The Problem
Lynch syndrome is under-recognized

The Solution
Collaboration between Surgeon, Pathologist and Genetic Counselor

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Learning Objectives

- Recognize that Lynch Syndrome is an important clinical problem
- Understand why Lynch Syndrome is under-recognized
- Discuss proposal for molecular analysis of all CRCs
Hereditary Colon Cancer Syndromes

- **Sporadic** (65%–85%)
- **Familial** (10%–30%)
- Rare CRC Syndromes:
  - MYH Polyposis
  - PJS, Cowden’s, JPC
- Lynch Syndrome (2-3%) (HNPCC)
- Familial Adenomatous Polyposis (<1%)

Adapted from Burt RW et al. *Prevention and Early Detection of CRC*, 1996
Warthin Syndrome?

- Aldred Scott Warthin MD PhD
- University of Michigan, Dept of Pathology 1895-1931

A little field, well tilled

Sir William Osler

- Seamstress’ story
- Family G
Lynch Syndrome

- Updated Family G
- Described other families
- Popularized recognition of Hereditary Colon Cancer Syndromes

Henry T. Lynch, M.D.
Professor of Medicine and Prev Medicine
Creighton University School of Medicine
Lynch- Family G, Branch I

I

Progenitor
Site Unknown
d. 60 y

II

Uterus, NOS
d. 55 y

III

Endometrium 55 y
Cecum 57 y
d. 75 y

IV

1
Colon, NOS 43 y
Colon, NOS 67 y
d. 85 y

2
Ascending Colon 70 y

3
Cecum 56 y
d. 82 y

V

1
Endometrium 49 y
d. 62 y

2
Cecum 58 y

3
Cecum 28 y
Rectum 45 y

- Affected Individual
- Female
- Male
- Deceased Individual
- Individual No.
- Cancer Site and Age at Diagnosis
- Age at Death
What’s in a name?

Cancer Fraternities
Cancer Family Syndrome
Hereditary Site Specific Colorectal Cancer
Lynch Syndrome
  Lynch Syndrome I- colon only families
  Lynch Syndrome II- colon, uterus, others
Hereditary Nonpolyposis Colorectal Cancer (HNPCC)
Muir-Torre Syndrome (with skin manifestations)
Turcot Syndrome (with brain tumors)
Genetic Features of Lynch Syndrome

- Familial Clustering of CRC
- Autosomal Dominant
  - Vertical Transmission
  - 50% risk in siblings/children
- Due to germline mutation in one of 4 DNA mismatch repair genes

Genes involved:
- Chr 2: MSH2, MSH6
- Chr 3: MLH1
- Chr 7: PMS2
Lynch Syndrome Results From Failure of Mismatch Repair (MMR) Genes

- Base pair mismatch
- Normal DNA repair
- Defective DNA repair (MMR+)

Molecular Diagnosis of Lynch CRCs
Loss of MMR proteins by IHC
or MSI by PCR
IHC for MMR Proteins
MMR System Repairs Replication Looping

Normal Tissue

Tumor Tissue

Addition of nucleotide repeats

Southern Blot

---CACACACA---

Shortened
Clinical Features of Lynch Syndrome

- High CRC risk - up to 80%
- Few adenomas with high cancer propensity
- Early onset - 44 yrs
- Proximal location - 65%
- Synchronous/Metachronous CRC - 20%
- Other cancers
Incidence of CRC in Lynch Syndrome vs Sporadic

Voskuil, Int. J. Cancer, 1997
Cancer Risks in Lynch Syndrome

Aarnio M et al. *Int J Cancer* 64:430, 1995

- Colorectal: 78%
- Endometrial: 43%
- Stomach: 19%
- Biliary tract: 18%
- Urinary tract: 10%
- Ovarian: 9%

% with cancer vs. Age (years)
Clinical Management of Lynch Syndrome
Gene Carriers or At Risk Family Members

- Colonoscopy, age 25 years, or 10 years younger than earliest diagnosis, repeat annually
- Colectomy for CRC or high risk adenoma
  - Type? Subtotal colectomy with IRA vs segmental resection
  - IHC of biopsy useful in pts < 50 yrs to tailor initial resection
- Other tumor screening
  - Endometrial, ovarian, gastric, small bowel, urinary tract
  - Prophylactic hysterectomy/oophorectomy
CASE STUDY: 40 YEAR OLD FEMALE PRESENTS FOR COLONOSCOPY

Colon cancer
Tubulovillous adenoma w/HGD
Melanoma

IHC – absence of MSH6
Germline testing – deleterious MSH6 mutation
Evolution of identification of Lynch syndrome

1. Amsterdam criteria
2. Bethesda criteria
3. Universal IHC screening on newly diagnosed CRC
Pathologist

- MSI-H histology

Primary Care/Surgeons

- Family History

Genetics

- MSI-H + family history = discussion of possible Lynch syndrome
Lynch syndrome estimates

- Estimated population incidence of Lynch syndrome is 1 in 370.
  (Based on the 2.8% incidence of Lynch syndrome in newly diagnosed colon cancers and the penetrance of Lynch syndrome being about 50%)
- That equals 829,800 out of 307,006,550 Americans
- Estimate that no more than 1.2% (10,000/829,800) of individuals with Lynch syndrome are aware of their diagnosis

Cancer Prevention Research, 4(1):1-5; 2011
Identification of Lynch Syndrome Makes a Difference

- Significant data indicate that an early diagnosis of Lynch syndrome followed by intense cancer surveillance and/or prophylactic surgery can prevent morbidity and mortality from LS cancers
  - 62% reduction in colon cancer risk in individuals with LS undergoing surveillance
  - 30% of women with LS who did not have risk reducing surgery had developed endometrial cancer and 5.5% developed ovarian cancer over a 10 year period

Genetics in Med 35-41, 2009
Amsterdam criteria
Amsterdam Criteria

- 3 first-degree relatives with CRC
- 2 or more generations
- 1 CRC by age 50
Amsterdam Criteria II

Other cancers, may be substituted for colon cancer in making the diagnosis

- endometrial cancer
- ovarian cancer
- gastric cancer
- hepatobiliary
- small bowel
- transitional cell ca of renal pelvis or ureter
Limitations of Amsterdam criteria

Amsterdam criteria

- requires careful assessment of family history
- Gathering:
  - 3 generation family history
  - Age of cancer diagnosis
  - Individuals tend to know the first degree history well but 2\textsuperscript{nd} and 3\textsuperscript{rd} degree less
- Family history inconsistently recorded by clinicians
- Inherited pattern may not be evident due to different types of cancer
Limitations of Amsterdam criteria

- <50% of Lynch syndrome families meet Amsterdam clinical diagnostic criteria

AND

- 50% of families meeting Amsterdam don’t have Lynch syndrome. They likely represent Familial Colorectal Cancer Syndrome X involving only risk for colon cancer
Bethesda criteria
Revised Bethesda criteria

- Perform MSI and/or IHC
  - CRC diagnosed in patient under age 50, OR
  - Synchronous/metachronous LS tumors, regardless of age, OR
  - CRC with MSI-H histology diagnosed in patient under age 60, OR
  - CRC in patient with \( \geq 1 \) first-degree relative with LS cancer, one cancer diagnosed \(<\) age 50, OR
  - CRC in patient with \( \geq 2 \) first- or second-degree relatives with LS cancer, regardless of age
MSI-H Histology

- Tumor infiltrating lymphocytes, OR
- Crohn’s-like lymphocytic reaction, OR
- Mucinous/signet-ring differentiation, OR
- Medullary growth pattern
Microsatellite Instability

- 12-20% of all CRC exhibit MSI and abnormal IHC\(^1\)-\(^5\)
  - 75% of these do not have LS but rather acquired hypermethylation of the \(MLH1\) promoter which silences gene expression \(^6\)-\(^7\)
  - Associated with somatic mutation V600E in the BRAF gene
    - Correlates with better prognosis (versus those with MSS CRC)
    - May benefit less from 5-FU-based chemotherapy
- Almost all LS-related CRCs exhibit MSI and/or abnormal IHC
- MSI and IHC analysis show >94% concordance\(^8\)

Genetic Testing for Lynch Syndrome
Using IHC on Tumor Tissue

Tumor IHC Testing
- MSH2/6/PMS2 absent
- MLH1 absent

B-RAF Tumor Testing
- Wild Type
- Mutant

Genetic Testing For Lynch Syndrome

All present
- No Further Testing MSS Sporadic

MSI Sporadic
But...

- 25% of patients with LS did NOT meet Bethesda guidelines
- 56% of patients with LS were diagnosed ≥50
- 35% of patients with LS did not have the characteristic histologic features

1 J Clin Oncol 2008;26:5783-5788.
Universal IHC/MSI testing
“Recently, IHC and/or MSI screening of all colorectal cancer and endometrial cancer regardless of age at diagnosis or family history, has been implemented at some centers to identify individuals at risk for Lynch syndrome.”

Currently done at:
- University of Colorado Hospital - Mandated to start at VA nationwide
- Ohio State within 18 months
- Huntsman Cancer Institute
- Cleveland Clinic
- Etc...

EVALUATION OF GENOMIC APPLICATIONS IN PRACTICE AND PREVENTION (EGAPP) Working Group Recommendations

- Offer screening for Lynch syndrome to all individuals at time of CRC diagnosis
- Focus on improved outcome in unaffected relatives who can then be tested
- MSI sensitivity 89%, IHC sensitivity 83%

Support for EGAPP Recommendations

- Universal testing:
  - Detects nearly twice as many cases of Lynch syndrome vs. targeting those diagnosed < age 50
  - Has cost-effectiveness ratio comparable with other preventive services (less than $25,000 per LY saved – same as colonoscopy every 10 years in adults ≥50 yrs)

- Findings support EGAPP recommendations

- Testing strategies using IHC have the most favorable cost-effectiveness ratios

Response to Screening

- MSI/IHC considered a screening test
- Screening well received by both patients/family members and medical professionals
- 90% of individuals screened through a research study proceeded with genetic counseling but a substantial decrease in follow-up was seen in a clinical setting
- Individuals may or may not choose to proceed with genetic testing following discussion

Pitfalls

- Exclusions
  - Any tumor exposed to XRT should not have MSI since there is contraction and expansion of MSI due to XRT
- Loss to follow-up
- Identifying low risk patients?
Informed Consent

- There is a general consensus that informed consent is not required for MSI/IHC analysis of biopsied or resected tumor samples.
  - Patient already has diagnosis of CRC
  - MSI or IHC not definitive for genetic diagnosis
  - Directed by patient care decisions
  - Have option to get genetic testing

- Some institutions include a fact sheet with pre-surgery packet
Solution:

Universal IHC/MSI testing on all CRC

Pathologist
- MSI-H histology
- MSI-H + family history = discussion of possible Lynch syndrome

Genetics
Notified of molecular results
Contacts provider for patients likely to have LS

Primary Care/Surgeons
- Family History

Review/Order IHC on CRC biopsy results
Anyone < 50