Detection of subclinical autonomic neuropathy in constipated patients using a sweat test

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Abstract

Chronic idiopathic constipation may be the result of an autonomic neuropathy. This hypothesis was tested in 23 constipated patients and 17 age matched controls, using the acetylcholine sweat spot test devised to test autonomic integrity in diabetes. Acetylcholine (0.01%) was injected in the dorsum of the foot painted with a mixture of starch and iodine. Active sweat glands appeared on the surface of the skin as small black dots which were photographed and counted, using a grid with 60 subareas. Two measurements were made: the number of dots per unit subarea (sweat spot test score) and the % number of abnormal subareas (with less than six spots). These two parameters were correlated. The median sweat spot test score was 9.53 in patients and 13.92 in controls (p=0.0001), the receiver operating characteristic curve showing that a score of 12 delimited normal and abnormal subjects. Increasing age was correlated with a low score in patients, probably because of prolonged symptoms. Seventy per cent of patients and one control had a borderline or abnormal number of subareas. These results suggest that idiopathic constipation is associated with a degree of autonomic denervation. The sweat spot test is an easy, inexpensive method to test this hypothesis and deserves a place in the clinical assessment of slow transit constipated patients.

(Gut 1992; 33: 1539–1543)

An intact autonomic nervous supply to the bowel is essential for normal function. Damage to one or both branches of the autonomic nervous system as a result of trauma, neoplasia, diabetes, infectious diseases, or neuronal degenerative diseases frequently leads to various gut motility (and secretory) problems, the most common being chronic constipation.

For many patients with severe chronic constipation no cause can be found, hence the condition is labelled as idiopathic. Chronic idiopathic constipation, however, is often associated with other motility disturbances involving multiple organs, suggesting an underlying systemic disease of the autonomic nervous system.

Previous attempts to detect an early autonomic neuropathy in these patients, based on the conventional cardiovascular tests suggested by Ewing and Clark, have given inconsistent results, probably because of their poor sensitivity and specificity. Sym pathetic denervation has also been assessed by investigating sweating ability in patients with unexplained gut dysmotility using the thermoregulatory sweat test and quantitative sudomotor axon reflex test. These tests require complex procedures, however, and specialised equipment which are not always readily available. A new test of autonomic integrity, the acetylcholine sweat spot test, has recently been claimed to be more sensitive than conventional tests, and reliable for quantification of the severity of autonomic nervous system damage.

The aim of this study is to investigate whether a subclinical impairment of the autonomic nervous system is present in patients with slow transit idiopathic constipation, and whether it has a possible role in the pathogenesis of this motility disorder.

Methods

PATIENTS

Twenty three patients (21 women, median age 42 years, range 28 to 78) with severe chronic idiopathic constipation entered the study. Severe chronic constipation was defined as constipation without bowel obstruction, lasting for more than 18 months, not responding to high fibre diets, with less than two bowel movements per week and without an alternating bowel habit (diarrhoea and constipation). Each patient had a proctogram to exclude Hirschsprung’s disease and an integrated dynamic defecography to identify anismus or other relevant ano rectal abnormalities. A radioopaque marker test showed that all patients had delayed colonic transit, which was confirmed in 11 patients by a radioisotope transit study. Nine patients had a partial or total colec tomy for constipation before this study.

CONTROLS

Seventeen age matched healthy volunteers (12 women, median age 42 years, range 28–69) served as controls. The exclusion criteria for the selection of controls were constipation or other gastrointestinal problems.

The acetylcholine sweat spot test was carried out injecting 0.1 ml of a 1% solution of acetylcholine (Miochol – CooperVision Ltd, Southampton, UK) intradermally using a 25 gauge needle in a marked area of the dorsum of the foot which had previously been painted with a 2% alcholic iodine solution and a suspension of maize starch in arachis oil (100 g in 100 ml). The reaction between sweat, iodine, and starch could be seen about four to five minutes later as fine black dots corresponding to functioning sweat glands (Fig 1). The marked skin area was photographed and the number of dots counted
unaware of the diagnosis. The final value obtained for statistical analysis corresponded to the median of the three observed values for each patient. The test took about 10 minutes to complete and counting the dots on the photo required 10 to 20 minutes.

As suggested by Ryder and coworkers the occurrence of five (8.3%) or more subareas with less than six dots was chosen as the criterion for abnormality. The mean number of dots/subarea (sweat spot test score) was also recorded for each subject.

**STATISTICAL ANALYSIS**

The Mann-Whitney U test was used in the comparison of the sweat spot test scores between patients and controls, and Fisher’s exact test for the analysis of abnormal subareas. The effect of a confounding factor such as age, and the relationship between the number of abnormal subareas and the sweat spot test score were analysed using the Spearman correlation coefficient (rho) with correction for ties.

The reliability of the method was also investigated by measuring the level of agreement between observers using the K statistic. The subjects were allotted to three arbitrarily defined categories for both the sweat spot test score (>11, 11–9.5, <9.5) and the percentage number of abnormal subareas (0, 0–0.33, >0.33). This was believed necessary, because random agreement between three observers for only two categories – that is, normal/abnormal – is very likely to occur, making results difficult to evaluate.

The sensitivity, specificity, and efficiency of the sweat spot test were calculated for both the sweat spot test score and the percentage number of abnormal subareas, and receiver operating characteristic curves were drawn for both parameters.

**Results**

The median sweat spot test score in patients (9.53, range 1.20 to 13.4) was significantly lower than in controls (median 13.92, range 9.95 to 21.5, p=0.0001, Fig 2). In controls the distribution of dots on the skin surface appeared to be uniform (Fig 1) whereas patients had a more patchy distribution (Fig 3).

There was no significant difference in age between the group of 23 patients (median age 42, range 28–78) and the 17 controls (median age 42, range 28–69). In the control group there was no correlation between age and the sweat spot test score while in patients there was a moderate but significant negative correlation, older patients having a lower sweat spot test score (rho=-0.45, p=0.035, Spearman correlation coefficient).

The number of abnormal subareas – that is, those subareas containing fewer than six dots – were also analysed for differences between the two groups. Because of blurring of the photographic image caused by focusing difficulties in some subjects, the number of subareas where counting was possible was less than the maximum of 60. In order to compare the number of

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**Figure 1:** Photograph of acetyl choline stimulated sweat glands in the dorsum of the foot painted with a maize starch/iodine solution. The artifact in the middle is the point of injection. Control subject, sweat spot test score 16.00. Note the even distribution of glands.

**Figure 2:** Sweat spot test response (SST score, number of sweat glands per unit area) in 23 constipated patients (closed circles) and 17 controls (open circles). The short horizontal line is the median in each group, p<0.0001, Mann-Whitney U test.
When the patients were analysed as a separate group, the correlation between the sweat spot test score and the % number of abnormal subareas was also significant (p=0.0001, rho=−0.91).

In the 21 subjects in whom the number of sweat glands were counted by three independent observers, the correlation between their observations were assessed. The K value was 0.81 (z=11.83, p<0.0001) for the agreement on the sweat spot test score and 0.78 (z=10.65, p<0.0001) for the agreement on the abnormal subareas.

The receiver operating characteristic curve is illustrated in Figure 5. The highest efficiency (83%) occurs with a sweat spot test score of 12 for the demarcation of normal from abnormal subjects (sensitivity 0.83, specificity 0.82). For the abnormal subareas the highest efficiency was with a percentage of abnormal areas of 0.033 or more (sensitivity 0.65, specificity 1.0).

Discussion
The aetiology of chronic idiopathic constipation is uncertain, and attempts to cure it have given poor results. Surgical treatment of severe chronic constipation not responding to medication had an inconsistent and unsatisfactory outcome, resulting sometimes in intractable diarrhoea requiring a definitive terminal ileostomy or in persistence of abdominal pain, bloating, and discomfort.10,11

The mechanisms underlying constipation have been investigated. Involvement of endogenous opioids and their receptors in the regulation of colonic transit was proposed by Kreek12 but further investigation failed to confirm this. The observation that most of the patients with chronic idiopathic constipation, like the population in our study, are women, has suggested the possible role of a disturbed hormonal environment in the pathogenesis13 but these findings have not been substantiated.14 psychological factors have also been indicated as a possible cause for some cases of idiopathic constipation,15 but there is no strong evidence to support this. Degeneration of intrinsic enteric neurones has also been suggested as a possible cause,16 but it might be the secondary result of long lasting bowel distension and laxative abuse.

Impairment of the autonomic nervous system was proposed as a cause of bowel motility dysfunction as early as 1928.17 The observation that constipation is frequently associated with other subclinical or clinically evident motility disorders of the bladder, oesophagus, ileum18,19 and that it is a common bowel disturbance in diseases involving the autonomic nervous system suggests the existence of a relationship between autonomic neuropathy and constipation. An early neuropathy may be present without major disturbances such as orthostatic hypotension, impotence or urinary incontinence, but it may be enough to disturb the fine regulation of gut motility, causing slow transit constipation. The cardiovascular tests1 conventionally used for detecting sympathetic autonomic neuropathy may not be adequate to detect a mild, otherwise asymptomatic neuropathy.20
Impairment of sweating is associated with various manifestations of autonomic neuropathy with and without the presence of other symptoms such as postural hypotension. Sweat regulation has been used as a test of autonomic nervous system function in diabetics since 1942. Camilleri and Fealey, using the thermoregulatory test and a quantitative sudomotor axon reflex test, showed in eight patients with gastrointestinal problems that an idiopathic autonomic denervation may be an aetiological factor in non-organic intestinal dysmotility. All these patients had patchy anhidrosis and two had positive cardiovascular tests.

Interpretation of the Sweat Spot Test Response

The intradermal injection of acetylcholine is effective in stimulating sweat glands because acetylcholine, and not noradrenaline, is the post ganglionic sympathetic neurotransmitter in sudomotor fibres. In the presence of exogenous acetylcholine a normal sweat response should therefore be expected even if nerve degeneration precludes adequate transmitter release. Normal response to the test, however, still depends on an intact nerve supply. Acetylcholine is known to be cotransmitted with vasoactive intestinal polypeptide at the synapse between the sympathetic nerve and the sweat gland, vasoactive intestinal polypeptide probably being responsible for the local vasodilatation which is necessary for a normal sweat response. We suggest that the presence of acetylcholine alone will not provide an adequate stimulus for the sweat response in the event of nerve degeneration, as the full response requires the combined action of both transmitters, provided that the receptors remain intact after nerve degeneration.

There is no evidence to suggest that sweat glands and their receptors remain unchanged after nerve degeneration. In fact, the sweating response to acetylcholine is reported to diminish gradually (one to two years) after denervation.

It is, therefore, possible that in the presence of an autonomic neuropathy of long duration the number of working sweat glands decreases and their distribution becomes irregular. In the original sweat spot test the foot was chosen because, impairment being more likely to occur in longer nerve fibres, the lower extremities would be the first to be involved in diabetic autonomic neuropathy. Although this theory has not been tested in other forms of autonomic neuropathy, we have assumed that the same holds true in cases where constipation was associated to a degree of autonomic neuropathy.

The sensitivity of the sweat spot test derived from the receiver operating characteristic curve is rather low at 83%. The most likely explanation is that these constipated patients are a heterogeneous group including patients without autonomic nervous system dysfunction, whose loss of peristaltic activity might instead be the result of intrinsic nerve damage or other motility disorder. A score of 12 on the sweat spot test response appears to be the delimiting number between normal and abnormal subjects in this population. The curve for abnormal subareas has a limited sensitivity, but the specificity is high at 1, for a scoring of 3% of abnormal subareas, which is the demarcation set by Ryder as a departure from normality.

This study suggests some degree of idiopathic autonomic denervation in patients with slow transit constipation. As with Ryder's findings in diabetics with autonomic neuropathy, most patients with slow transit constipation had a patchy anhidrosis and a number of working glands significantly below normal controls. In these patients, even when the number of glands was normal, their distribution was not uniform.

The significant decrease in sweat spot test score with increasing age would indicate that age is an influential factor in the response to the sweat spot test. There is a reported reduction both in the number of active sweat glands and in the secretory output of each gland with increasing age, a phenomenon which is not the result of a failure of the autonomic control of sweating but partly to trophic changes which occur with age in human skin. The correlation observed between age and loss of sweating ability, however, is only significant in subjects who are more than 70 years old, and only one of our patients...
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was over 70. In the present study, controls were matched for age with the patient group, and as with Ryder's normal subjects, no relationship was shown between age and sweat spot test score in the control group. The decrease in spot density with age observed in patients with constipation is not therefore an age related factor, but is most probably related to the duration of the symptoms as most had been constipated for many years.

The loss of sudomotor function seen in constipated patients in this study would at first indicate a degree of impairment of the sympathetic nervous system which would result in the loss of inhibitory tone on gut smooth muscle. This would consequently lead to increased motility because parasympathetic activity would be unopposed by diminished sympathetic activity. Increased motility does not necessarily result in increased propulsion, however. Conspitigers drugs such as opioids increase motility while inhibiting propulsive activity, and abnormal hypermotility patterns have been reported to occur in the irritable bowel syndrome and constipation. We can speculate that increased excitability of the smooth muscle in the bowel because of failure of the inhibitory sympathetic modulation of parasympathetic activity could result in increased mixing and segmental activity of the colon with impaired coordination of peristaltic activity.

If an autonomic neuropathy is present, it is unlikely that a selective sympathetic impairment would occur in the presence of normal parasympathetic function. The decreased sweating function and slow transit constipation seen in our patients could be a consequence of a mild cholinergic dysautonomia, rather than a sign of a more prominent sympathetic failure. Several reports of extreme cases of selective cholinergic dysautonomia identify ileus (or constipation) and loss of sweating as two of the symptoms experienced by those patients.

The extreme complexity of the enteric nervous system with its non-adrenergic non-cholinergic neurones and its connections with the autonomic nervous system and higher neural centres precludes a simple explanation for the effects of autonomic neuropathy in constipation.

In conclusion, idiopathic constipation may well be an early manifestation of an otherwise asymptomatic systemic neuropathy. The acetylcholine sweat spot test is an easy, inexpensive, time-saving test for detecting autonomic neuropathy. It is reliable for the quantification of the severity of nerve damage and deserves a place in routine clinical assessment of slow transit constipated patients.

The authors wish to thank Dr Robert Ryder for kindly showing us the intricacies of the sweat spot test, Dr Julian Roberts for providing data on intestinal transit time, and Dr Tom Koekoek for help with the statistics.

9 Roberts JP, Waldeon DW, Newell MA, Garvie NW, Williams NS. Comparison of radiopaque markers and radioscintopic assessment of whole bowel transit time in constipated patients. Gut 1990; 31: A617
10 Kamm MA. The surgical treatment of severe idiopathic constipation. Int J Colorect Dis 1987; 2: 279-85
14 Preston DJ, Lennard-Jones JE. Severe chronic constipation in young women: idiopathic slow transit constipation. Gut 1986; 27: 41-8
16 Wald A, Hinds JP, Caruana BJ. Psychological and physiological characteristics of patients with severe idiopathic constipation. Gastroenterology 1989; 97: 932-7
23 Dole HH, Feldberg W. The chemical transmission of secretory impulses to the sweat glands of the cat. J Physiol 1934; 82: 121-4
Detection of subclinical autonomic neuropathy in constipated patients using a sweat test.
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Gut 1992 33: 1539-1543
doi: 10.1136/gut.33.11.1539