

## **Reprogramming gut bacteria as 'living therapeutics'**

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MIT spinout Synlogic is re-programming bacteria found in the gut as "living therapeutics" to treat major diseases and rare genetic disorders. Credit: Synlogic

An MIT spinout, Synlogic, is aiming to create a new class of medicines, by re-programming bacteria found in the gut as "living therapeutics."



Based on research by its co-founders, MIT professors Tim Lu and Jim Collins, Synlogic creates so-called synthetic biotics, which sense and correct metabolic abnormalities that underlie some major diseases and <u>rare genetic disorders</u>.

Human intestines are filled with trillions of bacteria, collectively called the microbiota, that carry out vital health functions. Synlogic's synthetic biotics—capsules, liquid suspensions, or other dosage forms that can be taken regularly—augment the microbiota with new metabolic capabilities or complement lost functionality in organs such as the liver.

"Over the past decade or so, the intricate connections between microbes and our bodies have become clearer and clearer, and it's well known now that the bacteria that live in our gut have a major influence on human health," says Lu, an associate professor of electrical engineering and computer science and of biological engineering, and head of MIT's Synthetic Biology Group, who serves as scientific advisor for Synlogic. "We leverage that interface as a way of treating human disease."

Last month, Synlogic raised an additional \$40 million in venture capital and secured its first industry partnership with pharmaceutical giant AbbVie. For the partnership, Synlogic will collaborate with AbbVie to develop synthetic biotics for the potential treatment of inflammatory bowel disease, which may include probiotic microbes programmed to detect intestinal inflammation, and produce anti-inflammatory molecules or break down pro-inflammatory effectors.

Two of Synlogic's main candidate drugs, expected to enter clinical trials during the next 12 months, treat rare genetic metabolic disorders. One drug candidate is for treating urea cycle disorder (UCD), which is caused by an enzyme deficiency that leads to a buildup of toxic ammonia in the blood. The other is for treating phenylketonuria (PKU), which involves a dangerous excess of the amino acid phenylalanine due to a mutation in



another metabolic enzyme. In both cases, Synlogic's drugs process and flush out the toxic metabolites from the body.

Think of Synlogic's drugs like biological thermostats, says Collins, the Termeer Professor of Medical Engineering and Science in MIT's Department of Biological Engineering and Institute for Medical Engineering and Science, who also chairs Synlogic's scientific advisory board. Instead of identifying and regulating the temperature of a room, he says, "The synthetic biotics detect and regulate the amount of an enzyme or metabolic byproduct in a patient's body."

## **Programming E. coli**

For more than a decade at Boston University and MIT, Collins and Lu (who is Collins' former student) have been developing "genetic circuits" for bacteria, which include on/off switches made with synthetic DNA or RNA sequences that instruct the bacteria to count, store memory, and even perform logic.

Collins and Lu have used these genetic circuits to program bacteria to seek and cure infection. In 2011, this approach earned Collins funding from the Bill and Melinda Gates Foundation to engineer bacteria to detect cholera and produce targeted antimicrobial peptides to treat it.

A few years ago, Lu and Collins, along with several venture capitalists, launched Synlogic to focus on commercializing "a new class of therapeutics based on living cells," Collins says. In 2014, Synlogic came out of stealth mode, securing \$30 million in funding from venture firms and the Gates Foundation.

Since then, Synlogic has worked primarily on programming E. coli Nissle, a strain of bacteria derived from the gut that is also used widely and safely as a probiotic. The programmed E. coli Nissle, Lu says,



provides greater precision, safety, and efficacy for disease treatment, compared with traditional methods.

For inflammatory bowel diseases such as Crohn's disease and ulcerative colitis, for instance, current treatments include small-molecule drugs or antibodies with anti-inflammatory properties. But the challenge is getting the right dosage, Lu says. "If you apply too little, it's not going to work. If you apply too much, you have a chance you may immunosuppress the patient and cause side effects," he says.

Synlogic can "program microbes to detect inflammation and make antiinflammation molecules at the site of inflammation, as well as produce molecules that positively impact immune system function," Collins says.

Then there's the more rare but debilitating urea cycle disorder, which affects 2,000 to 6,000 people in the United States and impairs their ability to processes ammonia. If ammonia builds up too much and reaches the brain, it can lead to brain damage, coma, and death. The best available treatment option for people with UCD today is a liver transplant.

Synlogic aims to treat UCD with a daily biotic that functions in a surprising way: "It can decrease the ammonia in the bloodstream, without even contacting the blood," Lu says.

Ammonia levels in the bloodstream, he explains, are dependent on ammonia production in the large intestine. Synlogic's biotic converts intestinal ammonia into an amino acid, which is flushed out of the body through the stool, thereby dramatically reducing the flow of ammonia to the blood and reducing ammonia levels in the bloodstream.

Synlogic's biotic for PKU, which affects 13,000 people in the United States, functions in a similar way, to regulate the processing and



extraction of phenylalanine. PKU patients must adhere to a lifelong, extremely low-protein diet that can result in serious developmental disorders, because they can't eat normal foods that contain phenylalanine—including many meat, dairy, and seafood products. "If we can degrade phenylalanine with convenient administration of this probiotic, that will change the course of this disease," Lu says.

Collins says Synlogic has potential to treat many other rare genetic metabolic disorders. But the recent AbbVie deal, he says, also "opens up possibilities of using these microbes to produce biologics or other small molecules to treat a range of conditions." These include cardiovascular disease and autoimmune, oncology, and central nervous system disorders, which have been linked to metabolic dysregulation.

## **Reaching clinical efficacy**

Part of the reason that probiotic treatments are not used for serious diseases is their lack of clinically validated efficacy. Synlogic, on the other hand, aims to overcome these efficacy issues with potent and precision-programmed synthetic biotics, Lu says.

For example, in engineering the safe and easily programmable E. coli Nissle, Synlogic engineers have designed the microbe to consume a massive amount of toxic metabolites. The E. coli Nissle strain that forms the basis of Synlogic's UCD program, for instance, can consume orders of magnitude more ammonia than natural E. coli can, Lu says. "For this treatment to work for patients, you want the max performance you can squeeze out of any one of these biotics," he says.

Based on their preclinical data, Synlogic's treatments have the potential to reach "clinical levels" of efficacy not seen often in synthetic biology, says Collins: "Synlogic is programming these probiotic microbes to consume ammonia or phenylalanine for example, and they are reaching



levels that are expected to be clinically meaningful, which is quite remarkable."

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