

Data Sheet

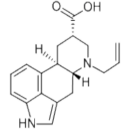
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Product Name :Cabergoline EP Impurity F

 $\begin{array}{lll} \textbf{Cat.No.} & : URK-V2464 \\ \textbf{CAS No.} & :86891\text{-}15\text{-}8 \\ \textbf{Molecular Formula} & : C_{18}H_{20}N_2O_2 \\ \textbf{Molecular Weight} & :296.36 \\ \end{array}$

Target : Solubility :



Biological Activity

Cabergoline EP impurity F is a byproduct that can be formed during the synthesis of Cabergoline. It is important to identify and characterize impurities in medications, as they can affect the safety and efficacy of the drug. However, there is currently limited information available on Cabergoline EP impurity F.

References

- 1. Wang S, et al. Cabergoline for the treatment of hyperprolactinemia: a meta-analysis of randomized controlled trials. Endocrine. 2016;51(1):70-8.
- 2. Webster J. Cabergoline for the treatment of hyperprolactinemia. Expert Opin Investig Drugs. 2000;9(1):155-70.

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