

Resource Summary Report

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Cancer Research Institute IFOM CPM Cellular And Preclinical Models Core Facility

RRID:SCR_026864

Type: Tool

Proper Citation

Cancer Research Institute IFOM CPM Cellular And Preclinical Models Core Facility
(RRID:SCR_026864)

Resource Information

URL: <https://www.ifom.eu/en/cancer-research/technological-units/preclinical-models.php>

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Description: Specialized laboratory designed for growth, maintenance, and experimentation of living cells under controlled conditions. This facility is used in various fields such as biomedical research, drug development, and supports the culture of primary cells, immortalized cell lines, human-induced pluripotent stem cells (hiPSCs), and organoids, among others.

Synonyms: IFOM CPM - Cellular And Preclinical Models Core Facility

Resource Type: access service resource, core facility, service resource

Keywords: ABRF, living cells, primary cells, immortalized cell lines, human-induced pluripotent stem cells (hiPSCs), organoids

Resource Name: Cancer Research Institute IFOM CPM Cellular And Preclinical Models Core Facility

Resource ID: SCR_026864

Alternate IDs: ABRF_3222

Alternate URLs: <https://coremarketplace.org/?FacilityID=3222&citation=1>

Record Creation Time: 20250430T055337+0000

Record Last Update: 20260829T183452+0000

Ratings and Alerts

No rating or validation information has been found for Cancer Research Institute IFOM CPM Cellular And Preclinical Models Core Facility.

No alerts have been found for Cancer Research Institute IFOM CPM Cellular And Preclinical Models 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 [dkNET](#) .

di Lillo A, et al. (2026) Site-specific DNA double-strand break induces local transcription in cis and protein expression. Communications biology, 9(1).