



NIOSH Reference: TN-09086  
Mfr. Reference: 111196

Centers for Disease Control  
and Prevention (CDC)  
National Institute for Occupational  
Safety and Health - ALOSH  
1095 Willowdale Road  
Morgantown, WV 26505-2888

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January 16, 1997

Mr. Robert E. Brennan  
Scott Aviation  
A Figgie International Company  
225 Erie Street  
Lancaster, New York 14086-0622

Dear Mr. Brennan:

The National Institute for Occupational Safety and Health (NIOSH) has reviewed your request dated November 11, 1996. This request was, in part, for approval of the Scott Aviation, thirty-minute, open-circuit, pressure-demand, 2216 psig Industrial Air-Pak, self-contained breathing apparatus (SCBA) listed in your standard application form.

Approval TC-13F-401 is granted to cover the Scott Aviation, thirty-minute, open-circuit, pressure-demand, 2216 psig Industrial Air-Pak, self-contained breathing apparatus (SCBA) listed in your standard application form. The respirator models affected by this approval are listed in your standard application form. A copy of the NIOSH test report is enclosed. The cautions and limitations listed on your approval label and in your instruction manual, copies of each enclosed, apply to this approval. Approval is granted for English language only on all documentation. It is the manufacturer's responsibility to correctly translate materials desired in languages other than English.

In making renewals or repairs, parts identical with those furnished by the manufacturer under the pertinent approval shall be maintained.

The approved assembly consists of the Scott Aviation, parts indicated by the approval label draft plus the assembly matrix. These parts are to be marked with the indicated numbers in a legible and permanent manner (marking cannot be removed without evidence of its previous presence).

Page 2 - Robert E. Brennan

The use of this approved device in combination with any other additional respirator components not covered under this approval, renders this certification invalid.

The enclosed approval label draft designs are to be used in preparing the approval labels. Label TC-13F-401 shall be prepared for use in the manual and on the harness assembly. Final copies of your labels and user's instructions must be submitted to NIOSH for concurrence before production.

Only those assemblies where approval numbers are assigned apply to this approval action. Production approval labels cannot include information on unapproved configurations.

Your quality control plans for this respirator were reviewed by NIOSH. On the basis of that review, your quality control plans are accepted as a part of this approval.


This certificate of approval is not an endorsement of the respirator by NIOSH, and such endorsement shall not be stated or implied in advertisements or other publicity. However, you may publicize the fact that the product has met the requirements of Title 42, Code of Federal Regulations (CFR), Part 84.

Any changes you wish to make to this respirator and accompanying documentation shall be submitted, and a modification of this approval shall be granted (including changes in materials, product design, manufacturing processes, labels, instruction manuals and quality control programs) before any changes are made. (Reference: Title 42, CFR, Part 84.35.)

Page 3 - Mr. Robert E. Brennan

Please submit samples of respirator packaging, bearing all required labels, instructions, and markings, for our approval, before adopting them. Please send us one production sample to be made a part of the record of this approval. We shall retain several other items as additional record material. All other material will be discarded unless we are otherwise advised by you within 7 days.

Sincerely yours,



Richard W. Metzler, Chief  
Certification and Quality  
Assurance Branch

Division of Respiratory Disease Studies

Enclosures