

'Wildly heterogeneous genes'

September 15 2013

Cancer tumors almost never share the exact same genetic mutations, a fact that has confounded scientific efforts to better categorize cancer types and develop more targeted, effective treatments.

In a paper published in the September 15 advanced online edition of *Nature Methods*, researchers at the University of California, San Diego propose a new approach called network-based stratification (NBS), which identifies cancer subtypes not by the singular mutations of individual patients, but by how those mutations affect shared genetic networks or systems.

"Subtyping is the most basic step toward the goal of personalized medicine," said principal investigator Trey Ideker, PhD, division chief of genetics in the UC San Diego School of Medicine and a professor in the departments of Medicine and Bioengineering at UC San Diego. "Based on patient data, patients are placed into subtypes with associated treatments. For example, one subtype of cancer is known to respond well to drug A, but not drug B. Without subtyping, every patient looks the same by definition, and you have no idea how to treat them differently."

Recent advances in knowledge and technology have made it easier (and less expensive) to sequence individual genomes, especially in the treatment of cancer, which is fundamentally a disease of genes.

But genes are "wildly heterogeneous," said Ideker. It is in combination, influenced by other factors, that mutated genes cause diseases like cancer. Every patient's cancer is genetically unique, which can affect the



efficacy and outcomes of clinical treatment.

"When you look at patients' data at the level of genes, everybody looks different," said Ideker. "But when you look at impacted <u>biological</u> <u>networks</u> and systems, groupings do appear. No genes are mutated in exactly the same place, but the mutations do appear in the same <u>genetic</u> <u>pathways</u>."

Specifically, the scientists looked at <u>somatic mutations</u> – present in tumors but not healthy tissues – in data from lung, uterine and ovarian cancer patients compiled by The Cancer Genome Atlas, an on-going National Institutes of Health-funded program to gather and catalogue the genomes of thousands of cancer patients.

Ideker said the NBS approach has immediate clinical value. Genome sequencing of cancer patients is rapidly becoming a standard part of diagnosis. Clinicians can use NBS, he said, to better match treatment to cancer subtype. And by chronicling treatment outcomes, funneling those results back into the database, they can further refine and improve cancer therapies, making them as personalized as the individuals themselves.

More information: *Nature Methods*, DOI: 10.1038/nmeth.2651

Provided by University of California - San Diego

Citation: 'Wildly heterogeneous genes' (2013, September 15) retrieved 1 May 2024 from <u>https://medicalxpress.com/news/2013-09-wildly-heterogeneous-genes.html</u>

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