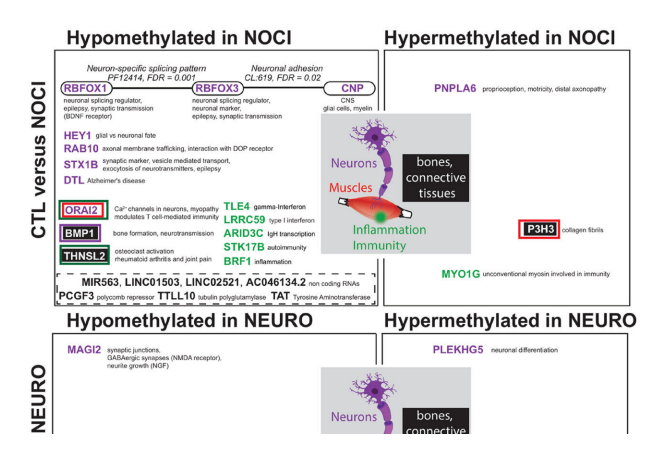


Relieving pain by mapping its biological signatures

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Genes with DMRs identified in the comparison groups are shown: CTL vs. NOCI in the upper panel, CTL vs. NEURO in the middle panel, and NOCI vs. NEURO in the lower panel. For each comparison, genes with DMRs were segregated according to the direction of the methylation changes observed: hypomethylation in the left and hypermethylation in the right column. Genes that belong to a functional network as detected by STRING 47 appear surrounded and linked together by a line, with the corresponding description and code as well as the false discovery rate (FDR). If there is no line between two genes, the



interaction between these genes is not detectable by the STRING program. This may either reflect a true absence of connection or insufficient knowledge. Functional networks may imply Homo sapiens (HSA) pathways either based on cellular processes (Kyoto Encyclopedia of Genes and Genomes, KEGG), or based on molecular reactions (Reactome), a gene ontology term (GO), functionally related protein family (INTERPRO), conserved protein domain (PF), and STRING cluster (CL). Genes with a functional link to the nervous system are shown in purple, those linked to the muscle in red, those linked to the bone and connective tissue (referred to as bone) in white, and those linked to the inflammatory system or immunity in green. For genes linked to two different systems, two different colors were used, one for the gene name and the other for the surrounding frame. Genes with no links to the above systems are shown as dashed squares. Credit: DOI: 10.1016/j.jpain.2021.09.001

Many people are confronted with chronic pain that can last for months or even years. How to best treat chronic pain? First, pain must be categorized for the right treatment to be prescribed. However, is that it is very challenging for patients to define their pain, its intensity or even its location using questionnaires. To overcome this difficulty, scientists from the University of Geneva (UNIGE) have joined forces with the research department of the Clinique romande de réadaptation (CRR) in Sion to carry out a complete epigenomic analysis of patients, making it possible to find the epigenetic signatures specific to each pain category. Thus, a simple blood test would make it possible to define which pain the person suffers from and, in the future, to prescribe treatment accordingly and to observe whether the biomarkers modified by the pain return to normal. These results can be read in the *Journal of Pain*.

Chronic <u>pain</u> is classified into two main categories: nociceptive pain—defined by the activation of receptors at the end of nerve fibers and found in osteoarthritis, burns or infections—and <u>neuropathic pain</u>, which is caused by damage to nerve structures, such as pain caused by



shingles. In order to classify which pain the patient suffers from, they fill in several questionnaires and quantify pain intensity of using assessment scales. However, this is very subjective and time-consuming.

Blind genome analysis

"At the CRR, we treat many people suffering from chronic diseases," explains Bertrand Léger, a researcher at the CRR and last author of the study. "We joined forces with UNIGE scientists to carry out a complete epigenomic study and define specific biomarkers for each type of pain, in order to be able to categorize the various types of pain quickly and reliably."

To do this, the Geneva team carried out an analysis of the entire genomes of 57 patients: 20 with no pain, 18 with nociceptive pain and 19 with neuropathic pain. "The aim was to start without any prior hypothesis to probe the genome as a whole and identify all the biomarkers involved in pain," explains Ariane Giacobino, the study's coauthor and a professor in the Department of Genetic Medicine and Development at UNIGE Faculty of Medicine.

Specific and potentially reversible biomarkers

Unexpectedly, not only did the scientists identify very striking epigenetic signatures of pain, but there was no overlap between nociceptive and neuropathic pain. "This total absence of similarities between the two categories of pain is very surprising, because intuitively, we might think that the difficulty in defining one's pain comes from a similarity in the epigenetic signature. We could prove that it absolutely not the case," notes Ariane Giacobino.

Indeed, the biomarkers specific to nociceptive pain are expressed by the



genes of the opioid system—involved in emotion, reward and pain—as well as by the genes of inflammation, specific to irritation. Conversely, the biomarkers for neuropathic pain are linked only to genes of the GABA system, the neurotransmitters of the central nervous system.

"Now that these epigenetic signatures are clearly defined, a <u>simple blood</u> test will make it possible to define the type of pain the person is suffering from and prescribe the appropriate treatment," says Bertrand Léger. The treatment will thus no longer target the symptoms, but the very root of the problem. And finally, since epigenetics is characterized by the fact that the expression of a gene is durably modified, the right treatment may return it to normal. "We could imagine monitoring the reversion of pain by observing, from an epigenetic point of view, whether the biomarkers return to normal, and adapt the treatment accordingly," concludes Ariane Giacobino.

More information: Ludwig Stenz et al, Genome-wide epigenomic analyses in patients with nociceptive and neuropathic chronic pain subtypes reveals alterations in methylation of genes involved in the neuro-musculoskeletal system, *The Journal of Pain* (2021). DOI: 10.1016/j.jpain.2021.09.001

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