Practical Pathology and Genetics of Hirschsprung Disease
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The only obligate diagnostic feature of the malformation, known as Hirschsprung disease (HSCR), is absence of ganglion cells in the myenteric and submucosal plexuses of the terminal rectum. Hypoganglionosis (paucity, but not complete loss of ganglion cells), segmental aganglionosis that does not involve the terminal rectum, and various forms of neuronal dysmophogenesis in which ganglion cells are present, but dysplastic, should not be regarded as HSCR, even though the clinical presentation may be similar. It is very likely that the pathogenesis of these conditions is distinct from HSCR. The pathology of some of these other conditions is reviewed elsewhere.1-3

Submucosal Biopsies
Aganglionosis can be diagnosed by evaluation of either submucosal or myenteric ganglia. Since, in either instance, the pathologist must establish absence of ganglion cells, adequate sampling is essential and the recognition of ancillary histopathological findings that correlate with aganglionosis can be helpful. In most centers, initial diagnosis of HSCR is based on histopathological study of suction rectal biopsies. Diagnostic biopsies must be taken at least 1-1.5 cm proximal to the anorectal squamo-columnar junction, measure >2 mm in diameter, and include sufficient submucosa. When properly oriented and sectioned adequately (75-100 levels), H&E-stained, paraffin-embedded sections are generally sufficient to exclude the presence of submucosal ganglion cells and suggest the diagnosis of HSCR. The presence of hypertrophic nerve fibers (>40 μm diameter) is observed in many, but not all cases, and helps establish the diagnosis. Similarly, the frequent, but not invariable, presence of abnormally thick and numerous AChE-stained nerve fibers in the muscularis mucosa +/- lamina propria can be used to confirm the diagnosis, particularly when neither ganglion cells nor hypertrophic nerves are observed.

The hypertrophic nerves that exist in most patients with HSCR arise from extrinsic autonomic and sensory fibers, which enter along with vessels from perirectal region and project for a finite distance rostrally. It is the number and diameter of these fibers that increases in HSCR, giving rise to the “hypertrophic” nerves that are frequently, but not always, observed in the myenteric plexus, submucosa, and mucosa of HSCR patients. Because these fibers only project for a finite distance proximal to the rectum, hypertrophic innervation may not be observed in biopsies taken rostrally in long-segment disease. Furthermore, extrinsic nerve hypertrophy of the rectum may not be observed in patients with combined deficiency of intrinsic ganglion cells and other peripheral ganglia (more common with long-segment HSCR) or very premature infants with delayed extrinsic innervation.

The average submucosal biopsy may be 3 mm across and the average distance between submucosal ganglia varies with age, but is probably about 500 microns and contains 2-7 ganglion cells, each of which measures 20-30 microns in diameter. With 3-5 um-thick paraffin sections, ganglia are missed in many random sections. Therefore, it is important to evaluate a lot (>75). With an adequate, well-oriented, not crushed, H&E-stained biopsy one can be fairly confident about the diagnosis of HSCR simply by careful examination of each section. In this situation, the presence of certain ancillary features of HSCR can be quite helpful, but is not required. The problem is that very often the biopsies are suboptimal due to paucity of submucosa, crush, or both. It is in these cases, where AChE-staining can be particularly valuable.
**AChE staining and interpretation**

AChE staining and interpretation

Histochemical staining for AChE activity is a useful adjunct for the diagnosis of HSCR. The procedure is only performed with frozen sections and therefore requires a second suction biopsy, if paraffin sections are also going to be evaluated. The traditional protocol for AChE staining requires approximately 90 min, but rapid procedures have been developed that require 5-10 minutes. AChE staining in the rectum of normal children includes staining of nerves in the submucosa and small fibers in the muscularis mucosa. Usually the latter are confined to the inner half of the muscularis mucosa. If fibers are positively stained in the lamina propria, they are extremely thin and few. Most, but not all, of the rectal submucosal biopsies from HSCR patients contain more densely packed, large, AChE-positive fibers through the full thickness of the muscularis mucosa. In addition, prominent AChE-positive fibers are often present in the lamina propria. The latter finding should not be relied upon too heavily because hypertrophic nerves are confined to the muscularis mucosa in biopsies from a significant subset of HSCR patients, particularly young infants.

**Histochemical diagnostic approach to HSCR**

Considerable variation exists in the approaches taken by clinicians and pathologists to diagnose HSCR. In particular, the number of biopsies, number and section thickness, and use of AChE-staining differ between groups. In some parts of the world, diagnosis of HSCR is based solely on histochemically stained frozen sections of suction rectal biopsies; paraffin sections are not required. The latter protocols generally stain separate sections from each biopsy for AChE and one or two markers of ganglion cell bodies, such as lactate dehydrogenase (LDH) and succinate dehydrogenase (SDH). A potential added value of this histochemical battery is that it may disclose subtle forms of enteric dysganglionosis (e.g., ganglion cell dysmaturity), apart from HSCR, which may explain the HSCR-like symptoms in a patient with ganglion cells.

With regard to diagnosis of HSCR, either the H&E/paraffin-based or histochemistry/frozen-based technique is valid, but experience with a given method is undoubtedly the most important variable that influences diagnostic accuracy. The pathologist must be secure with what constitutes adequate specimen, slides, orientation, etc. Above all, he/she should have the confidence to distinguish diagnostic from equivocal findings and clearly communicate the results to the clinician. In some instances, repeat biopsy is indicated, particularly if the specimen is inadequate or equivocal results are obtained.

**Intraoperative Seromuscular Biopsies**

Many surgical approaches to HSCR are currently employed with a trend toward one-step procedures that are often transanal. In other instances, diagnosis based on suction rectal biopsy is followed by a two-stage procedure that begins with placement of an ostomy proximal to the aganglionic segment. Intraoperative seromuscular biopsies are important to determine that ganglion cells are present at the level where the ostomy or anastomosis will be placed. During ostomy placement, aganglionic gut is frequently not biopsied intraoperatively. However, whether the definitive surgery is done in one or two stages, it is helpful if the transitional zone between aganglionic and ganglionic gut is mapped intraoperatively prior to resection of any bowel. Generally this is accomplished by sequential seromuscular biopsies, from distal-to-proximal.

Seromuscular biopsies should be a minimum of 1 cm in length and extend for a depth of 3-5 mm, so as to include the longitudinal and most of the circular layers of the muscularis propria. Proper orientation of the biopsy for frozen sections greatly facilitates sampling and identification of ganglion cells. The goal is to cut perpendicular to the serosal surface, thereby visualizing the both muscle layers and their interface in the histological sections. With a well-oriented biopsy,
two-to-five sections are generally sufficient to confirm/exclude aganglionosis. Recognition of ganglion cells in usually not difficult, although inflammation sometimes obscures their cytological features.

**Analysis of HSCR Resections**

The pathologist’s goals in analyzing resected aganglionic gut from an HSCR patient are to confirm the diagnosis of HSCR, map the transitional zone, and assess the integrity of the nervous system at the proximal end of the resection. Confirmation that the distal gut is aganglionic is straightforward. A map of the transition zone can be completed by sampling multiple areas along the length of the resected segment to document the presence/absence of submucosal and myenteric ganglia. Some pathologists prepare rolls from longitudinal strips of the entire length of the resection.\(^7\) I prefer to use transverse sections because the circumferential interface between aganglionic and ganglionic gut is often irregular, which will not be apparent in any single longitudinal strip.\(^7\) Full-circumference sections are needed, particularly if the aganglionic zone appears extend within two cm of the proximal resection margin. Construction of a decent map may require repeated sampling sessions.

Evaluation of the integrity of the proximal gut is the most challenging aspect of HSCR pathology. In principle, the genetic etiologies that produce distal aganglionosis may have more subtle effects on the number, distribution, circuitry, and/or differentiation of proximal neurons. Certainly, most HSCR patients harbor a transitional zone of variable length, in which the density of myenteric ganglia is obviously less than in normal gut. However, mild-to-moderate changes in neuronal density are extremely difficult to diagnose by routine analysis and yet may correlate with continued HSCR-like symptoms postoperatively. Incorporation of hypoganglionic transitional zone reportedly increases the likelihood of persistent post-surgical constipation that may necessitate a repeat operation. However, the pathological criteria used to define the transitional zone were not provided in these studies.

In addition to hypoganglionosis, submucosal hyperganglionosis (intestinal neuronal dysplasia, type B; IND) has been reported in the proximal gut of HSCR patients. IND is a controversial form of submucosal hyperganglionosis that has been defined entirely based on histochemically stained biopsies.\(^8\) Similar changes have been observed as isolated “neuropathy” in some children with HSCR-like symptoms, as well as in contexts of other primary disorders of dysmotility, including the transitional zone of HSCR. The significance of Hirschsprung-associated IND is hotly debated in the literature; some authors have advocated screening for IND with frozen sections at the time of surgery so as to extend the resection proximal to the affected area. However, no compelling data exists to suggest that IND-like changes in HSCR either predict a poor outcome or should be managed any differently from isolated HSCR.

A variety of other poorly understood histopathological findings are observed in aganglionic bowel and or the transitional zone. Some of these are summarize in Table 1. While most of these findings have no established clinical and/or genetic significance, their potential correlations with specific genetic defects and / or post-operative complications have not been adequately studied.
TABLE 1: MISCELLANEOUS HISTOPATHOLOGICAL FINDINGS IN HSCR

<table>
<thead>
<tr>
<th>Histopathological Finding</th>
<th>Location</th>
<th>References</th>
</tr>
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<tbody>
<tr>
<td>Eosinophilic neural infiltrates</td>
<td>Aganglionic and transitional zones; transmural including nerve plexuses</td>
<td>9</td>
</tr>
<tr>
<td>Submucosal arterial fibromuscular dysplasia</td>
<td>Transitional zone</td>
<td>10</td>
</tr>
<tr>
<td>Loss of c-kit immunoreactive interstitial cells of Cajal</td>
<td>Conflicting data</td>
<td>11-13</td>
</tr>
<tr>
<td>Peripheral nerve pattern of laminin expression</td>
<td>Aganglionic and transitional zones</td>
<td>14</td>
</tr>
<tr>
<td>Loss of nNOS innervation in muscularis propria</td>
<td>Aganglionic zone</td>
<td>15-17</td>
</tr>
</tbody>
</table>

Skip Lesions and Zonal Aganglionosis

It is important to be aware of two rare forms of congenital aganglionosis that deviate from the classic pattern, in which the distal rectum and uninterrupted contiguous bowel are devoid of nerve cell bodies. In the intestinal tracts of persons with “zonal” (“segmental”) aganglionosis, ganglion cells are present in the distal rectum, but are absent from a proximal segment of gut. In contrast, “skip areas” are ganglion cell-containing segments of large intestine, flanked proximally and distally by aganglionic gut.

Zonal aganglionosis is considered to be an acquired lesion (disruption) that results when ganglion cells (or their precursors) in a fully colonized segment of gut die due to ischemic, viral, immunologic, or other types of injury. The aganglionic segment can occur in small or large intestine. In some cases, a specific etiology is suggested by history (e.g., necrotizing enterocolitis) or other pathological findings (e.g., viral cytopathy). Alternatively, it has been suggested that zonal aganglionosis might result from failure of vagal and sacral crest cells to converge in the gut wall. At the time this hypothesis was introduced, it was less certain that sacral crest cells naturally adopt an enteric neural fate. Given recent evidence that a subset of colonic neurons derive normally from the sacral crest, the proposal is more tenable. However, evidence for neuron formation in the distal colon does not occur in experimental models in which hindgut colonization by vagal crest cells is prevented (discussed above).

Some colleagues and I reviewed the subject of skip areas in 1995 and found that only eleven cases had been reported. Since then, I have been informed of several other cases that were encountered by colleagues, and I suspect the entity may be more common than the literature might suggest. With one exception, skip areas are located in the large intestine and bracketed by aganglionic areas that invariably include both the distal rectum and appendix plus variable lengths of contiguous large and small intestine. Pathologists and surgeons must be cognizant of skip areas, and not use biopsies of the appendix as a means to diagnose total colonic aganglionosis since relatively large skip areas can be present in which ganglion cells exist. In at least one patient, the skip area was recognized, preserved, and used to establish a functional anastomosis between small intestine and anus.

Ultrashort-segment Hirschsprung Disease

Ultrashort-segment HSCR is a controversial entity that has been defined differently by various investigators. Initially, the term “ultrashort-segment HSCR” was reserved to describe patients with clinical and radiological findings similar to those of HSCR, but with ganglion cells in their rectal biopsies. Others reserved the diagnosis for only those biopsies that contain a normal density of ganglion cells and demonstrate an HSCR-like AChE-staining pattern. In either case, the underlying assumption is that an extremely short aganglionic zone in the distal rectum, inferior to
the submucosal biopsy site, is responsible for obstructive symptoms. Another possibility is that such patients have distal hypoganglionosis, which is extremely difficult to document because ganglion cells are normally sparse or absent in the terminal 2-3 cm of rectum.

Patients who fulfill revised criteria for “ultrashort-segment HSCR” probably constitute a heterogeneous set of disorders that may include aganglionosis, hypoganglionosis and anal achalasia (internal sphincter achalasia). In those patients with aganglionosis, the HSCR cannot be established by suction rectal biopsy because the aganglionic myenteric plexus does not extend proximal to the zone of physiological submucosal hypoganglionosis (2-3 cm from the dentate line). In patients that do not have aganglionosis, anatomic changes may not be evident and the diagnosis is based on physiological or histochemical criteria. Regulation of anal sphincter tone is a complex process, which is influenced by both the central and peripheral nervous systems. Heightened sphincter tone could be caused by psychogenic, myogenic, or neurogenic etiologies. A major stimulus for sphincter relaxation is nitric oxide (NO) release by efferent nerve fibers in the smooth muscle. Reduced nicotinamide-adenine dinucleotide phosphate (NADPH)-diaphorase activity (nitric oxide synthase), a marker for NO-producing nerve fibers, has been observed in the internal sphincter of patients with anal achalasia. In some cases, the symptoms resolve after sphincter myotomy.

Immunohistochemistry and the diagnosis of HSCR
An unbelievable number of papers have been published which essentially describe some neuron-specific marker that the authors suggest could be used to facilitate the diagnosis of HSCR. Although most of these antibodies are fairly specific, none is used widely in practice because pathologists who regularly search for ganglion cells in H&E-stained sections or LDH/SDH histochemically stained sections are quite good at discriminating neuron cell bodies without the need for special stains. False positive diagnoses result either from inadequate sampling or observer inexperience.

As opposed to most of the published immunohistochemical targets which are expressed in ganglionic gut, a marker specifically expressed in aganglionic gut would be much more valuable. The fundamental problem in the diagnosis of HSCR is sampling and the fact that the primary feature is absence of ganglion cells, which are sparsely distributed. A reliable marker is needed that detects changes related to the absence of ganglion cells. AChE histochemistry does this to some degree, but the hypertrophic nerve fibers revealed by this technique represent a quantitative change that is subject to individual interpretation. Also, as noted above, abnormal AChE staining is not present in all cases and is age-dependent. Recently, laminin immunohistochemistry was shown to highlight changes in submucosal nerves that may distinguish aganglionic bowel from ganglionic bowel. However, the sensitivity and specificity of this method have not been established.

Molecular genetics and HSCR
Hirschsprung disease (HSCR, intestinal aganglionosis) affects an estimated 1:5000 liveborn infants. The actual incidence of this malformation may much higher since intestinal aganglionosis is frequently associated with other anomalies, and many embryos with multiple defects might die in utero. It is now clear that most, if not all, cases of HSCR have a genetic basis and that HSCR is a complex multigene disorder characterized by incomplete penetrance and variable associated anomalies. Mutations in more than 11 different genes have been implicated in the pathogenesis of HSCR (Table 2); many were first recognized in murine models for this condition. However, according to most estimates, mutations in one or more of these genes only can be detected in about half of all HSCR cases. Therefore, other genetic and/or environmental factors are likely to be involved. RET is the gene in which mutations are most frequently
detected in patients with non-syndromic HSCR. In addition, recent studies indicate that non-coding polymorphisms (base-pair differences that do not affect protein structure and which are present in >1% of the normal population) in the proximal portions of the RET gene pose a significant risk for HSCR, possibly by reducing RET expression.²⁰-²²
TABLE 2: SUSCEPTIBILITY GENES IN ISOLATED AND SYNDROMIC HSCR

<table>
<thead>
<tr>
<th>SYNDROME</th>
<th>GENETIC DEFECT</th>
<th>% WITH HSCR</th>
<th>COMMON PHENOTYPIC FEATURES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Isolated HSCR(^{a})</td>
<td>RET</td>
<td>100%</td>
<td>TTF-1 mutations have been associated with hypothyroidism and cleft palate, but the only reported HSCR patient with a TTF-1 mutation had isolated HSCR.</td>
</tr>
<tr>
<td></td>
<td>EDNRB&lt;sup&gt;b&lt;/sup&gt;</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>EDN3&lt;sup&gt;c&lt;/sup&gt;</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>TTF-1</td>
<td>NE</td>
<td>TTF-1</td>
</tr>
<tr>
<td>Syndromic HSCR</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MEN2A</td>
<td>RET&lt;sup&gt;c&lt;/sup&gt;</td>
<td>&lt;1%&lt;sup&gt;23&lt;/sup&gt;</td>
<td>Medullary thyroid carcinoma, pheochromocytoma, parathyroid hyperplasia</td>
</tr>
<tr>
<td>Smith-Lemli-Opitz</td>
<td>DHCR7</td>
<td>16%&lt;sup&gt;24&lt;/sup&gt;</td>
<td>Growth retardation, pedal syndactyly, mental retardation, hypospadius, dysmorphic facies</td>
</tr>
<tr>
<td>Down</td>
<td>Trisomy 21</td>
<td>2-9%&lt;sup&gt;25&lt;/sup&gt;</td>
<td>Prominent epicanthal folds, upslanting palpebrae, hypotonia, mental retardation, flat midface, single transverse palmar crease</td>
</tr>
<tr>
<td>Waardenburg-Shah</td>
<td>SOX10, EDN3&lt;sup&gt;b&lt;/sup&gt;, EDNRB&lt;sup&gt;b&lt;/sup&gt;</td>
<td>100%&lt;sup&gt;d&lt;/sup&gt;</td>
<td>Deafness, piebaldism, other neurological deficits</td>
</tr>
<tr>
<td>Mowat-Wilson</td>
<td>ZHFX1B</td>
<td>62%&lt;sup&gt;26&lt;/sup&gt;</td>
<td>Abnormal facies, cardiac malformations, mental retardation, genitourinary anomalies</td>
</tr>
<tr>
<td>Haddad</td>
<td>PHOX2B&lt;sup&gt;e&lt;/sup&gt;</td>
<td>100%&lt;sup&gt;f&lt;/sup&gt;</td>
<td>Congenital central hypoventilation, neuroblastoma</td>
</tr>
<tr>
<td>Goldberg-Shprintzen</td>
<td>KIAA1279</td>
<td>100%</td>
<td>Microcephaly, various brain malformations, cleft palate</td>
</tr>
<tr>
<td>X-linked hydrocephalus</td>
<td>LICAM</td>
<td>NE</td>
<td>Cerebral aqueductal stenosis, hydrocephalus, absent corpus callosum</td>
</tr>
<tr>
<td>Cartilage-Hair Hypoplasia&lt;sup&gt;g&lt;/sup&gt;</td>
<td>RMRP</td>
<td>NE</td>
<td>Skeletal dysplasia, sparse blond hair, immunodeficiency, anemia</td>
</tr>
<tr>
<td>Bardet-Biedl</td>
<td>BBS1, BBS2, BBS4, BBS6&lt;sup&gt;h&lt;/sup&gt;, BBS7, BBS8</td>
<td>2%&lt;sup&gt;27&lt;/sup&gt;</td>
<td>Obesity, retinal degeneration, polydactyly, gonadal and renal malformation</td>
</tr>
<tr>
<td>Kauffman-McKusick</td>
<td>MKKS&lt;sup&gt;i&lt;/sup&gt;</td>
<td>NE</td>
<td>Polydactyly, congenital heart defect, hydrometrocolpos</td>
</tr>
</tbody>
</table>

<sup>a</sup>Since HSCR most often presents shortly after birth features of some syndromes (e.g., MEN2A) may not be obvious at the time of diagnosis.
<sup>b</sup>Isolated HSCR is more commonly associated with heterozygous mutations; Waardenburg-Shah syndrome is more commonly associated with mutations in both alleles.
<sup>c</sup>Missense mutations affecting one of three cysteine codons (Cys609, Cys 618, or Cys620)<sup>32</sup>.
<sup>d</sup>HSCR is a diagnostic feature that differentiates Waardenburg-Shah syndrome from other variants of Waardenburg syndrome.
<sup>e</sup>Congenital central hypoventilation syndrome has also been associated with RET, GDNF, and EDN3 mutations in rare patients, but concurrent HSCR has only been associated with PHOX2B mutations<sup>49</sup>.
<sup>f</sup>HSCR is a diagnostic feature of Haddad syndrome, but PHOX2B mutations also occur in patients with isolated congenital central hypoventilation<sup>49</sup>.
<sup>g</sup>Many patients have been presented with a HSCR and a variety of limb defects different from cartilage-hair hypoplasia<sup>37</sup>.
<sup>h</sup>BBS<sub>6</sub> and MKKS are the same gene<sup>50, 51</sup>.
<sup>i</sup>NE, not established.
The gene products encoded by all established “HSCR genes” are either expressed by neural crest-derived enteric neural progenitors or by adjacent cells in the colonization pathway that progenitors follow. Many of these gene products are intercellular signaling components that influence proliferation, survival, migration, and/or differentiation of enteric neural crest cells and, in some cases, other cell types as well. Recent research using avian, zebrafish, and murine models is providing insight into when, where, and how these gene products influence enteric neurodevelopment. Although much is unknown about the molecular and cell biology of each HSCR gene product in normal enteric neurodevelopment, it appears that defects in these genes lead to intestinal aganglionosis by interfering with events at very different points in the colonization route that neural crest cells negotiate en route to colonization of the entire gut. At the same time, studies murine and human studies are beginning to elucidate complex interactions that exist between polymorphic alleles of these genes.

It is important to realize that intestinal aganglionosis is a malformation, analogous to other birth defects like absent radii, cleft lip, congenital heart malformations, etc. HSCR can occur as an isolated anomaly or in the context of multiple malformations, which in many instances constitute recognizable syndromes. Genotype-phenotype correlation in patients with HSCR is often not sufficient to suggest mutation in a particular candidate gene because considerable phenotypic overlap exists among HSCR patients with mutations in different genes. However, complete clinical evaluation and family history should be obtained as these details sometimes provide clues to the nature of the genetic defect.

A particularly challenging problem in the field of Hirschsprung genetics is the phenomenon of "incomplete penetrance". Incomplete penetrance refers to the fact that only a subset of individuals who carry a particular mutation in one of the genes listed in Table 2 actually exhibit the HSCR phenotype (intestinal aganglionosis). Other persons, even immediate family members, with the same mutation are unaffected. Incomplete penetrance is observed in many other human genetic disorders and is usually attributed to one of three variables - genetic modifiers, environmental modifiers, or stochastic events. Substantial evidence has been gathered to suggest that genetic modifiers are responsible for some of the variable penetrance of HSCR mutations. In some cases, interactions between alleles of two different genes listed in Table 2 determine whether enteric neurodevelopment is perturbed. For example, subtle polymorphisms that alter the nucleotide sequence of the \textit{RET} gene, but not the amino acid sequence of the \textit{RET} protein, have been shown to influence penetrance of a particular missense mutation in the \textit{EDNRB} gene. In addition, genetic loci that influence penetrance of \textit{RET} mutations, but do not correlate with any of the genes listed in the table, have been mapped. Thus, HSCR is a complex genetic disorder that challenges the comprehension of scientists and clinicians, let alone the families with an affected child.

At this time, the genetic data listed in Table 2 has limited practical value. Mutational analysis for many of the genes listed is not easily obtained and does not affect clinical management or counseling, except in rare contexts. One possible exception is testing for MEN2A mutations of the \textit{RET} gene in patients with sporadic HSCR. Despite the fact that most patients with MEN2A do not have HSCR, some studies have reported relatively high rates (2.5-5%) of MEN2A-associated mutations in patients with what initially appears to be sporadic HSCR. Early diagnosis of MEN2A allows better surveillance for neoplasia in the affect individual and screening of relatives who may also carry the same mutation.
References


Pediatric Inflammatory Bowel Disease: New Insights and Controversies

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Introduction

The category of “inflammatory bowel disease” (IBD) has been traditionally defined as encompassing two entities, ulcerative colitis (UC) and Crohn’s disease (CD). The etiology of these two conditions is as yet unknown; therefore, there is no gold standard for their diagnosis. The pathogenesis of both UC and CD is not yet well defined but involves an interplay of generic susceptibility, abnormalities in the gut lining epithelium, imbalances between beneficial and pathogenic gut bacteria, and host immune response dysregulation (1). Diagnosis is achieved by (1) eliminating known causes of acute, self-limited enteritis and colitis (e.g., infections, allergies, medication-related injury) and (2) recognizing clinical, imaging, endoscopic, and histologic features characteristic of these two conditions. Although the classic forms of pediatric UC and CD are extensively described in the pathology literature, recent advances in imaging techniques, the increasing use of upper gastrointestinal endoscopy and colonoscopy to evaluate these conditions, the use of video capsule endoscopy to directly visualize the entire small intestine, and the implementation of diagnostic procedures early in the course of disease have resulted in the recognition of new findings and deviations from “classic” teachings that may confound the diagnosis of UC and CD and even raise the possibility of other types of IBD. In this presentation, the findings of classic UC and CD in the colon will be reviewed, but the emphasis is on atypical and newly described aspects detected in mucosal biopsies that may cause diagnostic difficulties in pediatric patients.

Histologic Features of Mucosal Colitis

Distinguishing an acute self-limited colitis (ASLC) from IBD is important because of the long-term psychological and medical management aspects of IBD. This distinction may be problematic in some cases for the following reason. The usual cases of ASLC are diagnosed by a combination of clinical, epidemiologic, and/or microbiological findings. Only the unusual cases with a prolonged clinical course, atypical laboratory findings, and/or negative infectious workups eventuate in colonoscopy. Colonic mucosal biopsy specimens from such patients may contain indeterminate findings that are difficult to categorize with certainty. The histologic features of active and chronic colitis in mucosal specimens are summarized below, followed by a discussion of the distinction between ASLC and IBD.

Active colitis

Activity is characteristic of both ASLC and IBD. The features of activity are well known:
1) Neutrophils in the lamina propria;
2) Cryptitis and crypt abscesses;
3) Erosions and ulcers.

Marginated neutrophils (i.e., neutrophils confined to the endothelial layer of small vessels) may be the result of the bowel cleansing process or the colonoscopic procedure itself; it is, therefore, not included as a manifestation of active colitis. These findings may be focal or diffuse; in my opinion, their pattern of distribution cannot reliably be used to distinguish infectious colitis from IBD. Inflammatory pseudomembranes, increased eosinophilia of the lamina propria, and mucin-depleted (‘withered”) crypts associated with a relative paucity of neutrophils are findings most suggestive of an ischemic injury or infection with toxin-producing bacteria such as C. difficile and enterohemorrhagic E. coli (2-6).

**Chronic colitis**

The histologic features of chronic colitis are summarized below:

1) Increased mononuclear cell inflammation in the lamina propria;
2) Crypt distortion and atrophy;
3) Surface villiform change;
4) Basal plasmacytosis in colonic mucosa distal to the cecum;
5) Basal lymphoid aggregates;
6) Paneth cells identified distal to the transverse colon.

Determination of increased mononuclear cell inflammation in the lamina propria is the most subjective and probably the least reliable criterion of chronic injury. This analysis is hampered by the fact that mononuclear cells are normally more numerous in the right than in the left colon, by variations in section thickness that may give the false impression of increased cellularity, and by the fact that increased mononuclear cell inflammation is typical of the resolving phase of infectious colitides (5,7). Precision in diagnosis is important because the pathologist’s diagnosis of “increased mononuclear inflammation” in colonic mucosa may be translated by clinicians to mean IBD (7). In most cases of established IBD, other features to support chronicity are present, particularly if multiple specimens are submitted. Basal plasmacytosis distal to the cecum is an excellent marker of chronicity but may be absent in the early phase of IBD and may diminish over time in inactive disease (8).

These changes of chronicity, however, are not synonymous with IBD. They may develop with the healing of any severe mucosal injury such as necrotizing enterocolitis of newborn infants or the colitis associated with Hirschsprung’s disease. The diagnosis of IBD, therefore, must be supported by appropriate clinical, imaging, endoscopic, and, in some cases, serologic data. If the pathologist has no information about the patient, it is best to diagnose such specimens descriptively as “findings compatible with healed injury” rather than “chronic inactive colitis” because the latter phrase will typically be interpreted as IBD by clinicians.
Changes of chronicity may be focal or diffuse within a mucosal specimen or among specimens. In addition, finding even a single abnormality among those summarized earlier is sufficient to support a diagnosis of chronic injury. Multiple abnormalities secondary to chronicity may be infrequent in pediatric IBD, particularly in the earlier phases of disease (see later discussion) and/or with limited mucosal sampling (9).

In both ASLC and IBD, inflammatory changes may be focal or diffuse within a mucosal specimen. In IBD, active inflammation is distributed throughout the lamina propria, whereas it tends to be confined to the upper half of the lamina propria in ASLC (7); however, this feature may not be evident if only one specimen is available for diagnosis. As expected, the major discriminator between ASLC and IBD is the presence of unequivocal features of chronicity in initial or follow-up biopsy specimens (3-5,7,10). Once chronicity is established, the clinician can correlate this finding with the remainder of the patient’s profile to exclude known causes of chronic injury (such as Yersinia infections and certain parasitic diseases) before establishing a diagnosis of IBD.

Artifacts That May Confound the Interpretation of Colonic Mucosal Inflammation

The colonoscope itself may be associated with minor mucosal damage. Since biopsies are obtained as the last procedure of the endoscopy, the colonoscope may have had time to irritate the mucosa, resulting in vascular congestion as well as edema and fresh hemorrhage in the lamina propria. When unaccompanied by epithelial damage or an inflammatory infiltrate, these findings, which are most prevalent in the distal colon, are best interpreted as iatrogenic in origin (2,11,12).

The colonic cleansing procedure may also induce iatrogenic lesions. Oral non-absorbable lavage agents and isotonic saline enemas are gentle and do not injure colonic mucosa; therefore, they are favored for use in infants and young children (13). The cleansing process, however, may be prolonged, resulting in poor compliance and inadequate cleaning. Therefore, a more vigorous but shorter cleansing protocol using sodium phosphate (taken orally and/or as enemas), oral magnesium citrate, and/or bisacodyl enemas and suppositories is often employed, particularly in older children and adolescents. These agents may produce mucosal lesions.

Sodium phosphate preparations (and, to a lesser extent, magnesium citrate) may produce aphthous ulcers with these characteristics:

1) Endoscopy: Discrete, small (1-3 mm), sharply defined erosions, surrounded by an erythematous halo. Most frequent in the rectosigmoid. Adjacent mucosa typically unremarkable;
2) Histology: Either an erosion overlying a lymphoid follicle or a focal, superficial, ischemic-type lesion with mucin-depleted crypts, modest active inflammation, and fibrinous exudate (14).
The prevalence of this change varies from 6% to 24% in recent clinical series (13-16). In a pediatric patient being evaluated for the cause of diarrhea, the differential diagnosis of these findings includes infectious colitis, drug-related injury (e.g., NSAID’s), and low-grade IBD, particularly Crohn’s disease. Recognition of the possible iatrogenic origin of the finding plus integration with all clinical data should permit correct categorization.

Another very common effect of oral/enema sodium phosphate preparations is the development, in otherwise unremarkable mucosa, of discrete foci of basally localized neutrophil cryptitis associated with apoptosis and, in some cases, an infiltrate of eosinophils. This constellation of findings is unaccompanied by crypt destruction (crypt abscesses), crypt architectural distortion, or increased mononuclear cell inflammation and macrophages in the surrounding lamina propria (13,15-17).

In the strictest sense, this constellation of findings does represent a “focal active colitis”. Given the fact that it is typically iatrogenic in origin, however, I recommend using a more descriptive term such as “non-specific basal cryptitis”, with a comment that such changes may be the result of bowel cleansing agents. The reason for this usage is that, based on my experience, the unqualified term “focal active colitis” will be interpreted by clinicians as representing either an infectious colitis or IBD, especially Crohn’s disease (7). As discussed by Xin and colleagues (18), focal active colitis that is likely caused by disease is characterized by involvement of a longer length of the crypts by inflammation than in iatrogenic injury, by the development of crypt abscesses, and by an increased concentration of lymphocytes and macrophages (or even mucin granulomas) in the adjacent lamina propria. Some biopsies may contain changes intermediate between typical iatrogenic effect and the classic features of true focal active colitis just described. In this situation, a descriptive diagnosis is again justified to alert the clinician that the differential diagnosis for the histologic findings includes infections, IBD, and bowel prep effect.

**Inflammatory Bowel Disease**

Accurate categorization of IBD cases as UC and CD has become increasingly important for several reasons:

1) Surgery is considered curative for UC but not for CD;
2) Patients with UC but not CD are candidates for creation of ileoanal pouches;
3) More specific treatments to treat the two conditions are emerging. For example, infliximab (Remicade), a chimeric monoclonal antibody to tumor necrosis factor-alpha has been successful in inducing remission in many patients with therapy-resistant CD. Proper patient selection is crucial because use of infliximab has been associated with a significant risk of serious infections, including reactivation of latent tuberculosis (19).
Clinical features

IBD is common in young patients: about 10% of cases develop before age 15 years and 30% of cases before the age of 25 years (20). It develops more commonly in Caucasians than in other races, and recent data from this country suggest that CD is becoming more common than UC (21). The clinical features of pediatric UC and CD are, in general, similar to those in adults. Features unique to pediatric IBD, however, are delayed puberty and growth failure. The latter is twice as prevalent in CD as in UC, is related more to disease activity than to corticosteroid therapy, and is especially associated with ileal disease (20,22). A family history of IBD is noted in 10% to 30% of patients (21).

Unlike the situation in adults, pediatric patients with certain congenital disorders may present with signs and symptoms that mimic IBD. The major conditions having this presentation include:

1) Glycogen storage disease, type Ib: terminal ileitis/colitis; perianal disease (mimics CD);
2) Chronic granulomatous disease: chronic active colitis with non-necrotic granulomas (mimics CD);

These entities are characterized by immunodeficiencies and impaired processing of infectious agents. In infants and young children with chronic colitis, these conditions should be ruled out; true idiopathic IBD is rare before the age of 2-3 years (21).

The findings of several recent studies also support the notion of an increased risk of IBD in celiac disease, with rates of IBD 3 to 8 times greater than in controls (23). This association suggests that patients not responding to adequate treatment for celiac disease should be evaluated for ileocolitis; conversely, treated IBD patients with persistent diarrhea should raise the possibility of concurrent but unrecognized celiac disease. The various possibilities can be sorted out by clinical, endoscopic, histologic and serologic findings. For example, increased villous intraepithelial lymphocytes are not characteristic of the duodenitis associated with IBD; their presence should at least raise the possibility of undiagnosed celiac disease (23,24).

Natural history

Pediatric IBD is characterized by exacerbations and remissions, but the natural histories of UC and CD are somewhat different (22). At first presentation, UC is characterized by more extensive endoscopic and histologic disease in children than adults. In recent series, pancolitis has been documented 50% or more of children compared to 8% to 20% of adults; conversely, proctitis or localized left-sided disease is present in approximately 25% of children but in up to half of adults at initial evaluation (20-22,25-27). Over time, progression to more extensive disease, including pancolitis, occurs in 50% to 70% of children with initially limited disease (compared to 10% to 30% of adults), especially those with an early age of onset and/or severe disease (20,28-31). At least one relapse will have
been documented in about 90% of pediatric UC patients after 10 years of follow-up, and approximately one-third will have had a colectomy, chiefly because of unresponsiveness to medical therapy, prepubertal growth failure or fulminant colitis (“toxic megacolon”) (20). Since by current definition UC is a disease confined to the colon, proctocolectomy is curative, and the patients are excellent candidates for creation of an ileoanal pouch.

Pediatric CD first presents as small bowel disease in 40% of patients, as ileocolitis in 30%, and colitis in 30%. Of interest, CD tends to present in the colon in very young patients, with ileal disease rarely documented before the age of 8 to 10 years. Ileal disease then increases in prevalence throughout adolescence. The reason for this finding is unclear, but may be related to immunologic changes associated with lymphoid follicle development and enlargement (Peyer’s patches) in the second decade of life (32).

Early surgery is much more common than in UC: up to 50% of pediatric patients with CD will have surgery for complications of the disease within 5 years of diagnosis and up to 90% will have an operation within 15 years (22). Surgery is performed for reasons like those for UC and for additional CD-related indications including treatment of intestinal fistulas and abscesses, gut obstruction, and intractable perianal disease (20,33). Since CD is a condition that may involve the entire gastrointestinal tract, surgery is limited to removal of clinically significant lesions and is usually not curative. Asymptomatic recurrences have been documented in up to 70% of patients within one year of surgery, and symptomatic recurrent disease is common within five years. Recurrences typically occur proximal to sites of previous anastomoses or ostomies. Patients with CD are usually not considered good candidates for ileoanal pouches because of the risk of recurrent disease in the pouch (20,22).

Pathologic findings

The pathologic features of UC and CD are identical in children and adults and have been codified in many recent reviews (6,7,34). The distinguishing features of classic UC and colonic CD are summarized in Table 1.

Classic UC begins in the rectum and, if it progresses, does so in a continuous retrograde fashion. Although by definition UC is confined to the colon, it may be accompanied by superficial mild non-specific mucosal inflammation in the terminal ileum (“backwash ileitis”) (see later discussion). At microscopy, the inflammatory process in UC is superficial (confined to the mucosa and submucosa) and is diffuse within the mucosa of the involved segment (34).

In contrast, colonic CD typically begins as localized right-sided or multifocal disease and progresses in a patchy fashion, with “skip areas” of uninvolved mucosa. Characteristic histologic findings include granulomas, deep or transmural inflammation (often characterized by the presence of lymphoid aggregates in the muscularis propria and/or at the muscularis/serosal interface), deep mural fissures or fistulas, and, in a minority of patients, a necrotizing or giant cell vasculitis. In the mucosa, the inflammatory lesions are often focal rather than diffuse (34,35).
Although diagnostically important, granulomas are not invariably present in otherwise typical cases of Crohn’s colitis. Even with serial sectioning, granulomas are detected in only approximately one-third of mucosal biopsy specimens from patients with Crohn’s colitis. In children with Crohn’s colitis, granulomas tend to be more common in the rectosigmoid than elsewhere in the colon. Also, prospective data suggest that the prevalence of granulomas decreases with increasing duration of disease, perhaps due to the effects of medical therapy; thus, granulomas may be more often detected in children than in adults (36).

The sarcoid-like granulomas characteristic of CD must be distinguished from foreign body-type granulomas and from the non-specific mucin granulomas that may be present in both UC and CD. Mucin granulomas are typically adjacent to or in direct contact with inflamed or ruptured crypts, tend to be poorly formed, and often contain giant cells (2,34). Their true nature can be determined by detection of intracytoplasmic mucin using stains such as the alcian blue-PAS with diastase pretreatment.

In mucosal biopsy specimens from untreated patients, the presence of non-mucin granulomas and focality of colitis are the best discriminators for CD (6,7). Focality of colitis in the rectosigmoid in children, however, should be interpreted with caution since it may be present at the onset of UC in this population (see later discussion). In adults, villiform surface change, crypt architectural distortion and crypt epithelial mucin depletion are said to be more common in UC than CD (6,7), but I have not found these to be particularly helpful distinguishing characteristics in pediatric IBD. The pathologist can offer the largest amount of useful information if, at the onset of pediatric IBD, colonoscopy (rather than flexible sigmoidoscopy) with protocol sampling of even endoscopically unremarkable mucosa is performed. Of 42 pediatric patients ultimately proven to have IBD in one recent study, 10 had normal rectosigmoid biopsy specimens. Additional, more proximal, sampling confirmed a diagnosis of CD in 60%; the remaining four patients were later classified as either UC or indeterminate colitis (26). Finally, as noted earlier, ileal disease is rare in CD before the age of 8 to 10 years. Therefore, untreated idiopathic colitis without distinguishing features (granulomas, focality) in very young patients can only be tentatively classified as UC; some of these cases may declare themselves to be CD later in their clinical course by the development of unequivocal ileal involvement (27,32).
Table 1
Classic Distinguishing Features of Ulcerative Colitis (UC) and Crohn’s Colitis (CC)

<table>
<thead>
<tr>
<th></th>
<th>UC</th>
<th>CC</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Gross features</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Isolated right-sided</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>colitis</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rectal involvement</td>
<td>Yes</td>
<td>Variable</td>
</tr>
<tr>
<td>Distribution</td>
<td>Diffuse</td>
<td>Diffuse or focal</td>
</tr>
<tr>
<td>Method of extension</td>
<td>Continuous(^{(a)})</td>
<td>Often discontinuous (“skip areas” of normal colon)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Involvement of gut</td>
<td>No(^{(b)})</td>
<td>Common</td>
</tr>
<tr>
<td>proximal to colon</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fistulas</td>
<td>No</td>
<td>Occasional</td>
</tr>
<tr>
<td>“Creeping” serosal fat</td>
<td>No</td>
<td>Common</td>
</tr>
<tr>
<td>Thickened bowel wall</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Strictures</td>
<td>Rare</td>
<td>Occasional</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Microscopic features</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Inflammation confined</td>
<td>Yes</td>
<td>Uncommon</td>
</tr>
<tr>
<td>to mucosa and submucosa</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Transmural inflammation</td>
<td>No(^{(c)})</td>
<td>Common</td>
</tr>
<tr>
<td>Fissuring ulcers</td>
<td>No(^{(c)})</td>
<td>Yes</td>
</tr>
<tr>
<td>Fistulas</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Sarcoid-like granulomas</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Distribution of</td>
<td>Diffuse</td>
<td>Focal or diffuse</td>
</tr>
<tr>
<td>inflammatory changes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>in mucosal specimens</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vasculitis</td>
<td>No</td>
<td>Yes</td>
</tr>
</tbody>
</table>

\(^{(a)}\)Recent studies document appendiceal or cecal involvement in some cases of localized left-sided UC
\(^{(b)}\)Possible upper gastrointestinal involvement by otherwise classic UC is a subject of current debate (see text for details)
\(^{(c)}\)May be present in fulminant colitis/toxic megacolon
Atypical Features and Controversial Aspects of Pediatric IBD

Unusual features in rectosigmoid mucosal biopsies at the onset of pediatric UC

At first presentation and before therapy, the majority of adult patients with UC (>90%) will have diffuse active colitis, usually with features of chronicity, in rectosigmoid mucosal specimens (25,37). Initial rectosigmoid specimens in children ultimately shown to have UC, however, demonstrate focal colitis and/or the absence of chronic changes in approximately one-third of patients and are completely normal in 4% to 8% (25,37,38).

These atypical findings are not specifically related to the patients’ ages at the onset of colitis (although they are predominantly found in patients younger than 10 years), the duration of symptoms before endoscopy, the symptoms themselves, or the ultimate evolution of UC (i.e., development of diffuse distal disease, proximal progression over time) (9,25,38). The reasons for these findings are unknown. One suggestion is that children may be evaluated earlier in the course of UC than adults (25-37); however, it is also clear that changes of chronicity may develop within a few weeks or months of symptom onset (5,9).

This presentation of distal UC with focal disease and a paucity or lack of features of chronicity in pediatric patients raises several diagnostic possibilities and stresses the need for a complete evaluation of the patient. First, it should be recognized that ulcerative colitis is not excluded by these findings (18,25,37,38). Second, they may represent a non-relapsing, infectious-type colitis, which often is patchy and may have rectal sparing (39). Crohn’s disease also enters the differential diagnosis; detection of focal proximal colitis, granulomas, ileal disease or perianal disease would support that diagnosis.

The predictive value of focal active colitis for development or recognition of CD once the confounding conditions discussed in the preceding paragraph have been eliminated has recently been examined. In a cohort of 29 pediatric patients with focal active colitis, 8 (28%) developed CD; most of the remainder had either infectious colitis or remained idiopathic (18). In contrast, focal active colitis in adults evolved into a diagnosis of CD over time in fewer than 15% of patients (40,41). One possible reason for the difference in outcome between the two populations is that unlike the case in adults, colonoscopy in children is typically performed for evaluation of abdominal pain, diarrhea or hematochezia rather than cancer surveillance, thus creating a bias towards detection of inflammatory diseases. As in children, many cases of “focal active colitis” (particularly those with minor abnormalities) in adults are likely secondary to effects of bowel cleansing agents such as sodium phosphate (18,40,41).
Effects of medical therapy on the histology of UC in colonic mucosal biopsy specimens

The classic teaching has been that quiescent UC heals with fixed morphologic changes that permit continued recognition of the colonic mucosa as injured. In 1993, however, Odze and colleagues demonstrated that medical therapy of left-sided UC with topical 5-aminosalicylic acid caused reversion of colonic mucosa to a normal appearance in 64% of patients (42). Since that time, several authors have confirmed and extended this observation. The results of these studies document that in patients with established extensive or pancolitis receiving contemporary medical therapy, histologic diffuse disease has become focal within the colon in up to 54% of patients and the rectum has become unremarkable in up to 34% on one or more occasions during follow-up (8,43-47). These results were not related to the duration of disease or the type of therapy employed (systemic or topical) (47).

Rectal sparing and focal colitis are typical of CD. Thus, to avoid diagnostic confusion, it is important to know the medication history of patients with presumed UC in whom such findings are documented. In both children and adults with chronic IBD, unfortunately, such information is often not available at the time of biopsy specimen interpretation. In this situation and in the absence of granulomas, focality and rectal sparing should be described but not interpreted, with the comment that prior medical therapy may have affected the histologic findings.

Appendiceal involvement as a “skip lesion” in UC

In several retrospective studies of colectomy specimens, the authors have attempted to determine the prevalence of appendiceal inflammation in patients with UC of various extents. Appendiceal involvement by UC must be distinguished from incidental acute appendicitis and has been defined as a lesion confined to the mucosa with architectural and cellular features of chronicity that may or may not be accompanied by activity (48). Not unexpectedly, the prevalence of chronic appendicitis in patients with pancolitis (whether due to UC or CD) is high, occurring in approximately 60% of cases (48,49).

Of greater interest is the prevalence of this finding in UC patients with less than pancolitis (i.e., with at least cecal/right-sided sparing in the colectomy specimen). In this group, appendicitis has been reported in 15% to almost 100% of pediatric and adult patients, giving rise to the notion of appendicitis as a “skip lesion” in otherwise classic UC (48,50-53). In a follow-up study of ileoanal anastomoses performed on patients with appendicitis as a presumed “skip lesion” of UC, there were no cases with outcomes suggesting that a diagnosis of CD had been missed (54).

A criticism of these studies relates to their retrospective nature (49,55). The extent of sampling of the cecum and right colon has not been performed in a standardized fashion, so that the extent of sampling has been either limited or unknown. Also, information about prior medical therapy and its possible effects on damaged mucosa has not been provided. Thus, whether or not this portion of the colon has been truly “normal”
at all times is debatable. However, appendiceal involvement has been documented in some cases that clearly represent otherwise localized left-sided UC (52). At present, an apparent appendiceal “skip lesion” should not be construed as evidence of CD without additional supportive clinical, imaging and/or histologic data.

**Cecal periappendiceal orifice inflammation as a “skip lesion” in UC**

Another proposed “skip lesion” of UC is endoscopic and histologic colitis confined to the periappendiceal cecal mucosa in patients with left-sided UC. Strict criteria to exclude CD are needed to support this interpretation: no granulomas, perianal disease, ileal disease or other documented colonic skip lesions should be present (55). The gross periappendiceal mucosal endoscopic findings vary from erythema and tissue granularity and friability to erosions and ulcers (55).

In several recent prospective endoscopic and biopsy surveys of patients with left-sided UC, the prevalence of periappendiceal localized colitis has varied from 54% to 75% (55-58). Biopsies examined from the colon between the left side and the periappendiceal area were normal in most instances (55-57) but on occasion demonstrated microscopic foci of colitis in endoscopically normal mucosa (58). In the study by Yang and colleagues, the prevalence of focal periappendiceal disease was the same in patients with and without prior medical therapy (55), and in an 8-year follow-up of their patients, d’Haens et al detected no features of CD (56). Although there are some issues with the interpretation of the histology in some of these series, the weight of evidence from these prospective studies supports the idea that the endoscopic periappendiceal cecal patch in children and adults does represent a skip lesion of UC and in isolation, therefore, should not be interpreted as evidence of CD.

**The conundrum of backwash ileitis in UC**

A perennial problem in both pediatric and adult IBD at the imaging, endoscopic and pathologic levels is the separation of ileitis due to CD from non-specific terminal ileal inflammation in patients with UC (“backwash ileitis”). Part of the problem relates to the lack of a modern consensus-derived gross and microscopic definition of backwash ileitis (34, 59). A contemporary provisional diagnosis of backwash ileitis in UC has been provided by Heuschen et al: a 5.0 cm or longer segment of terminal ileal mucosal inflammation with no granulomas. In resection specimens, no mural inflammation should be identified (60).

In a recent pediatric series, backwash ileitis (not precisely defined) was identified in 22% of colectomy specimens with pouchitis. Its presence did not predict any excess of problems with ileoanal pouches (such as refractory pouchitis and pouch failure) (61). A combined imaging, endoscopic and histologic study of backwash ileitis (not precisely defined) published in 2003 gave a prevalence of 39% in 18 pediatric patients with UC (62). In the absence of a standardized definition, such studies do not contribute to our understanding of this condition and its precise relation to IBD in children. A recent study by Haskell and colleagues, primarily in adults, has, however, extended our knowledge of...
terminal ileal inflammation in patients with UC in a study of 200 resection specimens in which CD had been rigorously excluded. In this cohort receiving contemporary medical treatment, ileitis was found in a minority of patients (17%), was typically mild and confined to the mucosa, was commonly (but not invariably) associated with pancolitis, and was not associated with an increased risk of ileoanal pouch complications compared to control patients lacking ileal inflammation in their resection specimens (59).

Based on the available data and in the absence of granulomas, ileal mucosal specimens with inflammatory changes should be diagnosed descriptively. In addition to being related to UC itself, such changes could be due to concurrent infection in patients on immunosuppressive therapy for their disease or to drug effects (e.g., NSAID’s). In addition, since some cases are associated with sparing of or minimal activity in the right colon, the term “backwash” is best avoided in the diagnosis, given the absence of definitive information on the pathogenesis of the ileal findings (59,60).

**Upper gastrointestinal tract involvement in inflammatory bowel disease**

Upper gastrointestinal inflammatory lesions of the esophagus, stomach, and/or duodenum in patients with CD have been well described in the literature. Among the possible findings, granulomas, focal duodenitis and focal gastritis in mucosal biopsy specimens are most closely associated with or are the best predictors of colonic CD (63-65). Focal (or “focally enhanced”) gastritis is defined as a localized and sharply defined area of gastritis, with active and chronic inflammation and gland injury, that is surrounded by normal mucosa and is easily identified at low-power inspection (64-67).

In contemporary medical practice, upper gastrointestinal endoscopy with biopsy in often performed before the institution of therapy in pediatric patients with newly diagnosed colonic IBD (both UC and CD). This procedure has lead to some interesting findings that are summarized in Table 2. The protocols and the extent of histologic sampling have not been uniform in these studies, but most are prospective in nature.

Although the gross and microscopic abnormalities are not always described in detail, an intriguing finding in these studies is that the overall prevalence of endoscopic and histologic inflammatory lesions in the esophagus, stomach and duodenum is roughly equal in patients with newly diagnosed and typical colonic UC and CD. When known causes of such inflammation (such as reflux esophagitis and Helicobacter pylori-associated gastritis) are excluded, there still remains a high prevalence of non-specific lesions, particularly H. pylori-negative diffuse gastritis, in both conditions (67-71). Whether this gastritis is incidental or related to IBD is unknown, but the important point is that the mere presence of upper gastrointestinal inflammatory lesions can no longer be used to automatically categorize a patient as having CD.

The presence of granulomas in upper intestinal mucosal biopsies is highly specific for a diagnosis of CD (see Table 2). Their prevalence has varied from 14% to 60% in recent pediatric series, and they are most often found in gastric mucosa (67-71). In contrast, they are much less common in the stomach of adults with CD (a prevalence of
only 5% in the study of Parente and colleagues) (66). In children, they are often detected in gastric and duodenal specimens even when synchronous colonic mucosal biopsies are negative for this finding (67,69,70).

Although focal gastritis can be seen in a minority of UC patients (8% to 21%) and non-IBD controls (2%-19%), it is more common in patients with CD (43% to 65%) with a calculated positive predictive value for colonic CD of 70% to 80% (66,67,71). All examples of gastritis, whether focal or diffuse, should be carefully evaluated to exclude Helicobacter infections as well as allergic and chemical-type injuries. As in the colon, “IBD-associated gastritis” is a diagnosis of exclusion.

In summary, upper endoscopy can be helpful in classifying colonic IBD. Biopsy specimens should be obtained from grossly normal as well as abnormal mucosa to detect treatable localized conditions, such as H. pylori-associated gastritis, and to detect granulomas and focal gastritis (66-68,70,71). In recent pediatric studies of the stomach in IBD, 70%-87% of patients had various inflammatory changes in antral biopsies despite the fact that only a minority (<40%) had endoscopically detectable mucosal changes (chiefly erythema and small ulcers) (67,71). The recent recognition that diffuse, non-Helicobacter chronic gastritis is common in patients with classic UC raises an interesting question: Is it a manifestation of UC? Perhaps adding credence to this possibility is a small number of pediatric and adult cases of “ulcerative enteritis” developing in patients with classic UC, typically after colectomy for treatment of the colitis. In these patients, as recently reviewed by Rubenstein and colleagues, the enteritis is diffuse, confined to the mucosa, lacks granulomas, and responds to the usual medical therapies for UC (72). Clarification of the exact nature of this enteritis (as well as the non-specific gastritis) will require additional prospective studies.

Use of wireless capsule endoscopy will likely increase the detection of small-bowel lesions in patients with IBD. For example, in a recent pediatric study of 20 patients aged 10 to 18 years suspected of having small-bowel CD but negative imaging studies of the gut as well as negative upper and lower endoscopy, 10 (50%) had multiple erosions/ulcers in the jejunum and ileum compatible with CD (73). It will be interesting to determine the rate of similar lesions in pediatric patients thought to have UC.
# Table 2
Upper Gastrointestinal Findings in Pediatric Patients with Crohn’s Disease and Ulcerative Colitis

<table>
<thead>
<tr>
<th>Series</th>
<th>Findings</th>
<th>Crohn’s Disease</th>
<th>Ulcerative Colitis</th>
<th>Controls</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kundhal et al.</td>
<td>Non-specific gastritis</td>
<td>92%</td>
<td>75%</td>
<td>None</td>
</tr>
<tr>
<td></td>
<td>Focal gastritis</td>
<td>52%</td>
<td>8%</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Granulomas</td>
<td>60%</td>
<td>0%</td>
<td></td>
</tr>
<tr>
<td>Tobin et al.</td>
<td>Esophagitis</td>
<td>72%</td>
<td>50%</td>
<td>91%&lt;sup&gt;(a)&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>Gastritis</td>
<td>92%</td>
<td>69%</td>
<td>27%</td>
</tr>
<tr>
<td></td>
<td>Duodenitis</td>
<td>33%</td>
<td>23%</td>
<td>9%</td>
</tr>
<tr>
<td></td>
<td>Granulomas</td>
<td>40%</td>
<td>0%</td>
<td>0%</td>
</tr>
<tr>
<td>Abdullah et al.</td>
<td>Abnormal endoscopy&lt;sup&gt;(b)&lt;/sup&gt;</td>
<td>64%</td>
<td>50%</td>
<td>None</td>
</tr>
<tr>
<td></td>
<td>Abnormal histology&lt;sup&gt;(b)&lt;/sup&gt;</td>
<td>82%</td>
<td>71%</td>
<td></td>
</tr>
<tr>
<td>Ruuska et al.</td>
<td>Endoscopic and/or histologic inflammatory lesions&lt;sup&gt;(b)&lt;/sup&gt;</td>
<td>80%</td>
<td>75%</td>
<td>23%&lt;sup&gt;(c)&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>Granulomas</td>
<td>25%</td>
<td>0%</td>
<td>0%</td>
</tr>
<tr>
<td>Sharif et al.</td>
<td>Focal gastritis</td>
<td>65%</td>
<td>21%</td>
<td>2.5%</td>
</tr>
<tr>
<td></td>
<td>Diffuse, non-H. pylori gastritis</td>
<td>34%</td>
<td>50%</td>
<td>15%</td>
</tr>
<tr>
<td></td>
<td>Granulomas</td>
<td>14%</td>
<td>0%</td>
<td>0%</td>
</tr>
</tbody>
</table>

<sup>(a)</sup>Controls = patients with reflux esophagitis  
<sup>(b)</sup>Includes esophagus, stomach and duodenum  
<sup>(c)</sup>Mixture of non-IBD patients
Summary

Because of the complexity of diagnosing and accurately classifying patients with pediatric IBD, it is clear that any one examination technique, such as histologic analysis of colonic mucosal biopsies, will seldom give a precise diagnosis. Rather, integration of all clinical, imaging, laboratory and morphologic data is necessary to achieve diagnostic accuracy. Pathologists, who usually lack the full range of clinical information, should not feel pressured to render specific diagnoses based on examination of limited colonic mucosal material. Our task is first to determine whether or not colitis is present. If it is, we can state whether it is active, chronic or both and, in untreated patients, give its distribution (focal or diffuse). Finally, the presence or absence of more specific features such as granulomas should be mentioned. Our reports will then be an important contribution to the clinician who must integrate all the data to obtain the most appropriate diagnosis.

References


ENTEROPATHIES OF INFANCY

PIERRE RUSSO, MD

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INTRODUCTION

Acute diarrhea, in the industrialized world, is usually self-limited, infectious in nature, and rarely requires biopsy evaluation. Chronic diarrhea can cause failure to thrive, frequently results in intestinal biopsy and may require parenteral nutrition and other forms of therapy.

Protracted diarrhea has replaced the older term “intractable diarrhea” in the GI literature and denotes a syndrome of severe diarrhea, usually occurring in infants less than a year of age and requiring aggressive nutritional and sometimes immunosuppressive management. Though not strictly synonymous with chronic diarrhea, the two are frequently used interchangeably. Several recent reviews on the subject of protracted diarrhea in childhood are available[1-4].

Disorders resulting in chronic diarrhea of childhood can be grouped into several major pathophysiologic categories:

I. Congenital transport disorders and disaccharidase deficiencies
II. Disorders of enterocyte development
III. Inflammatory and Immune-mediated disorders
IV. Systemic disorders and Tumors
V. Motility disorders
VI. Endocrine disorders – congenital adrenal hyperplasia
VII. Infections – “blind loop”, Giardia, HIV
VIII. Anatomic disorders – Malrotation, short gut, lymphangiectasia
IX. Pancreatic disorders – Cystic fibrosis, Shwachman disease

During the past twenty years, enteric infections and food intolerance have been replaced by rarer congenital conditions, partly due to greater awareness of these entities as well as due to improvements in diagnosis and management
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>ETIOLOGY</strong></td>
<td>n(%)</td>
<td>n(%)</td>
<td>n(%)</td>
</tr>
<tr>
<td>Enteric infection</td>
<td>18 (48)</td>
<td>4 (12)</td>
<td>2 (3)</td>
</tr>
<tr>
<td>Food intolerance</td>
<td>8 (22)</td>
<td>3 (10)</td>
<td>10 (17)</td>
</tr>
<tr>
<td>Autoimmune enteropathy</td>
<td>2 (5)</td>
<td>8 (25)</td>
<td>7 (12)</td>
</tr>
<tr>
<td>Structural enterocyte defects</td>
<td>2 (5)</td>
<td>7 (22)</td>
<td>16 (26)</td>
</tr>
<tr>
<td>Celiac disease</td>
<td>1 (2.5)</td>
<td>0 (0)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Eosinophilic enteropathy</td>
<td>1 (2.5)</td>
<td>1 (3)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Lymphangiectasia</td>
<td>1 (2.5)</td>
<td>1 (3)</td>
<td>2 (3)</td>
</tr>
<tr>
<td>Motility disorders</td>
<td>2 (5)</td>
<td>3 (9)</td>
<td>16 (26)</td>
</tr>
<tr>
<td>Munchausen syndrome by proxy</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td>1 (15)</td>
</tr>
<tr>
<td>Unknown</td>
<td>3 (7.5)</td>
<td>5 (16)</td>
<td>7 (11)</td>
</tr>
</tbody>
</table>


In evaluating intestinal biopsies, it is helpful to remember that disorders resulting in severe diarrhea tend to cluster in different age groups:

**Infant**
- Microvillus inclusion disease
- Autoimmune enteropathy
- Milk and soy protein intolerance
- Protracted infectious enteritis
- Hirschsprung’s disease
- Congenital transport defects
- Munchausen syndrome by proxy

**Toddler**
- Irritable colon of childhood
- Protracted infectious enteritis
- Giardiasis
- Sucrase-isomaltase deficiency
- Tumors
- Celiac disease
- Inflammatory bowel disease

Modified from Vanderhoof J.A. p.32-42 ch 3 in Pediatric Gastrointestinal Disease, Wyllie R. And Hyams J.S., eds W.B. Saunders, 1999

The type of diarrhea can also offer clues: Osmotic diarrhea results from an increased osmotic load in the distal small bowel and colon, with an osmotic “gap” in stool electrolytes, and usually stops when oral feeding is discontinued. It is usually associated with disaccharidase deficiencies and transport disorders, decreased absorptive surface (microvillous inclusion disorder) and hypermotility disorders. On the other hand, secretory diarrhea is characterized by an exudate of mucous, protein or blood, and a variable response to withholding of oral feeds.
Disorders usually associated with secretory diarrhea include congenital fluid transport disorders, inflammatory disorders and increased secretion of gastrointestinal activating substances.

In evaluating intestinal biopsies, disorders that cause protracted diarrhea in early life can be broadly divided into those in which villus architecture is maintained, and those characterized by villus atrophy:

**Normal villous morphology**

- Congenital chloride diarrhea
- Congenital carbohydrate malabsorption
- Congenital sucrose isomaltase deficiency
- Congenital bile acid malabsorption

**Lipid-filled enterocytes**

- Abetalipoproteinemia
- Chylomicron retention disorder

**Villous atrophy**

- **Pauci-inflammatory**
  - Microvillous inclusion disease
  - Epithelial dysplasia
- **Inflammatory**
  - Autoimmune enteropathy
  - IPEX
  - Enterocolitis with dysmorphic features
  - Congenital immunodeficiency disorders
  - Milk intolerance
  - Gluten-sensitive enteropathy

Most transport disorders and diasaccharidase deficiencies do not result in appreciable histologic changes on intestinal biopsies. In fact, normal findings on an intestinal biopsy should prompt the clinician to search for one of these entities.
Lipid transport disorders such as Abetalipoproteinemia, homozygous hypolipoproteinemia and chylomicron retention disease (Anderson disease) are characterized by accumulation of lipid within the enterocytes and clinical characteristics, such as fat malabsorption, low levels of serum lipids, failure to thrive in childhood, neurologic and visual problems resulting from malabsorption of fat-soluble vitamins. These conditions are associated with disorders of apolipoproteins, which reside on the surface of chylomicrons. Apolipoproteins native to the intestine are apolipoprotein (apo) A-I, apo A-IV, and apoB, which has two forms: apo B-100 and apo B-48, both encoded by the same gene located on chromosome 2[5].

Abetalipoproteinemia is an autosomal recessive disorder characterized by the absence of Microsomal Triglyceride Transfer protein (MTP), responsible for assembly of lipoprotein particles, and for the proper folding of ApoB, preventing its premature degradation[6]. Patients have diarrhea and fat malabsorption usually appearing within the first few months of life, with acanthocytosis, and deficiencies in fat-soluble vitamins resulting in retinitis pigmentosa and neurologic symptoms. Serum levels of cholesterol and triglycerides are typically low, and do not rise after a fatty meal. Fat-filled enterocytes are noted on intestinal biopsies of fasting patients, which on electron microscopy are irregular in size and generally non-membrane-bound. No lipid is noted in the extracellular space. Chylomicron retention (Anderson) disease is similar to abetalipoproteinemia in its gastrointestinal manifestations and impact on growth, though acanthocytosis is usually absent and neurologic and ocular abnormalities are much less severe. Also, in contrast to abetalipoproteinemia, serum fasting triglyceride levels are normal and hypocholesterolemia is less marked. The basis for the disorder appears to be an inability to export chylomicrons from the enterocyte into lacteals[7]. Fat-filled enterocytes are also seen in small bowel biopsies. On ultrastructure, lipid droplets are more uniform in size than the droplets
in abetalipoproteinemia, and some appear to be bounded by the membranes of the endoplasmic reticulum[8].

**Microvillus Inclusion Disease (MVID)**

Initially described by Davidson et al in 1978, and subsequently by investigators in other parts of the world, MVID is characterized by refractory secretory diarrhea occurring within the first week of life[9, 10]. There is a roughly equal sex incidence. Birth history is usually unremarkable, without evidence of polyhydramnios, as is seen in congenital chloride diarrhea. Tests of small intestinal function are diffusely abnormal[11]. The pattern of inheritance appears to be autosomal recessive[10].

Small bowel biopsies are characterized by severe villus atrophy, with little or no crypt hyperplasia, and without significant inflammation. The pathognomonic features are present on ultrastructural study and include absent or small stubby microvilli, vesicular structures located towards the apex of the enterocytes containing microvilli, and granules containing dense amorphous material. Even without an electron microscope, this disease can be strongly suspected on paraffin sections by the absence of a distinct brush border on the periodic-acid-Schiff (PAS) stain, and by the presence of PAS-positive diastase-resistant densities at the apex of the enterocytes. Similar observations are noted using immunohistochemical staining for alkaline phosphatase, CEA, and more recently, with anti-CD 10, a membrane-associated neutral peptidase[12]. Microvillus inclusions have also been reported in the colon, gallbladder and renal tubular epithelium in these patients[11]. These inclusions can sometimes be difficult to detect, and several biopsies may be required to identify them. It appears that MVID represents a spectrum of entities, ranging from cases with typical early presentation and diagnostic inclusions
on EM, to cases in which the clinical presentation is atypical, or cases in which inclusions either
cannot be demonstrated (microvillus dystrophy), or the inclusions are “atypical”[13-16].

In a multicenter survey of 23 patients with MVID, the one-year survival was less than
25%[10]. Medical therapy in these patients has generally proven to be ineffective and small
intestinal transplantation currently is the best treatment[10, 17, 18]. Instances of apparent
resolution of the disease with improvement of the mucosal features on long-term TPN have been
reported[19].

**Tufting Enteropathy (Epithelial dysplasia )**

First described by Reifen et al, then by Goulet and colleagues, patients with tufting
enteropathy present in the neonatal period with a watery diarrhea. Prenatal history is uneventful
and the disease appears to be inherited in an autosomal recessive fashion, as suggested by the
finding of other affected siblings and frequent parental consanguinity[20, 21].

Its histological hallmarks are severe villus atrophy with the formation of “tufts” of
rounded, tear-drop shaped enterocytes which appear to shed into the lumen. There may be a mild
increase in the lamina propria inflammatory cells, but intraepithelial lymphocytes (IELs) do not
appear to be significantly increased. Transmission electron microscopy has revealed a normal
brush border. Abnormalities in immunohistochemical staining for basement membrane laminin
and heparan sulfate proteoglycan, ultrastructural irregularities of desmosomes, and abnormal
distribution of α2β1 integrin have been observed in patients with this disorder, suggesting that
the early onset of the disease and its characteristic histomorphological changes may result from
abnormal cell-cell and cell-matrix interaction[22, 23]. Dietary manipulations, antibiotic and
immunosuppression have proven ineffective in these patients, and long-term TPN and small-
bowel transplantation are the current treatment options for these patients[11].
A probably related disorder resulting from the absence of intestinal epithelial α6β4 integrin has been described. The salient features are severe watery diarrhea in the first two weeks of life in association with pyloric atresia, with extensive sloughing of the epithelium observed on biopsies[24].

**Autoimmune Enteropathy**

The term “autoimmune enteropathy” (AIE) was proposed by Unsworth and Walker-Smith to describe a syndrome of protracted diarrhea with the presence of autoantibodies against gut epithelium[25].

More than 100 cases have since been reported in the past twenty years[26], most cases occurring in infancy or the first year of life. There is a strong male preponderance, family history of other affected siblings, frequent extra-intestinal involvement and various circulating autoantibodies[27, 28]. However, this entity has also been reported in older children[29] and even adults[30-32], in whom it may be responsible for a proportion of cases referred to as “refractory sprue”. Intestinal biopsies in most of the reported cases are characterized by severe villus atrophy and variable crypt hyperplasia. Concomitant colitis and gastritis are present in the majority of cases. Marked inflammatory destruction of intestinal crypts with extensive apoptosis is a feature noted in many of the cases. These findings are similar to those noted in severe intestinal graft-versus-host disease, and suggest an abnormal immune-mediated attack against intestinal epithelium.

The hallmark of this entity is the presence of circulating anti-enterocyte antibodies manifest as positive staining in a linear pattern along the apex and baso-lateral border of the
The antibodies are predominantly IgG, and have been described as complement-fixing, though IgM and IgA have also been described[33]. Antibodies reacting against mucus or goblet cells have also been described in occasional patients with an apparently related entity, and intestinal biopsies in these cases have shown a marked depletion of goblet cells[29, 34].

Extra-intestinal disease in these patients includes insulin-dependent diabetes, glomerulopathy and hemolytic anemia, as well as pulmonary and dermatologic manifestations[26]. Numerous autoantibodies have been detected in these patients, including anti-smooth muscle, anti-endoplasmic reticulum, anti-mitochondrial, anti-parietal, anti-adrenal, as well as anti-DNA and anti-ANA antibodies.

AIE has been associated with with several immunodeficiencies and autoimmune diseases:

**Autoimmune Enteropathy: Associated Conditions**

- IPEX Immunodysregulation / polyendocrinopathy / enteropathy / X-linked.
- Immunoosseous dysplasia, Schimke type
- Usher syndrome (hyperinsulinism with enteropathy and deafness)
- Autoimmune Polyendocrinopathy, Candidiasis and Ectodermal dysplasia

Perhaps the most well-recognized such association is with the X-linked syndrome of Immunodysregulation, Polyendocrinopathy and Enteropathy (IPEX) syndrome, initially described by Powell[35]. This syndrome has been related to mutations of the FOX P3 gene (Xp11.23-Xq13.3)[36, 37], and is the human equivalent of *scurfy* in the mouse[38]. The protein encoded by the gene, Foxp3, is expressed in regulatory CD4+/CD25+ T cells and is essential for immune homeostasis[39]. Scurfy mice are characterized by failure to thrive, overexpression of
CD4+/CD8+ lymphocytes, extensive multiorgan inflammatory infiltration and over production of cytokines[40].

Mortality in cases of AIE has been high, occurring in up to one-third of reported cases, but more frequent in the older descriptions. Diet changes and steroids alone have been largely ineffective, and the use of immunosuppressants appears to be necessary in most cases. Cyclosporine has induced a remission in several studies, though some remained resistant. A positive response to Tacrolimus has been reported in patients refractory to other therapies[41, 42].

“Syndromatic Diarrhea”

Girault et al described a group of patients with dysmorphic features, consisting of a prominent forehead, broad nose, hypertelorism and wooly, easily removable abnormal hair (trichorrhexis nodosa), in association with intractable diarrhea and immunodeficiency, which they termed “syndromatic intractable diarrhea”, because of the constellation of extra-intestinal manifestations[43]. Most of the patients reported were of Middle Eastern origin, and there was parental consanguinity and a history of similarly affected siblings. The immunodeficiency was characterized by impaired T-cell and antibody responses despite normal immunoglobulin levels. Intestinal biopsies revealed moderate villous atrophy without a conspicuous increase in inflammatory cells.

Other syndromes associated with early severe diarrhea include:

- Epidermolysis Bullosa with Pyloric atresia
- Tricho-hepato-enteric syndrome
- Infantile Systemic Hyalinosis
- Phosphomannose Isomerase Deficiency (CDG type Ib)
- Hypohidrotic Ectodermal Dysplasia
- Progressive Familial Cholestasis, PFIC I (ATP8B1)
- Lysinuric Protein Intolerance

Hair abnormalities can be seen in association with chronic diarrhea in other disorders, typically in *Menke’s* disease, Zinc deficiency and Lysinuric protein intolerance. Distinct hair shaft anomalies have been reported in two siblings as the tricho-hepato-enteric syndrome, in which thin sparse hair (trichomalacia) was seen in association with hypertelorism, chronic diarrhea, a progressive hypermethioninemia and a hemochromatosis phenotype[44].

**Congenital Immunodeficiencies**

Over 50% of children with immunodeficiencies present with chronic diarrhea, and the gastrointestinal tract is the second most common site of infection in these cases, after the pulmonary system. Immunodeficiency states may predispose to various abnormal intestinal immune reactions because of increased intestinal permeability, resulting in heightened exposure to dietary or microbial antigens.

**Primary Immunodeficiency Disorders with Prominent Gastrointestinal Manifestations**

**Antibody Deficiencies**
- X-linked agammaglobulinemia
- Common variable Immunodeficiency
- IgA deficiency
- Immunodeficiency with elevated IgM (hyper-IgM syndrome)
- Transient hypogammaglobulinemia of infancy
- IgG subclass deficiency

**Combined Immunodeficiencies**
- Severe combined immunodeficiency
- Major histocompatibility complex Class II deficiency

**T-Cell Disorders**
- Wiskott-Aldrich syndrome
- Ataxia telangiectasia
- DiGeorge syndrome

**Disorders of Phagocyte Functions**
- Chronic granulomatous disease
- CD11 / CD18 Leukocyte adhesion molecule deficiency
- Hermansky Pudlak syndrome
- Glycogen storage type IB

Other Disorders
- Chronic mucocutaneous candidiasis
- Autoimmune Polyendocrinopathy/candidiasis/ectodermal dystrophy
- IPEX (Immunodeficiency, Polyendocrinopathy, Enteropathy, X-linked)

*Common Variable Immunodeficiency (CVID)* is one of the more common primary immunodeficiencies. Chronic diarrhea is frequent in these patients, and other gastrointestinal manifestations include chronic active hepatitis and hemorrhagic gastritis[45, 46]. Histologic findings in the intestines fall into four major categories: infections (frequently *Giardia*), nodular lymphoid hyperplasia (NLH), celiac-like villous atrophy, and a severe Crohn’s-like enterocolitis. There is an absence of germinal follicles in the lymphoid tissues of the GI tract, and near absence of plasma cells in the lamina propria.

Chronic diarrhea refractory to a gluten-free diet in association with villus atrophy has been observed in patients with CVID, and major differentiating features from celiac disease include a crypt-destructive inflammatory process in association with prominent apoptosis, reminiscent of autoimmune enteropathy[47]. An accumulation of foamy macrophages in the lamina propria, resembling Whipple’s disease or chronic granulomatous disease, has been noted[47].

*IgA deficiency* is the most common primary immunodeficiency, with a prevalence of approximately 1 in 300[48]. Gastrointestinal disorders include *Giardia* infection, celiac disease, chronic inflammatory bowel disease, atrophic gastritis with pernicious anemia, food allergies and nodular lymphoid hyperplasia[49]. There is a 10-20-fold increase in celiac disease in patients with IgA deficiency[50, 51], and the clinical presentation, histologic changes and response to gluten are similar to that of non-IgA-deficient patients, the distinguishing feature being an
absence of IgA-bearing plasma cells from intestinal biopsies in the former[52]. Chronic diarrhea may also result from a non-specific enteritis which does not respond to gluten withdrawal[53].

**Severe Combined Immunodeficiency (SCID)** is due to a genetically heterogeneous group of disorders, and is the most severe form of congenital immunodeficiency. Infections, severe protracted diarrhea, failure to thrive and malabsorption are the major clinical manifestations[54, 55]. Neonatal graft-versus-host disease may occur from transplacentally acquired maternal lymphocytes, from the use of non-irradiated blood products, and as a result of bone marrow transplantation, which until recently was the only effective therapy for these children. Intestinal biopsies are typically void of plasma cells, and germinal follicles are absent from the appendix. Chronic viral[56] and bacterial infections[55] may also cause chronic diarrhea.

**Chronic Granulomatous Disease** results from an inability to produce hydrogen peroxide, the oxidant molecules required to kill ingested microbes. Stomatitis, oral ulcers, esophageal and gastric strictures are relatively frequent complications, and intestinal involvement, most often due to infections, occurs in over 50% of patients[57]. A Crohn’s disease-like picture is typically found, with thickening of the bowel wall, stenoses, ulcers and crypt abscesses with large granulomas[58-62]; pigmented histiocytes can be found in otherwise relatively uninvolved mucosa.
### Characteristic Histologic Changes in Intestinal Biopsies in Primary Immunodeficiency States

<table>
<thead>
<tr>
<th></th>
<th>XLA</th>
<th>CVID</th>
<th>IgA deficiency</th>
<th>SCID</th>
<th>CGD</th>
<th>CD11/CD18 deficiency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Paucity of plasma cells in lamina propria</td>
<td>+</td>
<td>+</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nodular lymphoid hyperplasia</td>
<td>+</td>
<td>+</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Giardia</td>
<td>+</td>
<td>+</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Regional enteritis</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td></td>
<td></td>
<td>+</td>
</tr>
<tr>
<td>Foamy macrophages in lamina propria</td>
<td>+</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>+</td>
</tr>
<tr>
<td>Non-specific villous atrophy</td>
<td></td>
<td>+</td>
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<td></td>
</tr>
</tbody>
</table>

Abbreviations: XLA – X-linked Agammaglobulinemia; CVID – common variable immunodeficiency; SCID – severe combined immunodeficiency; CGD – chronic granulomatous disease

### REFERENCES:


Eosinophilic Disorders of Childhood

Eduardo Ruchelli, MD

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Introduction

Allergy and eosinophilia in the gastrointestinal (GI) tract are not equivalent. GI eosinophilia is associated with infections (parasites), nutritional deficiencies, inflammatory processes, neoplasias, and systemic disorders. However, in addition to eosinophilia, other histologic changes usually accompany the increased number of eosinophils in most of these conditions. In contrast, there is a group of GI disorders that are primarily or exclusively characterized by tissue eosinophilia in the absence of known causes for eosinophilia. These primary eosinophilic GI disorders include eosinophilic esophagitis (EoE), allergic proctocolitis, and eosinophilic gastroenteritis. Several lines of evidence support an allergic cause for these disorders but in many patients a specific allergen is not easily identified.

Baseline eosinophils are normally found in the GI tract, specifically in the stomach, small bowel, and large intestine. Eosinophils are present prenatally. They convey antiparasite immunity, release cytotoxic granule proteins and, by interacting with T-cells, aid in antigen presentation, proliferation, and maturation and regulate T-cell responses. Normal levels in the GI tract range from ~5 eosinophils/high-powered field (HPF) in the stomach antrum and fundus to ~35 eosinophils/HPF in the cecum. Geographic variations have been found in the level of eosinophils in the GI mucosa. In addition to their number (subject to the known geographic variation), factors that determine whether eosinophils found in the GI tract should be considered “abnormal” include, their location (superficial vs deep; intraepithelial vs lamina propria), the finding of free granules, and the presence of associated pathologic abnormalities, such as epithelial changes, neutrophilia, and lymphocytosis.

Food allergy is believed to affect 2% to 8% of children younger than 2 years of age. It has been estimated to be the most frequent cause of chronic diarrhea in infancy in the U.S. The foreign antigenic substances are mainly cow’s milk in the infant and egg and soy protein in the older child. There is evidence that minute amounts of substances in breast milk may act as allergens in the infant. Many of these allergic reactions are temporary and appear to wane with age. Cross-reactivity between substances causing gastrointestinal and respiratory allergic reactions has been demonstrated. Predisposing factors in the child include a family history of atopy, IgA deficiency, early exposure to the antigen, and preceding gastrointestinal infection, possibly even including infection by Helicobacter pylori.

Food allergy can be mediated by type I, type III, or type IV immune reactions, singly or in combination. Type I reactions are of immediate onset and are associated with
mast cell degranulation and elevated IgE antibodies. Though infrequently documented on biopsy samples mucosal reactions are characterized by edema and altered motility and permeability. Type III (immune-complex) reactions are not believed to play a major role in food allergy, though they have been reported on occasion. Type IV, or cell-mediated reactions, result in the production of increased levels of IL-4, IL-5, and interferon-gamma, eosinophil chemoattractants secreted by activated lymphocytes and by eosinophils in an autocrine fashion all of which modulate eosinophil recruitment to the gut, the salient histologic feature of allergic enteropathy. In general, allergic disorders affecting the gut are non-IgE-mediated.

A clinical classification of food hypersensitivity reactions has been recently proposed and is presented in Table 1.

**TABLE 1. Classification of Gastrointestinal Food Hypersensitivity.**

<table>
<thead>
<tr>
<th>IgE-mediated disorders</th>
<th>Mixed IgE and non-IgE-mediated disorders</th>
<th>Non-IgE-mediated disorders</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Immediate gastrointestinal hypersensitivity</td>
<td>• Eosinophilic esophagitis</td>
<td>• Dietary protein (allergic) colitis and proctitis</td>
</tr>
<tr>
<td>• Oral allergy syndrome</td>
<td>• Eosinophilic gastroenteritis</td>
<td>• Dietary protein enteropathy</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Celiac disease</td>
</tr>
</tbody>
</table>

*Source: Adapted from Sampson HA and Anderson JA.*

**Eosinophilic Disorders of the GI Tract**

**Eosinophilic Esophagitis**

Over the last decade, it has been recognized that not all cases that present with symptoms of GER and have associated esophageal eosinophilia represent gastroesophageal reflux disease. A number of studies have described a unique group of patients with eosinophilic esophagitis who present with symptoms that are otherwise indistinguishable from those secondary to reflux esophagitis but fail to respond to conventional antireflux therapy. The degree of eosinophilia in this patient population tends to be unusually severe, greater than 20 eosinophils per high-magnification microscopic field. The symptoms as well as the histologic abnormalities frequently improve with either steroid or restricted diet therapy. On the basis of these observations and the frequent extra-intestinal allergic symptoms (asthma, eczema, and chronic rhinitis) experienced by these patients, an allergic etiology for this type of esophagitis has been proposed. However, one of the difficulties in establishing the actual etiology of this form of eosinophilic esophagitis is that in many patients a specific allergen is not easily identified. This form of esophagitis is usually referred as EoE.
While esophageal histology is essential in making the diagnosis of EoE, in some cases, endoscopy may reveal either a ringed appearance or linear furrows; however, these visual features are not pathognomonic for EoE. In some patients, superficial eosinophilic microabscesses in the squamous epithelium give the esophagus an endoscopic appearance characterized by white punctate lesions or a white exudates, which may be initially misinterpreted as Candida esophagitis. The main features of EoE are listed in Table 2.

In patients with EoE, the histologic changes are similar to reflux esophagitis, but the number of intraepithelial eosinophils is usually higher, frequently more than 20 eosinophils per high-magnification microscopic fields. Liacouras et al demonstrated that the clinical and histologic features of allergic esophagitis might evolve over years. Some patients may actually remain asymptomatic and present late in the course of the disease with dysphagia secondary to strictures. When eosinophilic infiltration is not confined to the esophagus but involves other segments of the gastrointestinal tract, classic eosinophilic gastroenteritis should be considered. Whether EoE is a variant of eosinophilic gastroenteritis or represents a different process is not clear.

**TABLE 2. Eosinophilic Esophagitis.**

<table>
<thead>
<tr>
<th>Gastrointestinal symptoms</th>
<th>Vomiting and regurgitation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Epigastric and chest pain</td>
<td>Heartburn</td>
</tr>
<tr>
<td>Nausea</td>
<td>Dysphagia</td>
</tr>
<tr>
<td>Dysphagia</td>
<td></td>
</tr>
<tr>
<td>Extra-intestinal symptoms</td>
<td>Asthma</td>
</tr>
<tr>
<td>Eczema</td>
<td>Chronic rhinitis</td>
</tr>
<tr>
<td>24-hour, pH probe testing</td>
<td>Normal</td>
</tr>
<tr>
<td>Histology</td>
<td>Severe esophageal eosinophilia</td>
</tr>
<tr>
<td>Treatment</td>
<td>Unresponsive to anti-reflux therapy</td>
</tr>
<tr>
<td></td>
<td>Responsive to steroids and restricted diet</td>
</tr>
</tbody>
</table>

Because EoE has only recently been recognized, both acute and long-term complications have been difficult to assess. However, a review of the literature suggests that while EoE has been mainly documented in children, it also occurs in adults. Adults identified with this disorder often suffer from significant dysphagia and the development of esophageal strictures. Thus, if EoE is left untreated in childhood, it may cause future signs and symptoms as have been seen in adults.

While initial reports have demonstrated that the disease responds to corticosteroid therapy, either taken orally or by swallowed inhalation therapy, many studies have demonstrated a high rate of recurrence after discontinuation of therapy. Recently, patients have been found to respond to a food elimination diet using an elementary formula, which results in complete resolution of esophageal eosinophilia and its corresponding symptoms.
Allergic Proctocolitis

Infants with allergic proctocolitis typically present during the first 2 months of life with blood-streaked stools, which may lead to a consideration of an anal fissure or tear. Growth is usually not affected, and peripheral eosinophilia is variable. Breast-fed infants may present at an older age and have less severe symptoms. The diagnosis in an infant fed cow’s milk or commercial formulas is relatively straightforward. Biopsies are performed to rule out other causes of bloody diarrhea in the infant such as infections and Hirschsprung–associated colitis.

Endoscopic features include erythema, mucosal friability, focal erosions, and nodularity, the latter suggestive of nodular lymphoid hyperplasia. Disease is usually limited to the distal colon. Eosinophils may be observed in stool smears, which should also be obtained to detect parasitic infection. Increased numbers of eosinophils, frequently clustered and sometimes accompanied by neutrophils, are characteristically noted in all compartments of the mucosa in biopsies. The number of eosinophils is variable, reportedly ranging from greater than 6 to more than 20 per high power field, and is frequently a focal finding. Eosinophil numbers may also vary as a result of dietary withdrawal therapies often empirically attempted prior to obtaining the biopsies. Infiltration of crypt and surface epithelium is generally readily noted and is an important diagnostic feature. The mucosal architecture is well preserved. Significant crypt architectural distortion is distinctly unusual and, if present, should suggest other disorders. Nodular lymphoid hyperplasia is a frequent concomitant finding. The differential diagnosis includes parasitic infections such as *Dientamoeba fragilis*, which can be identified in stool smears. It should be pointed out that eosinophilic infiltrates can also be seen in cases of Hirschsprung disease and that, conversely, the clinical and radiologic features of allergic colitis can occasionally mimic aganglionosis. The treatment is dietary change to a hypoallergenic formula. The condition is temporary; most patients can tolerate milk after 2 years of age.

Eosinophilic Gastroenteritis

Eosinophilic gastroenteritis involves eosinophilic infiltration throughout the GI tract. It is a rare condition, the etiology of which is unknown, and is the most difficult to treat eosinophilic disorder of the GI tract. Eosinophilic gastroenteritis is, in the broadest sense, a histological observation based upon a mostly subjective impression of increased eosinophils, usually seen in a biopsy sample of the intestine or colon where eosinophils normally reside. Pediatric patients of all ages have been reported, as well as adults, presenting with a bewildering array of clinical manifestations. Peripheral eosinophilia may be, but not always, elevated. Corroborating tests such as the detection of allergen-specific IgE antibodies, by pinprick or by in-vitro radioallergosorbent testing (RAST), are of varying diagnostic usefulness. Numerous case reports document unusual manifestations such as pancreatitis, pyloric stenosis, abdominal mass, intestinal perforation, and duodenal ulcers.
This situation is complicated by the absence of well-defined criteria for an objective assessment of increased eosinophils, especially in the intestinal and colonic mucosa, by the fact that increased numbers of eosinophils can be observed in diseases other than allergic, and by the difficulty in substantiating in an unequivocal manner an allergic etiology in many cases. The difficulty in establishing the diagnosis by histologic examination results from the variability in normal eosinophil counts, as well as by the variability in eosinophilic infiltration even in established disease. Mucosal eosinophil counts in the normal intestine and colon vary according to site, age, geographic location, and even time of year. Establishing the diagnosis histologically is thus fraught with difficulty and should not be attempted without clinical and laboratory correlation.

A greater degree of confidence in the diagnosis is justified in the presence of marked eosinophilic infiltration or large clusters of eosinophils, with epithelial infiltration and damage. Furthermore, increased eosinophils in the gastric body and antrum, where eosinophil counts are usually low, may be a more reliable marker. Perhaps the greatest value of the biopsy is to identify causes other than allergy for the patient’s symptoms.

References:


Celiac disease: Recent observations and some reflections on older ones
Cyrus E. Rubin, MD, University of Washington, Professor Emeritus of Medicine and Pathology, Division of Gastroenterology, Seattle, WA

Selected Bibliography

   The classical clinical description of the “celiac affection” by the great British clinician Samuel Gee. Proposed that “errors in the diet may perhaps be a cause of the disease and that cow’s milk is the least suitable kind of food for these children.” (a prediction of later recognition of lactase deficiency in celiac disease). Cites a child that “wonderfully throve when fed upon a quart of the best Dutch mussels daily”. He concludes that “if a patient be cured at all, it may be by means of diet”.

2. Dicke WK. Doctoral Thesis, University of Utrecht, Netherlands
   The origin of the gluten-free diet.

   2 & 3. These two of a classic series of studies indicate that wheat, barley, and rye as well as oats induce steatorrhea in celiac disease.

   The first histologic observation of the flat lesion in celiac disease at laparotomy by JW Paully – a British physician.

   The first demonstration of the flat intestinal lesion by suction biopsy.

   More extensive observations on the flat lesion in celiac disease in children and idiopathic sprue in adults. This is the origin of the name celiac sprue.

Recognizes the characteristic, but non-diagnostic histology of the flat intestinal lesion of untreated celiac sprue. Therefore proposed “a second criteria for a firm diagnosis of celiac sprue – a well documented and dramatic clinical response after institution of a strict gluten-free diet.” These two criteria remain the cornerstones for the diagnosis of celiac sprue to this day.


Intolerance to gluten in non-celiacs is common and thus the use of a clinical response to a gluten free diet alone is diagnostically unreliable.


This is the first work describing the specificity and sensitivity of serum TTG for the diagnostic screening for celiac sprue.


11 &12 These two papers provide further proof of the superiority of serum TTG in screening for celiac disease.

**HISTOLOGIC PROOF OF INJURIOUS PROPERTIES OF WHEAT, BARLEY & RYE IN CELIAC SPRUE**


An instrument making it possible to biopsy the whole length of the bowel repeatedly while it remains in situ.

First real time histologic demonstration of small bowel injury by wheat in celiac sprue.


Only wheat, barley, and rye destroy small bowel villi in celiac sprue. Oats do not destroy the villi but only non-specifically cause active inflammation in both celiac sprue and normal controls.


Severity of malabsorption is proportional to length of injured small bowel.

SEEKING COMPLICATIONS OF CELIAC SPRUE BY SCREENING?


75% of cases of celiac sprue not responding to a gluten free diet have an aberrant T-cell population similar to that seen in T-cell lymphoma complicating celiac disease.


There were histologic patterns predictive of poor outcome: 1. development of collagenous sprue, 2. marked mucosal thinning, 3. gastric metaplasia, 4. gastric inflammation and lymphoma.


This is one of the best studies of NHL in celiac disease in an otherwise confusing literature. This paper indicates that early diagnosis of celiac disease by mass screening does not significantly reduce its impact on the general population; therefore it comes out against mass screening to detect lymphoma in celiac disease.


Early treatment of celiac disease with a GFD does not decrease risk for autoimmune disease.

“Loss of response to a gluten free diet (refractory sprue) and ulcerative jejunitis are complications of celiac disease that may progress to enteropathy-associated T-cell lymphoma (EATL). Both conditions are characterized by the presence of nonlymphomatous monoclonal populations that show clonal identity with the lymphoma itself is also present in the enteropathic mucosa.”

DERMATITIS HERPETIFORMIS (DH)

   The first description of the flat small intestinal lesion in Dermatitis Herpetiformis (DH).

   Description of the varied small intestinal lesion in a series of DH patients despite almost uniform lack of malabsorption.

   Demonstration that the intestinal lesion in DH is related to gluten.

   Demonstration that a strict gluten free diet of long duration cures the skin lesion in dermatitis herpetiformis.

   Demonstration that rare patients with a normal proximal bowel in DH also carry the gene for celiac sprue in a latent form that can be made to penetrate by a high gluten diet.

GENERAL REFERENCE

   A generally excellent statement but it does not describe the genetic identity of DH and celiac sprue nor does it recognize that serum TTG is the method of choice for screening for celiac sprue because it is reproducible easily and objectively and has a high sensitivity and specificity.

A fabulous book on the origin of human societies and the domestication of plants being the initial step in progression from a primitive hunter-gatherer society to a more complex, sedentary agricultural society. The initial sedentary agricultural society first occurred 8000 BCE in the Fertile Crescent of southwestern Asia, where wild cereals grew prolifically after the end of the Pleistocene era, at the end of the ice age. Emmer wheat and barley were the first plants in the world to be domesticated from the wild cereals in the Fertile Crescent. They remain a major source of nutrition worldwide even though they are two of the three are the toxic grains in celiac disease!


Family study revealing the celiac gene to be a dominant with incomplete penetrance.