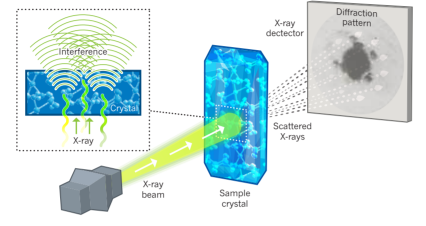


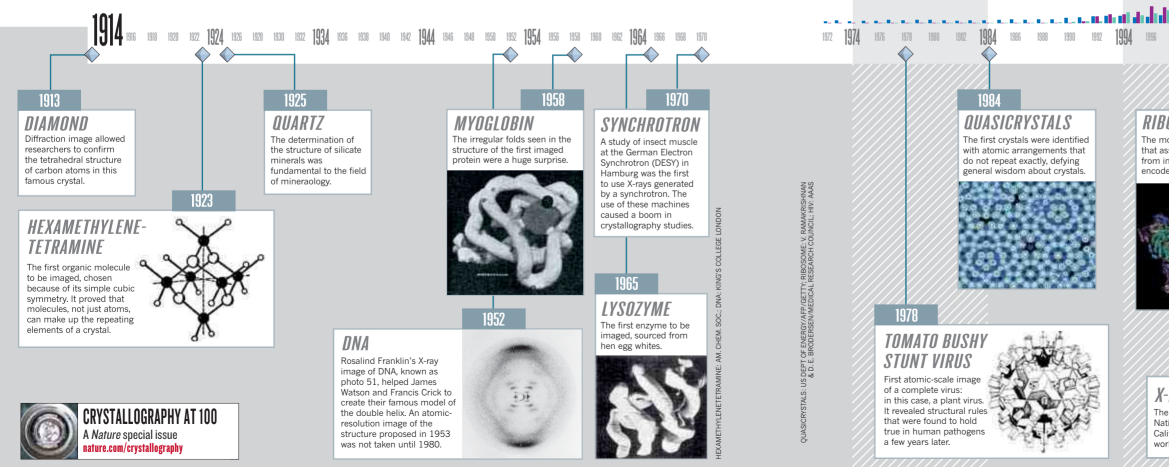
ATOMIC SECRETS *100 YEARS OF CRYSTALLOGRAPHY*

In 1914, German scientist Max von Laue won the Nobel Prize in Physics for discovering how crystals can diffract X-rays: a phenomenon that led to the science of X-ray crystallography. Since then, researchers have used diffraction to work out the crystalline structures of increasingly complex molecules, from simple minerals to high-tech materials such as graphene and biological structures, including viruses. With improvements in technology, the pace of discovery has accelerated: tens of thousands of new structures are now imaged every year. The resolution of crystallographic images of proteins passed a critical threshold for discriminating single atoms in the 1990s, and newer X-ray sources promise images of challenging proteins that are hard or impossible to grow into large crystals.



BIRTH OF AN IDEA

Von Laue hit on the idea that when X-rays passed through a crystal, they would scatter off the atoms in the sample and then interfere with each other like waves passing through a breach in a shore wall. In some places, the waves would add to each other; in others, cancel each other out. The resulting diffraction pattern could be used to back-calculate the location of the atoms that scattered the original X-rays. Von Laue and his colleagues proved his theory in 1912 with a sample of copper sulphate.



BERNHARD RUPP/GARLAND SCIENCE/TAYLOR & FRANCIS H. M. BERMAN *PROTEIN SCI.* **21**, 1587–1596 (2012), WITH UPDATES FROM WORLDWIDE PROTEIN DATA BANK

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WORLDWIDE PROTEIN DATA BANK/ RYSTALLOGRAPHY OPEN DATABASE

10.000

5.000

1976

GOING UP

models of some structures

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GETTING CLEARER Better techniques for both imaging and

interpreting data have allowed researchers

1986 1988 1990

The Worldwide Protein Data Bank has been collecting resolved

entries. Other databanks, including the Crystallography Open

Database (COD), include structures of everything from minerals

instructions into its database for how to print three-dimensional

to metals and small biological molecules. The COD is now adding

structures of proteins since 1971, and now holds nearly 100.000

1992

see finer details in some structures and

tackle ever more complicated molecules.



