



1. For devices without the direct mark, the user may obtain the UDI from the table below by referencing the corresponding information marked on the physical device.
  - a. Unique Device Identification: Policy Regarding Compliance Dates for Class I and Unclassified Devices, Direct Marking, and Global Unique Device Identification Database Requirements for Certain Devices (July 25, 2022)
2. This information can be reprinted from the product web page on the Henry Schein website at [www.HenrySchein.com](http://www.HenrySchein.com)

**Medical Device description:**

*ALM RETR, 2-3/4 INCH, SPREAD 2-1/4 INCH*

**UDI Codes cross-matching summary table**

| <b>HS REF</b> | <b>GTIN NUMBER<br/>(01)</b> | <b>LOT NUMBER<br/>(10)</b> | <b>MANUFACT. DATE<br/>(11)</b> |
|---------------|-----------------------------|----------------------------|--------------------------------|
| 104-7432      | 00304040016374              | 246401                     | 2018-02                        |
| 104-7432      | 00304040016374              | 246402                     | 2018-02                        |
| 104-7432      | 00304040016374              | 246403                     | 2018-02                        |
| 104-7432      | 00304040016374              | 246404                     | 2018-02                        |