

Product Name: Temozolomide Revision Date: 12/07/2022

## **Product Data Sheet**

# **Temozolomide**

Cat. No.: B1399

CAS No.: 85622-93-1
Formula: C6H6N6O2
M.Wt: 194.15

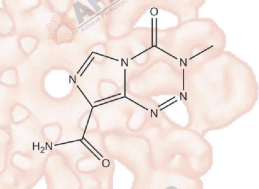
Synonyms:

Target: DNA Damage/DNA Repair

Pathway: DNA Alkylating

Storage: Store at -20°C, sealed storage, away

from moisture and light



## Solvent & Solubility

insoluble in EtOH; insoluble in H2O; ≥29.61 mg/mL in DMSO

In Vitro

Preparing Stock Solutions	Solvent  Concentration	1mg	5mg	10mg
	1 mM	5.1507 mL	25.7533 mL	51.5066 mL
	5 mM	1.0301 mL	5.1507 mL	10.3013 mL
	10 mM	0.5151 mL	2.5753 mL	5.1507 mL

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

# **Biological Activity**

Shortsummary	DNA methylating, chemotherapeutic agent			
IC <sub>50</sub> & Target		SI Control of the Con		
	Cell Viability Assay			
	Cell Line:	SK-LMS-1 leiomyosarcoma (MGMT-/p53+), Ewing sarcoma A-673 and		
	those benediction	GIST-T1 (both lines with MGMT+/p53- phenotype), and glioblastoma T98G		
In Vitro		(MGMT+/p53+)		
	Preparation method:	The solubility of this compound in DMSO is >6.6mg/mL. General tips for		
		obtaining a higher concentration: Please warm the tube at 37°C for 10 minutes		
		and/or shake it in the ultrasonic bath for a while. Stock solution can be stored		

		below -20°C for several months.		
		Delow -20 C for Several months.		
	Reacting conditions:	62.5, 125, 250 and 500 μM; 72 h		
	Applications:	In SK-LMS-1 cells, Temozolomide inhibited proliferative activity of SK-LMS-1		
	E La	cells. A-673 cells was most sensitive to temozolomide, the effect was time- and		
		dose-dependent. Preincubation of Ewing sarcoma cells with O6-benzylguanine		
		potentiated the cytotoxic effect of the alkylating agent and reduced viability of		
	chare perfective	tumor cells. GIST-T1 cells were insensitive to temozolomide.		
	Animal experiment			
	Animal models:	PARP1 wild - type (WT) and PARP1 knock - out (KO) mice		
	Dosage form:	68 mg/kg; once daily for 5 days; orally administrated		
	Applications:	In PARP1 WT mice, temozolomide significantly lowered concentrations of		
		NAD+ in the liver when compared with the control group (by 22%, p = 0.02). In		
In Vivo		the livers of PARP1 KO mice, there was also a statistically significant reduction		
	40.	in NAD+ in the temozolomide - only group when compared with the control (by		
	the Unitrovin	22%, p = 0.03).		
	Other notes:	Please test the solubility of all compounds indoor, and the actual solubility may		
	Activie fee	slightly differ with the theoretical value. This is caused by an experimental		
		system error and it is normal.		

### **Product Citations**

See more customer validations on www.apexbt.com.

### References

[1] Khusnutdinov RR1, Boichuk SV2. Mechanisms of Sensitivity of Soft Tissue Sarcoma Cells to Temozolomide. Bull Exp Biol Med. 2017 Jul 18.

[2]. Almeida GS1, Bawn CM1, Galler M1, et al. PARP inhibitor rucaparib induces changes in NAD levels in cells and liver tissues as assessed by MRS. NMR Biomed. 2017 May 22.

### Caution

#### FOR RESEARCH PURPOSES ONLY.

#### NOT FOR HUMAN, VETERINARY DIAGNOSTIC OR THERAPEUTIC USE.

Specific storage and handling information for each product is indicated on the product datasheet. Most APExBIO products are stable under the recommended conditions. Products are sometimes shipped at a temperature that differs from the recommended storage temperature. Shortterm storage of many products are stable in the short-term at temperatures that differ from that required for long-term storage. We ensure that the product is shipped under conditions that will maintain the quality of the reagents. Upon receipt www.apexbt.com



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