The New World of Genomics

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Learning Objectives

- To define genomic medicine
- To review early triumphs, and potential limitations, of genomic medicine
  - Guiding treatment
  - Risk assessment
  - Early detection/prevention
  - Diagnostics
- Applied case example: Genomic medicine approaches to hereditary breast-ovarian cancer
Genomic Medicine aka Personalized Medicine aka Precision Medicine

- An approach to customize medical care to an individual’s unique genetic makeup
- Every cell in the body contains DNA, molecules inherited from our parents that determine how the body looks and functions
- Sections of the DNA that contain information needed to make proteins are known as genes
- Variations in the DNA are associated with rare and common disease
  - Pathogenic and protective variants
- Many, if not most, human diseases have a genetic component
Precision Medicine is most simply:

- Human genetics and genomics information integrated into clinical medicine.

Precision Medicine more conceptually:

- Is a paradigm shift in our concept of human disease
- Instead of viewing disease based only on clinical information we define it based on its genetics/genomics
- We are literally rewriting the textbooks of medicine

From the top:

"Tonight, I'm launching a new Precision Medicine Initiative to bring us closer to curing diseases like cancer and diabetes — and to give all of us access to the personalized information we need to keep ourselves and our families healthier."

— President Barack Obama, State of the Union Address, January 20, 2015
Announcement of the NIH Precision Medicine Initiative

“Doctors have always recognized that every patient is unique, and doctors have always tried to tailor their treatments as best they can to individuals. You can match a blood transfusion to a blood type — that was an important discovery. What if matching a cancer cure to a genetic code was just as easy, just as standard? What if figuring out the right dose of medicine was as simple as taking our temperature?”

President Obama, January 30, 2013

“...because something called precision medicine... gives us one of the greatest opportunities for new medical breakthroughs that we have ever seen!”

Image courtesy of NIH
Francis Collins, Director of the National Institutes of Health, has been a key architect of the PMI

The Precision Medicine Initiative Cohort Program – Building a Research Foundation for 21st Century Medicine

Precision Medicine Initiative (PMI) Working Group Report to the Advisory Committee to the Director, NIH

September 17, 2015
<table>
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<th>Precision Medicine Initiative Details</th>
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<tr>
<td>• Large scale prospective cohort study</td>
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<td>• One million or more American</td>
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<tr>
<td>volunteers</td>
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<tr>
<td>• Broad diversity, all life stages</td>
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<td>• Begin enrollment in 2016</td>
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<tr>
<td>• Study genomic, environmental and gene-</td>
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<td>environment interactions</td>
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<td>• More precise preventive care</td>
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<td>• Better risk estimates and biomarkers for range of rare and common diseases</td>
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<td>• Fuel new targeted therapies e.g. cancer, heart disease</td>
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<tr>
<td>• Health data from EHRs and surveys</td>
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<td>• Physical exam</td>
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<td>• Blood sample</td>
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<td>• Whole genome sequencing</td>
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<td>• Home and mobile health (mHealth) sensors to correlate body measurements, calorie consumption and environmental exposures with health outcomes</td>
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<td>• Participants will control how their data is shared and used in research</td>
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<td>• Empower participants to improve own health</td>
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Precision Medicine Initiative – Integration of Genetic/Genomic Counseling

- Genetic counseling has integral role
  - Informed consent and specimen collection
  - Complexity of return of genomic based results
  - Decision making process
  - Improving health outcomes
- Service delivery models need to evolve

The UK announced the 100,000 Genomes Project in 2013

The 100,000 Genomes Project

The project will sequence 100,000 genomes from around 70,000 people. Participants are NHS patients with a rare disease, plus their families, and patients with cancer.

The aim is to create a new genomic medicine service for the NHS - transforming the way people are cared for. Patients may be offered a diagnosis where there wasn’t one before, in time, there is the potential of new and more effective treatments.

The project will also enable new medical research. Combining genomic sequence data with medical records is a ground-breaking resource. Researchers will study how best to use genomics in healthcare and how best to interpret the data to help patients. The causes, diagnosis and treatment of disease will also be investigated. We also aim to kick-start a UK genomics industry. This is currently the largest national sequencing project of its kind in the world.

Understanding genomics

Our Head of Engagement, Vivienne Parry, explains more about genomics in this film courtesy of our partners at Health Education England.

Take a look at our infographics to find out more about genomics.
The Explosion of Human Genetics and Genomics in Clinical Practice

Driven by high throughput sequencing
- 10 years ago – challenging to routinely sequence anyone
- 5 years – 2-3 genes possible (HCM 5 – 8 genes)
- 3 years – gene panels emerging (5-20 genes)
- 2 years – larger panels (10-40; pan-cardio – 84)
- 1 year ago – clinical exome emerges
  - exome = 19,000 – 20,000 genes encoding proteins
- Now – selective clinical exome sequencing:
  - 3 or more affecteds
  - negative panel
  - insurance coverage

Sequencing a genome
- Genome refers to the entire DNA structure
  - 6 billion letters of code distributed unequally amongst 46 chromosomes
  - 3 gigabytes of data
  - ~20,000 protein coding genes (2% of the DNA code)

- Cost has dropped considerably
  - 2001: $100 million per genome
  - 2010: $30,000
  - 2015: ~$1,000
Ending the diagnostic odyssey, with and without treatment ramifications
Doctors Sift Through Patients’ Genomes To Solve Medical Mysteries
By Rob Stein
http://www.npr.org/sections/health-shots/2012/09/25/160957147/doctors-sift-through-patients-genomes-to-solve-medical-mysteries

North County Twins Cured After Whole Genome Sequencing By Chris Chan
The use of WGS in the NICU provided differential diagnoses in a 50 hour time period.
WGS can shorten the time to diagnosis and quicken the move toward targeted treatment and genetic and prognostic counseling.

Sci Transl Med 2012
RESEARCH ARTICLE

GENOMICS

Noninvasive Whole-Genome Sequencing of a Human Fetus

Jacob O. Kitzman, Matthew W. Snyder, Mario Ventura, Alexandra P. Lewis, Ruolan Qiu, LaVone E. Simmons, Hilary S. Gammill, Craig E. Rubens, Donna A. Santillan, Jeffrey C. Murray, Holly K. Tabor, Michael J. Bamshad, Evan E. Eichler, Jay Shendure

Analysis of cell-free fetal DNA in maternal plasma holds promise for the development of noninvasive prenatal genetic diagnostics. Previous studies have been restricted to detection of fetal trisomies, to specific paternally inherited mutations, or to genotyping common polymorphisms using material obtained invasively, for example, through chorionic villus sampling. Here, we combine genome sequencing of two parents, genome-wide maternal haplotyping, and deep sequencing of maternal plasma DNA to noninvasively determine the genome sequence of a human fetus at 19.5 weeks of gestation. Inheritance was predicted at 2.8 × 10^5 parental heterozygous sites with 98.1% accuracy. Furthermore, 39 of 44 de novo point mutations in the fetal genome were detected, albeit with limited specificity. Subsampling these data and analyzing a second family trio by the same approach indicate that parental haplotype blocks of ~300 kilo-base pairs combined with shallow sequencing of maternal plasma DNA is sufficient to substantially determine the inherited complement of a fetal genome. However, ultra-deep sequencing of maternal plasma DNA is necessary for the practical detection of de novo mutations genome-wide. Although technical and analytical challenges remain, we anticipate that noninvasive analysis of inherited variation and de novo mutations in fetal genomes will facilitate prenatal diagnosis of both recessive and dominant Mendelian disorders.

Sci Transl Med 2012

MEDICINE

Whole-Genome Sequencing: The New Standard of Care?

Liam R. Brunham and Michael R. Hayden

Whole-genome sequencing may dramatically alter medicine, but there are obstacles to broad implementation.

Science 1 June 2012: 336 (6085), 1112-1113.

Science June 2012
Having the Genetic Code is not enough

- >99% of the DNA sequence identical
  - Several million genetic “variants” per person
  - Some variants (aka mutations) increase susceptibility to a specific disease
  - Some variants provide protection
  - Most variants remain beyond our understanding

- Large scale genomic studies have failed to identify all variants for common disease
  - 30+ variants (Type 2 diabetes; RR 1.2-1.4) explain 10% of disease heritability
  - Heart disease, schizophrenia, HTN

The current state of genomic risk assessment for CHD

The current state of genomic risk assessment for CHD

EGAPP Recommendations:
Genomics profiling to assess CV risk

- Found insufficient evidence to recommend testing for the 9p21 genetic variant or 57 other variants in 29 genes to assess risk for cardiovascular disease (CVD) in the general population, specifically for heart disease and stroke

- Found that the magnitude of net health benefit from use of any of these tests alone or in combination is negligible

- Discourages clinical use unless further evidence supports improved clinical outcomes

Genetics in Medicine 2010
The New World of Genomics

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Genomic Medicine – Early Triumphs

• **Diagnosis**: for some types of heart disease, detection of a gene mutation can lead to diagnosis and treatment to prevent sudden cardiac death before any symptoms occur

• **Guiding treatment**: instead of classifying cancers by the tissue where first detected, now beginning to categorize by genomic characteristics
  • select individualized treatments based on different somatic tumor signatures

• **Early detection/prevention**: individuals with *germline* BRCA mutations at higher risk of developing breast (and other) cancers
Early Triumphs - Diagnosis

- >6,000 inherited (Mendelian) diseases
- Collectively afflict more than 25 million Americans
- Large scale sequencing, ever decreasing cost, increasing data accumulation
  - As of Feb 2015, 2,937 genes underlying 4,163 Mendelian phenotypes discovered

Early Triumphs - Guiding Treatment

- Match advanced tumor alterations with specific targeted therapies, clinical trials
  - FoundationOne®: 315 somatic tumor pathway genes plus 28 common gene rearrangements
  - Doesn’t predict response to chemotherapy or recurrence (e.g. Oncotype)
- OSU CCC prospective single-arm trial*
  - 37 breast tumors
    - 192 tumor alterations, median of five per patient
    - 97% matched with at least one FDA-approved treatment or clinical trial
    - 65% of patients (n=24) with an FDA-approved breast cancer therapy

*2014 San Antonio Breast Cancer Symposium
Early Triumphs – Detection and Prevention

Familial 30%
Sporadic 60%
Hereditary 7-10%

Total Sporadic and Hereditary Cancers

Hereditary Breast and Ovarian Cancers

BRCA1 52%
BRCA2 32%
Other Genes 16%

Highly Penetrant Gene Variants

- Hereditary breast and ovarian cancer (HBOC)
- Caused by BRCA1 or BRCA2 germline mutation
  - Repairs double stranded DNA breaks
- Incidence
  - 1 in 500 women, in the general population
  - 2% of all individuals of Ashkenazi Jewish ancestry
  - 25% of all Ashkenazi Jewish women with ovarian cancer
Features That Indicate Increased Likelihood of *BRCA* Mutation

- Multiple cases of early onset breast cancer
- Ovarian cancer (with family history of breast or ovarian cancer)
- Breast and ovarian cancer in the same woman
- Bilateral breast cancer
- Ashkenazi Jewish heritage
- Male breast cancer along with female breast cancer in a family

HBOC-associated cancer risks

- Increased lifetime risk of multiple tumor types
  - Breast cancer (females): 50-85%
  - Contralateral breast cancer risk (female) depends on age of 1st diagnosis
  - Breast cancer (males): 6-16%
  - Ovarian cancer: 20-40%
  - Other tumors: pancreatic, melanoma, prostate
### Pathological Features of HBOC Cancers

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<th><strong>Breast cancer</strong></th>
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<tr>
<td>• <em>BRCA1</em>: ~80% are ER/PR/Her2/neu (triple) receptor negative</td>
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<td>• <em>BRCA2</em>: more likely to be ER/PR (+)</td>
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<td>• Prognosis appears to be the same as for sporadic breast cancer</td>
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<td>• Predominantly papillary serous</td>
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<td>• Can be mucinous but not as often</td>
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<tr>
<td>• Not typically associated with tumors of low malignant potential or borderline tumors</td>
</tr>
<tr>
<td>• Prognosis may be better than for sporadic ovarian cancer</td>
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### Sometimes used in treatment decisions: genomic based PARP inhibitors

| • PARPs work at molecular level to repair single strand DNA breaks |
| • Inhibition of PARP leads to the accumulation of defects, which leads to double-stranded breaks |
| • *BRCA*-associated tumors lack effective homologous-recombination DNA repair |
| • Results in selectively-induced cytotoxicity in tumor cells while sparing normal cells in patients with *BRCA*-associated tumors |
| • Dec 2014: FDA approved Lynparza (olaparib) for women with advanced ovarian cancer associated with defective *BRCA* genes |
Applications of genomic tumor analysis

• Ruth: 42 year old female recently diagnosed with metastatic breast cancer
• Age at diagnosis alone = NCCN referral!
• Some of the alterations in Ruth’s breast tumor:
  • ERBB2 amplification
    • Trastuzumab is effective in a subset of ERBB2 amplified breast cancers
  • BRCA2 617delAT (present in >80% of tumor cells)
    • May also identify true germline (hereditary) mutations

Ruth’s pedigree

Key
- Breast CA
- Ovarian CA
Misconceptions Regarding Family History

• Cancer on the father’s side doesn’t count
  – Half of all women with HBOC inherited it from their father

• Ovarian cancer not a factor in breast cancer risk
  – Ovarian cancer is an important indicator of hereditary risk, although it is not always present

• The most important thing in the family history is the number of women with breast cancer
  – Age of onset of breast cancer is more important than the number of women with the disease

Clinical Testing – Cancer Panels
Impact of results: medical management

Ruth tests positive for germline *BRCA2* 617delAT
- Recommend oophorectomy (w/ fallopian tubes)
- Eligible for specific clinical trials e.g. PARP Inhibitors

Ruth’s daughters (both *BRCA2* mutation negative)
- General population risk, follow ACS guidelines
- Cannot pass familial *BRCA2* mutation to children

- Ruth’s sister (*BRCA2* mutation positive)
  - Consider increased breast cancer screening +/- chemoprevention OR mastectomy and ovarian cancer screening OR oophorectomy (after child-bearing,<40)
Impact of results: medical management

• Ruth’s other sister (mutation status unknown)
  – Recommend screening as if mutation positive, until proven otherwise through testing
  – Same for other *at-risk* females in family

• Ruth’s father (obligate carrier)
  – Annual clinical exam; increased awareness
  – Annual prostate cancer screening
  – Follow ACS guidelines

All Disease is Complex Disease

• Predisposed to certain diseases because of our genes
• Not only genes that determine our health
• Microbiome: ecosystem of microorganisms that live on and in the human body
  – Degree influenced by environmental factors
  – Some degree of genetic influence of the host
• Lifestyle, habits, environment may cause some genes to be switched on and off, or even altered
  • Epigenetics: stable and heritable changes in gene expression NOT caused by changes in the DNA code