Esophageal Cancer Cases

Quiz Questions:

Relationship between proto-oncogene and oncogene?

Examples:

Ras

HER2/neu

MYC

Tumor Suppressor Genes

FAP Prophylactic Surgery

- Total proctocolectomy with ileoal pouch
- Total colectomy with ileorectal anastomosis
- Proctocolectomy with end ileostomy

Lynch Syndrome Prophylactic Surgery

- No role for prophylactic colon surgery
- Prophylactic hysterectomy with BSO

BRCA 1/2 Prophylactic Surgery

- Bilateral mastectomy reduces lifetime risk 90%
- Surveillance is a reasonable option
 - Yearly mammogram
 - Yearly breast MRI

BRAC 1/2 Prophyactic Surgery

- Prophylactic BOS age 35-40 or after childbearing
- Reduces risk of ovarian cancer 80%
- Surveillance not as effective
 - Transvaginal ultrasound
 - CA-125 screening

MEN 2A/2B or FMTC Prophylactic surgery

- Timing of thyroidectomy depends upon risk category
- Highest risk: Throidectomy within first year of life
- High risk: Thyroidectomy by age 5 or if calcinonin elevated
- Moderate risk: surveillance starting age 5
 - physical exam
 - neck ultrasound
 - serum calcitonin

FAP

Median age of dx 39

- Duodenal and ampullary tumors
- Gastric polyps
- Thyroid tumors
- Desmoid tumors

FAP screening

Colonoscopy age 10-12 EGD for duodenal polyps at age 20-30 CT 1-3 years after colectomy and q5 yers in those with family hx of desmoids

Lynch

Amdterdam Criteria

(Bethesda Criteria)

Mean age dx colon cancer 44-61 Predominant right side colon cancer Lifetime penetrance 82%

Lynch Other Cancers

- Endometrial
- Stomach
- Ovarian
- Urinary tract
- Biliary Tract
- Small bowel
- CNS

Lynch Screening

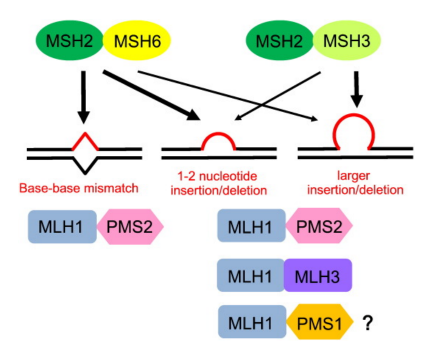
Colonoscopy q1-2 years staring age 20-25

Women with Lymch have 25-60% lifetime risk of endometrial cancer 45-12% lifetime risk of ovarian cancer Male: 1.2% risk of breast cancer (0.1% in general populations)

MMR and MSI

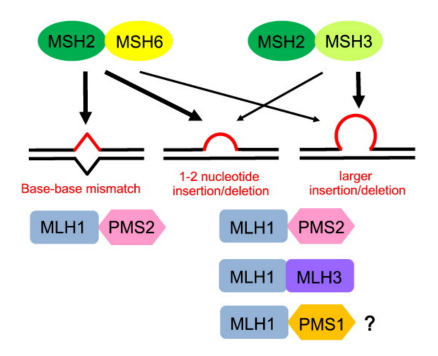
Greater than 90% of LS tumors are MSI-high (MSI-H) and/or lack expression of at least one of the MMR proteins by IHC. Ten percent to 15% of sporadic colon cancers exhibit abnormal IHC and are MSI-H most often due to abnormal methylation of the MLH1 gene promoter, rather than due to LS. Mutant BRAF V600E is found in many sporadic MSI-H CRCs and is rarely found in LS-related CRCs. There are some tumors that will have MLH1 methylation but lack a BRAF PV.

DNA Mismatch Repair Proteins

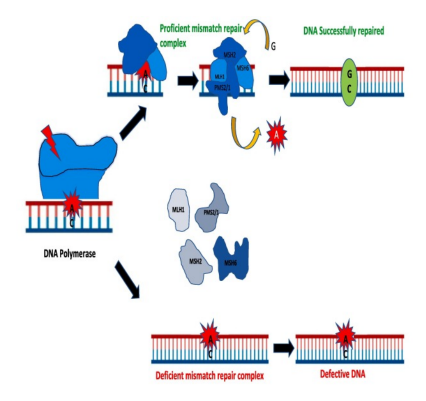


DNA Mismatch Repair Proteins

MLH1 PMS1 MSH6 MSH2 MSH5



DNA Mismatch Repair Proteins



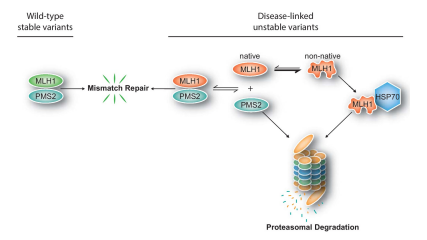
Mismatch Repair Proteins in Lynch Syndrome

Lynch Syndrome can be caused by loss of expression of:

- MLH1
- PMS1
- MSH6
- MSH2
- MSH5

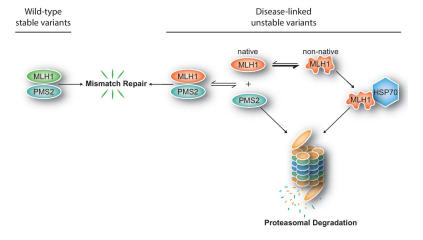
MLH1 and PMS Dimer in Mismatch Repair

wild-type MLH1 and PMS2 form a stable heterodimer



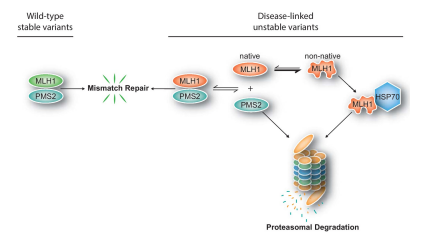
MLH1 and PMS Dimer in Mismatch Repair

Mutant MLH1 fails to form a stable heterodimer



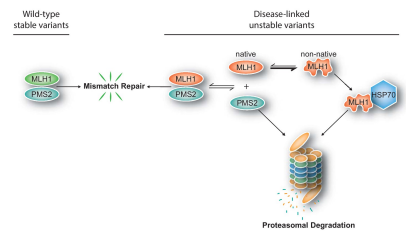
MLH1 and PMS Dimer in Mismatch Repair

Mutant MLH1 fails to form a stable heterodimer \rightarrow PMS2 is degraded



MLH1 and PMS Dimer in Mismatch Repair

If MLH1 is mutated \rightarrow PMS2 protein is *not* detected



Colon Polyposis: >10 adenomatous polyps

- Classical FAP
- Attenuated FAP (AFAP)
- MUTYH-associated polyposis (MAP)
- \bullet Colonic adenomatous polyposis of unknown etiology (CPUE)

Colon Polyposis: >4 hamartomatous polyps

- Puetz-Jaghers
- Juvenile Polyposis Syndrome
- Cowden Syndrome/PTEN Hamartoma Tumor Syndrome

Serrated Colon Polyps

5 serrated polyps/lesions proximal to the rectum, all being 5 mm in size, with 2 being 10 mm in size OR > 20 serrated polyps/lesions of any size distributed throughout the large bowel, with 5 being proximal to the rectum

Muir-Torre Syndrome

Muir-Torre syndrome refers to individuals with LS who have LS-associated skin findings of sebaceous adenomas/carcinomas or keratoacanthomas.

BRCA 1/2

BRCA1: 87% ifetime risk of breast cancer and 40-60% risk of ovarian vanver

BRCA2: 80% ifetime breast cancer 1nd 15-27% ovarian cancer - Elevated risk of prostate, pancreas, melanoma

Male carriers of BRCA2: 8.9% lifetime risk of breast cancer

BRCA risk factors:

- Breast cancer dx before age 40
- Bilateral breast vcancer
- Breast and ovarian cancer
- 2 family members with breast cancer < age 50
- Family history of breast and ovian
- Family history of male breast cancer

Attenuated FAP

 ~ 30 polyps 70% penetrance by age 80 - mean age at dx 50-55

Li-Fraumeni

Mutation of TP53 tumor suppressor gene

- Breast cancer 90% by age 50
- Sarcoma
- Leukemia
- Brain tumors
- Andrenocortical carcinoma

Breast cancer in Li-Fraumeni

Mastectomy favored to avoid radiation therapy Bilateral prophylactic mastectomy recommended

p16 = CDKNA2A mutation

- Increased risk of melanoma
 - Familial Atypical Multiple Mole Melanoma (FAMMM)
 - Familiarl llAtyopical Multiple Mole-Pancreatic Carcinoma (FAMMMPC)
- Melanoma penetrance 58-92% by 80
- Pancreatic cancer penetracne 17% by age 75

FAMMM

- Malignant melanoma in one or more first degree or seconddegree relatives
- High total body nevus count
- Nevi with certain features on microsopcy

Genetic testing not performed as only 50% of FAMMM harbor a mutation in CDKNA2A

Neurofibromatosis 1

Mutation in NF1 tumor suppressor gene

- Multiple neurofibromas
- Cafe au lait spots
- Lisch nodules (hamaartoma of the iris)

Risk of - NPNST - PHeochromocytoma - Astrocytoma - Leukemai

NF1 diagnosis

Two or more of the following 6 criteria:

- Six or more café-au-lait macules
- Two or more neurofibromas or one plexiform neurofibroma
- Axillary or inguinal freckling
- Optic glioma
- Two or more Lisch nodules
- Characteristic osseous lesions
- A first degree relative with NF1

Neurofibromatosis 2

NF2 gene

- Multiple neurofibromas
- Cafe au lait spots
- Bilateral vestibular shwannomas
- CNS tumors

Most affected develop bilateral schwannomas by age 30 with average age of death 26

Annual surveillance MRI stating age 10-12 and hearing evaluationm

PTEN

Cowden Syndrome Mutation in PTEN tumor suppressor gene

- Mucocutaneous facial lesions
- Macrocephaly
- Bilateral breast cancer
- Thyroid and endometrial tumors
- Hamartomatous polyposis of the GI tract

MEN1

- Mutation of MENIN tumor supressor
- Parathyroid
- Pitutiary
- Pancreatic islet cells

Hyperparathyoidism usually first presentaiotn Most common pancreatic tumor is non-functional

Dx by 2/3 of following:

- Parathyroid adenoma/hyperplasia
- Pancratic islet cell tumors
- Pituatiary tumors

MEN1 screening

Surveilance with serum prolactin, IGF-1, factin glucose and insuslin starting age 5 Calcium, chormogranin A, pancrea polypeptide glucaocon AP age 8 Serum gastrin starting age 20 Brain MRI starting age 5 Abdomeal CT/MRI stagting age20

MEN1 surgical treatment

Parathyroidectomy 3.5 gland or 4 gladn with autotransplantation

Pancreatic tumor resection if >2cm

Pituitary tumors resected via transsphenoidal

MEN2 Famly of Syndromes

- RET proto-oncogene
- Medullary thyroid cancer in almost 100%
- MEN2A
 - pheochromocytoma in 50%
 - Parathyroid hyperplasia in 20-30%

-MEN2B - pheochromocytoma in 50% - Megacolon - Marfinoid habitus - Ganglioneromas - Mucosal neuromas

MEN 2 Prophylactic surgery

Prophylactic total thyroidectomy

Testing for pheochromoyctoma prior with adrenal ectomy prior

Monitor calcitonin and CEA after thryoidectomy

Orientation Handbook



References