activity observed in our studies is the same as the 'duration response' because of the temporal association with the distension and neuropharmacological similarity of the activities. The nature of the distal inhibition has not been defined in our studies but is probably mediated through a non-adrenergic non-cholinergic pathway, similar to that responsible for lower oesophageal sphincter relaxation.

The different stimulation thresholds required for the proximal and distal responses also seem of interest. The proximal increase in activity was consistently initiated at a level of distension below that required for its perception. The distal inhibition, in contrast, did not occur until greater degrees of distension were induced and discomfort was felt. The proximal excitation may therefore be the more usual mechanism for clearing retained luminal contents which stimulate stretch receptors. The latter motor inhibition is probably not a mechanism for food transport under normal circumstances, although it might represent a subsidiary mechanism to encourage the clearance of larger boluses which would otherwise block the oesophagus.

Our patient studies appear to be the first to have applied intraluminal distension to the investigation of oesophageal transport problems and raise a number of issues for discussion. All the patients had severe chronic symptoms, which together with the prolonged oesophageal radionuclide transit, suggest that, in addition to inefficient primary peristalsis, the normal non-swallowing initiated mechanism for oesophageal clearance was impaired. The defective motor response to intraluminal distension in these people supports the idea of an abnormality of this secondary clearance mechanism. One group of patients (case numbers 3–5, 8, and 9) seem to belong to the spectrum of achalasia/diffuse oesophageal spasm as they show features of abnormal lower oesophageal sphincter relaxation or multi-peaked primary peristaltic waves or both. A diffuse abnormality of the intrinsic oesophageal nerves is well recognised in achalasia and presumably a similar
abnormality occurs in diffuse oesophageal spasm, although data in such disorders seem lacking.

A further group (case numbers 1–3) had normal standard manometry but a prolonged isotope transit. Abnormal clearance in the presence of normal primary peristalsis has been recognised by other workers but why this should occur has not been resolved. Studies comparing the amplitude of oesophageal contraction with the force acting on a bolus have shown the two are not related so that contractions sufficient to produce normally appearing peristaltic waves during oesophageal manometry are not necessarily sufficient to ensure adequate oesophageal transport. It is important to note that all three cases failed to show any abnormality of oesophageal function except prolonged supine transit and an abnormal distension response, so that standard manometry alone would have failed to recognise any abnormality. These patients presumably suffer from a defective intrinsically mediated oesophageal clearance response.

Patient number 6 had low pressure primary peristaltic activity. Though such activity makes interpretation of the distension response difficult, failure to initiate any proximal motor excitation again suggests an intrinsic neural abnormality to be present.

Finally, two of the patients (case numbers 5 and 9) showed delayed transit with normal distension responses. A possible reason for the lack of abnormality in these patients is that the distension technique used was too crude to show a more subtle peristaltic impairment. Studies using graded submaximal stimulation, at levels of intraluminal distension, more likely to occur after food ingestion, might have identified abnormalities in such patients.

In conclusion, it appears that investigation of the oesophageal motor responses to distension could be a useful adjunct to standard oesophageal manometry in the investigation of patients with suspected oesophageal clearance disorders, particularly when standard manometry is normal. Distension is simple to do, increases the duration of oesophageal manometry by only a few minutes and seems to provide useful information about oesophageal motility which cannot otherwise be obtained. Further studies of these responses, in patients with both structural and functional oesophageal dysfunction using graded submaximal distension are required so that the specificity and diagnostic potential of this technique can be more fully evaluated.

Dr Kendall is a WEG Knott Research Fellow of the British Digestive Foundation. Dr Thompson is a Wellcome Trust Senior Lecturer in Medicine.

We would like to thank Professor Lennard-Jones for help in preparing this manuscript and Miss E Walker for assistance in the studies.

References