Reading: Schneider, 2nd Edn. Chapter12

This 2-hour class on genetics and public health will use a case-based approach to help you learn about contemporary issues at the intersection of public health and genetics. Our overall goal is to encourage you to recognize the genetic aspect of public health problems. Just as learning about infectious organisms two centuries ago altered public health practices, from sanitation to immunizations, new knowledge and technologies in genetics are altering and will continue to impact public health practices. Genetic information influences health and disease across the life span, from preconceptional genetic counseling and fortification of flours with folate to improvements in our understanding of causes of death and disability, from newborn hearing screening where over half of congenital hearing loss is genetic to recognition of the familial risk factors inherent in, for example, Alzheimer disease, cancer, coronary artery disease and stroke.
An ongoing challenge for public health personnel is to incorporate current understanding of the science of health and disease in effective and ethical public health measures. Your own understanding of the relevance of the genetic components of your family health history to your own health and your willingness to think about these complex issues for society and public health are both part of your legacy. Think genetically.
Genetic Material (DNA), Packaged as Chromosomes, Encodes Proteins and Cellular Materials that Influence How Cells Grow and Develop.

The National Coalition for Health Professional Education in Genetics (www.nchpeg.org), a coalition of more than 120 health professional organizations, and the CDC (http://www.CDC.gov/genomics/training/competencies/default.html) have developed a set of competencies in genetics for health professionals and for the public health workforce. Review these competencies and continue to reflect upon them as you go through your MPH program.

The following have been identified as public health functions relevant to genetics:

- public health assessment
- evaluation of genetic testing
- development, implementation,
- evaluation of population interventions; and
- communication and information dissemination.

Critical issues include:

- partnerships and coordination
- ethical, legal and social issues; and
- education and training.


The following web sites may be useful for your further study.

- Genetic Alliance
- GeneReviews,
- Information for genetic professionals and on genetic conditions
- www.marchofdimes.com
- National Human Genome Research Institute
Birth defects:  Case 1 - Your sister has just found out at 16 weeks of pregnancy that she has a fetus with spina bifida. Describe the levels of the maternal child health pyramid that impact how this is handled.

The lecture (see Birth defects and the maternal child health pyramid.ppt) will challenge you to consider how the management and prevention of birth defects with a genetic component requires the interplay and cooperation of the various levels of public health service. You are encouraged to review the following web sites related to birth defects and folic acid and keep the following questions in mind.

Birth defects:

1) National Center for Birth Defects and Developmental Disabilities
   www.cdc.gov/ncbddd/

2) National Birth Defect Prevention Network
   www.nbdpn.org

Folic Acid:

1) Advocating for Folic Acid: A Guide for Health Professionals
   www.folicacid.net

2) National Council on Folic Acid
   www.folicacidinfo.org/about_us.php

Maternal Child Health Pyramid:
MCH Bureau Definitions of Core Public Health Services and Key Words

1. What are the needs of individuals with birth defects?
2. How do direct health care services help to meet these needs?
3. How do enabling services help to meet these needs?
4. How do population-based services help to meet these needs?
5. How does the public health infrastructure help to meet these needs?

Lecture - Birth defects and the maternal child health pyramid

Cancer: Case 2 - Your next-door neighbor tells you that 2 of her 4 daughters have recently been diagnosed with breast cancer. You all grew up together and are worried about the "cancer street." Describe how public health and genetic help you address risk assessment.

With the completion of the Human Genome Project inherited risk factors are increasingly being identified as contributors to common chronic diseases. As clinical testing strives to keep up with research advances in genetics, public health officials are recognizing the value of family history as an important screening tool. In this part of the lecture (see The Genetic Component of a Common Disease.ppt) we use the example of cancer and cancer clusters as a paradigm for the inherited genetic contribution to common diseases, and we discuss the incorporation of genetic information into public health investigations of these diseases.

After exploring these web sites
think about possible answers to the following questions:

1. Why is it important for a public health official to know about family health histories?
2. What are potential barriers that limit what public health investigators can learn about family health histories?
3. How is genetic susceptibility screening different than traditional public health screening tests like tuberculosis screening or smallpox screening with respect to:
   a. Disease symptoms (present/absent)
   b. Insurance, employment discrimination
   c. Who else is at risk?

Lecture - The Genetic Component of a Common Disease

Health manpower: Case 3 - You are a health planner and suddenly learn that there are no nutritionists in the state who have training in handling infants who are diagnosed on newborn screen with metabolic disease. Describe how you would address this need.

This lecture (see Health manpower and newborn screening.ppt - below) will take you through one public health geneticist's approach to this question. You are strongly encouraged to choose one of the current [for example: sickle cell and hemoglobinopathies, phenylketonuria (PKU), maple syrup urine disease (MSUD), homocystinuria, hypothyroidism, biotinidase deficiency, congenital adrenal hyperplasia (CAH) or medium chain acyldehydrogenase (MCAD)] conditions screened for in Virginia at birth and review the following web sites to answer the following questions. (Look at the conditions now included as the result of the 2005 General Assembly actions, the simplest way is to go to the VDH Genetics Program web site and look at What's new)

www.aap.org (Pediatrics 2000 Aug; 10692 pt 2)389-422. Screening the family from birth to the medical home. Newborn screening: a blueprint for the future - a call for a national agenda on state newborn screening programs)

http://genes-r-us.uthscsa.edu (National Newborn Screening and Genetics Resource Center, 2000 National NBS report)

www.geneticalliance.org (national coalition of genetic support groups, useful for getting information on a particular genetic condition by going to the particular condition's support group's web page)

www.marchofdimes.com (look for information sheets for parents)

1. Why is the condition screened for at birth?
2. How many children on average are born annually with this condition in Virginia and in the United States?
3. How is the condition treated?
4. What are the issues involved in informed consent/dissent for newborn screening?
5. What needs to be in place for an effective newborn screening and follow-up system for this condition in Virginia?

Lecture - Health manpower and newborn screening