Resource Summary Report

Generated by <u>dkNET</u> on Apr 16, 2025

SNPALYZE

RRID:SCR_009401 Type: Tool

Proper Citation

SNPALYZE (RRID:SCR_009401)

Resource Information

URL: https://www.dynacom.co.jp/english/product/snpalyze_e/

Proper Citation: SNPALYZE (RRID:SCR_009401)

Description: Software application (entry from Genetic Analysis Software)

Resource Type: software resource, software application

Keywords: gene, genetic, genomic, ms-windows, (98/me/nt4.0/2000/xp)

Funding:

Resource Name: SNPALYZE

Resource ID: SCR_009401

Alternate IDs: nlx_154640

Record Creation Time: 20220129T080252+0000

Record Last Update: 20250416T063546+0000

Ratings and Alerts

No rating or validation information has been found for SNPALYZE.

No alerts have been found for SNPALYZE.

Data and Source Information

Usage and Citation Metrics

We found 74 mentions in open access literature.

Listed below are recent publications. The full list is available at <u>dkNET</u>.

Yamaguchi N, et al. (2024) Autophagy-Related Gene ATG7 Polymorphism Could Potentially Serve as a Biomarker of the Progression of Atrophic Gastritis. Journal of clinical medicine, 13(2).

Chen R, et al. (2024) Association of IL13 polymorphisms with susceptibility to myocardial infarction: A case-control study in Chinese population. PloS one, 19(8), e0308081.

Amano S, et al. (2023) Prediction and association analyses of skin phenotypes in Japanese females using genetic, environmental, and physical features. Skin research and technology : official journal of International Society for Bioengineering and the Skin (ISBS) [and] International Society for Digital Imaging of Skin (ISDIS) [and] International Society for Skin Imaging (ISSI), 29(1), e13231.

Wang D, et al. (2022) Chemerin levels and its genetic variants are associated with Gestational diabetes mellitus: A hospital-based study in a Chinese cohort. Gene, 807, 145888.

Soflaei SS, et al. (2022) Association of Paraoxonase-1 Genotype and Phenotype with Angiogram Positive Coronary Artery Disease. Arquivos brasileiros de cardiologia, 119(4), 593.

Park JW, et al. (2022) Gene Dose-Dependent and Additive Effects of ABCG2 rs2231142 and SLC2A9 rs3733591 Genetic Polymorphisms on Serum Uric Acid Levels. Metabolites, 12(12).

Kondyarpu A, et al. (2021) Association of ISL1 polymorphisms and eosinophilic levels among otitis media patients. Journal of clinical laboratory analysis, 35(3), e23702.

Seo JE, et al. (2021) Association Between CLOCK Gene Variants and Restless Legs Syndrome in Koreans. Psychiatry investigation, 18(11), 1125.

Jeon YJ, et al. (2021) 3'-UTR Polymorphisms in Thymidylate Synthase with Colorectal Cancer Prevalence and Prognosis. Journal of personalized medicine, 11(6).

Bonella F, et al. (2021) Potential clinical utility of MUC5B und TOLLIP single nucleotide polymorphisms (SNPs) in the management of patients with IPF. Orphanet journal of rare diseases, 16(1), 111.

Fukunaga K, et al. (2021) Functional Characterization of the Effects of N-acetyltransferase 2 Alleles on N-acetylation of Eight Drugs and Worldwide Distribution of Substrate-Specific Diversity. Frontiers in genetics, 12, 652704.

Ohwaki A, et al. (2020) Placental Genetic Variants in the Upstream Region of the FLT1 Gene in Pre-eclampsia. Journal of reproduction & infertility, 21(4), 240.

Minn AKK, et al. (2020) Association study of long non-coding RNA HOTAIR rs920778 polymorphism with the risk of cancer in an elderly Japanese population. Gene, 729, 144263.

Oh J, et al. (2020) Association between Five Common Plasminogen Activator Inhibitor-1 (PAI-1) Gene Polymorphisms and Colorectal Cancer Susceptibility. International journal of molecular sciences, 21(12).

Kalantari S, et al. (2019) Single and multi-locus association study of TCF7L2 gene variants with susceptibility to type 2 diabetes mellitus in an Iranian population. Gene, 696, 88.

Matsusue A, et al. (2019) VNTR polymorphism in the monoamine oxidase A promoter region and cerebrospinal fluid catecholamine concentrations in forensic autopsy cases. Neuroscience letters, 701, 71.

Kobayashi H, et al. (2019) Adiponectin Receptor gene Polymorphisms are Associated with Kidney Function in Elderly Japanese Populations. Journal of atherosclerosis and thrombosis, 26(4), 328.

An HJ, et al. (2019) 3'-UTR Polymorphisms in the Vascular Endothelial Growth Factor Gene (VEGF) Contribute to Susceptibility to Recurrent Pregnancy Loss (RPL). International journal of molecular sciences, 20(13).

Ramezani S, et al. (2019) Association of PARP1 rs4653734, rs907187 and rs1136410 variants with breast cancer risk among Iranian women. Gene, 712, 143954.

Furukawa H, et al. (2018) Independent association of HLA-DPB1*02:01 with rheumatoid arthritis in Japanese populations. PloS one, 13(9), e0204459.