

- 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 <u>www.HenrySchein.com</u>

Medical Device description:

TISS FCP, 4-1/2IN(114 MM), 1 X 2 TTH, STD SERR HDL

UDI Codes cross-matching summary table

HS REF	GTIN NUMBER (01)	LOT NUMBER (10)	MANUFACT. DATE (11)
104-6618	00304040023105	AF2001	2020-01
104-6618	00304040023105	AF2003	2020-03
104-6618	00304040023105	AF2104	2021-04