## **Resource Summary Report**

Generated by NIF on Apr 22, 2025

# **NetworkX**

RRID:SCR\_016864

Type: Tool

## **Proper Citation**

NetworkX (RRID:SCR\_016864)

#### **Resource Information**

URL: https://networkx.github.io/

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**Description:** Software Python package for the creation, manipulation, and study of the structure, dynamics, and functions of complex networks.

**Resource Type:** software resource, data analysis software, network analysis software, data processing software, software application

Keywords: create, analysis, structure, dynamic, function, complex, network, data

**Funding:** 

Availability: Free, Available for download, Freely available

Resource Name: NetworkX

Resource ID: SCR\_016864

Alternate URLs: https://github.com/networkx/networkx

License: 3-clause BSD license

**Record Creation Time:** 20220129T080332+0000

Record Last Update: 20250422T055951+0000

## **Ratings and Alerts**

No rating or validation information has been found for NetworkX.

No alerts have been found for NetworkX.

#### **Data and Source Information**

Source: SciCrunch Registry

## **Usage and Citation Metrics**

We found 105 mentions in open access literature.

**Listed below are recent publications.** The full list is available at NIF.

Wei L, et al. (2024) Systems-level reconstruction of kinase phosphosignaling networks regulating endothelial barrier integrity using temporal data. NPJ systems biology and applications, 10(1), 134.

Skorupka A, et al. (2024) Detecting anomalies in graph networks on digital markets. PloS one, 19(12), e0315849.

Vishwanathan A, et al. (2024) Predicting modular functions and neural coding of behavior from a synaptic wiring diagram. Nature neuroscience, 27(12), 2443.

Nilsen AS, et al. (2024) Exploring effects of anesthesia on complexity, differentiation, and integrated information in rat EEG. Neuroscience of consciousness, 2024(1), niae021.

Wei L, et al. (2024) Systems-level reconstruction of kinase phosphosignaling networks regulating endothelial barrier integrity using temporal data. bioRxiv: the preprint server for biology.

Sadowski M, et al. (2024) Characterizing the genetic architecture of drug response using gene-context interaction methods. Cell genomics, 4(12), 100722.

Brempou D, et al. (2024) Using parenclitic networks on phaeochromocytoma and paraganglioma tumours provides novel insights on global DNA methylation. Scientific reports, 14(1), 29958.

Köhler CA, et al. (2024) Facilitating the Sharing of Electrophysiology Data Analysis Results Through In-Depth Provenance Capture. eNeuro, 11(6).

Wei L, et al. (2024) Using machine learning to dissect host kinases required for Leishmania internalization and development. bioRxiv: the preprint server for biology.

Xie X, et al. (2024) IL-1?-induced epithelial cell and fibroblast transdifferentiation promotes neutrophil recruitment in chronic rhinosinusitis with nasal polyps. Nature communications, 15(1), 9101.

Newsham I, et al. (2024) Early detection and diagnosis of cancer with interpretable machine learning to uncover cancer-specific DNA methylation patterns. Biology methods & protocols, 9(1), bpae028.

Gupta M, et al. (2023) Critical mechanistic features of HIV-1 viral capsid assembly. Science advances, 9(1), eadd7434.

Fan P, et al. (2023) Combination of antidepressants and antipsychotics as a novel treatment option for psychosis in Alzheimer's disease. CPT: pharmacometrics & systems pharmacology, 12(8), 1119.

Lu M, et al. (2023) Effect of intracranial electrical stimulation on dynamic functional connectivity in medically refractory epilepsy. Frontiers in human neuroscience, 17, 1295326.

Nanda P, et al. (2023) Multicellular growth as a dynamic network of cells. bioRxiv: the preprint server for biology.

Wu CG, et al. (2023) Extended regulation interface coupled to the allosteric network and disease mutations in the PP2A-B56? holoenzyme. bioRxiv: the preprint server for biology.

Wright SN, et al. (2023) Genome-wide association studies of human and rat BMI converge on synapse, epigenome, and hormone signaling networks. Cell reports, 42(8), 112873.

Iranzo J, et al. (2023) Protocol for comparing gene-level selection on coding mutations between two groups of samples with Coselens. STAR protocols, 4(1), 102117.

Collier M, et al. (2022) Pathogen transmission modes determine contact network structure, altering other pathogen characteristics. Proceedings. Biological sciences, 289(1989), 20221389.

Domingo-Fernández D, et al. (2022) Causal reasoning over knowledge graphs leveraging drug-perturbed and disease-specific transcriptomic signatures for drug discovery. PLoS computational biology, 18(2), e1009909.