



Dear Benjamin Bensal:

This e-mail provides confirmation that the annual registration for the following medical device establishment has been successfully completed for 2020:

Registration Number: 3002941807  
Owner Operator Number: 10043474  
KDL PRECISION MOLDING CORP  
11381 Bradley Ave  
Pacoima, CA 91331  
UNITED STATES

If you do not see a registration number assigned to the establishment and your establishment previously had one, please send an email to [reglist@cdrh.fda.gov](mailto:reglist@cdrh.fda.gov) and include the registration number you believe is assigned to your establishment. We will review and determine if a duplicate registration has been created for your establishment.

Your registration is valid until December 31, 2020. Registration for 2021 will be conducted between October 1 and December 31, 2020.

Please note that registering your device facility and listing your devices does not, in any way, constitute FDA approval of your facility or your devices.

Should you have any questions, please send an e-mail to the CDRH Registration and Listing Helpdesk at [reglist@cdrh.fda.gov](mailto:reglist@cdrh.fda.gov).

CDRH Registration and Listing Helpdesk  
Imports & Registration and Listing Team  
Division 2 Establishment Support  
Office of Regulatory Programs  
Office of Product Evaluation and Quality  
Center for Devices and Radiological Health  
U.S. Food and Drug Administration

Tel: 301-796-7400, Option 1

Email: [reglist@cdrh.fda.gov](mailto:reglist@cdrh.fda.gov)