Spectrum of malabsorption syndrome among adults & factors differentiating celiac disease & tropical malabsorption

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Background & objectives: Aetiology of malabsorption syndrome (MAS) differs in tropical and temperate countries over time; clinical and laboratory parameters may differentiate between various causes. This study was undertaken to investigate the spectrum of MAS among Indian adults and to find out the features that may help to differentiate between TM and celiac disease.

Methods: Causes of MAS, and factors differentiating tropical malabsorption (TM) from celiac disease (CD) were determined in 275 patients.

Results: Using standard criteria, causes in 275 patients [age 37.5±13.2 yr, 170, (61.5%) male] were, TM 101 (37%), CD 53 (19%), small intestinal bacterial overgrowth 28 (10%), AIDS 15 (5.4%), giardiasis 13 (5%), intestinal tuberculosis 7 (2.5%), strongyloidiasis 6 (2%), immunoproliferative small intestinal disease 5 (2%), Crohn’s disease 6 (2%), amyloidosis 4 (1.5%), intestinal lymphangiectasia 3 (1%) and unknown 22 (8%). On univariate analysis, patients with CD were younger than TM (30.6±12 vs. 39.3±12.6 yr, P<0.001), had lower body weight (41.3±11.8 vs. 49.9±11.2 kg, P<0.001), longer diarrhoea duration (median 36 inter-quartile range 17.8-120 vs. 24-months, 8-48, P<0.01), lower stool frequency (6/day, 5-8 vs. 8, 5-10, P<0.05), lower haemoglobin (9.4±3.2 vs. 10.4±2.7 g/dl, P<0.05), higher platelet count (2,58,000, range 1,35,500-3,23,500 vs. 1,60,000, 1,26,000-2,58,000/mm³, P<0.05), and more often had hepatomegaly (9/53, 17% vs. 4/101, 4%, P<0.01), and subtotal or partial villous atrophy (36/50, 72% vs. 28/87, 32%, P<0.001). Younger age (<35 yr), longer diarrhoea duration, higher platelet count and villous atrophy were significant on multivariate analysis.

Interpretation & conclusions: TM and CD are common causes of MAS among Indian adults. Younger age (<35 yr), longer diarrhoea duration, higher platelet count and villous atrophy were found to be associated with CD.

Key words Chronic diarrhoea - Crohn’s disease - small intestinal bacterial overgrowth - small intestinal diseases - tropical enteropathy - tropical sprue

Malabsorption syndrome (MAS) is a common condition in gastroenterology practice in tropics including India1. Aetiology of MAS in tropical areas differs from that in temperate countries and can be expected to vary over time1. In the past, tropical malabsorption (TM), popularly known as tropical sprue,
was a common cause of MAS in India. Epidemics of TM have been described in southern Indian villages in the past. Sporadic cases of TM have been described from other tropical countries such as Pakistan, Thailand and Malaysia and even in temperate region such as Puerto Rico and Britain. It is believed that, in recent years, with improvement in socio-economic status and sanitary conditions and increasing use of antibiotics, frequency of TM may have declined even in tropical countries in spite of contradictory evidence. Moreover, there could be considerable overlap between post-infectious MAS, which is a subgroup of TM, and post-infectious irritable bowel syndrome (IBS), a common condition in temperate countries. Celiac disease (CD), once thought to be uncommon in tropical countries including India, is being reported frequently as a cause of MAS among children and adults. Data on spectrum of MAS in Indian adults are however, scanty and contradictory.

It is difficult to differentiate between celiac disease and TM. Response to antibiotics, a criterion used to diagnose TM, may be misleading as patients with celiac disease may have secondary small intestinal bacterial overgrowth (SIBO) that at least temporarily may respond clinically to treatment with antibiotics. Thus, it has been proposed that celiac disease should not only be diagnosed using conventional criteria, and a serological test should be performed. Hence, there is a need to determine demographic, clinical and laboratory parameters that may help to differentiate TM from celiac disease in adults with MAS in tropical countries. This may help clinicians to assess the probability of celiac disease in a given patient with serological tests and empirical initiation of gluten free diet in patients with high probability of the disease even in tropical countries. This study was thus aimed to assess (i) the spectrum of MAS among Indian adults, and (ii) features that may help to differentiate TM and celiac disease among them.

**Material & Methods**

Consecutive patients with MAS attending the Luminal Gastroenterology Clinic of the Department of Gastroenterology at Sanjay Gandhi Postgraduate Institute of Medical Sciences, Lucknow, a tertiary referral center in north India, during a 10 yr period (from April 2000 to March 2010) were included. Patients were evaluated using a standard protocol, which was approved by the Institutional Ethics Committee.

**Protocol for evaluation for MAS:** Urine D-xylose, quantitative faecal fat measurement or faecal Sudan stain and duodenal biopsy were done to investigate MAS. Patients with abnormal results in at least two of three tests [urine D-xylose <1.0g/5g/5h, (faecal fat >7g/day or faecal Sudan stain >10 droplet/high power field) and any degree of villous atrophy on duodenal histology] were included in the study. Patients with abnormal results on one of the three tests and evidence supporting a specific cause of MAS were also included. Investigations for aetiology and complications of MAS followed a standard protocol. These included: (i) stool microscopy with various staining methods, (ii) haemogram, (iii) blood biochemistry, (iv) serological tests [human immunodeficiency virus (Enzyme-immunoassay, XIV chex, Qualigenes Diagnostics, Bangalore and Comb AIDS, Span Diagnostics, Surat, India)], (v) anti-endomysial antibody test (indirect immunofluorescence assay, Binding Site, UK), (vi) quantitative estimation of immunoglobulins (IgA, IgG and IgM) levels in serum, (vii) small bowel barium series, (viii) endoscopic duodenal or jejunal biopsies, which was interpreted by an experienced pathologist using a standard grading system, (ix) glucose hydrogen breath test (using 100 g glucose and a technique that has been described previously to diagnose SIBO) and duodenal biopsy to investigate SIbO.

Abdominal ultrasonography, bone marrow biopsy, and colonoscopy were done when deemed necessary on clinical grounds.

**Tests for absorption:**

- **D-xylose test** - Amount of D-xylose excreted in urine during a 5 h period following ingestion of 5 g of D-xylose was estimated using a colorimetric method. Excretion of <1.0 g D-xylose was considered abnormal.

- **Estimation of faecal fat** - Faecal fat was estimated by Van de Kamer’s technique after a three day fat load (75 g/day). Daily stool collection was done during the next three days, during which fat loading was continued. Mean daily stool weight and mean daily fat excretion was calculated. Faecal fat excretion >7 g/day was considered abnormal. Faecal fat was also tested by microscopic examination of the spot stool specimen stained with Sudan III stain. Faecal Sudan >10 droplets per high power field was considered as abnormal.

**Criteria for diagnosis:** Celiac disease: European Society of Pediatric Gastroenterology and Nutrition criteria, giardiasis and strongyloidiasis: demonstration on stool microscopy and/or duodenal biopsy; Crohn’s disease and intestinal tuberculosis: standard criteria described previously; SIBO: positive glucose...
Results of investigations for mucosal malabsorption:

**Urine D-xylose:** Urinary excretion of D-xylose, performed in 233/275 (85%) patients, was 0.53 \( \pm \) 0.30 g/5 g/5 h. It was abnormal in 212 of 233 (91%) patients with MAS.

**Faecal fat and faecal Sudan:** The mean faecal fat excretion, estimated in 167 of 275 (61%) patients, was 8.4 \( \pm \) 3.4 g/day. It was abnormal in 120 of 167 (72%) patients. The median faecal fat excretion by Sudan stain, evaluated in 209 of 275 (76%) patients, was 17 (range 3 - 50) droplets/HPF. It was abnormal in 168 of 209 (80.4%) patients.

**Results of histopathological examination of duodenal biopsy:** Duodenal biopsy was obtained in 226 (82%) patients with MAS; 17 (7.5%) had subtotal villous atrophy, 58 (25.7%) had partial villous atrophy, 45 (20%) had blunting of villi and one patient had normal villi; 217 (96%) had increased mononuclear infiltrate and 126 (55.8%) had increased intraepithelial lymphocytes (Fig. A-D).

Of the 275 patients, 148 (54%) had abnormal results in all the three tests (urine D-xylose, faecal fat either by Sudan or by Van de Kamer’s method and duodenal biopsy) and 94 (34%) had two of these tests abnormal; of them, urine D-xylose and faecal fat were abnormal in 25 (26%), faecal fat and duodenal biopsy were abnormal in 36 (38.2%) and urine D-xylose and duodenal biopsy were abnormal in 33 (35.1%) patients. Of the remaining 33 (12%) patients with only one of these three tests abnormal, included eight patients with SIBO (by GHBT), one with strongyloidiasis on stool microscopy, six with celiac disease with positive result on anti-endomysial antibody and subtotal or partial villous atrophy on duodenal biopsy, six with TM responding to treatment with tetracycline and folate, one intestinal tuberculosis with ileo-colic fistula on small bowel barium series responding to ATT, four with AIDS, two with IPSID on duodenal biopsy, two with histologically confirmed amyloidosis and three with Crohn’s disease diagnosed on imaging, endoscopy, histology and treatment response.

Aetiology of MAS

**Giardiasis:** Of the 13 (5%) patients with giardiasis [median age 42 yr, range 11-53 yr, 10 (77%) male], 4, 7 and 2 were diagnosed using stool microscopy,.
Histological examination of duodenal biopsies in patients with malabsorption syndrome. (A) Patient with tropical malabsorption showing blunting of villi with increased mononuclear infiltrate in lamina propria, (B) subtotal villous atrophy with crypt hyperplasia in a patient with celiac disease, (C) dense dysplastic lymphomononuclear infiltrate with blunting of villi in a patient with immunoproliferative small intestinal disease, (D) larvae of Strongyloides embedded in the mucosa (haematoxylin-eosin).

Histopathology and both tests, respectively. One patient had low level of serum IgA. All the patients responded to treatment with a nitroimidazole.

Intestinal tuberculosis: Of the seven (2.5%) patients with intestinal tuberculosis, [median age 39 yr, range 23-67 yr, 5 (71.4%) male], three had acid fast bacilli (AFB) on microscopic examination of smear, one grew AFB on culture and showed its DNA by polymerase chain reaction. Three other patients had features of ileo-cecal tuberculosis on small bowel barium series including ileo-colic fistula in one and jejunal stricture in another, multiple abdominal lymphadenopathy on computerized tomography scan and positive Mantoux test. All the patients responded to ATT.

Strongyloidiasis: Of the six (2%) patients with strongyloidiasis [median age 30 yr, range 18-70 yr, 5 (83%) male], larvae of strongyloides were detected on microscopic examination of stool or histopathological examination of duodenal biopsy in three each (Fig. D). Two of them were receiving corticosteroids and one had hypogammaglobulinemia. Peripheral blood eosinophilia was absent in four patients. Two patients died of septicemia. Two other patients with AIDS also had larvae of strongyloides on stool microscopy.

Acquired immunodeficiency syndrome (AIDS): Fifteen (5.4%) patients (median age 36 yr, range 30-57 yr) had AIDS. Patients with AIDS were more often male than those with MAS due to other causes [13/15 (87%) vs. 154/260 (59%), \( P<0.03 \)]. Patients with AIDS (n=15) and intestinal lymphangiectasia (n=2) had lower absolute lymphocyte count than MAS due to other causes (median 1296/mm\(^3\), range 360-2520 vs. 1775/mm\(^3\), range 176-6120, \( P<0.01 \)). Stool microscopy revealed larvae of strongyloides in two, Isospora belli
cyst in one, eggs of ascaris in one, both Hymenolepis nana and Cryptosporidium in one. Four patients had history of sexual promiscuity, four had undergone prior surgery and three had history of blood transfusion.

**Intestinal lymphangiectasia:** Three patients with intestinal lymphangiectasia were diagnosed on duodenal biopsy and responded to treatment with median chain triglycerides.

**IPSID and hypogammaglobulinemia:** Five (2%) patients (age 33 yr, range 27-38 yr, all male) were diagnosed IPSID on duodenal biopsy (Fig. C). All presented with chronic small bowel diarrhoea and weight loss. Three patients had been treated with ATT in the past by community physician. One patient had a mass in the right iliac fossa and two had digital clubbing. Three (60%) patients had SIBO on glucose hydrogen breath test and one had Giardia on stool microscopy. Three patients had increased level of serum alkaline phosphatase [median 130 (range 63 to 420 U/l)]. Three patients were successfully treated with tetracycline alone for early disease. Of the other two with advanced disease, one died of explosive diarrhoea and shock while receiving anti-cancer chemotherapy and the other declined further treatment.

**Amyloidosis:** Of the 275 patients with MAS, four (1.5%) (median age 47 yr, range 23 to 61 yr, three male) had presence of amyloid on rectal biopsy (n=2), duodenal biopsy (n=1) or abdominal fat pad aspirate (n=1) using Congo red staining.

**Crohn’s disease:** Of the 275 patients with MAS, six (2%) [median age 36 yr (range 22 to 53 yr), 5 male] had Crohn’s disease. Two of them had ileo-colic fistula. Two patients had been treated with ATT in the past. Two patients had SIBO on GHBT. Five patients responded to treatment and one died of sepsis.

**Tropical malabsorption and celiac disease:** Tropical malabsorption (n=101) was the commonest cause of MAS followed by celiac disease (n=53). Forty of 49 patients (82%) with celiac disease, who underwent test for anti-endomysial antibody, had positive result. Of the remaining nine patients, three had low level of serum IgA, one had anti-tTG antibody and other had anti-gliadin antibody in serum. The clinical and demographic details of TM and celiac disease patients are shown in Table I. On univariate analysis, patients with celiac disease were younger than TM, had lower body weight (41.3+11.8 vs. 48.9+11.2 kg, P<0.001) though height was comparable (156.9 ± 11.9 vs. 158.5 ± 9.6 cm), longer diarrhoea duration, lower stool frequency, lower haemoglobin levels, higher platelet count, hepatomegaly, subtotal or partial villous atrophy or blunted villi (Tables I and II). The best cut-off value of age that differentiated TM from celiac disease was 35 yr (area under curve 0.68, sensitivity 70%, specificity 61%). The best cut-off value of haemoglobin that differentiated TM from celiac disease was 10 g/dl (area under curve 0.62, sensitivity 63%, specificity 56%) and that of platelet count was 200 X 10³/mm³ (area under curve 0.62, sensitivity 62%, specificity 77%). Patients with celiac disease more often had haemoglobin value less than 10 g/dl than those with TM [30 (57%) vs. 37 (37%), P=0.017] and had platelet count more than 200 x 10³/mm³ [28 (53%) vs. 29 (29%), P=0.003]. However, only one patient with celiac disease had thrombocytosis (platelet count 600 X 10³/mm³). Younger age (<35 yr), longer diarrhoea duration, higher platelet count and villous atrophy were significant on multivariate analysis (Table III). GHBT, performed in 52 (51.5%) of 101 patients with TM, revealed SIBO in 12 (23%) patients.

Two patients with strongyloidosis and one with IPSID died. Of the remaining patients, nine with idiopathic MAS did not respond to treatment. Frequency of recurrence after successful treatment was comparable among patients with celiac disease and TM (two patients each) during a follow up period of 12.5 ± 15.3 and 10.8 ± 14 months, respectively.

**Discussion**

The present study showed that TM and celiac disease are common causes of MAS among Indian adults. Other causes included SIBO, AIDS, giardiasis, hypogammaglobulinemia, intestinal tuberculosis, strongyloidiasis, IPSID, Crohn’s disease, amyloidosis and intestinal lymphangiectasia. In 8 per cent patients, the cause for MAS remained unknown.

Aetiological spectrum of MAS is expected to change in developing countries with time. Infective diseases and TM were believed to be common causes in the past, with non-infective causes such as celiac disease being rare in several tropical countries. Epidemics of TM were reported from southern India and tropical countries including South East Asia, Caribbean, the Mexico, the Middle East, parts of Africa, Hong Kong and Malawi. The current data show that TM continues to be a major cause of sporadic MAS in Indian adults. This is in accordance with another study from southern India. A study from north India has shown celiac disease to be a common
cause of MAS in Indian adults and children. In a recent study from Delhi, though tropical sprue was one of the major causes of malabsorption, celiac disease was commoner. However, the difference in spectrum of MAS from the current study might be explained by age group of patients included and inclusion criteria in addition to geographical factors. The study from Delhi and the present study showed that celiac disease was more common among patients with malabsorption at younger age than those at older age. Hence, inclusion of a large proportion of young patients might explain high frequency of celiac disease in that study. In the present study, only 7 per cent patients were in the age group 12-18 yr, a proportion much lower than the Delhi study (30%). It is noteworthy that all these studies, including the present one are hospital-based study and, therefore, the findings cannot be generalized to the community.

There is a need to study the clinical and basic laboratory parameters to differentiate between celiac disease and TM as the two conditions clinically mimic closely. Celiac disease usually manifests at a younger age, and may present without diarrhoea but with atypical symptoms. Due to a lack of awareness among physicians about celiac disease in tropical countries, it may remain undiagnosed for a long period. On the other

| Table I. Demographic, clinical and laboratory parameters of patients with celiac disease and tropical sprue |
|--------------------------------------------------|--------------------------------------------------|
| Celiac disease (n=53) | Tropical malabsorption (n=101) |
| Age (yr, mean ± SD) | 30.6 ± 12 | 39.3 ± 12.6*** |
| Age >35 yr | 16 (30%) | 61 (60%)*** |
| Gender (female) | 29 (54.7%) | 43 (42.6%) |
| BMI (kg/m², mean ± SD) | 16.4 (4.1) | 19.4 (4.1)*** |
| Duration of diarrhoea (mo, median, IQR) | 36 (18,120) | 24 (8, 48)*' |
| Stool frequency/day (median, IQR) | 6 (5,8) | 8 (5, 10)*' |
| Antitubercular treatment in the past | 11 (21%) | 14 (14%) |
| Oedema | 6 (11%) | 12 (12%) |
| Glossitis | 7 (13.2%) | 10 (9.9%) |
| Hepatomegaly | 9 (17.0%) | 4 (4%)*' |
| Ascites | 2 (3.8%) | 2 (2%) |
| Splenomegaly | 2 (3.8%) | 1 (1%) |
| Clubbing | 4 (7.5%) | 2 (2%) |
| Lost ankle jerk | 2 (3.8%) | 4 (4%) |
| Haemoglobin (mean ± SD) | 9.4 (3.2) | 10.4 (2.7)*' |
| Platelet counts (>200 x 10³/mm³, median, IQR) | 258 (136, 324) | 160 (126, 258)*' |
| Serum alkaline phosphatase (IU/l, median, IQR) | 142 (127, 209) | 130.5 (87, 190) |
| Serum albumin (g/dl, median, IQR) | 2.3 (1.7, 2.8) | 2.3 (1.8, 3.6) |
| Serum iron (µg/dl, median, IQR) | 54.4 (39, 105) | 60 (35, 77) |
| Total iron binding capacity (µg/dl, median, IQR) | 283 (251, 310) | 281 (249, 335) |
| Stool weight (g/day, median, IQR) | 542 (391, 767) | 475 (366, 650) |
| Faecal fat*' (g/24 h, median, IQR) | 7.4 (6.4, 9.5) | 7.8 (6.6, 9.0) |
| D-Xylose*' (g/5 g/5 h, median, IQR) | 0.5 (0.3, 0.6) | 0.5 (0.3, 0.6) |

BMI, Body mass index; IQR, Inter quartile range. *Faecal fat was done in 45/53 (85%) and 96/101 (95%) patients with celiac disease and tropical sprue respectively. *Urine D-xylose was done in 42/53 (79%) patients with celiac disease and 94/101 (93%) with tropical malabsorption. Normal range of various parameters: haemoglobin:>11.5 g/dl; platelet counts: 150-400×10⁰/mm³; serum alkaline phosphatase: 35-150 U/l; serum albumin: 3.5-5.5 g/dl; serum iron: 60-150 µg/dl; total iron binding capacity: 270-380 µg/dl; faecal fat: >7g/day or faecal Sudan: >10 droplet/HPF; D-Xylose: >1g/5g/5 h

*P<0.05, ***<0.001 compared to celiac disease patients
hand, TM is a disease among adults, may start acutely after an episode of gastrointestinal infection and may be partly treated with antibiotics before reaching a tertiary healthcare centre. Hence, patients with celiac disease are expected to have long-lasting clinical or subclinical MAS before being finally diagnosed as compared to patients with TM. Hence, our findings of younger age, lower body mass index, longer duration of diarrhoea, lower haemoglobin and more marked villous atrophy are quite expected.

The current study showed that platelet count was higher among patients with celiac disease than those with TM. Thrombocytosis, defined as a platelet count >600 × 10^3/µl, is common in patients with celiac disease and may result from inflammatory mediators or from reversible hyposplenism\(^27\). As high as 60 per cent patients with untreated celiac disease are reported to have thrombocytosis\(^28\). However, only one patient with celiac disease in the current series had platelet count 600×10^3/µl. This might be related to the fact that platelet count may be lower in Indian population\(^29\). Also, associated deficiency of folate and vitamin B12 due to malabsorption might have resulted in a lower platelet count\(^30\). Folate and vitamin B12 deficiency and megaloblastic anaemia are expected to be more common in patients with TM\(^31\) with consequent lower platelet count. By ROC curve, a cut-off value of platelet count more than 200×10^3/µl differentiated celiac disease and TM with 62 per cent sensitivity and 77 per cent specificity. The finding of this study suggests that there was an observed difference between celiac disease and TM during initial assessment.

Some causes of MAS detected in the present study are unusual in occurrence. Of the 13 patients with giardiasis, 12 were immunocompetent. Intestinal tuberculosis, though often associated with biochemical malabsorption, clinical malabsorption is uncommon\(^32\). Malabsorption in intestinal tuberculosis might result from associated SIBO, secondary amyloidosis, lymphatic blockage and diffuse mucosal disease\(^33\). Interestingly, half of the patients had MAS due to strongyloidiasis in absence of immunodeficiency and two-thirds did not have eosinophilia, a parameter by which clinicians suspect strongyloidiasis\(^34\). Absence of eosinophilia in hyperinfection with strongyloides has been reported\(^35\). Five patients had IPSID as a cause for MAS. This is in accordance with an earlier study from southern India in which 1.5 per cent patients with chronic diarrhoea and MAS had IPSID\(^36\). Hypogammaglobulinemic sprue was another cause of MAS that has been uncommonly reported from India\(^37\).

### Table II. Histological parameters of patients with celiac disease and tropical malabsorption

| Parameter                        | Celiac disease (n=53) | Tropical malabsorption (n=101) |
|----------------------------------|-----------------------|--------------------------------|
| Subtotal villous atrophy         | 12 (24)               | 3 (3.4)***                     |
| Partial villous atrophy          | 18 (36)               | 19 (22)                        |
| Blunting of villi                | 2 (4)                 | 6 (7)                          |
| Any villous atrophy              | 36 (72)               | 28 (32)***                     |
| Mononuclear infiltrate           | 47 (94)               | 84 (97)                        |
| Increased intra-epithelial lymphocytes** | 37 (74)   | 44 (51)**                      |

Values are no. (%)

*Duodenal biopsy was obtained in 50 patients with celiac disease and 87 with tropical malabsorption

\(^{P^*}<0.01, \text{ ***}^{<0.001}\) compared to celiac disease patients

### Table III. Result of multivariate analysis of factors differentiating between celiac disease and tropical malabsorption

| Factor                        | Crude OR (95% CI) | Adjusted OR (95% CI) | P (Wald’s test) | P (LR-test) |
|-------------------------------|-------------------|----------------------|-----------------|-------------|
| Age >35 vs age <35 yr         | 3.53 (1.482, 6.728) | 2.739 (1.072, 6.999) | 0.0353          | 0.0316      |
| Presence vs absence of hepatomegaly | 0.202 (0.065, 0.851) | 0.258 (0.056, 1.184) | 0.0814          | 0.074       |
| Duration of diarrhoea in months | 0.9926 (0.9862, 0.9991) | 0.9927 (0.986, 0.9994) | 0.0329          | 0.0321      |
| Stool freq/24 h               | 1.11 (0.996, 1.237)  | 1.107 (0.972, 1.262) | 0.126           | 0.1092      |
| Haemoglobin                   | 1.154 (1.016, 1.311) | 1.113 (0.950, 1.303) | 0.1851          | 0.1804      |
| Any form of villous atrophy   | 0.184 (0.043, 0.268)  | 0.099 (0.032, 0.308)  | < 0.001         | <0.001      |
| Platelet count                | 0.9957 (0.9921, 0.9994) | 0.9932 (0.9867, 0.9998) | 0.0446          | 0.0323      |
Many of these unusual causes of MAS may not be suspected even in the tropics resulting in fatal outcome and inappropriate treatment with anti-tuberculosis treatment. Hence, awareness about these diseases is required before clinicians can suspect and diagnose these conditions.

The present study provides important data that may have implications on two conditions, namely TM and post-infectious irritable bowel syndrome (IBS). TM may provide useful models to study post-infectious IBS\(^6\). SIBO has been proposed as a condition overlapping both post-infectious IBS and TM\(^8\). Thus, our findings may serve as a useful reference point for future research in this area.

In conclusion, the present study show that TM and celiac disease are common causes of MAS among Indian adults. There were several other unusual causes, awareness about which is required before the clinicians can suspect and diagnose. On univariate analysis, patients with celiac disease were younger than TM, and inappropriate treatment with anti-tuberculosis in the upper gut in patients with malabsorption syndrome from the tropics. BMC Gastroenterol 2003; 3 : 9.

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