

60 Utahns are among landmark large-scale genome sequencing study

October 27 2010

Just seven months after University of Utah geneticists took part in a <u>landmark study</u> that sequenced for the first time the genome of an entire Utah family, U of U researchers have taken part in another historic study that is the first large-scale genome sequencing project – 179 people representing three continents – and 60 Utahns played a major role in this study, too.

Published Wednesday, Oct. 27, 2010, in *Nature*, the study demonstrates how quickly the science of genome sequencing is expanding – first from individuals, then to families, and now to large groups of people representing distinct geographic and ethnic populations, according to Lynn B. Jorde, Ph.D., professor and chair of human genetics in the U of U School of Medicine and a senior investigator and co-author on the study. Rapid-fire technological advances and highly sophisticated computers are giving geneticists the ability to sequence the genomes of increasingly larger groups of people at lower costs, and this study is an example of the research that breakthroughs in technology enable. It highlights as well the significant role Utahns have played in groundbreaking genetics studies.

"This study provides the first large-scale inventory of human genetic diversity," Jorde said. "It also is a good example of how Utahns have helped researchers worldwide. Genetics studies keep coming back to Utah."

The study is part of the 1000 Genomes Project Consortium – an



international collaboration of hundreds of geneticists from dozens of laboratories with the goals of cataloging the range of genetic diversity found in people worldwide and characterizing 95 percent of the most common genetic mutations found in humans. Along with being the first large-scale genome-sequencing project, the study made several important findings, including confirmation of earlier work of Jorde's lab group and scientists at the Institute for Systems Biology in Seattle that gave the first direct estimate of the rate of genetic mutation in humans; providing evidence of recent natural selection in new genes; discovering a number of genetic mutations that occur often enough to be considered among the most common in humans; and identifying new "mobile elements" – DNA sequences that randomly reshuffle in the genome.

Earlier this year, Jorde, who is on the 1000 Genomes Project steering committee, was part of the team that was the first to sequence the genome of an entire family – two parents and two children who live in Utah. As part of that study, published in March in Science, he estimated the rate at which genetic mutations are passed from generation to generation at 60 – meaning each parent passes 30 genetic mutations to their offspring. Most gene mutations are harmless, but understanding the rate at which mutations are passed among generations is an essential part of understanding the human biological clock, according to Jorde. To confirm his estimated mutation rate, which was half of what had been estimated previously by indirect methods, researchers in the current study sequenced the genomes of two families of three people each.

"We were delighted that the mutation rate estimate obtained from the 1000 Genomes Project was exactly the same as our estimate," Jorde said.

In the large-scale sequencing phase of the project – in which the genomes of 60 Utahns representing Northern Europe; 60 Nigerians representing Africa; and 30 Chinese and 30 Japanese people



representing Southeast Asia were sequenced – the researchers identified a number of new genes that appear to have undergone recent natural selection. That mirrors another study from Jorde's lab, published in Science in July of this year, that showed evidence of genetic natural selection within the past 5,000 to 10,000 years that enables the people of Tibet to thrive in the low-oxygen environment of extremely high altitudes. Finding further proof of natural selection in a broader diversity of people in the current study provides evidence to answer another basic question about humans: are we still evolving?

"These studies provide an unequivocal answer – yes," Jorde said.

The 60 Utahns in the project were part of a number of families who 30 years ago volunteered for a U of U study investigating how genetic variations are passed in families.

As they sequenced the genomes of those three population groups, the researchers also discovered a number of genetic mutations that occur often enough – defined as being found in greater than 1 percent of people – to be considered common gene mutations. With the discovery of these new gene mutations, more than 15 million DNA variations have been identified in the human genome and the researchers believe the 1000 Genomes Project has reached its goal of identifying 95 percent of common genetic mutations. That's a major step that provides a reference of genetic variation for researchers worldwide trying to unravel the genetic bases of countless diseases, according to Jorde.

"People now have a catalog for comparing their research results," he said. "Our ultimate goal is to use this information to understand the causes of disease."

In a third phase of the study, researchers including Jin chuan Xing, Ph.D., a postdoctoral fellow in Jorde's lab, looked at the DNA of 697



people and discovered new "mobile elements" – DNA sequences that pop up randomly in peoples' genomes. Although the function of mobile elements is unknown, they comprise half the human <u>genome</u> and are believed to help propel evolution. This was the first study to look at mobile elements in a large population of people, Xing said.

"We discovered a lot of new ones," he said. "This will help us look at how mobile elements assemble."

Provided by University of Utah

Citation: 60 Utahns are among landmark large-scale genome sequencing study (2010, October 27) retrieved 4 May 2024 from <u>https://medicalxpress.com/news/2010-10-utahns-landmark-large-scale-genome-sequencing.html</u>

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