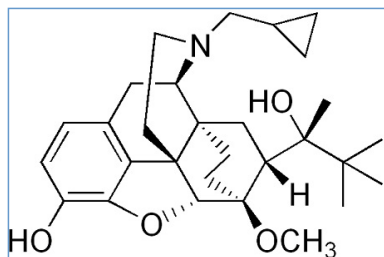


Stabilis



Buprenorphine Hydrochloride



Stabilité des préparations

0.3 mg/mL - 0.125 mL® = Par Pharmaceutical		Ethanol 95° 157,5 µL Sirop simple 217,5 µL	2-8°C		30			4587
0.3 mg/mL - 0.125 mL® = Par Pharmaceutical		Ethanol 95° 157,5 µL Sirop simple 217,5 µL	22-27°C		30			4587





















Bibliographie

	Type	Source
4587	Revue	Rochani A, Nguyen V, Becker R, Kraft W, Kaushal G. Stability-indicating LC-MS Method for Determination of Stability of Extemporaneously Compounded Buprenorphine Oral Syringes for Neonatal Abstinence Syndrome. J Pediatr Pharmacol Ther 2021 ; 26, 4: 395-404.



Dictionnaire

 Antalgique	 Solution buvable
 Stabilité des préparations	 Contenant
 Origine	 Excipient
 Température	 Conservation
 Durée de stabilité	 Biosimilaire
 Données conflictuelles	 Bibliographie
 Seringue PP orale	 Flacon injectable
 A l'abri de la lumière	 Jour
 Bibliographie	 Dictionnaire