A celery gene may protect roses from disease

Researchers from the North Carolina State University, NC, USA, have undertaken a study intended to overcome petal blight in roses – one of the most common post-harvest diseases which significantly shortens their longevity.

Several fungal plant pathogens secrete mannitol – an agent that interferes with the plant’s defense mechanisms. Many plants naturally produce mannitol dehydrogenase (MTD) which catabolizes mannitol of fungal origin. It is not well determined whether roses produce this enzyme, but even if they do, the amount is not sufficient to protect them from the unfavorable effect of mannitol. MTD has already been demonstrated to improve defense mechanisms, for instance in petunias. Therefore, North Carolina State University horticultural scientists, Dr. John Dole and Dr. John Williamson, introduced the MTD celery gene into the rose’s genome to improve the mannitol-catabolizing capacity. Genetically modified roses growing in the North Carolina State University greenhouses appear and smell just like the conventional roses. In the nearest future, tests will be conducted to find out if they have become more resistant to the petal blight.

Mick Kulikowski, the editor of News Services of the North Carolina State University, summarized this innovative study with a short biotech rhyme: “Roses are red, Celery is green, Roses last longer with a celery gene”.

Sources

15th European Congress on Biotechnology

The Biotechnology Association of Turkey, along with the European Federation of Biotechnology (EFB), will organize the 15th European Congress on Biotechnology of the European Federation of Biotechnology – ECB15, which will be held in Istanbul, September 23-26, 2012. The organizers intend to attract top scientists, decision and policy makers, high-tech and start-up companies, entrepreneurs, investors and students from Europe, Asia and the Middle East.

The main theme of the 15th European Congress on Biotechnology is Biocrossroads – analyzing the impact of the life sciences industry in addressing humanity’s great challenges. It will be covered by four parallel sessions: Health & Medicine – focusing on the latest achievements in the field of biotechnology to improve existing treatments and put forward novel approaches, with a special focus on the development of new innovative and targeted therapies; Industrial Biotechnology – a multidisciplinary approach including applied biocatalysis, biochemical engineering, microbial physiology and systems biology – all focused on innovation in industrial biotechnology and the development of the Knowledge-Based Bio-Economy; Plant & Environmental Biotechnology – exploring the benefits and challenges derived from the application of biotechnologies in agriculture and their key role in providing the biomass the world needs; Systems Biology & Technology – novel “omics” and technology-based approaches for dramatically increasing and improving capacities for experimentation.

Deadline for abstract submission is April 1, 2012. A number of abstracts will be selected for oral presentations. All other qualifying abstracts will be selected for poster presentations. Further information is available on the website: www.ecb15.org.

Environmental Microbiology and Biotechnology Conference 2012

From April 10-12, 2012, University of Bologna, Italy, will host the conference Environmental Microbiology and Biotechnology in the Frame of the Knowledge-Based Bio and Green Economy (EMB2012). The event is organized by the Environmental Biotechnology section of the European Federation of Biotechnology (EFB) in cooperation with the Federation of European Microbio-
logical Societies (FEMS). The program will be especially focused on the advances in sustainable decontamination of polluted habitats, purification and reuse of water resources, biofixation of climate change-inducing gases and the production of bio-based chemicals, materials and fuels from bio-waste. The organizers will address several priorities of the Europe 2020 strategy with the intention to contribute toward constructing a more sustainable and competitive knowledge-based economy in Europe and in the Partner Countries. Abstract submission is open until February 1, 2012. Selected papers will be invited for the full publication in New Biotechnology.

More details about the conference may be found at: www.unibo.it/EMB2012.

**FDA approved first cell therapy against wrinkles**

In June 2011, United States Food and Drug Administration (FDA) approved the Biologics License Application for laViv (azficel-T) – the first personalized cell therapy. The product was developed by Fibrocell Science, Inc. (Exton, PA), a biotechnology company that focuses on the development of personalized autologous cell therapies for aesthetic, medical and scientific applications. laViv is intended to improve the appearance of moderate to severe nasolabial fold wrinkles (smile lines) in adults.

laViv is an autologous cellular product composed of fibroblasts suspended in Dulbecco’s Modified Eagle’s Medium (DMEM) without phenol red. Firstly, patient’s own dermal fibroblasts are harvested from behind the ear through the skin biopsy. Next, the cells are aseptically expanded using standard tissue culture procedures. The recommended course of laViv administration is a series of three treatments, typically three to six weeks apart. The process for production of laViv takes approximately 11 to 22 weeks.

Investigators involved in the clinical trials of laViv point out that people who can afford this type of cosmetic treatments may be especially attracted to the idea that their own cells are used in this approach rather than artificial fillers. Moreover, the investigators claim that the effect seems to be longer lasting in comparison to the other fillers available.

Noteworthy, laViv is only the cellular therapy product approved in the United States so far. The two other FDA approved products have strictly medical use: Cartilgel (autologous cultured chondrocytes) developed by Genzyme (Cambridge, MA) is used to repair knee injuries and Provenge (sipuleucel-T), manufactured by Dendreon Corporation (Seattle, WA), is used to treat metastatic hormone-refractory prostate cancer.

**Sources**


**Individualization of cancer treatment using personalized tumorgrafts**

*Molecular Cancer Therapeutics* published recently the results of a cutting edge pilot study of a treatment guided by personalized tumorgrafts in patients with advanced stage of cancer. In a study led by Manuel Hidalgo, Director of the Clinical Research Programme at the Spanish National Cancer Research Centre (CNIO), 14 patients with advanced solid cancers were evaluated in a personalized tumor model created by implanting fragments of tumor tissue in immunodeficient mice, which were treated with 63 anticancer drugs, both as single agents and in different combinations. Using this approach, an effective treatment regimen in the xenograft model was identified for 12 patients. 11 of them received 17 prospectively guided treatments, and 15 of these treatments resulted in durable responses. No effective treatments were found in 2 patients. The authors reported a remarkable correlation between drug activity in the model and clinical outcome, both in terms of resistance and sensitivity.

The results presented by Hidalgo and colleagues are very promising, however there are several limitations in the application of this approach in clinics. Firstly, the procedure requires fresh tumor tissue, which is difficult or impossible to obtain in many patients. Moreover, it usually takes 6-8 months to grow a tumor in mice and obtain graft, and it can be achieved in around 60% of cases. Additionally, in some cases, an effective treatment can-
not be found. Another obvious limitation is the cost of this type of personalized treatment. Nevertheless, the group of Manuel Hidalgo will conduct further research to develop methods which would allow to solve these problems and to make this technology widely available for patients.

The study is conducted in collaboration with Johns Hopkins University in Baltimore, USA, Hospital de Madrid, Spain, and the company Champions Oncology, Baltimore.

Sources

Mission Therapeutics Ltd, a new British spin-out company, raised £6 million

A team led by Professor Stephen Jackson together with the Cancer Research Technology (Cancer Research UK’s commercial arm) and the University of Cambridge announced launching Mission Therapeutics Ltd, the new spin-out company.

Mission Therapeutics will translate the newest cell biology research achievements on DNA repair from Professor Jackson’s laboratory at the Gurdon Institute, University of Cambridge, into drugs which are intended to improve the management of life-threatening diseases, including cancer. Professor Jackson claims that these drugs could also improve the effectiveness of existing cancer treatments, such as radiotherapy and certain chemotherapies. The company will principally exploit novel research on ubiquitin pathways that control cellular responses to DNA damage. More information about the research interest and projects conducted in Professor Stephen Jackson’s laboratory can be found at: www.gurdon.cam.ac.uk/~jacksonlab. Interestingly, the company has raised over £6 million from a venture capital syndicate led by Sofinnova Partners company, together with Imperial Innovations (technology commercialization and investment company), SR One (the corporate venture capital arm of GlaxoSmithKline) and Roche Venture Fund. Mission Therapeutics will be based on the Babraham Research Campus, Cambridge, UK.

Source

Science Multimedia Center

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Science Multimedia Center is a valuable source of scientific news prepared in approachable and visually attractive manner. Such innovative way of presenting science may certainly serve as a valuable tool and an inspiration to the academic teachers and science popularizers.

All the recordings can be accessed at: www.sciencemag.org/site/multimedia.