

Resource Summary Report

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Binding and Expression Target Analysis

RRID:SCR_005396

Type: Tool

Proper Citation

Binding and Expression Target Analysis (RRID:SCR_005396)

Resource Information

URL: <http://cistrome.org/BETA/>

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Description: A software package that integrates ChIP-seq of transcription factors or chromatin regulators with differential gene expression data to infer direct target genes. BETA has three functions: (1) to predict whether the factor has activating or repressive function; (2) to infer the factor's target genes; and (3) to identify the motif of the factor and its collaborators which might modulate the factor's activating or repressive function. BETA requires ~2GB RAM and 1h for the whole procedure. BETA may run on the web server at Cistrome or may be downloaded.

Abbreviations: BETA

Synonyms: BETA - Binding and Expression Target Analysis

Resource Type: production service resource, service resource, data analysis service, analysis service resource, software resource

Defining Citation: [PMID:24263090](#)

Keywords: transcription factor, chromatin regulator, transcriptome, chip-seq, cistrome, gene expression, target gene, motif, differential gene expression

Availability: Registration required, Open unspecified license

Resource Name: Binding and Expression Target Analysis

Resource ID: SCR_005396

Alternate IDs: OMICS_00515

Record Creation Time: 20220129T080230+0000

Record Last Update: 20260821T042637+0000

Ratings and Alerts

No rating or validation information has been found for Binding and Expression Target Analysis.

No alerts have been found for Binding and Expression Target Analysis.

Data and Source Information

Source: [SciCrunch Registry](#)

Usage and Citation Metrics

We found 455 mentions in open access literature.

Listed below are recent publications. The full list is available at [NIF](#) .

Foidart P, et al. (2026) Whole-genome doubling drives immune evasion by silencing antigen presentation. *Cancer cell*, 44(7), 1418.

Wuergezhen D, et al. (2025) An eGFP-Col4a2 mouse model reveals basement membrane dynamics underlying hair follicle morphogenesis. *The Journal of cell biology*, 224(2).

Parihar AK, et al. (2025) Adaptive responses of large-seeded lentils across diverse Indian climates. *Heliyon*, 11(3), e42184.

Vora M, et al. (2025) Genome-wide analysis of Smad and Schnurri transcription factors in *C. elegans* demonstrates widespread interaction and a function in collagen secretion. *eLife*, 13.

Li B, et al. (2025) Genetic insights into cardiac conduction disorders from genome-wide association studies. *Human genomics*, 19(1), 20.

Chambers TL, et al. (2025) Methylome-proteome integration after late-life voluntary exercise training reveals regulation and target information for improved skeletal muscle health. *The Journal of physiology*, 603(1), 211.

Citu C, et al. (2025) Identification and catalog of viral transcriptional regulators in human diseases. *iScience*, 28(3), 112081.

Tang Q, et al. (2025) p63 co-opts the skin Krt8-to-Krt5 transition for enamel organ development. *bioRxiv : the preprint server for biology*.

Harvey C, et al. (2025) Evaluation of a biomarker for amyotrophic lateral sclerosis derived from a hypomethylated DNA signature of human motor neurons. *BMC medical genomics*, 18(1), 10.

Londono-Correa D, et al. (2025) Crystallized and fluid cognitive abilities have different genetic associations with neuropsychiatric disorders. *Research square*.

Houten R, et al. (2025) Digital Versus Paper-Based Consent from the UK NHS Perspective: A Micro-costing Analysis. *PharmacoEconomics - open*, 9(1), 27.

Wei Q, et al. (2025) Enhancing the performance of SSVEP-based BCIs by combining task-related component analysis and deep neural network. *Scientific reports*, 15(1), 365.

Lai Y, et al. (2025) Transcriptomic analysis reveals the function of m6A regulators in aged cochlea. *Brazilian journal of otorhinolaryngology*, 91(3), 101578.

Li Z, et al. (2024) EstroGene2.0: A multi-omic database of response to estrogens, ER-modulators, and resistance to endocrine therapies in breast cancer. *bioRxiv : the preprint server for biology*.

Ismaeel A, et al. (2024) Coordinated Regulation of Myonuclear DNA Methylation, mRNA, and miRNA Levels Associates With the Metabolic Response to Rapid Synergist Ablation-Induced Skeletal Muscle Hypertrophy in Female Mice. *Function (Oxford, England)*, 5(1), zqad062.

Liu M, et al. (2024) Mapping the causal associations of cytokines with sarcopenia and aging traits: Evidence from bidirectional Mendelian randomization. *Journal of cachexia, sarcopenia and muscle*, 15(3), 1121.

Gu W, et al. (2024) A MTA2-SATB2 chromatin complex restrains colonic plasticity toward small intestine by retaining HNF4A at colonic chromatin. *Nature communications*, 15(1), 3595.

Mao G, et al. (2024) Unveiling the bioinformatic genes and their involved regulatory mechanisms in type 2 diabetes combined with osteoarthritis. *Frontiers in immunology*, 15, 1353915.

DiCiaccio B, et al. (2024) ZBTB7A is a modulator of KDM5-driven transcriptional networks in basal breast cancer. *Cell reports*, 43(12), 114991.

Valenzuela-Fuenzalida JJ, et al. (2024) The Association between Anatomical Variants of Musculoskeletal Structures and Nerve Compressions of the Lower Limb: A Systematic Review and Meta-Analysis. *Diagnostics (Basel, Switzerland)*, 14(7).