

Massive genome study unlocks secrets of how cancers form

February 6 2020, by Sara Hussein



A patient undergoes a mammogram to detect early signs of breast cancer

A massive, decade-long study sequencing the genomes of dozens of cancers has revealed the secrets of how tumours form and may pave the way for better and more targeted treatment.

The Pan-Cancer Project [brought together](#) over 1,300 researchers globally to tackle the mammoth task of sequencing the genomes of 38 types of cancer in nearly 2,800 patients.

Their work produced a host of new discoveries—from the number and location of so-called driver mutations that push cells to reproduce uncontrollably, to the surprising similarities between cancers found in different types of tissue.

The results were published Thursday in nearly two dozen papers in *Nature* and other Nature Research journals and represent the largest and most comprehensive study of whole cancer genomes ever.

"With the knowledge we have gained about the origins and evolution of tumours, we can develop new tools and therapies to detect cancer earlier, develop more targeted therapies and treat patients more successfully," said Lincoln Stein, a member of the project steering committee, in a statement issued by the Ontario Institute for Cancer Research.





















Among the key findings of the work is the massive variety in cancer genomes, said Peter Campbell of the Wellcome Sanger Institute, another steering committee member.

"The most striking finding is just how different one person's cancer genome is from another person's," he told AFP.

The study found thousands of combinations of mutations in individual cancers, as well as over 80 processes that cause the mutations, some of them age-related and others inherited or linked to lifestyle factors such as smoking.

WHO global cancer report

Most prevalent cancers, 2018

MEN			WOMEN		
		Cases			Cases
Lung		1,368,524	Breast		2,088,849
Prostate		1,276,106	Colorectal		823,303
Colorectal		1,026,215	Lung		725,352
Stomach		683,754	Cervix uteri		569,847
Liver		596,574	Thyroid		436,344
Bladder		424,082	Corpus uteri		382,069
Oesophagus		399,699	Stomach		349,947
Non-Hodgkin lymphoma		284,713	Ovary		295,414
Kidney		254,507	Liver		244,506
Leukaemia		249,454	Non-Hodgkin lymphoma		224,877

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Source: WHO report on Cancer/IARC Global Cancer Observatory

A massive decade-long study has sequenced the genomes of 38 types of cancer, revealing secrets about how tumours form

'Exciting themes'

But within the enormous variety there were "exciting themes", Campbell said.

For example, the work found the early development of some cancers can occur decades before diagnosis, sometimes even in childhood.

"This shows that the window of opportunity for early intervention is much wider than we expected," Campbell said.

The research also found that patterns of mutations, and where they occur, can help identify the approximately 1-5 percent of cancers that

cannot be identified through regular diagnostics.

A sequenced genome can even reveal occasional misdiagnosis of a cancer type.

Most work on sequencing the cancer [genome](#) has focused on the approximately two percent known as the protein-coding genes.

But the Pan-Cancer Study sequenced entire genomes, uncovering new cancer-causing driver mutations in the other 98 percent, known as non-coding genes.



Gene sequencing helps map out the many types of mutations that can cause cancer

The researchers found enormous variation in the number of mutations in a given cancer, from very few in some cancers seen in children, to up to 100,000 in lung cancer samples.

And in around five percent of cases, no known driver mutations were found at all, implying there are mutations that have not yet been identified.

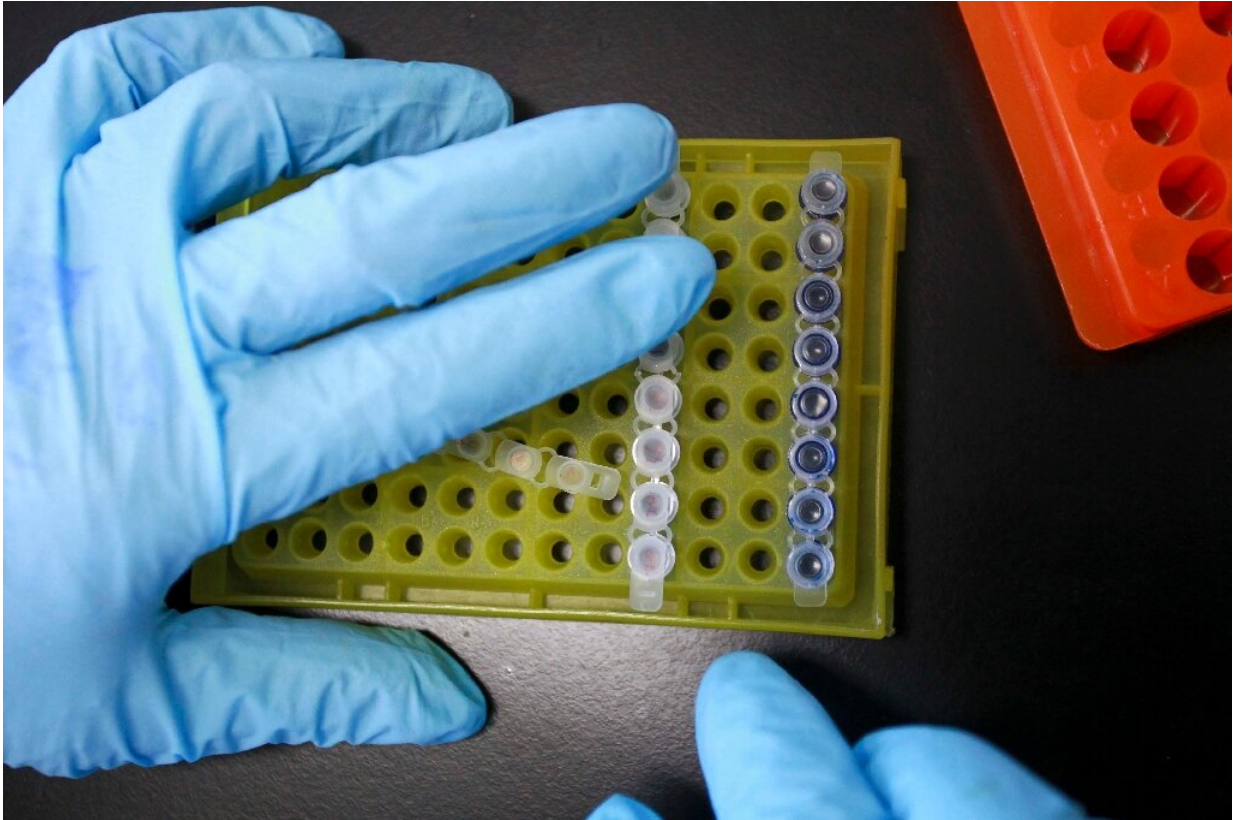
'Enormous spectrum'

The sequencing helps map out the many types of mutations—from changes in single DNA letters to much larger insertions or deletions of genetic code—that can cause cancer.

It also revealed that cancers in different parts of the body are sometimes much more alike than had been thought.

"We may have a type of breast cancer and prostate cancer where the driver mutations are similar," said Joachim Weischenfeldt, a co-author and associate professor at the University of Copenhagen.

"This means that the patient with prostate cancer may benefit from the same treatment as the one you would give the breast cancer patient," he said in a statement issued by the university.



Most work on sequencing the cancer genome has focused on the approximately two percent known as the protein-coding genes

In practical terms, the findings will help identify difficult-to-diagnose cancers, allow more targeted treatment based on the specific driver mutations behind a particular cancer, and potentially allow earlier diagnosis of developing tumours.

"We are finding that [cancer](#) represents the far end of an enormous spectrum of change," said Campbell.

"If we can understand the forces at play in our normal organs as we age, what causes [mutations](#) to accumulate, what causes some clones to expand and others to fade, what lifestyles do to tilt this balance, then we can

think about ways to intervene early, with a view to preventing or slowing the emergence of untreatable cancers."

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