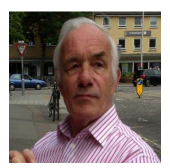
Histology of gluten related disorders

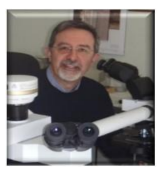
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Michael N Marsh



Vincenzo Villanacci



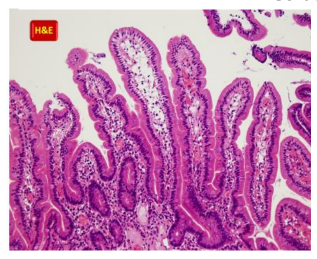
Amitabh Srivastava

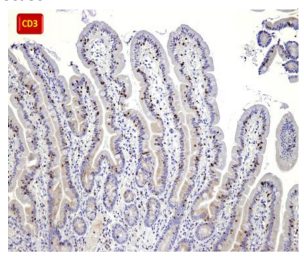
Gluten related disorders range a inflammatory disorders of the small intestine malabsorption characterized bv ingestion of gluten in individuals with a genetic background. Clinical certain presentation can vary from full-blown to subtle malabsorption and atypical symptoms. Diagnosis currently relies on clinicopathologic studies including mucosal biopsy, serologic tests, and the effects of a diet free of gluten on the symptoms. Mucosal pathologic features are also variable, ranging from mild abnormalities, including intraepithelial lymphocytosis, to completely flat mucosa (1). Since there is no specific biomarkers for non-coeliac gluten sensitivity a combination of clinical and histology would play an improtant in identifying such individuals. Classification of mucosal pathology in gluten-sensitive enteropathy has been a subject of controversy among pathologists and needs to be revised according to the current understanding of the disease (2-4).

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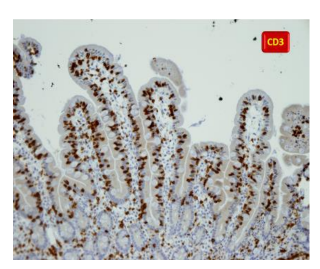
Celiac disease



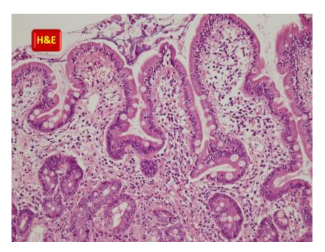


Normal (10X)



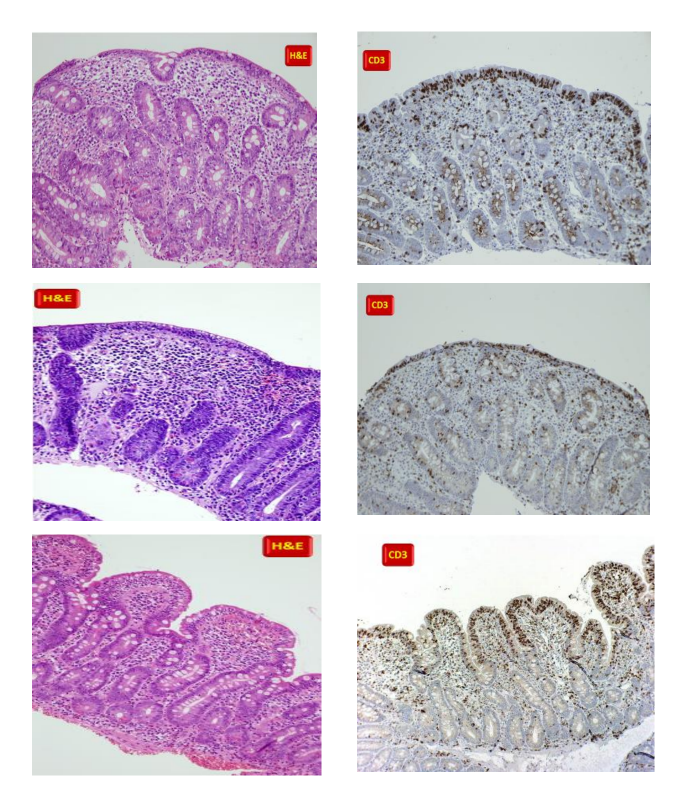


Type 1-2 Marsh (10X)



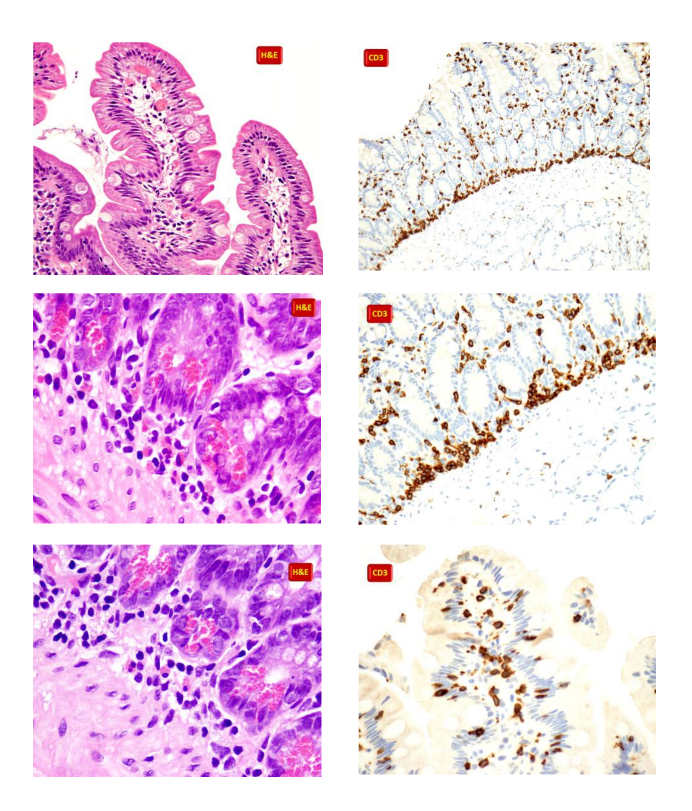


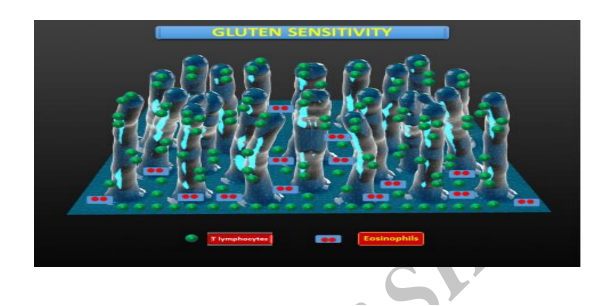
Type 3 Marsh (10X)

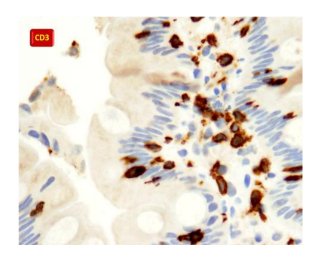


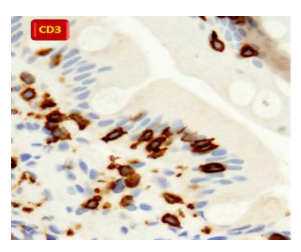
Type 3 Marsh (10X)

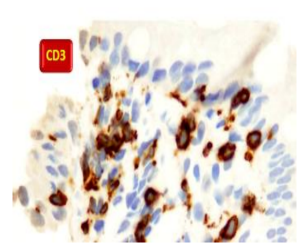
Non Celiac Gluten Sensitivity

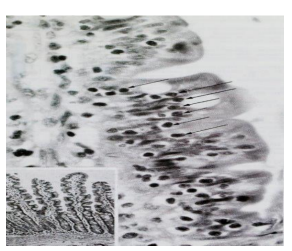












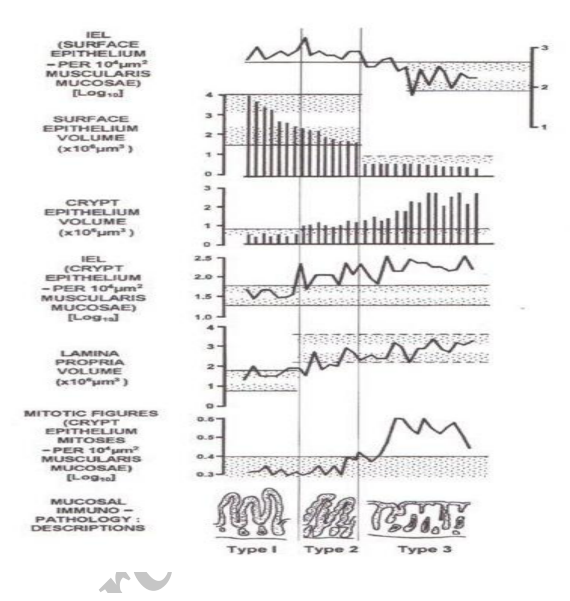


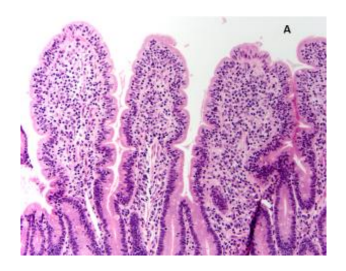
Diagram. These data (Marsh MN et al, unpublished) illustrate the underlying immunopathologic features of celiac musosa as it becomes flat.

It is based on the computerised morphometric technique devised and employed in our laboratory, all measurements ultimately dependent on an invariant "comparator" of a constant test area of muscularis mucosae.

It refers not only to the crypts, but also the important changes occurring within the lamina propria: each stage has its characteristic changes, thus adding to the validity of the Marsh classification.

Overall, the inclusion of so many controlled numerical data in one diagram provides an almost dynamic view across the mucosa as it is becoming flat. We need further descriptive accounts, related to this underlying format, based on the evolutionary alterations in the production of inflammatory molecules and gene activations during the temporal development of the severe lesion.

This micrograph shows a Marsh II lesion, with well-preserved villi, and hypertrophied crypts. The epithelium is heavily infiltrated with small, non-mitotic lymphocytes. This specimen actually came from a first degree relative, and was one of the original findings which lead to the formulation of the Marsh Classification.



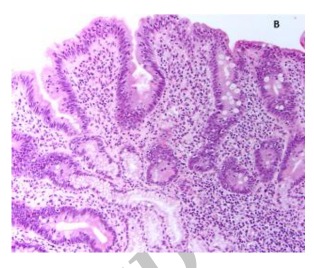


Figure. A variety of diseases can mimic celiac disease on histology. Tropical sprue (A) may show marked intraepithelial lymphocytosis (IEL). The lack of significant villous atrophy in the presence of significantly increased IELs should raise suspicion for tropical sprue. Total or subtotal villous atrophy with increased IELs mimicking celiac disease can also be seen in patients with immune deficiency disorders, as seen here in a patient with common variable immunodeficiency (B).

References

- 1. Marsh MN, Johnson MW, Rostami K. Mucosal histopathology in coeliac disease: a rebuttal of Oberhuber's sub-division of Marsh III. Gastroenterol Hepatol Bed Bench 2015;8:99-109.
- 2. Marsh MN. Studies of intestinal lymphoid tissue: XIII Immunopathology of the evolving celiac sprue lesion. Pathol Res Pract 1989; 185:774-77.
- 3. Villanacci V. The histological classification of biopsy in celiac disease: time for a change? Dig Liver Dis 2015;47:2-3.
- 4. Rostami K, Aldulaimi D, Holmes G, Johnson MW, Robert M, Srivastava A, et al. Microscopic enteritis: Bucharest consensus. World J Gastroenterol 2015; 21:2593-604.