Flagship Results Paper from the MedSeq Project Published in Annals of Internal Medicine

The final flagship results paper from the 4-year, \$10-million NIH-funded MedSeq Project, the first-ever randomized trial conducted to examine the impact of whole genome sequencing (WGS) in healthy primary care patients, was published June 26, 2017, in the *Annals of Internal Medicine*.

This paper is of note because it is the first randomized clinical trial of "end-to-end" WGS in the day-to-day practice of medicine, enrolling healthy adults and delivering sequencing reports to their primary care physicians without intermediation by a geneticist or genetic counselor. Additionally, this is the first study to comprehensively analyze and report nearly 5,000 disease-



associated genes, including monogenic disease risks, carrier variants, pharmacogenomic associations and polygenic risk estimates. A remarkable 22% of the patients had new monogenic disease risk results - higher than previously suspected or documented. Moreover, overall, patients did not exhibit anxiety or distress about



Senior author and director of the Genomes2People Research Group, Robert C. Green, MD, MPH (left) and first author and assistant professor, Jason Vassy, MD, MPH, SM.

their findings, and the actions of the primary care doctors in response to the sequencing were rated by genetics experts and were found to be largely appropriate. Finally, this new paper reports the first estimates of potential economic burden of WGS in healthy adults on the health care systems, showing that resulting consultations, clinical tests and other downstream health care costs were modestly increased for the patients who received WGS in comparison to the control group.

Several mainstream media and scientific outlets covered the announcement of this exciting paper; read on for links to those articles in addition to links to

the newly published <u>MedSeq paper</u>, <u>press release</u>, <u>video</u> of first author Dr. Jason Vassy, and a <u>complete list</u> of MedSeq Project publications, which investigators have previously published from this implementation study.

NPR All Things Considered

Routine DNA sequencing may be helpful and not as scary as feared

Wired

You can get your whole genome sequenced. But should you?

Science

One in five 'healthy' adults may carry diseaserelated genetic mutations

MIT Technology Review

DNA testing reveals the chance of bad news in your genes

The Scientist

The consequences of sequencing healthy people

The Washington Post

Scientists discover DNA might not be that useful as part of your annual checkup

Reuters Whole genome sequencing not ready for routine use: study

STAT News

In healthy patients, genome sequencing raises alarms while offering few benefits

Bio-IT World

Sequencing healthy patients reveals that many carry rare genetic disease risks

CardioBrief

Does whole genome sequencing have a role in primary care?

Science Daily

Sequencing finds rare genetic disease risk in one out of five healthy adults

GenomeWeb

Studies make case for sequencing healthy individuals, but questions remain

Front Line Genomics Questioning the value of genome sequencing in healthy people

Xconomy

<u>New study: DNA tests for healthy people have</u> <u>"uncertain value"</u>

American Journal of Managed Care (AJMC)

Genome Sequencing Reveals Rare Disease Risk in 1 of 5 Generally Healthy Adults