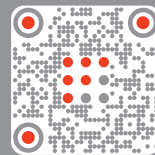




Pro 7 Comprehensive

Seven key areas analyzed to personalize health and wellness recommendations to patients



Register/Order

Analyze seven key areas of health

What are SNPs and what is NutriGenomics?

Single nucleotide polymorphisms (SNPs) are the most frequent type of DNA variation found in humans. Characterization of some SNPs may help predict the risk of developing certain diseases and understand an individual's response to certain foods and drugs.

NutriGenomics studies common genetic variations, such as SNPs, related to unique body responses to nutrition. Everyone is genetically unique when analyzing markers such as these. This allows providers to better personalize medicine by tailoring nutritional treatment to each patient.

Research shows patients are more motivated to follow healthier treatment, diet, and lifestyles when the advice provided is personalized and based on their genetic profile.^{1,2}



The GX Sciences Pro 7 Comprehensive Panel is an innovative, easy to read, comprehensive genetic test focusing on seven key areas of health.

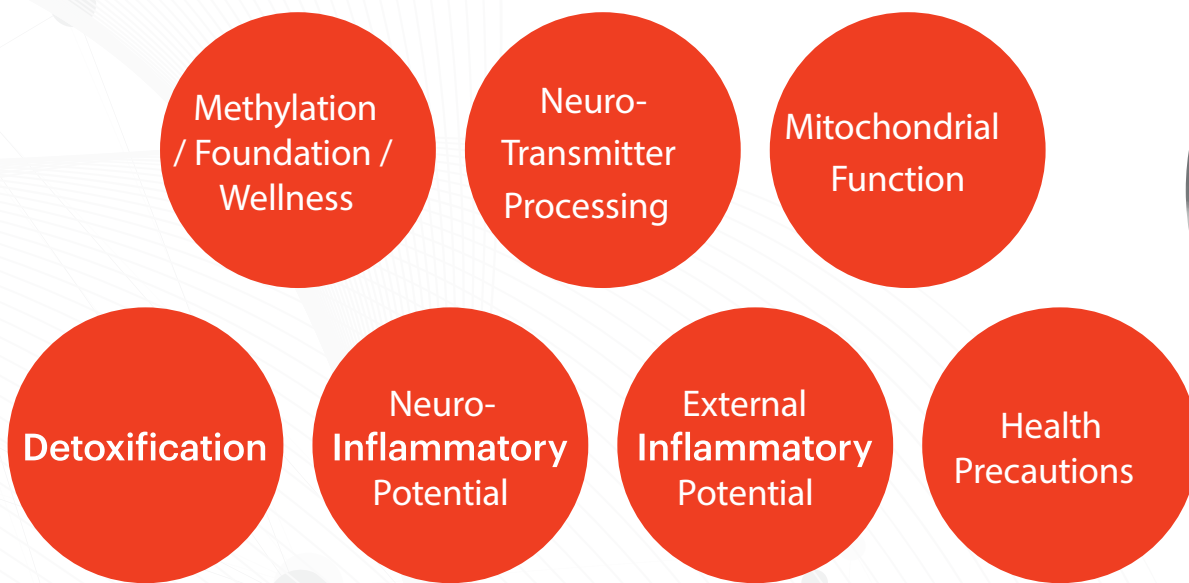


- **Optimize overall health**
Ideal for healthy patients who want to further optimize and maintain their health and wellness.
- **Personalize supplementation**
The report associates each genetic variation, categorizes them, and recommends easy to read therapeutics and formulas based on APIs.
- **Lifestyle and laboratory recommendations**
Easy to read categorical recommendations for the provider to consider in consulting and prescribing.

Personalize health & wellness

Seven categories analyzed

The genetic variations analyzed with Pro 7 are associated with seven key categories of health, hand selected by our experts, that provide nutritional recommendations, health information, and more based on scientific DNA validation of a patient's specific needs in these key areas:



The GX Sciences Pro 7 Comprehensive report includes:



Categorized tables of genetic analysis and explanation



Supplement, prescription, lifestyle, and lab recommendations



Easy to read summary of personalized recommendations

Results are shared securely with the healthcare professional through the GX Sciences online portal.

How it works

Kit contents:

- 1x requisition form
- 1x instructions card
- 1x buccal swab
- 1x ID label sticker
- 1x containment bag
- 1x prepaid return shipping bag

1. Collect the DNA sample (cheek swab) according to the included instructions.

2. Complete the requisition form and any additional instructions. These items must accompany the sample.

3. Submit the sample and forms to the GX Sciences laboratory via prepaid return shipping bag.

4. Results are provided via online portal typically in 5-10 business days after receipt of the sample.

About Us

GX Sciences is an advanced genetic and biomarker testing company/laboratory based in Austin, TX that empowers providers with the most comprehensive Nutrigenomic and Pharmacogenomic testing and blood analysis interpretation platform available in the North American market.

We specialize in:

- Scientifically validating each patient's specific nutritional, laboratory, and lifestyle recommendations.
- Suggested health precautions based on the patient's individual DNA findings.
- Blood analysis via micro-sampling that offers an easy and less painful way for providers to obtain biomarker results, such as Vitamin D levels, with a simple finger prick.
- Customized, secure, and private software specifically designed by and for medical professionals.
- Typical average in-house lab turnaround times of just 5-10 business days.

Together we create the future of personalized medicine.

References

1. Goordazi MO. Lancet Diabetes Endocrinol. 2018;6(3):223-236.
2. Ordovas JM, et al. BMJ. 2018;13;361:bmj.k2173.

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