

GIT PATHOLOGY







ORAL CAVITY

Congenital disorders: hare-lip and cleft palate



xincidence 1 : 950 newborns

Iateral cleft – isolated or complete

➡ fusion defect of the first branchial arch –maxilar process with fronto-nasal lateral process genetic x acquired (environmental) unilateral x bilateral

- cheiloschisis (upper lip) complete/incomplete
- 눡 gnathoschisis (jaw)
- ⇒palatoschisis (hard palate)
- 눡 uranoschisis (soft palate)
- ⇒staphyloschisis (uvula)

medial, oblique, transverse cleft (rare)





сору

Salivary glands



*3 pairs of large major salivary glands, numerous small ones
*serous / mucinous
*secretory units → glandular ducts
*double cell layer – external myoepithelia
*tumors mostly in the parotid gland, in adults usually epithelial tumors



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Salivary gland tumors

🗶 parotid ⇒cca 75%, mostly benign (70-85%) submandibular ➡ 40% malignant minor salivary glands **⇒**50% malignant *****sublingual mostly malignant

Histologic types



- Selected types
- >> pleomorphic adenoma (benign mixed tumor)
- * adenolymphoma (Warthin tumor)
- oncocytoma
- mucoepidermoid carcinoma
- adenoid cystic carcinoma
- malignant mixed tumor

Pleomorphic adenoma (mixed tumor of salivary glands)

- former name "myxochondroepithelioma"
- benign epithelial tumor
- × 80% in the parotid gland, most common parotid tu
- any age, mostly middle-late adult age
- slow-growing firm mass
- well-demarcated, but capsule incomplete
- *frequent recurrences after incomplete resection
- Iow risk of malignant transformation (4%), in long duration, recurrences

Pleomorphic adenoma



× micro:

- histologic diversity
- ductal and myoepithelial tumor cells
- epithelial elements in strands or sheets
- myxoid to chondroid stroma produced by myoepithelia
- \Rightarrow often penetrates the capsule \Rightarrow protuberances



Pleomorphic adenoma





Pleomorphic adenoma



Warthin's tumor (cystadenolymphoma)



- 5-10% of total salivary gland tumors, benign
- ✗ M> F, 6th-7th decade
- Iower pole of the parotid gland
- Iow recurrence rate, malignant transformation (ca, malignant lymphoma) highly uncommon
- risk factors:
 - ⇒ smoking (8x), radiation, EBV
- origin? (heterotopic salivary tissue in a LN; reactive epithelial proliferation + lymphocytic infiltration)
- histology:
 - cystic or cleftlike spaces with double-layered epithelial lining, dense lymphoid stroma (usually + germinal centers)

Warthin's tumor (cystadenolymphoma)





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Warthin's tumor (cystadenolymphoma)





Tonsillitis chronica



xrecurrent chronic inflammation

*acute exacerbations - enlarged, red, swollen painful tonsils

xgross:

- mostly purulent exudate with necrotic epithelial cells and bacteria in crypts forming semi-firm foul-smelling debris
- ⇒pseudomembranous tonsillitis (diff. dg. x EBV)

Tonsillitis chronica



complications: local, distant

- ⇒ phlegmonous acute tonsillitis (necrosis and ulceration, penetration of bacteria into the interstitium → inflammation may progress into retrotonsillar stroma)
- abscessi (tonsillar, peritonsillar, retropharyngeal) + spread
- distant rheumatic fever, glomerulonephritis

micro:

reactive hyperplasia of lymphoid tissue, lacunae filled with neutrophils, debris and bacteria, local fibrotisation

Tonsilla palatina chronic purulent inflammation



3

- 1 squamous cell epithelium in the lacuna
- 2 detritus with neutrophils
- 3 germinal centers
- 4 ulceration

MAR C

3



mostly epithelial, less commonly mesenchymal lesions

*****sequence

normal tissue – hyperplasia – dysplasia – CIS – invasive carcinoma

risk factors

smoking, alcohol, their combination; irradiation, HPV, betel, other chronic irritation



Squamous cell carcinoma 😤



elevated/ulcerated firm lesion

Esophageal diverticula



- diverticulum an acquired outpouching of the esophageal wall involving all the layers of the wall (true diverticulum)
- ★ Zenker's diverticulum (pharyngoesophageal) most common type, located on the posterior wall in the pharyngoesophageal junction, weakening of m. constrictor pharyngis, developed by pulsion (forcible distension of the esophagus) → pulsion diverticulum
- Midthoracic diverticulum in the mid-chest; developed by traction (external forces pulling on the wall inflammation with scarring e.g.) → traction diverticulum
- Epiphrenic diverticula above the diaphragm
- Signs: dysphagia, regurgitation, foetor ex ore
- **Complications:** *putrid inflammation, ulceration, perfortion into mediastinum*

Esophageal varices



 congested and dilated submucosal veins in the distal third of the esophagus
 in portal hypertension
 porto-caval anastomoses
 complications - rupture with massive hemorrhage into the lumen, haematemesis, haemorrhagic shock

Esophageal varices - endoscopy





copy

Esophageal varices gross





Mycotic oesophagitis



x candida, aspergillus, mucor, cryptococcus

Superficial form low-level immunodeficiency, patients on broad-spectrum ATB or corticosteroids therapy, diabetics, pregnancy...

generalised form + secondary deep mycotic infections – high-grade immunodeficiency - AIDS, neoplasia (haemathologic), immunosuppression, debilitated patients

Mycotic oesophagitis



🗴 gross:

white to gray pseudomembranes with hemorrhagic bases after removal

micro:

 necrotic mucosa with mixed inflammatory infiltrate, numerous fungal organisms (pseudo/hyphae, yeasts)
 special impregnation/staining for detection of the fungi (Groccott, PAS, Giemsa)







Mycotic oesophagitis (Groccott silver impregnation)



Mycotic oesophagitis - detail



Mycotic oesophagitis - detail (PAS staining)



Reflux oesophagitis



- chemically caused inflammation in gastro-oesophageal reflux disease (sq. epithelium sensitive to acids)
- signs x pathology low correlation
 - Þ heartburn, dysphagia
- **x** gross:
 - ⇒ mucosal hyperemia, epithelial erosions, ulcerations, scarring, stenosis
- micro:
 - 3 reactive alterations of the squamous cell epithelium: basal zone hyperplasia, elongation of lamina propria papillae, inflammatory infiltrate with eosinophils.
- complication: Barrett's oesopagus (predisposition to maligancy)!

Reflux oesophagitis





Regular oesophageal epithelium

Reflux oesophagitis: basal zone hyperplasia (>20%), elongation of lamina propria papillae (into the superficial 1/3)

Barrett's oesophagus



- complication of reflux oesophagitis
- risk for the development of adenocarcinoma!
- ★ replacement of the normal squamous cell epithelium by columnar epithelium with goblet cells (= intestinal metaplasia) → risk of dysplasia
- ★ → oesophageal adenocarcinoma (so-called Barrett's carcinoma)

Barrett's oesophagus





1 regular oesophageal mucosa

2 metaplasia

3 gastro-oesophageal junction

4 cardia



Barrett's oesophagus

1 intestinal type epithelium 2 goblet cells 3 hyperemic lamina propria






Barrett's oesophagus PAS ALC staining – blue goblet cells





Barrett's oesophagus - dysplastic epithelium





- usually in the mid-third of the esophagus
- ✗ M>F, ≥45 yrs of age
- Risk factors:
 - carcinogenic substances in food (aflatoxin, nitrosamines), tobacco, alcohol, HPV, very hot beverages, chronic inflammation
- **Symptoms:**
 - dysfagia, weight loss, cachexia



× gross:

polypoid exophytic mass, ulcerative or diffuse infiltrative neoplasm

***** bad prognosis:

→ tendency to spread through submucosal lymph vessels → distant satellite tumor foci











STOMACH













***** two general types:

Content of the second secon

• gross:

hyperemic, oedematous mucosa with erosions, haemorrhage

• micro:

hyperemia, oedema, neutrophilic (foveolar) inflammatory infiltrate, erosions

Gastritis



chronic - chronic inflammatory changes leading to mucosal atrophy and epithelial metaplasia

- usually associated with Helicobacter pylori
- •micro evaluation:
 - » inflammatory infiltrate in lamina propria mucosae lymfoplasmocytic (grade of chronicity) + neutrophils (grade of activity)
 - » presence of HP (+/-) and quantitative analysis
 - *» presence of atrophy, intestinal metaplasia (complete, incomplete) and possible dysplasia*

Clinical-pathological classification of chronic gastritis

- Chronic non-atrophic gastritis (superficial) ("B")
- 2) Chronic atrophic gastritis
- I. Autoimmune gastritis ("A")
- II. Chronic multifocal atrophic gastritis
- 3) Special forms (chemical reactive, lymphocytic, eosinophilic, granulomatous)

Clinical-pathological classification of chronic gastritis

× Chronic non-atrophic gastritis (superficial)

Helicobacter pylori

- gross: antrum and body mucosa
- micro: superficial or deep inflammation, active chronic gastritis, lymphocyte and plasma cell response forming lymphatic follicles in the glandular area, final mucosal atrophy
- higher risk of developing NHL

Chronic non-atrophic gastritis - follicular



Chronic non-atrophic gastrits



Chronic active gastritis - grade of activity 2





intraepithelial neutrophils

Chronic non-atrophic gastritis Helicobacter pylori (Warthin-Starry)





Chronic non-atrophic gastritis Helicobacter pylori (Giemsa–Romanowski)



Chronic gastritis



× Chronic atrophic gastritis

1/ Autoimmune chronic atrophic gastritis ("A")

autoimmune, anti-parietal cell and anti-intrinsic factor antibodies, hypochlorhydria, association with vitamin B12 deficiency and pernicious anemia

- gross: mucosa of the gastric body and fundus atrophy
- *micro:* chronic non-active gastritis (severe mucosal atrophy with intestinal or pseudopyloric metaplasia, fibrosis)
- higher risk of developing adenocarcinoma!

Chronic gastritis



× Chronic atrophic gastritis

2/ Chronic multifocal atrophic gastritis (pangastritis)

Helicobacter pylori-associated
 low grade of inflammation (body + antrum)
 epithelial reactive changes, erosions
 uneven distribution of atrophic foci

Chronic atrophic gastritis (gastric body)





Chronic atrophic gastritis (gastric body)



Chronic gastritis - intestinal metaplasia





Chronic gastritis

Special forms:

Chemical (reflux) gastritis / reactive gastropathy (former "C")

bile reflux, duodenal reflux after partial gastrectomy, NSAIDs

micro: hyperemic, oedematous mucosa, foveolar hyperplasia, vasodilatation, little inflammatory response

* Lymphocytic, eosinophilic, granulomatous...

Reactive gastropathy (gastritis C)



Reactive gastropathy (gastritis C)





Hypertrophic gastropathy

* uncommon, large mucosal folds

- Ménétrier's disease (hyperplastic hypersecretory gastropathy with protein loss)
- hypersecretory gastropathy (hyperplasia of parietal and chief cells)
- glandular hyperplasia in Zollinger Ellison syndrome (in neuroendocrine tumors with gastrin production)

Gastric erosions



- definition:
 - Imited by m. mucosae, tiny superficial defects < 3 mm
- causes:
 - NSAIDs, alcohol, vomiting, stress, burns, infection
- Iocalisation:
 - antrum and body
- microcirculation disorder, capillary rupture
 complete regeneration within a few days

Peptic ulceration



- Ulcer definition: mucosal defect progressing through the m. mucosae into the submucosa or deeper
- risk factors/causes:
 - general: genetics, age, stress, alcohol, smoking
 - Iocal: gastric hyperacidity, HP gastritis, NSAIDs
- Iocalisation:
 - pylorus, lesser curvature, bulbus duodeni, (Meckel's diverticulum, stomic junction, GE junction)

Gastric ulceration



Acute ulcer:

- sharply demarcated defect 4 25mm; acute gastritis, severe stress (shock, trauma, burns), NSAIDs, severe hyperacidity
- Chronic ulcer:
 - slightly overhanging margins, radial adjacent mucosal folds
 - gross: smooth base
 - 4 histologic zones: 1) fibrinoid necrosis and cell debris active u. 2) mixed inflammatory infiltrate 3) granulation tissue 4) fibrous scar
 - complications: bleeding (overt, occult), penetration, perforation, scarring + obstruction, rare malignant transformation

Chronic peptic ulcer of the stomach





Perforated duodenal ulcer

391--1964



Chronic peptic ulcer of the stomach - basis

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- 1 surface cell detritus
- 2 narrow capillary lumina (compressed by proliferating tissue)
- 3 granulation tissue with inflammatory infiltrate and fibrous tissue
- 4 fibrinoid necrosis

Peptic duodenal ulcer - edges of the ulcer




Chronic peptic duodenal ulcer



Important gastric tumors



SEUDOTUMORS

non-tumorous polyps (inflammatory, hyperplastic, fundic gland polyps)

× EPITHELIAL

adenoma (in the setting of chronic gastritis/intest. metaplasia)

➡ malignant: carcinoma (adenoca, neuroendocrine ca, ...)

NON-EPITHELIAL

gastrointestinal stromal tumors (GISTs)
 lymphomas (NHL: MALT, DLBCL)

Gastric carcinoma



- most common malignant gastric tumor
- Iocation: antrum, pylorus, lesser curvature
- risk factors:
 - precursor lesions: chronic gastritis with intestinal metaplasia, infection with HP, intraepithelial neoplasia
- EBV, dietary carcinogenes (salted, smoked food), familial
 clinical features:
 - vomiting, abdominal discomfort, weight loss, anorexia
 Direct spread into adjacent organs/tissues
 Metastases: LN regional, distant (Troisier's supraclavicular LN), portal circulation (liver), peritoneal dissemination, lung, ovarian Krukenberg tumor in females.



Gastric carcinoma

Classification:

macroscopical:

 exophytic (polypous)
 excavated (ulcerated)
 infiltrative (linitis plastica)

 depth of invasion:

 early: only in mucosa and submucosa
 advanced: extended into the muscular wall

 histological type

WHO histological classification of gastric tumors



- 🗴 Tubular
- Papillary
- Mucinous
- Signet-ring cell
- Adenosquamous
- Squamous
- Undifferentiated
- × Neuroendocrine

Lauren's histological classification

× Intestinal:

- \Rightarrow 50%, HP chronic gastritis, \downarrow tendency
- intestinal metaplasia-connected, neoplastic tubular glands/papillary formations, columnar epithelium, expansive growth
- ⇒ > 50 yrs, M:F 2:1
- Diffuse:
 - ⇒ 30%, ↑ tendency
 - dissociated cells infiltrating singly / in small clusters into the stomach wall, signet-ring cells possible, reactive desmoplasia fibrosis (scirrhus)
 - Þ earlier age, M:F 1:1
- × Mixed

Gastric adenocarcinoma - exophytic growth



Gastric adenocarcinoma - intestinal type





1 normal gastric mucosa

2 tubopapillary adenocarcinoma

3 muscularis mucosae

line - sharp demarcation of the tumor from normal mucosa

Gastric adenocarcinoma (intestinal type) infiltration into lamina muscularis propria



1 tumor cells

2 smooth muscle cells

Gastric adenocarcinoma - diffuse type



Gastric adenocarcinoma - diffuse type





1 intestinal metaplasia of the mucosa

2 diffuse infiltration with signet-ring cells

Gastric adenocarcinoma - diffuse type



Gastric adenocarcinoma - diffuse type



Gastric adenocarcinoma - diffuse type detail (PAS)



Krukenberg tumor







Krukenberg tumor



Gastrointestinal stromal tumors (GISTs)



- mesenchymal tumors
- * arising from intestinal cells of Cajal (pacemaker cells controlling peristalsis)
- rigin anywhere in the GIT: predominantly the stomach and small intestine
- spindle-like or epitheloid cells, IHC CD117+,
- biologic behaviour prognosis :
 - according to mitotic rate, size, localization



Oesophageal GIST



Oesophageal GIST - detail (spindle-like cells, low malignancy)



Oesophageal GIST - detail (spindle-like cells, highly malignant)



Intestinal GIST - detail (spindle-like cells, highly malignant)



Intestinal GIST IHC CD117 positivity







INTESTINES

Normal mucosa of the small intestine



 villi to crypts height ratio 3:1 – 5:1
 standard number of intraepithelial lymphocytes: 40 IEL / 100 enterocytes

brush border – microvilli (PAS+, alkaline phosphatasis +)
 differentiated enterocytes



Normal villi of the small intestine



Malabsorption syndromes



- a group of symptoms resulting from an alteration in the digestion / absorption of nutrients mostly in the small intestine
- symptoms:
 - anorexia, diarrhea, steatorrhea, weakness, weight loss, abdominal distention,
 - growth disturbances, eczema, neurologic/psychologic disturbances, bleeding disorders, anaemia, tetany
- disturbance of:
 - digestion intraluminal, terminal in the brush border
 - mucosal absorption enterocytes abnormalities, reduced intestinal surface area
 - Iymphatic transport

Malabsorption syndromes

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- Classification
 - primary enterocytes' disorder (inborn, acquired)
 - *secondary cause apart from enterocytes*
- commonly mixed causes

Defects of mucosal absorption



*** Brush border enzymatic deficiency** (lactose intolerance – lactase deficiency)

Celiac disease gluten (gliadin)-sensitive enteropathy



- prevalence 0,5-1% in Caucasian Europeans
- associated with dermatitis herpetiformis Duhring, DM I., Sjögren sy, etc.
- immunological sensitivity to gluten (component of wheat)
- antibodies EMA, ARA, TG (non-specific antigliadin)
- genetic (HLA), immune, exogenous factors



- gluten-free diet necessary, sm. lifelong
- risk of malignant disease:
 - malignant lymphomas (T-cell), carcinomas of the small intestine
- × clinical
 - infancy (6-24 m.), adults 30-60 yrs; silent, latent
- symptoms:
 - ➡ irritability, diarrhoea, fatigue...
- endoscopy:
 - Ioss of mucosal folds, mosaic mucosal pattern, prominence of the submucosal vessels



- micro: most changes in the proximal part of the small intestine
- basic histologic features:
 - increased number of intraepithelial T-cells
 - inflammatory infiltrate (plasma cells, eosinophils, neutrophils, T-cells) in lamina propria mucosae
 - villous atrophy
 - reactive hyperplasia (elongation) of the crypts

Marsh classification 0-IV





- IEL specific activated CD8+ T-cell subpopulation
 direct cytotoxic activity – killing of
- enterocytes
- increased enterocytes' turnover
- non-specific histology diff. dg. alimentary allergies, viral infections, giardiasis, tropical sprue

Celiac disease Marsh I





Celiac disease Marsh IIIc





Celiac disease atrophic mucosa - detail

2

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1 Total villous atrophy

2 Increased number of intraepithelial lymphocytes and dedifferentiated enterocytes

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3 Goblet cell
Inflammatory bowel disease (IBD)



- idipathic, chronic relapsing inflammatory disorders of not completely known origin
 genetic predisposition, immunologic factors
 etiology:
 - ⇒ aberrant local immune response to exogenous stimulus (microbiome)→ increased transepithelial permeability → inflammation acceleration
- × Crohn disease
- **×** Ulcerative colitis
- Indeterminated colitis (10-15%)





common histologic features:

- 1) abnormal crypt architecture
- 2) crypt atrophy
- dense inflammatory infiltrate in lamina propria, basal plasmacytosis
- **4)** Paneth cell metaplasia in the left colon



Clinical features:

- recurrent attacks of diarrhea, abdominal pain, fever
- abrupt beginning, lasting days to weeks, symptom-free intervals,
- → common other AI diseases:
 - iritis, ankylosing spondylitis, erythema nodosum, PSC
- Gross:
 - ⇒ terminal ileum, or anywhere else in the GIT (oral-anus)
 - sharply demarcated , segmental lesions and skip lesions:
 - shallow \rightarrow longitudinal ulcers
 - wall stenosis and thickening, fissuring, fistulae



Histology:

- transmural inflammatory infiltrate
- formation of lymphatic follicles/germinal centres
- non-caseating granulomas (not always present) in submucosa, subserosa and regional lymph nodes
- ➡ fissuring, ulceration
- ➡ fibrosis



- **Complications:**
 - narrowed lumen, intestinal strictures, obstruction
 - malabsorption, protein loss
 - perforation, peritonitis, fistulae formation
 - ➡ hemorrhage
 - ⇒systemic AA amyloidosis
 - ⇒ carcinoma









Crohn's disease in the colon (transmural chronic inflammatory infiltrate)







Crohn's disease inflammatory infiltrate in the submucosa





Crohn's disease inflammatory infiltrate in the subserosa



Crohn's disease - granuloma in the submucosa



- 1 Multinucleated giant cell
- 2 Granuloma in the submucosa
- 3 Inflammatory infiltate in the submucosa
- 4 Muscularis propria with inflammatory infiltrate

Crohn's disease - inflammatory infiltrate



1 Lymphatic follicle in the subserosa

Ulcerative colitis Clinical features:



- relapsing attacks of bloody mucoid diarrhea, cramps, lower abdominal pain
- start rectum + sigmoid, continuous retrograde extension, may affect the entire colon (pancolitis)
- unclear etiology, autoimmune and genetic factors, variable triggers
- associated with systemic disorders (eye, skin, joint, bile tract primary sclerosing cholangitis)

×Gross:

hyperemia, oedema, flat ulcerations, regenerative hyperplastic mucosa forming pseudopolyps

Ulcerative colitis



×Micro:

- non-specific inflammatory infiltrate only in the mucosa and submucosa
- crypt abscesses, crypt destruction
- ➡ no granulomas, no skip lesions
- very little fibrosis, no mural thickening
- high risk of carcinoma development

Ulcerative colitis



Microscopic phases of the inflammation:

⇒**1. active**

- hyperemia, mixed inflammatory infiltrate, crypt abscesses
- ⇒2. healing
 - less neutrophils, no crypt abscesses, epithelial regeneration

⇒3. remission

 mucosal architectural disarray, atrophy, inflammatory changes sm. only in the rectum

Complications:

toxic megacolon, hemorrhage, perforation, peritonitis, carcinoma development

Ulcerative colitis - gross



Ulcerative colitis superficial inflammatory infiltrate





4 Mucosa ulceration

Ulcerative colitis - crypt abscess



Ulcerative colitis - dysplastic changes in the epithelium





Ulcerative colitis



Ulcerative colitis basal plasmacytosis



Ulcerative colitis basal plasmacytosis



Ulcerative colitis Paneth cell metaplasia in the left colon





Ulcerative colitis epithelial dysplasia

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2

3



- 1 Ulcer in the mucosa
- 2 Inflammatory infiltrate in the mucosa
- 3 LG epithelial dysplasia

Further types of enterocolitis

- » pseudomembranous
- ischemic
 - short-term decreased blood supply to the intestine (shock, trauma, surgery)
- microscopic (collagenous, lymphocytic)
 - chronic watery diarrhea, normal colonoscopy, associated with autoimmune diseases
- infectious
- postradiation
- × others

Pseudomembranous colitis



xetiology:

infection – bacterial (Clostridium difficile, Salmonella, Staph. aureus)

⇒antibiotic-associated

⇒uremia

×gross:

⇒greyish pseudomembranes on the mucosal surface, ulcers Micro:

fibrinous pseudomembrane with neutrophils, bacteria and macrophages, adherent to the necrotic mucosa





Endoscopy (copy)





Pseudomembranous colitis - detail





Pseudomembranous colitis (Clostridium difficile etiology)





Ileus – intestinal obstruction/disruption of the normal motility

*mechanic (strangulation, *dynamic obturation)

- ➡Adhesions
- **⇒**hernias
- **⇒**volvulus
- ⇒invagination
- **⇒**tumors
- obstruction (foreign body)
 congenital atresia
 meconial in cystic fibrosis

- paralytic: toxic-infective, drugs, peritonitis, postoperative
- vascular: hemorrhagic infarction
- myopathy, neuropathy
- Hirschprung' disease





Clinical features: signs of "acute abdomen" with acute pain, cramps, abdominal distention, nausea and vomiting, stop of the stool/gases passage
 Type / severity of the signs acccording to localisation + stage of obstruction

Intestinal wall in/above the obstruction:
 dilatation → inflammation → (peritonitis, sepsis)
 → mural necrosis → perforation → stercoral
 (fecal) peritonitis



Volvulus, bowel infarction



Gallstone ileus



Hemorrhagic infarction of the intestine





copy

result of intestinal ischemia
Hemorrhagic infarction of the intestine





Hirschprung' disease



Intestinal polyps



× Non-neoplastic polyps

hyperplastic polyp (<5 mm) minimal malignant potential, part of group of serrated lesions

⇒juvenile polyp - hamartoma; in children under 5 years, in the rectum, sporadic or part of juvenile polyposis sy (AD, haemorrhage, ↑ risk of ca)

⇒ Peutz - Jeghers hamartoma polyps + mucocutaneous hyperpigmentation; single/multiple (P-J syndrome - ↑ risk of pancreatic, pulmonary, ovarian, breast cancer)

Intestinal polyps



*Neoplastic sporadic adenomatous polyps

 ⇒ tubular adenoma (smaller, spheric, pedunculated)
 ⇒ villous adenoma (large, flat, sessile, often HG dysplasia and high malignant potential)
 ⇒ tubulovillous adenoma

Familial syndromes



1/ Familial hereditary polyposis syndromes
 ⇒ familial adenomatous polyposis (FAP):

 AD, mutation of suppresor APC gene, 100-2500 colonic adenomas, teenagers, 100% risk of cancer
 ⇒ Gardner syndrome: FAP variant, dental anomalies extraintestinal tumors: osteomas, gliomas, lipomas, fibromas
 ⇒ Peutz - Jeghers syndrome :

melanotic mucosal and cutaneous pigmentation with hamartomatous intestinal polyps



2/ Lynch syndrome

 (hereditary non-polyposis colorectal cancer, AD), DNA mismatch repair defect + increased rate of mutations; younger age, right colon.
 Increased risk of multiple tumors of the stomach, small intestine, liver, gallbladder tract, urinary tract, brain, skin, prostate.

Serrated lesions



special heterogenous group of polypous lesions, serrated (sawtooth, stellate) morphology, part of intraepithelial neoplasias

- precursors of perhaps one third of colorectal cancers
- classification: dysplastic, non-dysplastic

Classification of serrated lesions/polyps

➤ Non-dysplastic
 ⇒ hyperplastic polyp
 ⇒ sessile serrated adenoma/polyp
 ➤ With dysplasia
 ⇒ sessile serrated adenoma/polyp w. dysplasia
 ⇒ traditional serrated adenoma







Colon – hyperplastic polyp





Colon - juvenile polyp



Colon - hamartomatous P-J polyp





Adenomas





Polyposis of the colon

2 3

cm.





Tubular adenoma



- 1 Tubular adenoma with low grade dysplasia
- 2 Border between dysplastic and normal epithelium
- 3 Lamina muscularis mucosae
- 4 Normal intestinal epithelium

Tubular adenoma – low grade dysplasia



Tubular adenoma – high grade dysplasia





Villous adenoma







Villous adenoma





* high incidence in the Czech Republic and other developed countries ✗ 60 - 70 % in the rectum and sigmoid (50%) detectable by per rectum examination) Risk factors: lifestyle + diet, smoking, alcohol high intake: refined carbohydrates, fat, red meat decreased intake: unabsorbable vegetable fiber, protective micronutrients (vtamins A,C,E) *predisposing factors: genetic ⇒polyposis

ulcerative colitis



×Gross:

- exophytic, polypous
 - proximal colon long time asymptomatic
- endophytic, ulcers with heaped-up edges
 - distal colon early stenosis

中 annular

encircling lesions

⇒infiltrative

rare, linitis plastica type



×Micro:

tubular adenocarcinoma (most frequently)

⇒other adenocarcinoma types:

- cribriform comedo-type
- micropapillary
- medullary
- mucinous
- serrated
- signet ring cell

adenosquamous, spidle-like, squamous cell, undifferentiated



- **×**TNM classification and tumor progression:
 - pTis intraepithelial/intramucosal (100% 5-year survival, no metastases)
 - ⇒pT1 submucosa (90% survival)
 - pT2 into the muscularis propria (+LN metastases
 possible);
 - ⇒pT3 subserosa (+ metastases common), 35% survival in LN meta pT3N1

⇒pT4 transperitoneal/invasion into adjacent organ Distant metastases present - 8% survival.



Staging stage I: T1, T2, no meta stage II: T3, T4, no meta stage III: any T, LN meta, no distant meta (M0) stage IV: any T, any N, M1









- **1** Intestinal adenocarcinoma structures
- 2 Normal colonic epithelium
- 3 Muscularis propria





Adenocarcinoma structures
 Pericolonic fat and fibrous tissue





- 1 Perineural invasion of adenocarcinoma
- 2 Peripheral nerve
- 3 Pericolonic fat





1

Mucinous adenocarcinoma

Colorectal carcinoma - complications



*stenosis
 *obstructive ileus
 *hemorrhage (occult!, overt)
 *perforation
 *penetration
 *stercoral peritonitis



Peritoneal carcinomatosis widespread metastases in the peritoneum/ omentum





Appendix - normal





Appendix - periappendicitis





Appendicitis



- ★Causes: ?obstruction, stool stagnation → collapse of the draining veins → ischemia of the wall → bacterial proliferation → inflammation (catarrhal, phlegmonous)
- Trombosis of mesenteriolar veins \rightarrow ischemic necrosis of the appendiceal wall \rightarrow secondary bacteria invasion \rightarrow gangrenous inflammation
- **Complications:**
 - ⇒ peritonitis
 - periappendiceal abscess
 - ➡portal pyemia
 - ⇒adhesions





Phlegmonous appendicitis



Parasitic appendicitis - Oxyuriasis vermicularis (pinworm) in the lumen





Neuroendocrine tumors (NETs)



*arise from neuroendocrine cells of the gastro-enteropancreatic system (GEP-NET)

*histologic classification WHO 2010:

NET G1 (carcinoid)
NET G2
NEC G3 large cell or small cell type
compound adenoneuroendocrine carcinoma

Neuroendocrine tumors (NET)



* arise from neuroendocrine or precursor cells of the GIT mucosa

x mostly in the **ileum and appendix** (80%)

all NETs (except for a very few) are considered malignant in various grade





***classification** *depends on:*

Iocation
 type of the endocrine product

×gross:

small, round-shaped, flat nodules of yellowish colour, infiltrating the wall to different depth, superficially ulcerated or covered with normal mucosa, sometimes exophytic

GEP-NET



×micro:

trabecular, glandular structures - tubules, palisading or compound structure

- regular cells with clear cytoplasm and round or oval-shaped nucleus; slight nuclear polymorphism
- Iow mitotic activity
- chromogranin A in cytoplasm

GEP-NET



× possible production of various endocrine substances: serotonin, somatostatin, gastrin
 × serotonin active only locally in intestine → intestinal

- hypermotility with diarrhea
- **★** liver metastases → carcinoid syndrome:

cutaneous "flush" and cyanosis, nausea and vomiting, astmatic bronchoconstrictive attacks, endocardial fibrosis in right ventricle, hepatomegaly

Carcinoid of the appendix



Carcinoid of the appendix



Carcinoid of the appendix





THANK YOU FOR ATTENTION