Gastritis

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What is gastritis?

- A symptom complex
- Endoscopic appearance of the stomach
- Microscopic inflammation of the stomach
Gastritis: definitions

- Gastritis: inflammation associated with epithelial cell damage and regeneration
- Gastropathy: mucosal injury without inflammation
- Atrophy: loss of normal mucosal glands
- Metaplasia: change in epithelial cell types
gastritis

• There is not a close relationship between clinical symptoms and histologic gastritis
• Although gastritis may not produce symptoms, its complications do.
Gastritis: correlation of endoscopic and histologic findings

- 98 patients with endoscopic mucosal changes attributed to gastritis: 27 had normal biopsy
- 69 patients with normal endoscopic appearance: 63% had histologic evidence of gastritis
- Interobserver variability for some features of gastritis (Gastrointest Endosc 1995;42:420)
Classification of Gastritis and Gastropathy

Gastritides

Chronic Nonspecific
- Diffuse antral-predominant gastritis with *Helicobacter pylori*
- Multifocal atrophic gastritis with or without *H. pylori*
- Diffuse corporal atrophic gastritis

Infectious
- Viral
- Bacterial
  - *H. pylori* (see Chapter 48)
  - Others, including mycobacterial infection
- Fungal
- Parasitic

Granulomatous
- Crohn's disease
- Sarcoidosis
- Foreign bodies
- Infections
- Tumor-associated

Gastropathies

Distinctive Forms
- Collagenous
- Lymphocytic
- Eosinophilic

Miscellaneous
- Gastritis cystica profunda
- Graft-versus-host disease

Reactive (Erosive "Gastritis")
- Aspirin and other nonsteroidal anti-inflammatory drugs/other medications
- Alcohol
- Portal hypertensive gastropathy
- Cocaine
- Stress
- Radiation
- Bile reflux
- Ischemia
- Prolapse/hialtal hernia
- Trauma (e.g., gastric tubes)

Hyperplastic
- Me´ ne´ trier's disease and hyperplastic, hypersecretory gastropathy
- Zollinger-Ellison syndrome (see Chapter 31)
Chronic nonspecific Gastritis classification

- Nonatrophic - type B
  - H. pylori
  - Antrum > corpus
  - Diffuse antral predominant gastritis

- atrophic - type A
  - AUTOIMMUNE (body, fundus)
    - Diffuse corporal atrophic gastritis
    - Autoimmune metaplastic atrophic gastritis
  - MULTIFOCAL (H. pylori, antrum = corpus)
    - Multifocal atrophic gastritis
Chronic nonspecific gastritis

- Diffuse antral gastritis
- Diffuse corporal atrophic gastritis
- Multifocal atrophic gastritis
Natural history of H. pylori Infection

- H. pylori Infection (weeks → months)
  - Chronic Superficial Gastritis (years → decades)
    - Gastroduodenal Ulcer
    - Chronic Superficial Gastritis
    - Lymphoproliferative Disease
    - Chronic Atrophic Gastritis
      - Gastric Adenocarcinoma
H. Pylori gastritis

- Acute
- Chronic
  - Antral predominant gastritis, also called type B, nonatrophic, diffuse antral predominant gastritis
  - Atrophic gastritis, also called type A, multifocal atrophic gastritis, metaplastic atrophic gastritis
Clinical significance of H pylori gastritis

• Acute
  – May be symptomatic with epigastric pain, nausea and vomiting
• Antral predominant gastritis
  – Duodenal ulcer
• Atrophic gastritis
  – Intestinal metaplasia
  – Gastric ulcer
  – Gastric adenocarcinoma
Where to biopsy for H. pylori

- 2 from antrum
- 2 from gastric body
- 1 from incisura: site most likely to show atrophic gastritis and premalignant dysplasia
Where to biopsy for H. pylori
Endoscopic findings of H. pylori

- No distinct endoscopic pattern
- Normal
- Red streaks in antrum
- Erosions and ulcerations
- Hypertrophy, atrophy
Autoimmune metaplastic atrophic gastritis

- Immune response directed against parietal cells and intrinsic factor
- 3x more common in women
- Autosomal dominant disorder
- Northern European
- Associated with other autoimmune disorders: Hashimotos thyroiditis and vitiligo
Autoimmune metaplastic atrophic gastritis: endoscopic findings

- Appearance of multiple polyps
- Absent or inconspicuous rugae in body and fundus
- Submucosal blood vessels visible through thin atrophic overlying mucosa
- Usually no antral involvement
Chronic atrophic gastritis associated with pernicious anemia, fundus
Intestinal metaplasia

• Eventually atrophic glands are replaced by metaplastic epithelium
• H. pylori: Intestinal metaplasia develops at a rate of 1-2% per annum to yield a lifetime risk of 50-75%
Patchy intestinal metaplasia, paler than surrounding mucosa, in the antrum of 52 year old woman with dyspepsia. Test for H. pylori was positive
Intestinal metaplasia of the gastric antrum
H. Pylori and peptic ulcer disease

- Gastritis is found in virtually all patients infected with H. pylori
- In the United States, 80% of pts with DU and 60% with GU are associated with H. pylori
- Fewer than 20% of people with H. p ever develop PUD
- H. p. Rx with antibiotics dramatically decreases ulcer recurrance rate
Gastric adenocarcinoma
Gastritis and gastric cancer

- Adenocarcinoma
  - H pylori
  - autoimmune

- MALT: H. pylori

- Carcinoid: autoimmune
Gastritis and gastric cancer: H pylori and adenocarcinoma

- 2nd most common cancer worldwide
- in US: 8th cancer related mortality in men and 10th in women
- Decreased incidence of gastric adenocarcinoma in western populations parallels decrease in prevalence of H pylori
- H pylori infected individuals have 2-10x increased incidence of gastric cancer
- 36 and 47% of all gastric cancers in developed and developing countries respectively are attributable to H pylori
- Multifactorial
H. Pylori and Gastric adenocarcinoma

Diagram:

- H. pylori Infection in Early Childhood
- Chronic Active Gastritis
- Multifocal Atrophic Gastritis
- Incomplete Intestinal Metaplasia
- Dysplasia
- Invasive Gastric Carcinoma

Environmental Factors
Hypochlorhydria
Chronic gastritis and gastric adenocarcinoma

• Increased risk with intestinal metaplasia; not known if cancer arises from intestinal metaplasia or whether it represents a marker of increased risk
• Autoimmune gastritis: 3-18x increased risk of gastric adenocarcinoma
Gastritis and gastric cancer
MALToma

- Low grade B cell lymphoma
- Mucosal associated lymphoid tissue
MALToma

- associated with chronic H. pylori infection in more than 90% of cases
- Primary gastric lymphoma accounts for 3% of gastric neoplasms and 10% of lymphomas
- 50% of gastric lymphomas are MALT
- H. pylori induces mucosal inflammatory reaction, lymphoid follicles -> B cell monoclonal cells -> autonomous uncontrolled growth
• Gastric MALToma: dense monotonous lymphoid infiltrate in the lamina propria
Autoimmune gastritis and carcinoid tumors

- Loss of parietal cell mass → hypochlorhydria → G cell hyperplasia and hypergastrinemia → chronic stimulation of enterochromaffin like cells by gastrin
Gastric carcinoid
What to do about gastric cancer and gastritis: look for H pylori?

Unknown if treatment for H pylori decreases risk of gastric cancer

• Studies are difficult because of long cancer development process that may take several decades
What to do about gastritis and gastric cancer: look for H pylori?

- some studies show improvement in inflammation and intestinal metaplasia
- 2 studies show improvement in gastritis and superficial epithelial damage but no improvement in intestinal metaplasia or atrophy
- 1 study from China: healthy H. pylori carriers, treated and followed for 7.5 years. No overall decrease in gastric cancer. Subgroup of patients with no precancerous lesions on presentation did have decreased gastric cancer risk. (JAMA 2004;291:187)
What to do about gastritis and cancer risk: look for H. pylori

- If you find H. pylori: eradication should be considered because it is a carcinogen (ASGE)
- Insufficient evidence to recommend screening asymptomatic patients for H. pylori to prevent gastric cancer (up to date)
- Consider testing first degree relatives of patients with noncardia gastric cancer (Mayo)
What to do about gastritis and cancer risk: surveillance scope for dysplasia or cancer?  
ASGE guidelines

• Pernicious anemia: Single endoscopy should be performed to identify carcinoid and gastric cancer

• Endoscopic surveillance of gastric intestinal metaplasia has not been extensively studied in the United states and therefore cannot be uniformly recommended

• Patients at increased risk for gastric cancer due to ethnic background or family history may benefit from surveillance
What to do about gastritis and cancer risk: dysplasia

- Low grade dysplasia: surveillance EGD every 3 months for at least 1 year with topographic mapping biopsy strategy
- High grade dysplasia: consider endoscopic resection or gastrectomy
MALToma

- H. pylori therapy is useful in patients with localized, mucosal or submucosal, nonbulky, flat disease (without metastasis, LN, or diffuse large B cell lymphoma)
- Only 10% of lymphoma patients
- 50-90% complete remission
Infectious gastritis: CMV

- Immunocompromised patient
- Epigastric pain, fever, atypical lymphocytosis
- Endoscopic findings: congested, edematous mucosa, erosions, ulcerations, nodular mucosa
CMV gastritis
CMV gastritis: pathology

- Cytomegalic cells with intranuclear and intracytoplasmic inclusions of cytomegalovirus
Granulomatous gastritis: Crohn’s disease

- Crohn’s disease of the stomach is uncommon
- Almost always associated with intestinal disease
- Nausea, vomiting, epigastric pain, anorexia and weight loss
- Endoscopy: reddened mucosa, irregularly shaped ulcers, erosions, nodular lesions and cobblestone pattern
Crohn’s gastritis

• Severe nodular gastritis in 18 year old male with crohn's disease
Gastropathy

- Hemorrhagic or erosive lesions
- Caused by irritants such as medications or reduction in mucosal blood flow
- NSAIDs, alcohol, trauma, sepsis
- Disruption of normal protective barrier: mucin, bicarbonate, epithelium, PG
Gastropathy

- Endoscopic findings
  - Acute: may be diffuse (NSAIDs and alcohol) or confined to body and fundus (stress)
  - Chronic: usually antrum
NSAID gastropathy
NSAID gastropathy
gastropathy

• Pathology:
  – Acute: subtle changes
  – Chronic: foveolar hyperplasia, edema, increased smooth muscle fibers, vascular dilatation and congestion
  – Few inflammatory cells
Foveolar hyperplasia

- Tortuous, corkscrew appearance
- Marker of increased epithelial cell turnover
- Chemical injury and H pylori gastritis
Bile reflux gastropathy

- Often occurs after gastric surgery
- Asymptomatic or abdominal pain, emesis and weight loss
- Erosions, redness, bile staining of gastric mucosa
- Treatment: sulcralfate, ursodeoxycholic acid, surgery