



NDA 205737
NADA 141358

**DEEMED GRANTED -
MEDICAL GAS CERTIFICATION REQUEST**

Air Liquide Large Industries U.S. LP
Attn: Steve R. Miller
Sr. Director, Quality and Regulatory Affairs
2700 Post Oak Boulevard
Houston, TX 77056

Dear Mr. Miller:

Please refer to your April 22, 2013, request received on April 24, 2013, for certification of Oxygen, USP as a designated medical gas. You have requested to market Oxygen, USP for human and animal drug use.

A request for certification of a medical gas as a designated medical gas submitted under section 575(a)(1) of the Federal Food, Drug, and Cosmetic Act (FD&C Act) is deemed to be granted unless, within 60 days of the filing of the request, FDA finds that one or more of the bases for denying the request listed at section 575(a)(2) of the FD&C Act applies. FDA has made no such finding in connection with your request, and 60 days have passed since your request was filed. Accordingly, by operation of section 575(a)(2) of the FD&C Act, your request for certification of Oxygen, USP as a designated medical gas is deemed to be granted, and you now have in effect an approved new drug application (NDA 205737) and an approved new animal drug application (NADA 141358) for this gas.

If any of the information you have submitted in connection with your request becomes incomplete or inaccurate, please consult section IV.D of the draft guidance document entitled *Certification Process for Designated Medical Gases* (available at <http://www.fda.gov/downloads/Drugs/GuidanceComplianceRegulatoryInformation/Guidances/UCM332136.pdf>) for instructions on providing FDA with complete, up-to-date information. Please address any such communications to:

Food and Drug Administration
Center for Drug Evaluation and Research
Office of Pharmaceutical Quality
5901-B Ammendale Road
Beltsville, MD 20705-1266

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Please include the NDA and NADA numbers listed above at the top of the first page of any such communications.

If you have any questions, please call Michael Folkendt at (301) 796-1670 or by email at Michael.Folkendt@fda.hhs.gov.

Sincerely,

{See appended electronic signature page}

Charles J. Andres, Ph.D.
Business Process Improvement Manager
Office of New Animal Drug Evaluation
Center for Veterinary Medicine
FDA

Michael Folkendt
Associate Director for Regulatory Affairs
Office of Program and Regulatory Operations
Office of Pharmaceutical Quality
Center for Drug Evaluation and Research
FDA

This is a representation of an electronic record that was signed electronically and this page is the manifestation of the electronic signature.

/s/

MICHAEL M FOLKENDT
09/25/2015

CHARLES J ANDRES
09/28/2015

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