ACE information sheet

FDA Intended Use

Importing FDA-regulated products? As a customs broker, Livingston needs to know how the product being imported is intended to be used and/or why the product is being imported.

Importers should confirm that their invoices clearly show what the end use is going to be for the product so that the customs broker can enter the appropriate end use code. To ensure your descriptions are complete enough we have highlighted below the intended use descriptions required.

Biologics Intended Use

- 080.000 CBER-regulated final product ready for use
- 081.000 CSER-regulated product for processing into a medical device
- 082.000 Human cells, tissues, and cellular tissue based products for implant, transplant, infusion or transfer into human recipient
- 100.000 Importation for personal use
- 110.000 Import of biological drug or device for trade show
- 140.000 Standard import of a biological drug or device for non-commercial distribution in organization support program
- 150.007 Bulk drug substance for processing into a pharmaceutical product
- 155.000 CBER product for further manufacturer of a licensed biological product under a short supply agreement (21CFR 601.22)*
- 170.000 Import of biological product, drug, or device that is US goods returned to manufacturer
- 180.000 Import of biologic for NON-clinical research use only.
- 180.009 Import of biological or chemical for research and development into a pharmaceutical product
- 180.010 Import of a biological or chemical for research and development into a medical device
- 180.016 CBER product sample for testing or lot release
- 970.000 CBER-Import for Export
- 940.000 Compassionate use/emergency use
- UNK Unknown

Drugs Intended Use

- 080.000 For Human Medical Use as a Non-Food Product under Controlled Distribution PRE Prescription
- 130.000 For Consumer Use as a Non-food Product OTC Over the Counter
- 150.007 Active Pharmaceutical Ingredient/ bulk drug substance for processing into a pharmaceutical product
- 180.009 Chemical for research and development in a pharmaceutical product investigational new drugs, clinical trials or other human/animal ingestion
- 180.017 Chemical for research and development in a pharmaceutical product laboratory testing only-- no human/animal ingestion
- 970.000 Import for Export program
- 100.000 Importation for personal Use



- 155.009 Importation of a drug constituent part (drug product) for use in a medical product regulated under a
 device (CDRH) application type (e.g. for use in a PMA/510(k) drug-device combination product
- 150.017 Importation of a drug component (API) for use in a medical product regulated under device (CDRH) combination product
- 150.018 Active Pharmaceutical ingredient/ bulk drug substance to be used for pharmacy compounding
- UNK Unknown

Medical Devices

- 081.001 For Human Medical use as a medical device (standard import of a medical device, accessories, or components) regulated as finished devices, import of refurbished device, import of a reprocessed device
- 081.002 Refurbishing
- 081.003 Domestically manufactured device that is part of a medical device convenience kit
- 081.004 Foreign manufactured device that is part of a medical device convenience kit
- 081.005 Device for use in a drug/device combination product
- 081.006 Import of a medical device under enforcement discretion, applies to the following product codes: 800--UG, 86N--FF, 86N--FG, 80N--XQ, 90L--MB, 90L--MD only
- 100.000 Personal Use
- 110.000 Public exhibition or taking orders (includes trade shows)
- 140.000 Charitable organization use
- 081.007 Component for further manufacturing into a finished medical device
- 081.008 Device for use in a NDA/ANDA/BLA drug device combination product
- 170.000 Repair
- 180.010 Research & development as a medical device
- 180.014 Research & development for bench testing or non-clinical research
- 180.015 Research & development clinical investigation use
- 920.001 US manufactured medical device returned as over-stock, refund
- 920.002 US manufactured device sale to a third party
- 940.000 Compassionate use, emergency use device
- 950.001* Single use device for domestic reprocessing
- 950.002* Multi-use device for domestic reprocessing
- 970.000 Import for export (must be further manufactured or processed, and then exported)- device or accessory
- 970.001 Import for export (must be further manufactured or processed, and then exported)-component

NOTE: Conditional affirmations are required if applicable to the product being declared, for example, if the product requires premarket clearance 510k, then PM# must be provided.

- * annotates that additional information may be needed at the time of entry in order for FDA to make a final admissibility decision.
- UNK Unknown



Electronics

- 085.000 For Veterinary Medical Use as a Non-Food Product under Controlled Distribution
- 090.000 For Military Use as a Non- Food Product
- 100.000 For Personal Use as a Non- Food Product
- 110.000 For Public Exhibition or Display as a Non-Food Product
- 120.000 For Public Safety Use as a Non-Food Product
- 130.000 For Consumer Use as a Non- Food Product
- 140.000 For Charitable Organization Use as Non-Food Product
- 150.000 For Commercial Processing as a Non-Food Product
- 155.000 For Commercial Assembly as a Non-Food Product
- 170.000 For Repair of a Non-Food Product
- 180.000 For Research and Development as a Non-Food Product
- 970.000 For import for export
- 980.000 For other use

Contact Livingston

Have questions or need help with your FDA imports under ACE?

Contact your account executive,

write to us at: simplify@livingstonintl.com

or give us a call at 1-855-225-5548

