biotechne[®] TOCRIS

Print Date: Mar 5th 2024

Certificate of Analysis

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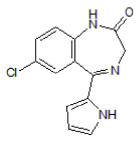
Product Name: Ro 5-3335

CAS Number: 30195-30-3

IUPAC Name: 7-Chloro-1,3-dihydro-5-(1*H*-pyrrol-2-yl)-2*H*-1,4-benzodiazepin-2-one

1. PHYSICAL AND CHEMICAL PROPERTIES

Batch Molecular Formula: Batch Molecular Weight: Physical Appearance: Solubility: Storage: Batch Molecular Structure: C₁₃H₁₀CIN₃O. 259.69 Off White solid DMSO to 100 mM Store at +4°C



2. ANALYTICAL DATA

HPLC:	Shows 99.9% purity				
¹ H NMR:	Consistent with structure				
Mass Spectrum:	Consistent with structure				
Microanalysis:	Carbon Hydrogen Nitrogen				
	Theoretical 60.13 3.88 16.18				

Found

Caution - Not Fully Tested • Research Use Only • Not For Human or Veterinary Use

60.08

3.87

bio-techne.com	North America	China	Europe Middle East Africa	Rest of World
info@bio-techne.com techsupport@bio-techne.com	Tel: (800) 343 7475	info.cn@bio-techne.com Tel: +86 (21) 52380373	Tel: +44 (0)1235 529449	www.tocris.com/distributors Tel:+1 612 379 2956

Catalog No.: 4694

16.11

Batch No.: 4

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Description:

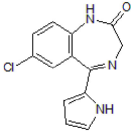
Ro 5-3335 is a core binding factor (CBF) inhibitor; preferentially kills human leukemia cell lines with CBF fusion proteins (IC₅₀ = 1.1 μ M). Represses RUNX1/CBF β -dependent transactivation in reporter assays and inhibits transcriptional regulation by RUNX1 and CBF β . Reduces leukemia burden in a mouse model. Attenuates RUNX1-dependent hematopoiesis in zebrafish embryos. Also a Tat antagonist; inhibits HIV-1 replication in vitro.

Physical and Chemical Properties:

Batch Molecular Formula: C₁₃H₁₀ClN₃O. Batch Molecular Weight: 259.69 Physical Appearance: Off White solid

Minimum Purity: ≥98%

Batch Molecular Structure:



References:

NH

Cunningham *et al* (2012) Identification of benzodiazepine Ro5-3335 as an inhibitor of CBF leukemia through quantitative high throughput screen against RUNX1-CBFß interaction. Proc.Natl.Acad.Sci.USA **109** 14592. PMID: 22912405.

Okuda et al (2001) RUNX1/AML1: a central player in hematopoiesis. Int. J. Hematol. 74 252. PMID: 11721959.

Cupelli *et al* (1995) The human immunodeficiency virus type 1 Tat antagonist, Ro 5-3335, predominantly inhibits transcription initiation from the viral promoter. J.Virol. **69** 2640. PMID: 7884917.

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4

Storage: Store at +4°C

Solubility & Usage Info: DMSO to 100 mM

Stability and Solubility Advice:

Some solutions can be difficult to obtain and can be encouraged by rapid stirring, sonication or gentle warming (in a 45-60°C water bath).

Information concerning product stability, particularly in solution, has rarely been reported and in most cases we can only offer a general guide. *Unless contradicted by product-specific protocols or instructions, our standard recommendations apply:

SOLIDS: Provided storage is as stated on the product label and the vial is kept tightly sealed, the product can be stored for up to 6 months from date of receipt.

SOLUTIONS: We recommend that stock solutions, once prepared, are stored aliquoted in tightly sealed vials at -20°C or below and used within 1 month. Wherever possible solutions should be made up and used on the same day.