

NIEHS study offers insight into why cancer incidence increases with age

By Robin Arnette

Scientists have known for years that age is a leading risk factor for the development of many types of cancer, but why aging increases cancer risk remains unclear. In a [study](http://www.ncbi.nlm.nih.gov/pubmed/24287154) (<http://www.ncbi.nlm.nih.gov/pubmed/24287154>) published in the February issue of the journal *Carcinogenesis*, researchers from the NIEHS Epidemiology Branch, [Zongli Xu, Ph.D.](#), and [Jack Taylor, Ph.D., M.D.](#), identified DNA methylation sites across the human genome that changed with age.

Researchers have suspected that DNA methylation, or the binding onto DNA of chemical tags called methyl groups, may be involved in the increase of cancer incidence with age. Methyl groups activate or silence genes by affecting interactions between DNA and the cell's protein-making machinery. Xu and Taylor demonstrated that a subset of those sites - the ones that become increasingly methylated with advancing age - are also disproportionately methylated in a variety of human cancers.

"You can think of methylation as dust settling on an unused switch, which then prevents the cell from turning on certain genes," Taylor said. "If a cell can no longer turn on critical developmental programs, it might be easier for it to become a cancer cell."

Age-related methylation and tumor growth

Xu and Taylor made the discovery using blood samples from participants in the [Sister Study](#), (<http://www.sisterstudy.niehs.nih.gov/English/index1.htm>) a nationwide research effort to find the environmental and genetic causes of breast cancer and other diseases. More than 50,000 sisters of women who have had breast cancer are participating in the study.

The researchers analyzed blood samples from 1,000 women, using a microarray that contained 27,000 specific methylation sites. Nearly one-third of the sites showed increased DNA methylation in association with age. They then looked at three additional data sets, from smaller studies that used the same microarray, and found 749 methylation sites that behaved consistently across all four data sets. As an additional check, they consulted methylation data from normal tissues and seven different types of cancerous tumors in [The Cancer Genome Atlas](#), (<http://cancergenome.nih.gov/>) a database funded by the National Cancer Institute and the National Human Genome Research Institute.

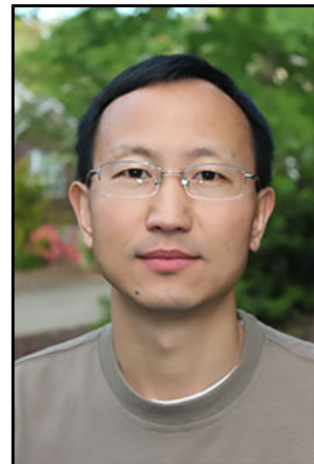
Taylor said that DNA methylation appears to be part of the normal aging process and occurs in genes involved in cell development. Cancer cells often have altered DNA methylation, but the researchers were surprised to find that 70-90 percent of the sites associated with age showed significantly increased methylation in all seven cancer types. Taylor suggests that age-related methylation may disable the expression of certain genes, making it easier for cells to transition to cancer.

Understanding how the environment affects methylation

The research also determined how fast these methylation events accumulate in cells. They occur at a rate of one per year, according to Xu.

"On your 50th birthday, you would have 50 of these sites [from the subset of 749] that have acquired methyl groups in each cell," Xu said. "The longer you live, the more methylation you will have."

For future work, Xu and Taylor want to examine more samples, using a newer microarray that will explore methylation at 450,000 genomic methylation sites. The additional samples and larger microarray, which will provide 16 times more genomic coverage, will allow them to address whether environmental exposures during adulthood or infancy affect methylation profiles. These additional studies will help scientists better understand why methylation happens as people march toward their retirement years.



Xu is a staff scientist in the NIEHS Epidemiology Branch. (Photo courtesy of Zongli Xu)



Taylor holds appointments in the NIEHS Epidemiology Branch and the Molecular Carcinogenesis Laboratory. (Photo courtesy of Steve McCaw)

DNA methylation is one of several epigenetic mechanisms that can control gene expression without changes in DNA sequence. This study is part of a broader research effort, funded by NIEHS, to understand how environmental and other factors affect epigenetic mechanisms in relation to health.

Citation: [Xu Z, Taylor JA.](#)

(<http://www.ncbi.nlm.nih.gov/pubmed/24287154>)

2014. Genome-wide age-related DNA methylation changes in blood and other tissues relate to histone modification, expression, and cancer. *Carcinogenesis* 35(2):356-364.

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