

Produktinformation



Forschungsprodukte & Biochemikalien
Zellkultur & Verbrauchsmaterial
Diagnostik & molekulare Diagnostik
Laborgeräte & Service

Weitere Information auf den folgenden Seiten! See the following pages for more information!



Lieferung & Zahlungsart siehe unsere Liefer- und Versandbedingungen

Zuschläge

- Mindermengenzuschlag
- Trockeneiszuschlag
- Gefahrgutzuschlag
- Expressversand

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Datasheet for WM2090-01-0005 WM2090 Viable Cells

Overview

| Description: | WM2090 Viable Cells - WM2090-01-0005 |
|--------------|--------------------------------------|
| Item No.: | WM2090-01-0005 |
| Size: | 5 x 1 million cells |
| Origin: | Human |

Product Details

| Background: | WM2090 is a metastatic human melanoma cell line. This cell line features the specific V600E (Val600Glu) mutation at codon 600 in the BRAF gene. This mutation causes constitutively active kinase activity and activation of MEK and ERK signaling pathway. WM2090 cells produce xenograft tumors when injected into immunocompromised mice. |
|--------------------|--|
| Synonyms: | Melanoma, patient derived tumor, tumor models, skin cancer, xenograft |
| Species of Origin: | Human |

Target Details

| Purity/Specificity: | Cells are sterile, validated by short tandem repeat profiling, and are tested as negative for mycoplasma. It is recommended that cell lines are tested for mycoplasma contamination and short tandem repeat (STR) profiling every 10 passages or each time a frozen seed stock is made. See cell culture protocol for additional details. |
|---------------------|---|
| Relevant Links: | Cell Line EULA |
| | Melanoma Cell Culture Protocol |

Application Details

| Application Note: | The key applications of these cell lines include genetic studies, xenograft production, drug testing, and drug target discovery. These cell line models can be used in various biological assays, and for identifying critical target genes, and cell signaling pathways. |
|-------------------|---|
| Assay Dilutions: | All assays should be optimized by the user. Recommended dilutions (if any) may be listed below. |



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Cell Line Data

| Cell Line: | Human Melanoma |
|-----------------|---|
| Product Type: | Viable Cells |
| Cell Viability: | Yes |
| Stage: | Metastasis |
| BRAF: | V600E |
| CDK4: | WT |
| C-Kit: | WT |
| N-RAS: | WT |
| PTEN: | WT |
| Paired: | No |
| Medium: | Tumor Specialized Media with 2% HI-FBS |
| Sub-culture: | Cells should be maintained between 30 – 95% confluence in tumor specialized medium with 2% FBS; split cultures 1:6 every 5 days using 0.25% trypsin/EDTA. |
| Incubation: | 36°C with 5% CO2 |

Formulation

| Physical State: | Frozen Cell Suspension |
|-----------------|---|
| Concentration: | 1.0 million cells/mL Count By Hemocytometer |
| Buffer: | None |
| Preservative: | None |
| Stabilizer: | None |
| | |

Shipping & Handling

| Shipping Condition: | Dry Ice |
|---------------------|---|
| Storage Condition: | Cells are frozen with 90% FBS/10% DMSO solution at about 5x10^6 cells/ml. Store vial in liquid nitrogen upon arrival. |
| Expiration: | Expiration date is two (2) years from date of receipt. |

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Images



Flask

Human melanoma tumor cells with known gene mutations, disease stage, STR, and RPPA profiling

Disclaimer

No test method can provide total assurance that the hepatitis B virus, hepatitis C virus, human immunodeficiency virus, or any other infectious agents are absent. Thus, all blood products, including purified proteins derived from human blood sources, should be handled at Biosafety Level 2 as recommended by the CDC\NIH manual entitled Biosafety in Microbiological and Biomedical Laboratories for potentially infectious human serum, blood specimens or proteins derived from same. Source material for the human blood product supplied to your facility has been tested for the detection of HIV antibody, Hepatitis B surface antigen, antibody to Hepatitis C, HIV 1 antigen(s), antibody to HTLV - I/II, and syphilis by FDA guidelines. All units were found to be nonreactive/negative for these tests. All human blood source material is collected in FDA licensed centers and is tested with FDA approved test kits.

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