The human genome is at last complete. Researchers have been working for decades toward this goal, and the Human Genome Project claimed victory in 2001, when it had read almost all of a person's DNA. But the stubborn remaining 8 percent of the genome took another two decades to decipher. These final sections were highly repetitive and highly variable among individuals, making them the hardest parts to sequence. Yet they revealed hundreds of new genes, including genes involved in immune responses and those responsible for humans developing larger brains than our primate ancestors. “Now that we have one complete reference, we can understand human variation and how we changed with respect to our closest related species on the planet,” says geneticist Evan Eichler of the University of Washington, one of the co-chairs of the Telomere-to-Telomere consortium that finished the genome.
In the horizontal bars below, each of the 22 numbered human chromosomes and two sex chromosomes (X, Y) are divided into regions (thin vertical stripes) of 1,000,000 bases, or nucleotides. Different bar lengths reflect the chromosomes’ varying physical lengths, as demonstrated by these chromosome-pair illustrations. Colors indicate regions that reached 50, 90 or 99 percent completion in each year.

In 2002, 40% of the genome flipped to 99% coverage. About 70% was completed at this level.

In 2003 chromosomes 1–5 and 12 received new coverage, whereas 13, 14 and 19–22 saw little change.

Completion from previous years is carried over in gray.
From 2004 to 2013 updates were relatively minor. 

In 2006 less than 1% of the genome flipped coverage status.

By 2009 about 80% of regions had been sequenced to 99% or better.

By 2013 about 8% of the genome was not sequenced at all.

In 2022 scientists added 251,330,203 bases for a totally gapless genomic sequence.

Credit: Martin Krzywinski; Sources: UCSC Genome Browser; “The Complete Sequence of a Human Genome,” by Sergey Nurk et al., in Science, Vol. 376; April 2022

Editor’s Note (7/22/22): The graphic in this article was edited after posting to correct the number of bases in a totally gapless genomic sequence in 2022.

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*Credit: Nick Higgins*

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