In accordance with the Paperwork Reduction Act of 1995, the Centers for Disease Control and Prevention (CDC) has submitted the information collection request titled Pulmonary Function Testing Course Approval Program to the Office of Management and Budget (OMB) for review and approval. CDC previously published a "Proposed Data Collection Submitted for Public Comment and Recommendations" notice on June 2, 2020 to obtain comments from the public and affected agencies. CDC received one non-substantial comment related to the previous notice. This notice serves to allow an additional 30 days for public and affected agency comments.

CDC will accept all comments for this proposed information collection project. The Office of Management and Budget is particularly interested in comments that:

(a) Evaluate whether the proposed collection of information is necessary for the proper performance of the functions of
the agency, including whether the information will have practical utility;
(b) Evaluate the accuracy of the agencies estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used;
(c) Enhance the quality, utility, and clarity of the information to be collected;
(d) Minimize the burden of the collection of information on those who are to respond, including, through the use of appropriate automated, electronic, mechanical, or other technological collection techniques or other forms of information technology, e.g., permitting electronic submission of responses; and
(e) Assess information collection costs.

To request additional information on the proposed project or to obtain a copy of the information collection plan and instruments, call (404) 639-7570. Comments and recommendations for the proposed information collection should be sent within 30 days of publication of this notice to www.reginfo.gov/public/do/PRAMain Find this particular information collection by selecting "Currently under 30-day Review - Open for Public Comments" or by using the search function. Direct written comments and/or suggestions regarding the items contained in this notice to the
Proposed Project

Pulmonary Function Testing Course Approval Program. (OMB Control No. 0920-0138, Exp. 11/30/2020) – Revision – National Institute for Occupational Safety and Health (NIOSH), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

NIOSH has the responsibility under the Occupational Safety and Health Administration’s Cotton Dust Standard, 29 CFR 1920.1043, for approving courses to train technicians to perform pulmonary function testing in the cotton industry. Successful completion of a NIOSH-approved course is mandatory under this Standard. In addition, regulations at 42 CFR 37.95(a) specify that persons administering spirometry tests for the national Coal Workers’ Health Surveillance Program must successfully complete a NIOSH-approved spirometry training course and maintain a valid certificate by periodically completing NIOSH-approved spirometry refresher training courses. Also, 29 CFR 1910.1053(i)(2)(iv), 29 CFR 1910.1053(i)(3), 29 CFR 1926.1153(h)(2)(iv) and 29 CFR 1926.1153(h)(3) specify that pulmonary function tests for initial and periodic examinations in general industry and construction performed under the
respirable crystalline silica standard should be administered by a spirometry technician with a current certificate from a NIOSH-approved spirometry course. NIOSH is requesting a three-year approval.

To carry out its responsibility, NIOSH maintains a Pulmonary Function Testing Course Approval Program. The program consists of an application submitted by potential sponsors (universities, hospitals, and private consulting firms) who seek NIOSH approval to conduct courses, and if approved, notification to NIOSH of any course or faculty changes during the approval period, which is limited to five years. The application form and added materials, including an agenda, curriculum vitae, and course materials are reviewed by NIOSH to determine if the applicant has developed a program which adheres to the criteria required in the Standard. Following approval, any subsequent changes to the course are submitted by course sponsors via letter or email and reviewed by NIOSH staff to assure that the changes in faculty or course content continue to meet course requirements. Course sponsors also voluntarily submit an annual report to inform NIOSH of their class activity level and any faculty changes.

Sponsors who elect to have their approval renewed for an additional five year period, submit a renewal application and supporting documentation for review by NIOSH staff to ensure the
course curriculum meets all current standard requirements. Approved courses that elect to offer NIOSH-Approved Spirometry Refresher Courses must submit a separate application and supporting documents for review by NIOSH staff. Institutions and organizations throughout the country voluntarily submit applications and materials to become course sponsors and carry out training. Submissions are required for NIOSH to evaluate a course and determine whether it meets the criteria in the Standard and whether technicians will be adequately trained as mandated under the Standard.

NIOSH will disseminate a one-time customer satisfaction survey to course directors and sponsor representatives to evaluate our service to courses, the effectiveness of the program changes implemented since 2005, and the usefulness of potential Program enhancements.

Application form changes consist of minor text edits that clarify questions and information, thereby reducing the need for applicants to contact NIOSH for guidance. In addition, parts of the forms were reformatted to reduce redundancy and increase clarity for applicants. Two of the forms have updated titles which reflect the purpose of the applications (initial sponsorship and sponsorship renewal forms).

The estimated annual burden to respondents is 147 hours. There will be no cost to respondents other than their time.
## Estimated Annualized Burden Hours

<table>
<thead>
<tr>
<th>Type of Respondents</th>
<th>Form Name</th>
<th>No. of Respondents</th>
<th>No. of Responses per Respondent</th>
<th>Average Burden per Response (in hours)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Potential Sponsors</strong></td>
<td><strong>Initial Application</strong></td>
<td>3</td>
<td>1</td>
<td>3.5</td>
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<tr>
<td></td>
<td><strong>Annual Report</strong></td>
<td>34</td>
<td>1</td>
<td>28/60</td>
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<tr>
<td></td>
<td><strong>Report for Course Changes</strong></td>
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<td>1</td>
<td>30/60</td>
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<tr>
<td></td>
<td><strong>Renewal Application</strong></td>
<td>13</td>
<td>1</td>
<td>6</td>
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<tr>
<td></td>
<td><strong>Refresher Course Application</strong></td>
<td>3</td>
<td>1</td>
<td>8</td>
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<tr>
<td></td>
<td><strong>One-Time Customer Satisfaction Survey</strong></td>
<td>32</td>
<td>1</td>
<td>12/60</td>
</tr>
</tbody>
</table>

Jeffrey M. Zirger,
Lead,
Information Collection Review Office,
Office of Scientific Integrity,
Office of Science,
Centers for Disease Control and Prevention.

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