Anomalies of peristalsis in idiopathic diffuse oesophageal spasm

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SUMMARY Oesophageal manometry revealed ‘interrupted peristalsis’—proximal sequential contraction with simultaneous pressure increases in the mid-oesophagus and sequential contraction distally—and abnormally slow distal propagation of peristalsis in seven of a consecutive series of 12 patients with idiopathic diffuse oesophageal spasm (IDOS). These two abnormalities occurred together in six patients. It is proposed that ‘interruption’ of peristalsis and ‘slow’ distal propagation are apparent rather than real, and that they reflect failure of normal luminal obliteration during oesophageal contraction. This argument is extended to the ‘aperistalsis’ of achalasia. It is also proposed that IDOS, achalasia, and related idiopathic motor disorders may share a common pathogenetic abnormality—namely, functional obstruction at the level of the lower oesophageal sphincter. According to this hypothesis, the presentation of the patient and the progression of his disease are determined by the degree and duration of that obstruction, and the functional reserve of the oesophageal body musculature.

Idiopathic diffuse oesophageal spasm is characterised by symptoms of chest pain, dysphagia, or both and by radiographic abnormalities ranging from tertiary waves to the classical ‘corkscrew’ oesophagus. Manometric abnormalities are variable; and there are no uniform, generally accepted criteria for manometric diagnosis. Most would agree, however, that the typical patient with IDOS has powerful, prolonged, repetitive, and simultaneous contractions in the lower half or two-thirds of the oesophageal body.\textsuperscript{1} Some,\textsuperscript{2,3} but not all, would insist that the absence of peristalsis—that is, axially sequential contraction—in this segment excludes IDOS. In such patients, the distinction between IDOS and ‘vigorous achalasia’ becomes blurred. Another apparent variant of IDOS, initially described by Pope\textsuperscript{4} and subsequently given, by Benjamin et al.,\textsuperscript{5} the appellation ‘nutcracker oesophagus’, is characterised by powerful, prolonged but uniformly sequential distal oesophageal contractions.

A review of manometric records obtained at the University of Vermont from patients with IDOS during the 18 month period from January 1979 to June 1980 revealed two additional interesting abnormalities. One, the occurrence of simultaneous pressure waves in the mid-oesophagus with sequential contraction below that level, hereafter referred to as ‘interrupted peristalsis’, has not to my knowledge been described before. The other, abnormally slow propagation of oesophageal contraction in the distal segment, was reported previously by us in certain patients with Parkinson’s disease.\textsuperscript{6} A description of these abnormalities and discussion of their interpretation form the basis of this report.

Methods

Patients with IDOS

Between January 1979 and June 1980, a diagnosis of idiopathic diffuse oesophageal spasm (IDOS), initially suspected on the basis of clinical and radiographic findings, was established by manometry in 12 patients. This group comprised nine women and three men. Ages ranged from 29 to 89 years, with a mean of 61.6 years. One patient presented with odynophagia alone. Each of the remaining 11 complained of dysphagia, which was associated with chest pain in five, eructation in two, regurgitation of food in one, and weight loss in one. None had symptoms of gastrooesophageal reflux. In the one
patient whose barium swallow was reported to be normal, manometry revealed extremely powerful (up to 470 mm Hg; 62.5 kPa), repetitive pressure waves, which were frequently simultaneous, in the lower half of the oesophageal body. Radiographic evidence of incoordination, ranging from prominent tertiary waves to a classical ‘corkscrew’ oesophagus, was present in the remaining 11. In addition, five had an obvious hiatus hernia, two a B (Schatzki’s) ring, two had an A ring, and one had a lower oesophageal diverticulum. Manometry in these patients revealed powerful simultaneous pressure waves, which were frequently prolonged, repetitive, or both, in the distal half to two-thirds of the oesophageal body. Intermittent sequential contractions were recorded from the distal oesophagus in 10 of the 12 patients. All showed sequential contraction in the proximal oesophagus.

**Comparison group**

In order to provide a frame of reference for the interpretation of measurements in patients with IDOS, data on the manometric characteristics of lower oesophageal contraction in 10 healthy subjects are presented. This group comprised nine men and one woman. Ages ranged from 19 to 38 years. None had oesophageal symptoms. All gave informed consent, and performance of manometric studies in them was approved by the University of Vermont Committee on Human Experimentation for the Medical Sciences.

**Oesophageal manometry**

Patients with IDOS were studied with a commercially manufactured tube assembly (Arndorfer Medical Specialties, Greenland, Wisconsin). This has four openings for pressure transmission at the same axial level of the assembly, with four additional openings spaced at 5 cm intervals above that point. Normal subjects were studied with a simple triple-lumen tube, with openings spaced at 5 cm intervals. Each lumen was perfused at a rate of 0.6 ml/min by a pneumohydraulic capillary infusion system (Arndorfer Medical Specialties). Pressures were measured by external transducers (Bell and Howell P 327–1), the outputs of which were recorded on a Gilson Macropolygraph multichannel recorder. Respiration and swallowing were monitored by pneumograph and myograph respectively.

In patients with IDOS, lower oesophageal sphincter pressures (LOSP) were measured simultaneously by the four sensors located at the same axial level of the assembly. A station withdrawal method was used. LOS pressures were expressed as the mean of the end-expiratory measurements obtained by the four sensors, with gastric baseline pressure being used as zero reference. Oesophageal contractile responses to ‘dry’ and ‘wet’ (5 ml boluses of water at room temperature) swallows were then recorded by openings spaced at 5 cm intervals.

In normal subjects, the responses to 10 ‘dry’ and 10 ‘wet’ swallows were recorded with pressure sensors 2, 7, and 12 cm above the LOS.

**Results**

Lower oesophageal sphincter pressures in the group of patients with IDOS tended to be high. They ranged from 2.0 to 7.58 kPa (15 to 57 mm Hg) with a mean of 3.57 kPa (31.5 mm Hg). It should be emphasised that these are end-expiratory pressures.
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which are almost invariably less than mid-respiratory and end-inspiratory pressures. LOS relaxation in response to swallowing occurred in all patients, but was judged to be incomplete in eight.

'Interrupted peristalsis'—defined as simultaneous rise in pressure at two or more proximal recording points with a later pressure rise at a distal recording point—was observed in seven of the 12 patients. In each it was an intermittent phenomenon. It is illustrated in Fig. 1. 'Slow distal propagation', also shown in Fig. 1, and defined as a time interval between onset of pressure rise at the two most distal recording points which exceeded the greatest value (2.4 s) recorded in any of the normal subjects, was observed in seven patients with IDOS. The maximum interval in these seven patients ranged between 3.4 and 6.4 seconds. Again, 'slow distal propagation', like 'interrupted peristalsis', was intermittent. It is important to note that six of the seven patients with 'slow distal propagation' also had 'interrupted peristalsis'.

Maximum values for distal oesophageal contractile amplitude, duration, and propagation time are shown in Fig. 2. Values for amplitude and duration exceeded, often by a substantial margin, the highest normal values in nine and 11 patients with IDOS respectively.

Discussion

The normal subjects referred to in this study tended to be younger than, and were not sex-matched with, the patients with IDOS. For this reason, it could be argued that they do not represent an ideal control group. However, results reported recently by Weihrauch et al. would suggest that age and sex per se have little effect on contractile function of the oesophagus; and the only age-related change noted by Hollis and Castell was a reduction in the amplitude of oesophageal contraction. Data from healthy individuals are included in this report primarily to provide a general frame of reference as an aid to the interpretation of measurements obtained in the patients with IDOS. This is important because normal values obtained in any one laboratory depend to a significant degree upon the methods and equipment used.

The primary purpose of this paper is to draw attention to two manometric abnormalities—'interrupted peristalsis' and 'slow distal propagation'—which were observed, almost always in association, in approximately half of a group of 12 patients with IDOS. At first sight, one might interpret the manometric phenomenon of 'interrupted peristalsis' to represent a contraction which passes sequentially along the proximal oesophagus, then occurs simultaneously in a segment of oesophagus, and then again becomes sequential at a lower level. Although our understanding of oesophageal peristalsis is limited, this sequence of events seems inherently improbable. An alternative explanation is presented in Fig. 3. The central concept is that 'aperistalsis'—that is to say, lack of axially sequential contraction—is apparent rather than real. It is proposed that the recording, during manometry, of a simultaneous rise in pressure at two or more points along the length of the oesophagus is due not to simultaneous onset.

Fig. 2 Maximum values for amplitude, duration, and propagation time of distal oesophageal contractions in controls (C) and patients with IDOS (P). Patients with 'interrupted peristalsis' are represented by the open circles.

Fig. 3 An explanation for 'simultaneous contraction'. I, to left of Figure, represents typical diffuse oesophageal spasm. II, in centre of Figure, represents spasm with 'interrupted peristalsis' and 'slow distal propagation'. III, to right of Figure, represents achalasia. See text for detailed explanation.
of contraction at those points but to the transmission of pressure generated at a proximal point to more distal points. This hypothesis requires the existence, either transient at the time of contraction, or persistent, of a cavity below the level of the proximal pressure sensor and containing within it the distal pressure sensor(s). In the first illustration (I) of Fig. 3, a contraction progresses normally from A to B. The oesophageal lumen below point B is open, so pressure rises simultaneously at B, C, and D. If true contraction at C and D is absent, the configuration of the pressure waves at B, C, and D will be similar. However, if significant contraction occurs at C and D, onset of pressure rise at B, C, and D, will remain simultaneous, but the duration of the pressure wave will increase progressively from B to D (hatched lines). Thus, the pressure wave recorded at any given point may be a composite, representing the summation of pressures generated not only at that point itself, but also above it, below it, or both. In section II of Fig. 3, sequential pressure rises are recorded at points A and B. A cavity exists below B, so that pressure rises simultaneously at points B and C. Pressure generated at B is not transmitted to D, as the oesophageal lumen in closed between C and D. However, if true contraction occurs at D, the onset of pressure rise at that point will be not only sequential, but will appear delayed relative to the pressure wave at point C, onset of which reflects contraction at point B. This hypothesis therefore explains not only ‘interrupted peristalsis’ but also ‘slow distal propagation’; and, if correct, it implies that peristalsis is not in fact ‘interrupted’ and propagation of contraction in the distal segment is not necessarily slow. The right half of section II of Fig. 3 depicts another phenomenon that may be observed if points B and C are in open luminal continuity. Pressure generated by contraction at point B causes a simultaneous rise in pressure at C. Subsequent contraction at C will, in turn, if the oesophageal lumen remains open, cause a simultaneous rise in pressure at B. Thus, the pressure wave at each point will have two peaks. An example of this rather commonly observed phenomenon is shown in Fig. 4. Section III of Fig. 3 depicts the situation in achalasia. If the entire oesophageal lumen is dilated and remains open, the pressure wave generated by contraction at the most proximal recording point (A) will be transmitted to B, C, and D. Thus, the entire oesophageal body will appear to contract simultaneously. If oesophageal muscle below point A retains the ability to contract, the pressure wave at distal recording points will appear abnormally long (hatched lines). Contractions will appear feeble, as pressure is dissipated in the large volume of the dilated oesophagus. This general hypothesis is supported by recent observations\textsuperscript{10,11} that peristalsis may ‘return’ after successful treatment of achalasia. Such treatment depends upon relief of distal obstruction (at the LOS) and is accompanied by a reduction of oesophageal calibre. The basic conclusion of this hypothesis is that ‘aperistalsis’ may be a figment of the manometrist’s imagination. The corollaries are that a simultaneous rise in pressure at two or more levels does not necessarily mean simultaneous contraction; that a pressure wave with two or more peaks does not necessarily indicate that the oesophagus at a given level has contracted more than once; and that an abnormally long pressure wave at

\[ \text{Fig. 4} \quad \text{The double-peaked pressure wave. Tracing from patient with IDOS. The interrupted lines join the beginning and end of pressure waves at the three recording points. After the 'dry' swallow (DS) pressure rises simultaneously at all three points. The first peak at 33 and 38 cm is due to pressure transmitted from 28 cm. The second peaks at 33 and 38 cm are sequential, and are due to independent contraction at those points. After the 'wet' swallow (WS), pressure rises sequentially at all three points, and is due to progressive contraction. Pressure generated by the unusually powerful contraction at 38 cm is transmitted upwards. It distorts the descending limb of the pressure wave at 33 cm, and causes a distinct second peak at 28 cm. In this case, the pressure wave ends simultaneously at the three points. These are not true repetitive contractions.} \]
a given level does not necessarily imply that the
duration of muscular contraction at that same level
is abnormal. I would emphasise, however, that the
duration of contraction is truly prolonged in at least
some patients with IDOS; and that true repetitive
contractions undoubtedly do occur, not only in the
lower oesophagus in certain patients with IDOS,
but also in the proximal oesophagus in some patients
with scleroderma. Such contractions probably
represent a response to incomplete oesophageal
emptying.

The relationship between achalasia, IDOS, and
other less well-classified idiopathic disorders of
oesophageal motility remains controversial. Arguing
from the assumptions discussed above, I would
propose that these disorders may share a common
pathogenesis. The basic postulates are that the
initial abnormality in all these disorders may be
functional obstruction at the level of the LOS; and
that abnormalities of motility in the oesophageal
body may be secondary consequences of this ob-
struction. Patients with achalasia have, by definition,
failure of LOS relaxation which, as determined by
manometry, may be partial or complete. Such
dysfunction of the LOS is usually present in 'vigorous
achalasia' and frequently present in IDOS. The
finding of 'normal' LOS relaxation in some patients
with IDOS means only that the LOS has relaxed to
the point where its luminal diameter is as great as
that of the recording catheter. Hence, normal
relaxation on manometry should not necessarily be
equated with normal opening of the LOS. As smooth
muscle undergoes hypertrophy in response to
obstruction, it is germane to note that oesophageal
muscular hypertrophy occurs in both achalasia and
IDOS. If the hypothesis is correct, the likeli-
hood that a patient will present with, or progress to,
florid achalasia will depend upon the magnitude of
resting LOS pressure, the degree and duration of
LOS dysfunction, and the stamina of the oesophageal
body. The following sequence can be envisaged. The
initial response to functional obstruction at the LOS
will be hypertrophy of the oesophageal muscle. So
long as the capacity for luminal obliteration during
contraction is retained, manometry will show force-
ful but sequential contractions, and the oesophagus,
may be characterised as a 'high squeezer' or 'nut-
cracker'. As this capacity is lost in the lower
oesophagus, manometry will reveal simultaneous
pressure waves distally, and may also show 'inter-
rupted peristalsis' and 'slow distal propagation'. As
loss of this capacity for luminal obliteration extends
upwards, manometric abnormalities will evolve
through the spectrum of typical diffuse oesophageal
spasm, vigorous achalasia, and achalasia. Progress-
ion through this spectrum may be halted at any
stage if the oesophageal body has adequate reserve,
or if the degree of LOS dysfunction decreases. In
this connection, it may be pertinent that sliding
hiatal hernia is very rarely associated with achala-
sia, but is common in IDOS. If one believes, as I do,
that the strength of the LOS, and its resting pressure
as recorded by infused-catheter manometry, depend
not only on the intrinsic sphincter, but also on the
external structures, such as diaphragmatic crura,
which normally buttress it, then the development of
a hernia in a patient with IDOS may reduce the
degree of functional LOS obstruction and help to
prevent progression to achalasia.

Patients with symptomatic IDOS often do not
respond well to medical therapy. In those whose
symptoms are severe and refractory, a long myotomy
may be performed. The results of this procedure are
frequently unsatisfactory, with benefit occurring in
approximately two-thirds of patients subjected to it. If
the hypothesis outlined above is correct, it
would be more logical to perform a short myotomy of
the type used in achalasia, as extension of the
myotomy above the LOS would be expected only to
reduce oesophageal propulsive force.

References

1. Creamer B, Donoghue FE, Code CF. Pattern of esoph-
2. DiMarino AJ, Cohen S. Characteristics of lower eso-
ophageal sphincter function in symptomatic diffuse esophageal
3. Mellow MH. Symptomatic diffuse esophageal spasm. Manome-
tric follow-up and response to cholinergic stimulation and cholinesterase inhibition. Gastro-
4. Pope CE II. Abnormalities of peristaltic amplitude and
force—a clue to the etiology of chest pain? In: Van-
trappen G, ed. Proceedings of the Fifth International Sympos-
um on Gastrointestinal Motility. Herentals, Belgium: Typo-
5. Benjamin SB, Gerhardt DC, Castell DO. High ampli-
tude peristaltic esophageal contractions associated with
chest pain and/or dysphagia. Gastroenterology 1979; 77: 478-83.
6. Kaye MD, Hoehn MM. Esophageal motor dysfunction in
Parkinson's disease. In: Vantrappen G, ed. Proceed-
ings of the Fifth International Symposium on Gastro-
intestinal Motility. Herentals, Belgium: Typo-
7. Weihrauch TR, Vallerins P, Alpers H, Ewe K. Assess-
ment of various factors influencing esophageal pressure
measurement. II. Significance of physiological factors in
8. Hollis J, Castell DO. Esophageal function in elderly


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