Quick Guide For NMPA UDID GDSN Testing

Quick Guide For Medical Device License Holders/General Agents To Conduct NMPA UDID GDSN Testing.

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1. **Submit the Authorization Statement to GS1 China.**
   
   1.1 Log into the *website of Authorization Statement*, click "Agree", complete the form, and click "Submit" to trigger the data subscription.

   ✔️ **Note:** when the sign “✅” appears, it means the submission is successful.

   1.2 E-mail to GS1 China contact (Yingxi Yang, yangyx@ancc.org.cn) to indicate the completion of step 1.1 and confirm the request for testing.

2. **Complete the Authorization process on the NMPA UDID portal.**

   2.1 Log into the *Production system of NMPA UDID*, select “中国物品编码中心” (GS1 China) as the authorized agency, upload the Authorization Letter, and complete the authorization.

   ✔️ **Note:** normally the Authorization in the UDID Production system will be auto-synced into the UDID Pre-production system within 72 hours and work for data testing.

3. **Make sure relevant medical device license information has been maintained in the UDID Pre-production system.**

   3.1 Log into the *Pre-production system* to view or maintain license information.

4. **Upload UDI data into local data pools based on NMPA UDID data requirements.**

   4.1 Please find the latest version of the NMPA UDID to GDSN Attribute Mapping document at the top of the webpage *here*.

5. **Publish UDI data to NMPA UDID’s Pre-production GLN (6907777445543).**

6. **Manage CIC messages.**

   6.1 Take correction actions based on the CIC messages of "Review" (if any) resulting from data validations. Please find the NMPA UDID CIC Message Mapping document at the top of the webpage *here*.

7. **Check UDI data in the Pre-production system of NMPA UDID.**

   7.1 Log into the *Pre-production system*. The medical device license holder/general agent’s user account in NMPA UDID’s Production system also works in the Pre-production system.