



SZABO SCANDIC

Part of Europa Biosite

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

SZABO-SCANDIC Handels GmbH

Quellenstraße 110, A-1100 Wien

T. +43(0)1 489 3961-0

F. +43(0)1 489 3961-7

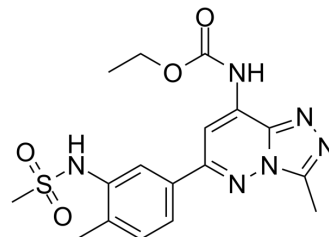
mail@szabo-scandic.com

www.szabo-scandic.com

[linkedin.com/company/szaboscandic](https://www.linkedin.com/company/szaboscandic) 

Bromosporine

Cat. No.:	HY-15815
CAS No.:	1619994-69-2
Molecular Formula:	C ₁₇ H ₂₀ N ₆ O ₄ S
Molecular Weight:	404.44
Target:	Epigenetic Reader Domain; Apoptosis; CDK; HIV
Pathway:	Epigenetics; Apoptosis; Cell Cycle/DNA Damage; Anti-infection
Storage:	Powder -20°C 3 years 4°C 2 years In solvent -80°C 2 years -20°C 1 year



SOLVENT & SOLUBILITY

In Vitro

DMSO : ≥ 51.7 mg/mL (127.83 mM)
 * "≥" means soluble, but saturation unknown.

	Solvent Concentration	Mass	1 mg	5 mg	10 mg
Preparing Stock Solutions	1 mM		2.4726 mL	12.3628 mL	24.7255 mL
	5 mM		0.4945 mL	2.4726 mL	4.9451 mL
	10 mM		0.2473 mL	1.2363 mL	2.4726 mL

Please refer to the solubility information to select the appropriate solvent.

In Vivo

- Add each solvent one by one: 50% PEG300 >> 50% saline
Solubility: 10 mg/mL (24.73 mM); Suspended solution; Need ultrasonic
- Add each solvent one by one: 10% DMSO >> 90% corn oil
Solubility: ≥ 2.5 mg/mL (6.18 mM); Clear solution

BIOLOGICAL ACTIVITY

Description

Bromosporine is a potent BET inhibitor with an IC₅₀ value of 2.1 μM for PCAF. Bromosporine can arrest cell cycle and induce apoptosis in cancer cells. Bromosporine exhibits excellent antitumor activity in xenograft mice model when combined with [5-FU](#) (HY-90006). Bromosporine can increase CDK9 T-loop phosphorylation in HIV-1 latency models, resulting the protection of reactivate HIV-1 replication from latency. Bromosporine can be used to research colorectal cancer, acute myeloid leukemia (AML) and AIDS^{[1][2][3][4]}.

IC₅₀ & Target

IC₅₀: 2.1 μM (PCAF)^[2]
 Apoptosis, BET, CDK9, HIV-1^{[1][3][4]}

In Vitro

Bromosporine (0-1000 nM; 72 h) synergistically inhibits cell growth in CRC cells with [5-FU](#) (HY-90006)^[1].

Bromosporine (various concentration; 48 h) causes a distinct increase in the cells arrested at G1 phase when combined with [5-FU](#)^[1].

Bromosporine (various concentration; 48 h) decreases the expressions of PARP, caspase 3, and 9^[1].

Bromosporine (0.1, 0.5 and 1 μ M; 6-10 days) inhibits AML cells in a dose-dependent manner^[3].

Bromosporine (2.5 μ M; 72 h) activates HIV-1 replication in vitro in latent HIV-1 J-Lat clone C11 cells^[4].

Bromosporine (1-50 μ M; 48 h) does not induce marked toxicity in primary CD4+ T cells^[4].

MCE has not independently confirmed the accuracy of these methods. They are for reference only.

Cell Proliferation Assay^[1]

Cell Line:	HCT116 and HT29
Concentration:	0, 30, 60, 120, 240, 480 and 1000 nM
Incubation Time:	72 h
Result:	Synergistically inhibited cell growth in CRC cells with 5-FU (HY-90006) (0-16 μ g/mL) and exhibited a dose-dependent manner.

Cell Cycle Analysis^[1]

Cell Line:	HCT116 and HT29
Concentration:	Various concentration
Incubation Time:	48 h
Result:	Caused a distinct increase in the cells arrested at G1 phase when combined with 5-FU (HY-90006).

Western Blot Analysis^[1]

Cell Line:	HCT116 and HT29
Concentration:	Various concentration
Incubation Time:	48 h
Result:	Elevated the level of apoptosis in both cell lines through cleavage of PARP, caspase 3, and 9.

Cell Proliferation Assay^[3]

Cell Line:	MV4;11, KASUMI-1, OCI-AML3 and K562
Concentration:	0.1, 0.5 and 1 μ M
Incubation Time:	6-10 days
Result:	Inhibited these AML cells in a dose-dependent manner.

Cell Cytotoxicity Assay^[4]

Cell Line:	PBMCs
Concentration:	1 μ M, 2.5 μ M, 5 μ M, 10 μ M, 25 μ M and 50 μ M
Incubation Time:	48 h

	<table> <tr> <td>Result:</td><td>Did not induce marked toxicity in primary CD4+ T cells with CC₅₀ over 10 μM.</td></tr> </table>	Result:	Did not induce marked toxicity in primary CD4+ T cells with CC ₅₀ over 10 μM.						
Result:	Did not induce marked toxicity in primary CD4+ T cells with CC ₅₀ over 10 μM.								
In Vivo	Bromosporine (100 mg/kg; i.p.; daily for 10 days) shows better antitumor activity than individual when co-treated with 5-FU (HY-90006)^[1]. MCE has not independently confirmed the accuracy of these methods. They are for reference only. <table> <tr> <td>Animal Model:</td><td>Female BALB/c nude mice (5-6 weeks; injected with 1 × 10⁶ cells/100 μL of HT116 cells)^[1]</td></tr> <tr> <td>Dosage:</td><td>100 mg/kg</td></tr> <tr> <td>Administration:</td><td>i.p.; daily for 10 days</td></tr> <tr> <td>Result:</td><td>Exhibited better antitumor activity than individual Bromosporine or 5-FU (HY-90006) when co-treated with the two agent.</td></tr> </table>	Animal Model:	Female BALB/c nude mice (5-6 weeks; injected with 1 × 10 ⁶ cells/100 μL of HT116 cells) ^[1]	Dosage:	100 mg/kg	Administration:	i.p.; daily for 10 days	Result:	Exhibited better antitumor activity than individual Bromosporine or 5-FU (HY-90006) when co-treated with the two agent.
Animal Model:	Female BALB/c nude mice (5-6 weeks; injected with 1 × 10 ⁶ cells/100 μL of HT116 cells) ^[1]								
Dosage:	100 mg/kg								
Administration:	i.p.; daily for 10 days								
Result:	Exhibited better antitumor activity than individual Bromosporine or 5-FU (HY-90006) when co-treated with the two agent.								

CUSTOMER VALIDATION

- Proc Natl Acad Sci U S A. 2019 Feb 19;116(8):2961-2966.
- Biochem Pharmacol. 2020 Jul;177:113946.
- Patent. US20180263995A1.

See more customer validations on www.MedChemExpress.com

REFERENCES

- [1]. Cheng X, et al. BET inhibitor bromosporine enhances 5-FU effect in colorectal cancer cells. Biochem Biophys Res Commun. 2020 Jan 22;521(4):840-845.
- [2]. El-Shershaby MH, et al. From triazolophthalazines to triazoloquinazolines: A bioisosterism-guided approach toward the identification of novel PCAF inhibitors with potential anticancer activity. Bioorg Med Chem. 2021 Jul 15;42:116266.
- [3]. Picaud S, et al. Promiscuous targeting of bromodomains by bromosporine identifies BET proteins as master regulators of primary transcription response in leukemia. Sci Adv. 2016 Oct 12;2(10):e1600760.
- [4]. Pan H, et al. The bromodomain and extraterminal domain inhibitor bromosporine synergistically reactivates latent HIV-1 in latently infected cells. Oncotarget. 2017 Oct 6;8(55):94104-94116.

Caution: Product has not been fully validated for medical applications. For research use only.

Tel: 609-228-6898

Fax: 609-228-5909

E-mail: tech@MedChemExpress.com

Address: 1 Deer Park Dr, Suite Q, Monmouth Junction, NJ 08852, USA