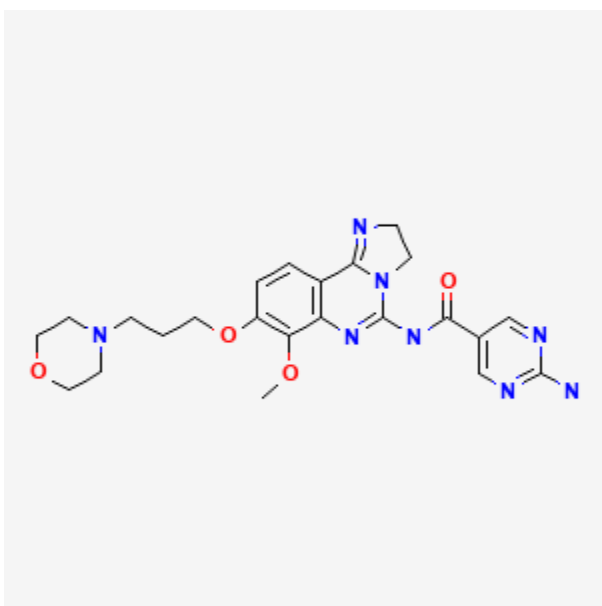




Copanlisib

Revised: November 15, 2023.

CASRN: 1032568-63-0



Drug Levels and Effects

Summary of Use during Lactation

Copanlisib has been removed from the US market. No information is available on the clinical use of copanlisib during breastfeeding. Because copanlisib's half-life is about 39 hours, it might accumulate in the infant. The manufacturer recommends that breastfeeding be discontinued during copanlisib therapy and for 1 month after the last dose.

Drug Levels

Maternal Levels. Relevant published information was not found as of the revision date.

Infant Levels. Relevant published information was not found as of the revision date.

Disclaimer: Information presented in this database is not meant as a substitute for professional judgment. You should consult your healthcare provider for breastfeeding advice related to your particular situation. The U.S. government does not warrant or assume any liability or responsibility for the accuracy or completeness of the information on this Site.

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Effects in Breastfed Infants

Relevant published information was not found as of the revision date.

Effects on Lactation and Breastmilk

Relevant published information was not found as of the revision date.

Substance Identification

Substance Name

Copanlisib

CAS Registry Number

1032568-63-0

Drug Class

Breast Feeding

Lactation

Milk, Human

Antineoplastic Agents

Enzyme Inhibitors

Signal Transduction Inhibitors