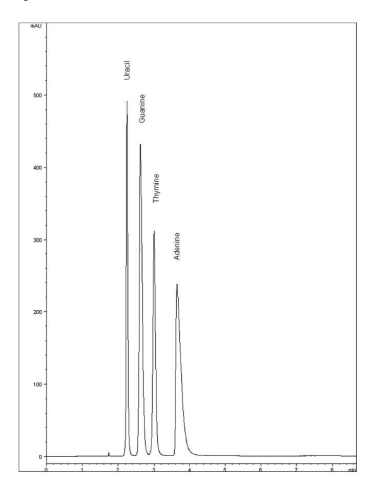


Nucleobases analyzed by HPLC - AppNote

Nucleotides Uracil, Guanine, Thymine, Adenine, Excellent Peak Shape and Resolution

This Method is easy to prepare, use and reproduce. Separation is accomplished under 100% Aqueous Conditions yet there is an alternate Selectivity. These bases may be difficult to retain on Columns with ordinary Silica that contain significant amounts of Silanols.



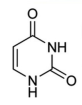
Printed from the Chrom Resource Center
Copyright 2024, All Rights Apply
MicroSolv Technology Corporation

9158 Industrial Blvd. NE, Leland, NC 28451 tel. (732) 380-8900, fax (910) 769-9435

Email: customers@mtc-usa.com

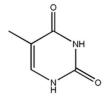
Website: www.mtc-usa.com

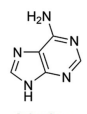




Uracil

Guanine





Thymine

Adenine

Peaks:

- 1. Uracil (U)
- 2. Guanine (G)
- 3. Thymine (T)
- 4. Adenine (A)

Method Conditions

Column: Cogent Diamond Hydride™, 4µm, 100Å

Catalog No.: 70000-75P Dimensions: 4.6 x 75mm

Mobile Phase: DI Water / 0.1% Acetic Acid

Temperature: 25°C
Injection vol.: 2.5μL
Flow rate: 1mL / minute
Detection: UV @ 254nm

Notes: Nucleobases (or Nucleotide Bases) are the parts of DNA and RNA that may be involved in pairing. The main Bases are Cytosine, Guanine, Adenine (DNA and RNA), Thymine (DNA) and Uracil (RNA). They are usually simply called "Bases" in Genetics.



Attachment

Nucleobases Analyzed by HPLC pdf Download File

Printed from the Chrom Resource Center Copyright 2024, All Rights Apply

MicroSolv Technology Corporation

9158 Industrial Blvd. NE, Leland, NC 28451 tel. (732) 380-8900, fax (910) 769-9435

Email: customers@mtc-usa.com

Website: www.mtc-usa.com