



NIOSH Reference: TN-17651  
Mfr. Reference: GER00299

Centers for Disease Control  
and Prevention (CDC)  
National Institute for Occupational  
Safety and Health (NIOSH)  
National Personal Protective  
Technology Laboratory (NPPTL)  
P.O. Box 18070  
Pittsburgh, PA 15236-0070  
Phone: 412-386-4000  
Fax: 412-386-4051  
March 25, 2011

Mr. Robert Brunell  
Quality Control Manager  
Louis M. Gerson Company, Inc.  
16 Commerce Boulevard  
Middleboro, Massachusetts 02346

Dear Mr. Brunell:

The National Institute for Occupational Safety and Health (NIOSH) has reviewed your request accepted on February 22, 2011. This request was for an extension of approval to TC-23C-1526, TC-23C-1579, TC-23C-1580, TC-23C-1581, TC-84A-3333, TC-84A-3332, TC-84A-3334, TC-84A-3335, and TC-84A-3336 to add private label versions of these 9000 series Half Mask Air Purifying Respirators with chemical cartridges, part numbers G01, G04, G05, and G02 or combination chemical and particulate cartridges, part numbers G02, G01, G03, G04 and G05 with the G95P P95 filter, for Libus of Buenos Aires, Argentina under the same part numbers, reference assembly matrix Rcr18AMj.xls.

This request is granted. Approvals are granted only for documentation written in the English language. It is the manufacturer's responsibility to correctly translate materials desired in languages other than English.

The CD enclosed with this letter contains the final private label respirators approval label. The cautions and limitations which apply to this approval are on the approval label. Only those assemblies affected by this request, or where new approval numbers are assigned, apply to this approval action. Production approval labels cannot include information on unapproved configurations.

The approved assemblies consist of the parts as listed on the approval label and the assembly matrix. Parts are to be marked with the numbers indicated on the approval label in a legible and permanent manner (marking cannot be removed without evidence of its previous presence).

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The manufacturer is responsible for properly packaging, labeling, and controlling the respirators produced under this private label approval. At a minimum, the items that must be controlled are the approved user instructions, all approval labeling, all approved packaging, use claims, marketing materials, and the respirator design and construction details. Any change to this NIOSH-approved respirator or approval documentation without prior notification and approval is a violation of this approval and renders this certification as invalid.

No changes may be made to any respirator and accompanying documentation without prior written approval of NIOSH. Requests for changes must be submitted to NIOSH and a modification of this approval must be granted before changes are made.

Sincerely yours,

A handwritten signature in blue ink, appearing to read "Heinz W. Ahlers". The signature is fluid and cursive, with the first name being the most prominent.

Heinz W. Ahlers  
Chief, Technology Evaluation Branch  
National Personal Protective Technology Laboratory

Enclosures



NIOSH Reference: TN-16693  
Mfr. Reference: GER000277

Centers for Disease Control  
and Prevention (CDC)  
National Institute for Occupational  
Safety and Health (NIOSH)  
National Personal Protective  
Technology Laboratory (NPPTL)  
P.O. Box 18070  
Pittsburgh, PA 15236-0070  
Phone: 412-386-4000  
Fax: 412-386-4051  
December 15, 2009

Mr. Robert Brunell  
Quality Control Manager  
Louis M. Gerson Company, Inc.  
16 Commerce Boulevard  
Middleboro, Massachusetts 02346

Dear Mr. Brunell:

The National Institute for Occupational Safety and Health (NIOSH) has reviewed your request dated November 15, 2009. This request was for extensions of approvals to TC-23C-1578, TC-84A-0170, and TC-84A-3173 to add private label versions of these air purifying respirators consisting of the model 9000 half mask facepiece, chemical cartridge, part number GO3, or particulate filter part number G70, or particulate filter part number XP100 for Libus of Buenos Aires, Argentina, under the same model and part numbers, as referenced on assembly matrix Rcr18AMag.xls.

This request is granted. Approvals are granted only for documentation written in the English language. It is the manufacturer's responsibility to correctly translate materials desired in languages other than English.

The CD enclosed with this letter contains the final respirator, cartridge and filters approval labels. The cautions and limitations which apply to this approval are on the approval labels. Only those assemblies affected by this request, or where new approval numbers are assigned, apply to this approval action. Production approval labels cannot include information on unapproved configurations.

The approved assemblies consist of the parts as listed on the approval labels and the assembly matrix. Parts are to be marked with the numbers indicated on the approval label in a legible and permanent manner (marking cannot be removed without evidence of its previous presence).

The manufacturer is responsible for properly packaging, labeling, and controlling the respirators produced under this private label approval. At a minimum, the items that must be controlled are the approved user instructions, all approval labeling, all approved packaging, use claims, marketing materials, and the respirator design and construction details. Any change to this NIOSH-approved respirator or approval documentation without prior notification and approval is a violation of this approval and renders this certification as invalid.



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No changes may be made to any respirator and accompanying documentation without prior written approval of NIOSH. Requests for changes must be submitted to NIOSH and a modification of this approval must be granted before changes are made.

Sincerely yours,



Heinz W. Ahlers  
Chief, Technology Evaluation Branch  
National Personal Protective Technology Laboratory