Butyrolactone I

CDK inhibitor

Cell permeable and selective inhibitor of the cyclin-dependent kinases CDK1/cyclin B, CDK2 and CDK5. Inhibits cell cycle progression at the G1/S and G2/M transitions. G1/S and G2/M transitions, in W138 cells, were inhibited, concurrent with inhibition of the phosphorylations of, respectively, retinoblastoma protein and histone H1. It is an important tool for probing the cellular roles of CDKs. Blocks Fas-induced apoptosis. Displays antitumor activity.

Citations: 15

View Online »

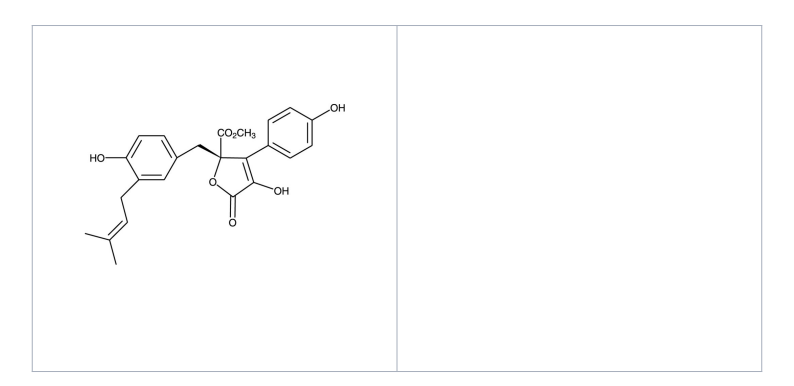
Ordering Information

Order Online »

BML-CC210-1000	1mg
BML-CC210-0200	200µg

Manuals, SDS & CofA

View Online »



Handling & Storage

Use/Stability As indicated on product label or CoA when stored as recommended. Stable for at least

1 year after receipt when stored, as supplied, at -20°C. Stock solutions are stable for up

to 3 months at -20°C.

Handling Protect from light.

Long Term Storage -20°C

Shipping Blue Ice

Regulatory Status RUO - Research Use Only

Product Details

Appearance White solid.

CAS 87414-49-1

Couple Target CDK

Couple Type Inhibitor

Formula $C_{24}H_{24}O_7$

MW 424.5

Purity ≥99% (TLC)

Solubility Soluble in 100% ethanol (25mg/ml) or DMSO(>10mg/ml).

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European Sales Office