Secondary oesophageal peristalsis in patients with non-obstructive dysphagia

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Abstract
Secondary peristalsis was investigated in 30 patients with non-obstructive dysphagia and 20 age matched controls. Oesophageal motility was recorded at 3 cm intervals along the oesophageal body. Primary peristalsis was tested with 5 ml water swallows. Secondary peristalsis was stimulated with 10 ml boluses of air and water injected in the mid-oesophagus and by distensions (5 seconds duration) with a 3 cm balloon at the same level. Primary peristalsis was normal in 19 of the 20 control subjects and in nine of the 30 patients with dysphagia; 11 patients had diffuse spasm and 10 had non-specific abnormalities of primary peristalsis. Secondary peristalsis was triggered significantly less frequently by air and water distension in dysphagia patients (median success rate of 10% for the air boluses and 0% for the water boluses) than in control subjects (50% and 30% respectively, p<0.005), and was abnormal in six of nine patients with normal primary peristalsis, nine of 11 patients with diffuse spasm and eight of 10 patients with non-specific motor abnormalities. The median frequency of balloon induced secondary peristalsis, however, was not significantly different in the two groups (0% controls, 40% non-obstructive dysphagia, p=0.22). For each stimulus, there were no significant differences in the response rate in the three subgroups of patients. The major pattern of failure of secondary peristalsis in response to the air and water boluses was the complete absence of any oesophageal response. The amplitude of complete secondary peristalsis triggered by the water boluses and the balloon was greater in the patients with dysphagia (p=0.03) than in normal subjects, while the amplitude of the secondary peristaltic responses triggered by the air boluses was similar in the two groups. Secondary peristaltic velocity was also similar in normal subjects and patients with non-obstructive dysphagia. Patients with non-obstructive dysphagia show a noticeable defect in the triggering of secondary peristalsis which may make an important contribution to the delayed oesophageal bolus transit and dysphagia seen in this condition.

Methods
SUBJECTS
We studied 30 patients with non-obstructive dysphagia (17 women and 13 men), ranging in age from 26–74 years (median 59±5 years). All complained of dysphagia as their predominant symptom but nine patients also had chest pain. Mechanical obstruction was excluded in all patients by either endoscopy (n=29) or barium swallow (n=26), or both. Patients taking medication that might affect oesophageal motility were asked not to take this from at least 48 hours before the study.

Twenty age matched healthy subjects (eight women and 12 men), ranging in age from 23–76 years (median 50 years) served as controls. Subjects were free of gastrointestinal symptoms, had no history of upper gastrointestinal surgery, and were not taking antacids regularly or any medication known to change oesophageal motor function. All patients and volunteers gave written informed consent and the study was approved by the Human Ethics Committee of the Royal Adelaide Hospital.
MANOMETRY
Oesophageal motility was recorded with a 13 lumen manometric assembly made from a 4.5 mm diameter silicon rubber extrusion that incorporated a 6 cm sleeve sensor. Side holes, spaced 3 cm apart, recorded pressures from seven sites along the oesophageal body starting at 2 cm above the proximal margin of the lower oesophageal sphincter. A side hole in the gastric fundus recorded gastric pressure and a side hole in the pharynx monitored swallowing. A 3 cm long silicon rubber balloon, which fitted tightly around the catheter assembly when deflated provided focal distension 12-5 cm above the lower oesophageal sphincter. An infusion port was located immediately above the balloon for the rapid injection of boluses of air or water. The catheter was fixed and maintained in position so that the mid portion of the sleeve sensor was located within the lower oesophageal sphincter. The oesophageal and gastric side holes and the sleeve sensor were perfused with degassed distilled water at 0.6 ml/minute by a low-compliance pneumohydraulic capillary infusion system.14 The pharyngeal side hole was perfused at 0.3 ml/minute. Pressures were sensed by external pressure transducers (Deseret Medica Inc, Park Davis & Co, Sandy, Utah, USA; model 38-8000-1) with output to a 12 channel polygraph recorder (Grass Instrument Co, Quincy, MA, USA; model 7D). Recordings were made at a paper speed of 5 mm/sec.

STUDY PROTOCOL
Subjects were studied after an overnight fast. The catheter was passed pernasally. After a 10 minute adaptation period, both primary and secondary peristalsis were tested in each subject.

Primary peristalsis was tested with 10, 5 ml water swallows. Each swallow was separated by an interval of 30 seconds. Secondary peristalsis was triggered by oesophageal distension with 10 ml boluses of air and water and by inflating the intraoesophageal balloon. Each stimulus was tested five times. Balloon distension was performed in only 23 of the 30 patients because of technical problems such as balloon rupture or intolerance of balloon distension due to chest discomfort. The air and water boluses were rapidly injected through the infusion port by hand. The 10 ml air bolus was injected within 0.5 second, while the 10 ml water bolus was delivered within 1.5 seconds. The balloon was inflated with 17 ml of air within 0.5 seconds to a diameter of 3 cm and sustained for 5 seconds before being quickly deflated over 0.5 seconds. The distending stimuli were given at least 15 seconds after any preceding primary peristaltic wave, and an interval of 20 seconds was allowed after each stimulus for any response to occur. This interval was selected as it is comparable to the latency of the secondary peristaltic response after physiological reflux events.15 During this time the subjects were instructed not to swallow. At the end of the 20 second period the subject was asked to perform a dry swallow that served to reduce the desire of the subject to swallow during the distension stimulus as well as to clear any residual air or water.

DATA ANALYSIS
The contraction amplitude at each recording site and the latency of the wave onset between adjacent recording sites were determined for both primary and secondary peristalsis. Amplitude was measured from basal end-expiratory intraoesophageal pressure to the peak of the pressure wave. The onset of the major upstroke of the pressure wave was used as the reference point for determination of the wave latency.

Primary peristalsis was classified as complete if a propagated pressure wave of ≥12 mm Hg at the proximal two oesophageal recording sites and ≥25 mm Hg in the distal five oesophageal channels, traversed all the recording sites.16,17 The minimum latency of wave onset between adjacent recording sites that defined peristaltic progression was set at 0.5 seconds, corresponding to a peristaltic velocity of 6 cm/sec.17 Criteria for failed peristalsis were either failure of a pressure wave ≥12 mm Hg in the proximal two recording sites and ≥25 mm Hg in the distal five channels to traverse all of the oesophageal recording sites or synchronous pressure waves occurring at two or more recording sites. In addition, no response to distension was judged to have occurred if a pressure wave ≥10 mm Hg was seen in less than two recording sites. For complete peristaltic responses, mean amplitude and velocity were calculated for the five distal recording sites corresponding to the distal 12 cm of the smooth muscle segment of the oesophageal body. Normal primary peristalsis was defined as the occurrence of eight or more complete peristaltic responses to the 10 water swallows.18

Secondary oesophageal motor responses to air and water boluses were typically characterised by a propagated pressure wave that traversed the entire oesophagus,19 and were analysed according to the criteria given above for primary peristalsis. The response to balloon distension, however, was different from the response triggered by the air and water injections as there were separate responses above and below the balloon.19 During balloon inflation, a sustained pressure wave was seen above the balloon with inhibition of motor activity below. After balloon deflation, the pressure wave above the balloon subsided and a motor response was seen below the balloon. Each component of the balloon response was analysed individually using the same criteria as for primary peristalsis.19

STATISTICAL ANALYSIS
The frequencies of successful primary and secondary peristalsis were determined for each stimulus in each subject. Differences
in response rates among stimuli were analysed using log linear modelling techniques and non-parametric tests for paired and unpaired data. Data for peristaltic amplitude and velocity and the pattern of oesophageal motor responses to distension were pooled and subjected to analysis of variance. Group data for response rates are expressed as median values (interquartile range) and those for peristaltic amplitude and velocity as mean (SEM).

Results

**PRIMARY PERISTALSIS**

The rate of successful primary peristalsis was significantly lower in the patients with dysphagia (50% (29%-90%)) than in the normal subjects (90% (80%-100%), p<0.005). Nineteen of the 20 normal subjects and nine (30%) of the 30 dysphagia patients exhibited normal primary peristalsis (Fig 1). Conversely, 21 (70%) of the 30 dysphagia patients had abnormal primary peristalsis, with a median response rate in this group of 40% (10%-50%). Eleven of the 21 patients were diagnosed as having diffuse oesophageal spasm and eight had non-specific motor abnormalities including two with high amplitude peristaltic contractions in the distal oesophagus. The mean contraction amplitude and propagation velocity for the primary peristaltic responses are summarised in the Table. Peristaltic amplitude was greater in the patients with dysphagia than the normal volunteers (p<0.0005) but the primary peristaltic velocities were similar in the two groups (p=0.7).

**SECONDARY PERISTALSIS**

**Air and water boluses**

Peristaltic response rates to air boluses were significantly lower in the patients with dysphagia as a group (10% (0%-20%)) than in the normal subjects (50% (20%-90%), p<0.002) (Fig 2). Response rates in the subgroups of patients with normal primary peristalsis (20% (0%-40%)), diffuse spasm (20% (0%-20%)), and non-specific motor abnormalities (0% (0%-20%)) were not statistically different and were each less than that in normal subjects. Using the fifth centile (20%) as the lower limit of normal, secondary peristaltic responses to air were abnormal in six of nine patients with normal primary peristalsis, nine of 11 patients with diffuse spasm, and eight of 10 patients with non-specific motor abnormalities.

Water boluses produced similar results with a median response rate of 0% (0%-20%) in patients with dysphagia compared with 30% (10%-80%) in the normal subjects (p<0.005). Response rates in the subgroups of patients with diffuse spasm (20% (0%-20%)) and non-specific motor abnormalities (0% (0%-20%))
The pattern of the motor responses for the air and water boluses are shown in Figure 3. In the normal subjects, 52% of the air boluses triggered a peristaltic response while 21% failed to trigger any response at all. A small proportion of air boluses triggered failed or synchronous responses. In the patients with dysphagia, however, only 18% of air boluses triggered secondary peristalsis and 43% produced no response. The proportion of failed and synchronous responses were similar in the patients and controls, and among the three subgroups of patients. Water boluses resulted in a similar patterns of responses in the two groups.

**Balloons distension**

The frequency of secondary peristaltic responses seen distal to the balloon after deflation is shown in Figure 4. The response rate below the balloon varied widely among subjects and the mean (SEM) values in the patients with dysphagia (20% (0-60%)) were not statistically different from those in the normal subjects (0% (0-70%); p=0.22). In the normal subjects, the response rates to balloon distension was significantly less than than for the air boluses (p<0.04) but similar to that for the water boluses. In the dysphagia patients, the secondary peristaltic response rate to balloon distension was greater than that for air (p<0.05) and water boluses (p<0.004).

The contraction amplitude of secondary peristalsis elicited by balloon distension was greater in the dysphagia patients than the normal subjects (p=0.005) while the propagation velocities were similar in the two groups (Table).

The pattern of balloon responses are shown in Figure 5. There were no significant differences in the proportions of the motor responses either above or below the balloon between the two groups nor among the three subgroups of patients.

**Discussion**

Non-obstructive dysphagia is a common indication for referral for oesophageal manometry as it is generally believed to be a result of oesophageal motor dysfunction. In a significant proportion of patients, however, manometric assessment of primary peristalsis fails to reveal any abnormality and the cause of the symptoms remains unclear. One possible explanation for this discrepancy is that testing of primary peristalsis is not sufficiently sensitive to detect subtle oesophageal motor abnormalities. This notion is supported by several studies showing radiological and scintigraphic abnormalities in patients with dysphagia and normal primary peristalsis.

Abnormal secondary oesophageal peristalsis is a potential mechanism that might account for these discrepancies, as secondary peristalsis is an important mechanism of oesophageal volume clearance. This study is the first assessment of secondary peristalsis in patients with non-obstructive dysphagia and shows that they commonly have defective secondary
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The mechanisms underlying defective secondary peristalsis in response to the air and water injection remain to be defined, but inferences are possible from the pattern of the responses and previous observations. As secondary peristalsis is a reflex response to oesophageal distension, the defect may lie either in the oesophageal motor nerve supply or muscles, or in oesophageal sensation, or perhaps both. In patients with abnormal primary peristalsis, it is possible that there is a defect in the efferent limb of the reflex arc, although the pattern of abnormal primary peristalsis did not have any significant bearing on the pattern of abnormal secondary peristalsis. Interestingly, in our study some patients with abnormal primary peristalsis had normal secondary peristalsis suggesting that the defect in primary peristalsis in these patients lies in the central control mechanisms rather than in the peripheral neural pathways or oesophageal muscle. In patients with both abnormal primary and secondary peristalsis, however, the defect is more difficult to localise.

In patients with normal primary peristalsis and abnormal secondary peristalsis, the efferent pathways seem to be intact and this suggests that there is a defect in the afferent limb of the reflex pathway with impaired oesophageal sensitivity to distension. This hypothesis has not been examined in patients with non-obstructive dysphagia. In patients with oesophagitis, however, Williams et al noted that the distension threshold required to trigger a motor response to prolonged oesophageal distension was higher than that in healthy controls. Deschner et al have reported that the distension threshold for symptoms was lower in patients with non-obstructive dysphagia but the volume required to elicit symptoms has been shown to be different from the volume required to trigger an oesophageal motor response.

Secondary peristaltic responses to balloon distension were not significantly different between normal subjects and patients with dysphagia. This was due largely to a low response rate in the normal subjects. We observed similar findings in a study of secondary peristalsis in gastro-oesophageal reflux disease and this supports our conclusion from other studies that balloon distension is an ineffective method of testing secondary peristalsis. The apparent discrepancy between the balloon responses and those to air and water boluses most likely reflects differences in the nature of the distending stimulus, the balloon providing a focal and fixed distension in contrast to the more diffuse and mobile stimulus with air and water boluses.

Although secondary peristalsis has not been formally examined in patients with non-obstructive dysphagia, recent studies have shown that oesophageal motor responses to prolonged distension are abnormal in these patients. These studies showed abnormal

Figure 5: Patterns of manometric response to balloon distension. The proportion of various patterns of manometric response are derived from pooled data of the individual responses. Responses have been divided into those occurring above and below the balloon. The responses above the balloon occurred during inflation and those below the balloon occurred after deflation. The patterns of response were similar in the two groups.
contractions in the distal oesophagus during balloon inflation in patients with non-obstructive dysphagia while the motor responses proximal to the balloon were similar to those in normal subjects. We also noted similar patterns of motor activity above the balloon in the normal subjects and the patients, but failed to detect any motor activity in the distal oesophagus during balloon inflation. A possible explanation may be the greater balloon diameter used in our study leading to a greater degree of distal inhibition. Alternatively, we used sudden and relatively brief balloon distension rather than gradual distension (1 ml/3 sec) which may have influenced the pattern of the response.

Evaluation of the manometric characteristics of the secondary peristaltic responses showed a higher mean peristaltic amplitude in the patients with non-obstructive dysphagia than in the normal volunteers, while the propagation velocity in the two groups was similar. The importance of this finding is not clear but it is interesting to note that a similar difference is seen for primary peristaltic amplitude and perhaps reflects a generally increased oesophageal contractile strength. Similar results were reported by Jacob et al who found higher primary peristaltic amplitudes in patients with dysphagia than in patients without dysphagia.1

In summary, we have shown a noticeable defect in secondary peristalsis in patients with dysphagia. This defect may well result in delayed bolus transit along the oesophagus and contribute to the development of the patient’s symptoms. Examination of secondary peristalsis may be useful in the diagnostic evaluation of patients with non-obstructive dysphagia.

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