

Resource Summary Report

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Stanford University Human Immune Monitoring Center Core Facility

RRID:SCR_018266

Type: Tool

Proper Citation

Stanford University Human Immune Monitoring Center Core Facility (RRID:SCR_018266)

Resource Information

URL: <http://himc.stanford.edu>

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Description: Core designed for immune monitoring services for clinical and translational studies. Goals include providing standardized, state-of-the art immune monitoring assays at RNA, protein, and cellular level, testing and developing new technologies for immune monitoring, archive, report, and mine data from immune monitoring studies. HIMC uses online database for integration of data from standard HIMC assays, along with de-identified clinical and demographic data.

Abbreviations: HIMC

Synonyms: Stanford University - Human Immune Monitoring Center, Human Immune Monitoring Center (HIMC)

Resource Type: access service resource, core facility, service resource, data or information resource

Keywords: Immune monitoring service, clinical study, translational study, RNA, protein, data, core facility, USEdit, ABRF

Availability: Open

Resource Name: Stanford University Human Immune Monitoring Center Core Facility

Resource ID: SCR_018266

Alternate IDs: ABRF_399

Alternate URLs: <https://coremarketplace.org/?FacilityID=399>

Record Creation Time: 20220129T080339+0000

Record Last Update: 20260728T043549+0000

Ratings and Alerts

No rating or validation information has been found for Stanford University Human Immune Monitoring Center Core Facility.

No alerts have been found for Stanford University Human Immune Monitoring Center Core Facility.

Data and Source Information

Source: [SciCrunch Registry](#)

Usage and Citation Metrics

We found 1 mentions in open access literature.

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

Mehl LC, et al. (2025) Environmental factors contribute to cancer therapy toxicity. bioRxiv : the preprint server for biology.