## PRP Section 8100

**Instructions to Providers Having a Quality Control Materials Review**

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Introduction

.01 A quality control materials (QCM) review is a type of peer review that is a study and appraisal by a QCM reviewer of an organization’s (hereinafter referred to as provider) system of quality control to develop and maintain QCM. The system represents the provider’s policies and procedures that the provider has designed and is expected to follow when developing the materials. The QCM reviewer’s objective is to determine whether the system is designed and whether the provider is complying with its system appropriately so that users of the materials, primarily CPA firms and their employees, have reasonable assurance to rely on the materials. The materials can be part or all of a firm’s documentation of its system, in the form of, for example, manuals, programs, and practice aids (forms and questionnaires). Users rely on the materials to assist them in performing and reporting in conformity with professional standards in conducting their accounting and auditing practices. In addition, one of the reasons that providers elect to have an independent review of their system of quality control for the development and maintenance of the QCM they have developed, and of the materials themselves, is to provide more cost-effective peer reviews for firms that have acquired or use such materials.

.02 The purpose of these instructions is to provide overall guidance to providers having a QCM review under the AICPA Peer Review Program (the program). Providers should be aware of their review responsibilities and requirements as discussed in PRP section 1000, Standards for Performing and Reporting on Peer Reviews, with an emphasis on paragraphs .154–.205 and paragraphs 17–22 of appendix A (as well as these instructions). In addition, all individuals at the provider involved in the review should read and become familiar with the standards, PRP section 2000, Peer Review Standards Interpretations; PRP section 3000, Other Guidance; and materials relative to the aspect of the review that most directly affects their role at the provider.

.03 An independent review of the system for the development and maintenance of QCM and the resultant materials (the QCM review) is required for certain providers (see PRP section 1000). In addition, a provider may have a QCM review voluntarily so that peer reviewers of user firms can place reliance on the QCM to reduce the scope of the review of the firm’s third-party materials.

.04 A QCM Review is intended to provide the QCM reviewer with a reasonable basis for expressing an opinion on whether, during the year under review

a. the provider’s system for the development and maintenance of the materials was suitably designed and was being complied with during the period under review to provide user firms with reasonable assurance that the materials are reliable aids to assist them in conforming with those professional standards the materials purport to encompass, and

b. the resultant materials are reliable aids.

.05 A QCM review encompasses judgmental review of all of the materials opined on in the report. The extent of review of each module or guide is based on the QCM reviewer’s assessment of risk, taking into consideration factors such as industries with higher inherent risk, new pronouncements and standards, and so on.

.06 QCM reviews are administered by the AICPA Peer Review Board’s National Peer Review Committee (NPRC). In addition, the QCM Task Force is involved in the administration and acceptance process. The task force’s involvement includes performing oversight reviews prior to acceptance, developing practice aids, and recommending enhancements to the guidance related to QCM reviews.

Prior to the Review

.07 Providers required to have a QCM review should have the review once every 3 years and should arrange to have the review administered by the NPRC. Providers should submit the Information Required to Schedule QCM Reviews form no less than 60 days prior to the commencement of the review. The QCM review should not commence until the provider and reviewer are informed that the selected QCM review team is approved to perform the review. It is the responsibility of the provider to verify that the QCM review team is qualified to perform the review, including ensuring that the QCM review team doesn’t have the following independence impairments:

- The reviewing firm uses materials developed by the provider as an integral part of its system of quality control.
- A QCM review team member was involved in the development of the provider’s materials.
The provider is an association to which the reviewing firm belongs.

Any other conflicts of interest.

.08 The provider and the QCM reviewer should agree on an appropriate date for the review to commence and the anticipated exit conference date. Ordinarily, the review should be performed within six months following the end of the year to be reviewed. In most circumstances, the applicable period should not change from one triennial review period to the next. In the event of substantial change in the system for the development and maintenance of the materials or in the resultant materials, the provider should consult with the NPRC to determine whether an accelerated review is warranted.

.09 The terms and conditions of the QCM review may be summarized in an engagement letter between the provider and the reviewing firm or association, if an association formed the QCM review team.

.10 A contact person should be designated as liaison to provide assistance to the QCM review team and should be available throughout the review.

.11 Provide the following to the QCM reviewer as soon as possible:

a. The quality control documentation, including the procedures for developing the materials (including distribution), ensuring the materials are current and relevant, determining the on-going qualifications of all parties involved in the development and maintenance of the materials, soliciting and evaluating feedback from users of the materials, policies regarding the issuance of updates to the materials, the method of updating, procedures undertaken to provide such updates (if such policies exists), and the procedures for monitoring compliance with the provider’s quality control policies and procedures

b. Documentation of the monitoring procedures performed by the provider during the review year

c. A list of the materials on which an opinion is to be expressed

d. A list of the personnel involved in the development and maintenance of the materials

e. A list of the external or guest authors and technical reviewers involved in the development and maintenance of the materials

f. Other information requested by the QCM reviewer

.12 Have the following available for the QCM review team when they perform the site visit:

a. Personnel information to the extent requested by the QCM reviewer

b. Documentation to support the qualifications and expertise of personnel involved in the development or maintenance of the materials, such as current résumés including titles or positions, relevant training, experience, industry expertise, CPE records, and so on.

c. Documentation to support the qualifications and expertise of external or guest authors and technical reviewers

d. Any communications relating to allegations, investigations, or litigation involving the provider, its personnel, or non-personnel contributors or reviewers (such as guest authors or technical reviewers) since the provider’s last review year-end

.13 The provider should provide a comfortable, adequate working area for the QCM review team and, if necessary, assist in coordinating accommodations for the QCM review team.

.14 The review of the provider’s quality control policies and procedures includes interviews of the provider’s personnel. The objective of these interviews is to provide corroborative evidence that certain policies and procedures have been properly communicated and are being complied with. The QCM review team may perform one-on-one interviews or focus groups. The QCM reviewer will arrange for the scheduling of interviews with selected members of the provider’s personnel. The provider should see that this schedule is communicated to the appropriate individuals and that they understand the importance and purpose of the interviews. The QCM review team will endeavor to have these discussions and interviews without disrupting the provider’s operations.
During the Review

.15 The designated liaison should meet with the QCM review team at the beginning of the review to orient them to the policies and procedures, introduce them to appropriate personnel, and provide them with a tour of the office.

.16 During the course of the QCM review, the QCM review team may find it necessary to discuss matters with the appropriate personnel. Provider personnel should be asked to be available to the QCM review team as necessary during the course of the QCM review.

.17 In addition, provider personnel may need to coordinate with AICPA staff to plan oversight procedures.

Completion of the Review

.18 A provider that has a QCM review should respond promptly to questions raised in the review in order to assist the QCM review team in reaching its conclusions. Prior to issuing its report or finalizing Finding for Further Consideration (FFC) form(s), if applicable, the QCM review team will communicate any matters documented on the Matter for Further Consideration (MFC) form(s), findings documented on the FFC form(s), deficiencies or significant deficiencies to be included in the peer review report, and the type of report to be issued through one or more exit conferences. The designated liaison should arrange for appropriate personnel to attend the exit conference. The exit conference may be attended by representatives of the NPRC, the QCM Task Force, the AICPA Peer Review Board, AICPA staff, or other board authorized organizations with oversight responsibilities.

.19 The QCM review team should communicate, if applicable, that the provider will be required to respond to the findings documented on the FFC form(s) and the deficiency(ies) or significant deficiencies included in the peer review report. The QCM review team should also communicate that the provider may be required, if applicable, to (1) take certain actions to correct the deficiencies or significant deficiencies noted in the report and (2) complete an implementation plan to address the findings noted in the FFC form(s). The QCM review team should discuss with the provider the implications of these steps on the acceptance and completion of the peer review. The exit conference is also the appropriate vehicle for providing suggestions to the provider that are not included in the report, FFC form(s), or MFC form(s).

.20 The provider will provide the QCM reviewer with written representations, at a minimum, relating to the following matters:

a. Situations where management is aware that its materials were used and substantially relied upon in an engagement that was later found to not comply with the applicable standards or regulations (auditing, review, reporting, and so on) in all material respects, when the materials were found to be an underlying cause of the engagement deficiencies.

b. Access to all sources of feedback, including user feedback.

c. Situations or a summary of situations where management is aware that its personnel or non-personnel contributors or reviewers (for example, guest authors or reviewers) have not complied with the rules and requirements of state board(s) of accountancy or other regulatory bodies, as applicable (including applicable licensing requirements in each state in which it practices if the provider is a firm or has employed CPA personnel), and if applicable, how the provider has or is addressing and rectifying situations of noncompliance.

d. Restrictions or limitations of CPA personnel or non-personnel contributors that impacts their ability to practice public accounting within three years preceding the current peer review year-end that were imposed by or agreed to with other regulatory, monitoring, or enforcement bodies (for example, the Public Company Accounting Oversight Board, Securities and Exchange Commission, U.S. Government Accountability Office, Department of Labor, any state board of accountancy or AICPA or state society professional ethics committee, or any other government agency).

e. Access to records and systems of control, including but not limited to, employee files of leased and per diem employees, records related to non-personnel contributors or reviewers, and so on.

f. Materials provided for review that are complete and represent the final version of the materials.

The written representations should be addressed to the QCM reviewer performing the peer review. Because the QCM reviewer is concerned with events occurring during the review period and through the date of his or her QCM review
report that may require an adjustment to the QCM review report or other review documents, the representations should be dated the same date as the QCM review report. See appendix A for an illustration of provider representations.

.21 Ordinarily any FFC forms should be responded to by the provider during the peer review, for example, during or immediately following the exit conference. This would allow the QCM reviewer to assist the provider in developing its responses and obtaining the necessary signatures on the FFC forms and allow the QCM reviewer to review the responses at that time, all of which will expedite the process. In some cases, the provider will choose to check the box on the FFC form that it agrees with the finding and will implement the reviewer’s recommendation. If the provider prefers to provide a description of the actions taken or planned to be taken (and timing), the QCM reviewer can provide assistance in ensuring that the responses are appropriate and comprehensive. However, it is also recognized that the provider may prefer to provide its final responses after it has had the opportunity to discuss them further internally, develop a plan of action and more formally respond. In either case, the completed FFC forms should be submitted to the QCM reviewer no later than two weeks after the exit conference, or by the peer review’s due date, whichever is earlier. FFC forms are then submitted by the QCM reviewer with the applicable working papers to the NPRC.

.22 The provider will receive a report on the QCM review within 30 days of the exit conference date. However, the provider should not publicize the results of the review or distribute copies of the report to its personnel, customers, or others, until it has been advised that the report has been accepted by the NPRC as meeting the requirements of the program.

.23 If the provider receives a report with a peer review rating of “pass with deficiencies” or “fail,” the provider should respond in writing to the deficiencies or significant deficiencies and related recommendations identified in the report. The letter of response should be addressed to the NPRC and should describe the actions planned (including timing) or taken by the provider with respect to each deficiency in the report. The provider should submit a copy of the report, and its letter of response, to the NPRC within 30 days of the date it received the report from the QCM reviewer. Prior to submitting the response to the NPRC, the provider should submit the response to the QCM reviewer for review, evaluation, and comment.

.24 If the provider receives a report with a review rating of “pass” or “pass (with a scope limitation),” a letter of response is not applicable, and the provider does not submit a copy of the report to the NPRC.

.25 Reviewers and providers should understand that professional judgment often becomes a part of the process and each party has the right to challenge the other on such matters. If, after discussion with the QCM reviewer, the provider disagrees with one or more of the findings, deficiencies, or significant deficiencies, the provider should contact NPRC staff for assistance in the matter. If the provider still disagrees after contacting the NPRC, the provider’s response on either the FFC form or in the letter of response, as applicable, should describe the reasons for such disagreement.

.26 The AICPA Peer Review Board encourages the provider to work with the QCM reviewer to develop recommendations that both parties believe will be effective in correcting the matters, findings, and deficiencies noted during the QCM review. Experience shows that improvement is more likely to occur when the provider’s responses describe specific actions to be taken. Therefore, a response limited to the provider’s comment that it will emphasize or reemphasize a policy or procedure should be combined with more specific actions.

.27 Once the QCM reviewer has finalized the QCM review workpapers and the report, the documents are due to the NPRC within 30 days of the exit conference. All QCM reviews undergo a technical review process. In addition, all QCM reviews are subjected to oversight by the QCM Task Force. The level of oversight is dependent on various factors. At a minimum, oversight encompasses NPRC staff performing on-site oversight during the fieldwork procedures, reviewing the QCM reviewer’s working papers, and reviewing a sample of the QCM materials opined upon in the report. The task force can judgmentally elect to perform additional oversight procedures as deemed necessary.

.28 Once technical review and oversight procedures are completed, QCM reviews are presented to the full NPRC with the task force’s recommendation for consideration and acceptance. QCM reviews are considered by the full NPRC during its regularly scheduled meetings or conference calls.
.29 Once the QCM review report and related documents are accepted by the NPRC, an acceptance letter is sent to the provider. The review results are posted to the AICPA website to make QCM review results easily accessible to firms that use the materials, their peer reviewers, and other interested parties.

.30 As part of the acceptance process, the provider may be requested to perform remedial, corrective actions related to the deficiencies or significant deficiencies noted in the QCM review report, in addition to those described by the provider in its letter of response. If a provider does not agree to perform the required actions, this will delay acceptance of the review. If a provider does not perform the required actions, this will delay completion of the peer review.

.31 The program is based on the principle that a systematic monitoring and educational process is the most effective way to attain high quality performance throughout the industry and CPA profession. Thus it depends on mutual trust and cooperation. The provider is expected to take appropriate actions in response to findings, deficiencies, and significant deficiencies identified with its system of quality control or its compliance with the system, or both. Based on the information on the FFC form(s), the provider may be required to have an implementation plan in addition to or as an affirmation of the plan described by the provider in its response to the findings on the FFC form(s). If a provider does not perform the required action in the implementation plan, it could jeopardize the provider’s ability to schedule future QCM reviews. For those providers that are required to obtain a QCM review, disciplinary actions will be taken for a failure to cooperate, failure to correct inadequacies, or when a provider is found to be so seriously deficient in its performance that education and remedial, corrective actions are not adequate.

Fees and Expenses

.32 The NPRC is authorized to establish fees to fund the administration of QCM and CPE peer reviews. Refer to the AICPA web site for the most current fee schedule.
Appendix A

Illustration of a Provider Representation Letter that has No Significant Matters to Report to the QCM Reviewer

October 31, 20XX

To the QCM reviewer:

We are providing this letter in connection with the quality control materials review of [name of provider] and the [insert the titles of the materials] as of the date of this letter and for the year ended June 30, 20XX.

We confirm, to the best of our knowledge and belief, that there are no known circumstances when our materials were used and substantially relied upon in an engagement that was later found to not comply with the applicable standards or regulations in all material respects when the above named materials were found to be an underlying cause resulting in the engagement deficiencies. We also confirm that we have considered all sources of feedback, including feedback from users. We have made you aware of any situations when management is aware that its personnel or non-personnel contributors or reviewers have not complied with the rules and requirements of state board(s) of accountancy or other regulatory bodies (as applicable) and how the provider has or is addressing and rectifying situations of noncompliance. We have also determined that none of our CPA personnel or non-personnel contributors or reviewers are subject to any restrictions or limitations that impacts their ability to practice public accounting within three years preceding the current peer review year end that were imposed by or agreed to with other regulatory, monitoring, or enforcement bodies. Further, we have provided the QCM reviewer with any other information requested and access to records and systems of control, including but not limited to, employee files of leased and per diem employees, files related to non-personnel contributors or reviewers, user feedback, and so on.

Sincerely,

[Name of Signatory]

[Name of Provider]

[The next page is 8201.]

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1 Should be dated the same date as the quality control materials review report.
2 For example, auditing, review, reporting standards, and so on. Consideration should also be given to regulatory guidance, such as the Employee Retirement Income Security Act, the Office of Management and Budget, the Department of Labor (DOL), and so on.
3 Including guest or external authors or reviewers.
4 Including applicable licensing requirements in each state in which it practices if the provider is a firm or has employed CPA personnel.
5 For example, the Public Company Accounting Oversight Board, Securities and Exchange Commission, U.S. Government Accountability Office, DOL, any state board of accountancy or AICPA or state society professional ethics committee, or any other government agency.
6 Letter should be signed by the appropriate party at the provider that has primary responsibility for the system to develop and maintain the materials.