DEPARTMENT OF HEALTH AND HUMAN SERVICES

45 CFR Parts 155 and 156

[CMS-9922-F]

RIN 0938-AT53

Patient Protection and Affordable Care Act; Exchange Program Integrity

AGENCY: Centers for Medicare & Medicaid Services (CMS), HHS.

ACTION: Final rule.

SUMMARY: This final rule revises standards relating to oversight of Exchanges established by states and periodic data matching frequency. This final rule also includes new requirements for certain issuers related to the collection of a separate payment for the portion of a plan’s premium attributable to coverage for certain abortion services.

DATES: This final rule is effective on [insert date 60 days after the date of publication in the Federal Register].

FOR FURTHER INFORMATION CONTACT:

Emily Ames, (301) 492-4246 or Marisa Beatley, (301) 492-4307.

SUPPLEMENTARY INFORMATION:

I. Background

A. Legislative and Regulatory Overview

Sections 1311(b) and 1321(b) of the Patient Protection and Affordable Care Act (PPACA) provide that each state has the opportunity to establish an Exchange. Section 1311(b)(1) of the PPACA gives each state the opportunity to establish an Exchange that facilitates the purchase of qualified health programs (QHPs) by individuals and families,
and provides for the establishment of a Small Business Health Options Program (SHOP) that is designed to assist qualified small employers in the state in facilitating the enrollment of their employees in QHPs offered in the small group market in the state.

Section 1313 of the PPACA describes the steps the Secretary of Health and Human Services (the Secretary) may take to oversee Exchanges’ compliance with HHS standards related to title I of the PPACA and ensure their financial integrity, including conducting investigations and annual audits.

Section 1321(a) of the PPACA provides broad authority for the Secretary to establish standards and regulations to implement the statutory standards related to Exchanges, QHPs, and other identified standards of title I of the PPACA.

Section 1321(c)(2) of the PPACA authorizes the Secretary to enforce the Exchange standards using civil money penalties (CMPs) on the same basis as detailed in section 2723(b) of the Public Health Service Act (PHS Act). Section 2723(b) of the PHS Act authorizes the Secretary to impose CMPs as a means of enforcing the individual and group market reforms contained in Part A of title XXVII of the PHS Act when a state fails to substantially enforce these provisions with respect to health insurance issuers.

Section 1303 of the PPACA, as implemented in 45 CFR 156.280, specifies standards for issuers of qualified health plans (QHPs) through the Exchanges that cover abortion services for which public funding is prohibited (also referred to as non-Hyde abortion services). The statute and regulation establish that, unless otherwise prohibited by state law, a QHP issuer may elect to cover such non-Hyde abortion services. If an issuer elects to cover such services under a QHP sold through an individual market Exchange, the issuer must take certain steps to ensure that no premium tax credit (PTC)
or cost-sharing reduction (CSR) funds are used to pay for abortion services for which public funding is prohibited.

As specified in section 1303(b)(2), one such step is that individual market Exchange issuers must determine the amount of, and collect, from each enrollee, a separate payment for an amount equal to the actuarial value of the coverage for abortions for which public funding is prohibited, which must be no less than $1 per enrollee, per month. QHP issuers must also segregate funds for non-Hyde abortion services collected through this payment into a separate allocation account used to pay for non-Hyde abortion services.

Section 1411(c) of the PPACA requires the Secretary to submit certain information provided by applicants under section 1411(b) of the PPACA to other federal officials for verification, including income and family size information to the Secretary of the Treasury.

Section 1411(d) of the PPACA provides that the Secretary must verify the accuracy of information provided by applicants under section 1411(b) of the PPACA for which section 1411(c) does not prescribe a specific verification procedure, in such manner as the Secretary determines appropriate.

Section 1411(f)(1)(B) of the PPACA requires the Secretary to establish procedures to redetermine eligibility on a periodic basis, in appropriate circumstances, including for eligibility to purchase a QHP through the Exchange and for advance payments of the premium tax credit (APTC) and CSRs.

Section 1411(g) of the PPACA allows the exchange of applicant information only for the limited purposes of, and to the extent necessary to, ensure the efficient operation
of the Exchange, including by verifying eligibility to enroll through the Exchange and for APTC and CSRs.

On October 30, 2013, we published a final rule entitled, “Patient Protection and Affordable Care Act: Program Integrity: Exchange, Premium Stabilization Programs, and Market Standards; Amendments to the HHS Notice of Benefit and Payment Parameters for 2014,” (78 FR 65046), to implement certain program integrity standards and oversight requirements for State Exchanges.

On March 27, 2012, we published a final rule entitled “Establishment of Exchanges and Qualified Health Plans; Exchange Standards for Employers,” (Exchange Establishment Rule (77 FR 18309), in which we codified the statutory provisions of section 1303 of the PPACA in regulation at 45 CFR 156.280, and established many standards related to Exchanges. On February 27, 2015, we published the Patient Protection and Affordable Care Act; HHS Notice of Benefit and Payment Parameters for 2016, final rule (80 FR 10750) (hereinafter referred to as the 2016 Payment Notice) providing guidance regarding acceptable billing and premium collection methods for the portion of the policy holder’s total premium attributable to non-Hyde abortion coverage for purposes of satisfying the statutory separate payment requirement.

On March 8, 2016, we published the Patient Protection and Affordable Care Act; HHS Notice of Benefit and Payment Parameters for 2017, final rule (81 FR 12204), in which we provided issuers the option to adopt a premium payment threshold policy to avoid situations in which an enrollee who owes only a de minimis amount of premium has his or her enrollment terminated for non-payment of premiums.

On November 9, 2018, we published a proposed rule entitled “Patient Protection...
and Affordable Care Act; Exchange Program Integrity” (83 FR 56015), which proposed to revise standards relating to oversight of Exchanges established by states and periodic data matching frequency and authority. It also proposed new requirements for certain issuers related to the billing and collection of the separate payment for the premium portion attributable to coverage for certain abortion services.

B. Stakeholder Consultation and Input

HHS has consulted with stakeholders on policies related to the operation of Exchanges. We have held a number of listening sessions with consumers, providers, employers, health plans, the actuarial community, and state representatives to gather public input, with a particular focus on risks to the individual and small group markets, and how we can alleviate burdens facing patients and issuers. We consulted with stakeholders through regular meetings with the National Association of Insurance Commissioners, regular contact with State Exchanges through the Exchange Blueprint process and ongoing oversight and technical assistance engagements, and meetings with Tribal leaders and representatives, health insurance issuers, trade groups, consumer advocates, employers, and other interested parties.

II. Provisions of the Proposed Rule and Analysis of and Responses to Public Comments

A. Exchange Establishment Standards and Other Related Standards Under the Affordable Care Act

1. Functions of an Exchange (§155.200)

We proposed to revise §155.200 to clarify that the Exchanges must perform oversight functions generally, and cooperate with oversight activities, in accordance with
section 1313 of the PPACA and as required under 45 CFR part 155. Section 155.200 describes the functions that an Exchange must perform. Section 155.200(c) specifies that the Exchange must perform functions related to oversight and financial integrity in accordance with section 1313 of the PPACA. HHS interprets this requirement broadly to include program integrity functions related to protecting against fraud, waste, and abuse, including functions not explicitly identified in section 1313 of the PPACA. We believe State Exchanges, including State Exchanges on the Federal Platform (SBE-FPs), have also generally interpreted this requirement broadly, as evidenced by their engagement in activities designed to combat fraud and abuse.

However, questions about the breadth of this function have arisen when Exchanges have sought to understand what uses and disclosures of personally identifiable information (PII) are permitted under §155.260. Specifically, we received questions about whether Exchanges are permitted under §155.260 to disclose applicant PII to government oversight entities, such as state departments of insurance, when investigating fraudulent behavior related to Exchange enrollments on the part of agents and brokers.

As noted in the proposed rule, we believe that use and disclosure of PII related to Exchange program integrity efforts, such as combatting fraud, currently fall under §155.200(c), but seek to make that position more clear. Therefore, we proposed to revise §155.200(c) to clarify that the Exchanges must perform oversight functions generally, and cooperate with oversight activities, in accordance with section 1313 of the PPACA.

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1 Section 155.260 limits an Exchange’s use and disclosure of PII when an Exchange creates or collects personally identifiable information for the purposes of determining eligibility for enrollment in a qualified health plan; determining eligibility for other insurance affordability programs, as defined in §155.300; or determining eligibility for exemptions from the individual shared responsibility provisions in section 5000A of the Internal Revenue Code. One of the permitted uses and disclosures is for the Exchange to carry out the functions described in §155.200.
and as required under 45 CFR part 155, including overseeing its Exchange programs, Navigators, agents, brokers, and other non-Exchange entities as defined in §155.260(b). We further explained that because this is a clarification and not a new function, we did not believe it would impose additional burdens on Exchanges, but instead would help resolve questions about the available tools and authority to enable Exchanges to effectively oversee and combat potentially fraudulent behavior.

After consideration of comments received, we are finalizing this provision as proposed, with one technical modification to remove a redundant term included in the proposed regulation text. The comments we received on this topic are summarized below, along with our responses.

**Comment:** All commenters on this topic supported the proposed amendment to §155.200(c) as it clarifies that oversight and transparency for all Exchanges is required with respect to determining eligibility for APTC and combatting fraud. Two commenters encouraged HHS to work closely with states once the proposal is finalized to ensure that individuals who are assisting consumers receive proper notice and training on the applicable compliance requirements and standards in their states. One commenter suggested that HHS solicit stakeholder feedback on the possibility of incorporating an additional level of collaborative issuer-Exchange oversight and verification prior to enrollment when the applicant’s coverage has been previously terminated for fraud.

**Response:** We remain committed to improving Exchange program integrity, including efforts related to combatting fraud, and appreciate commenters’ support for our clarification that Exchanges are permitted to use and disclose applicant PII to certain entities for these efforts. We agree that it is important for agents, brokers, Navigators,
and other assisters to understand the applicable standards in their state, and plan to work closely with states to ensure compliance. We continue to explore other pathways for combatting fraud in Exchanges and appreciate commenters’ recommendations.

We are finalizing the amendment to §155.200(c) as proposed, with one modification. We are removing the reference to assisters because it is redundant of the reference to non-Exchange entities. Non-Exchange entities are defined in §155.260(b) and include Navigators, non-Navigator assistance personnel, certified application counselors, agents, brokers, web-brokers and other individuals or entities who gain access to PII submitted to an Exchange or collect, use or disclose PII gathered directly from Exchange applicants or enrollees.

2. Verification process related to eligibility for insurance affordability programs (§155.320)

We requested comment on our proposed plans to expand the current scope of Medicare periodic data matching (PDM), which only identifies and notifies those dual enrollees receiving financial assistance, to also include the Exchange population not receiving financial assistance. Specifically, we proposed to add a new authorization compliant with Health Insurance Portability and Accountability Act of 1996 (HIPAA) (Pub. L. 104-191) standards to the single streamlined application to permit Exchanges using the federal platform to collect PHI in order to determine enrollees’ Medicare enrollment status. We also proposed to leverage the current attestation question on the single, streamlined application, for applicants to provide written consent permitting the Exchange to terminate their coverage if they are found later to be dually enrolled in Medicare and a QHP to expand the scope of Medicare PDM to the population not
receiving financial assistance. We will not finalize these proposed actions, but will continue to identify and notify dual enrollees receiving financial assistance as part of current Medicare PDM operations.

Under §155.330, Exchanges are required to periodically examine available data sources to identify whether enrollees on whose behalf APTC or CSRs are being paid have been found eligible for or are enrolled in Medicare, Medicaid, Children's Health Insurance Program (CHIP), or the Basic Health Program (BHP), if a BHP is operating in the service area of the Exchange. Individuals identified as enrolled both in Exchange coverage (with or without APTC or CSRs) and one of these other forms of coverage are referred to as dually enrolled consumers. Generally, if an individual is eligible for or enrolled in such other forms of coverage that qualify as minimum essential coverage (MEC) under section 5000A of the Code, the individual is not eligible to receive APTC or CSRs. For instance, if an individual is eligible for premium-free Medicare Part A or enrolled in Medicare Part A or Part C (also known as Medicare Advantage), all of which qualify as MEC, he or she is not eligible to receive APTC or CSRs to help pay for an Exchange plan or covered services.

The Secretary has broad authority under section 1321(a) of the PPACA to establish regulations setting standards to implement certain statutory requirements under title I of the PPACA, including with respect to the establishment and operation of Exchanges, the offering of QHPs through the Exchanges, the establishment of the risk adjustment and reinsurance programs, and such other requirements as the Secretary determines appropriate. Additionally, section 1411(g) of the PPACA allows the exchange of certain applicant information as necessary to ensure the efficient operation of
the Exchange, including verifying eligibility to enroll in coverage through the Exchange and to receive APTC or CSRs.

Furthermore, 45 CFR 155.430(b)(1)(ii) requires an Exchange to provide an opportunity at the time of plan selection for enrollees receiving and not receiving financial assistance to choose to remain enrolled in a QHP if he or she becomes eligible for other MEC, or to terminate QHP coverage if the enrollee does not choose to remain enrolled in the QHP upon completion of the redetermination process. As such, for plan year 2018 and thereafter, we added language to the existing single, streamlined application to support compliance with this requirement by all Exchanges using the federal platform. This new language allows all consumers, regardless of whether they are seeking financial assistance, to authorize the Exchange to obtain eligibility and enrollment data and, if so desired by the consumer, to end their QHP coverage if the Exchange finds during its periodic eligibility checks that the consumer has become eligible for or enrolled in other MEC, such as Medicare, Medicaid/CHIP, or BHP.

In addition, for plan years beginning with the 2020 plan year, we stated in the proposed rule our intention to add a new HIPAA authorization to the single, streamlined application used by Exchanges using the federal platform, which would meet HIPAA standards regarding how one’s protected health information (PHI) is collected and used. In the preamble to the proposed rule, we discussed using this proposed new HIPAA authorization to expand the current scope of Medicare PDM to individuals in the Exchange population who are not receiving financial assistance and who authorize the Exchanges using the federal platform to conduct certain PDM by requesting PHI from HHS such as their name, Social Security Number, Medicare eligibility or enrollment
status, and other data elements the Exchange may determine necessary, to allow the Exchange to determine whether the consumer is dually enrolled in Medicare and Exchange coverage. This HIPAA authorization would allow HHS to check Medicare enrollment databases for applicants regardless of whether they seek or receive financial assistance.

As we discussed in the preamble to the proposed rule, for consumers who request voluntary termination upon a finding of dual enrollment, the Exchange would terminate coverage after following the current PDM process outlined in §155.330(e)(2)(i), which requires Exchanges to provide notice of the updated information the Exchange found, as well as a 30-day period for the enrollee to respond to the notice. We emphasize again, because the Exchange cannot identify through this process those consumers who are eligible for, but not enrolled in premium-free Part A, we encourage all consumers who are 65 and older to apply with the Social Security Administration (SSA) to receive an eligibility determination with respect to Medicare.

We received multiple comments on this discussion regarding expanding the scope of Medicare PDM to the Exchange population not receiving financial assistance. After further consideration of the technical complexity of implementing a HIPAA authorization on the single, streamlined application and the potential burden on consumers to read, decipher, and agree to legal agreements many may find confusing, we will not pursue the addition of a new authorization to the single streamlined application. Instead, we will explore other means through which the Exchanges can expand the scope of Medicare PDM to the Exchange population that is not receiving financial assistance. A summary of these comments and our responses to those comments follow:
Comment: Most commenters generally supported HHS’s goal to reduce dual enrollment in Medicare and Exchange coverage, but cautioned HHS about the consequences of terminating QHP coverage for this population. Commenters noted that terminating Exchange coverage could: (1) interfere with the continuity of care, (2) create gaps in coverage, especially for those dual enrollees who have not yet enrolled in Medicare Part B, (3) cause other family members on the Medicare beneficiary’s policy to lose coverage, and (4) cause increased consumer confusion over their coverage options. Rather than terminating QHP coverage, commenters recommended targeted outreach and education to the Medicare eligible population to ensure this population fully understands the consequences of dual enrollment, the appropriate time to enroll in Medicare Part B to avoid financial penalties for delayed enrollment, and how access to their Medicare eligibility information intersects with QHP termination via Medicare PDM. One commenter recommended that we prevent all individuals with Medicare from enrolling in QHP coverage through screening at initial application.

Response: Given the technical complexity of implementing a HIPAA authorization on the single streamlined application and the potential burden it would place on consumers as consumers would be required to read, decipher, and agree to complex legal agreements that may be confusing for consumers, we are reconsidering our approach to expanding Medicare PDM to the Exchange population not receiving financial assistance. We are exploring other options to identify and notify this population of their dual enrollment in Medicare and Exchange coverage to ensure that this population is able to enroll in Medicare Part B at the appropriate time and without financial penalty.
For enrollees in Exchanges using the federal platform who are receiving financial assistance, the Exchanges will continue to end subsidies or QHP coverage for those consumers who permit the Exchange to do so in accordance with §155.330. For the Exchange population receiving financial assistance, terminating QHP coverage as part of Medicare PDM ensures that consumers are not enrolled in unnecessary duplicative coverage, reduces the potential for taxpayer financial liability related to possibly having to repay APTC at the time of federal income tax reconciliation, and also protects the integrity of the Exchange by ensuring enrollees no longer eligible for financial assistance do not receive these subsidies inappropriately.

HHS is also aware of concerns from stakeholders that consumers often do not know when they should contact the Exchange to end their QHP coverage after enrolling in Medicare. We believe this voluntary option to provide written consent for the Exchange to end a Medicare dual enrollee’s QHP coverage will alleviate some of the confusion consumers currently face when transitioning from Exchange coverage to Medicare as the Exchange provides information in the initial warning notice on how to end QHP coverage after enrolling in Medicare. Furthermore, in instances where the dual enrollee does not take action, the Exchange will automatically end coverage for the dual enrollee; thus, saving the enrollee time and reducing the risk of the consumer having to pay back some or all of the APTC received when they file their federal income taxes.

In addition, in response to commenter concerns about the consequences of termination of dually enrolled consumers’ coverage, we note that enrollees receiving financial assistance have 30 days to respond to their Medicare PDM notice before the Exchange takes action as specified in §155.330(e)(2)(i)(D). As we noted in the preamble
to the proposed rule, upon receiving the required notice, the enrollee could (1) return to the Exchange and terminate his or her QHP coverage, (2) revoke the prior authorization for the Exchange to terminate his or her QHP coverage in the event dual enrollment is found, so that he or she would remain enrolled both in the QHP and in Medicare, or (3) notify the Exchange that he or she is not eligible for, or enrolled in, Medicare. For enrollees who revoke their prior authorization for the Exchange to terminate their QHP enrollment where the Exchange finds the enrollee is eligible for or enrolled in Medicare, or who disagree that they are eligible for or enrolled in Medicare, the Exchange would only proceed to terminate the enrollee’s APTC and CSRs, and not his or her enrollment in QHP coverage through the Exchange, using the process specified in §155.330(e)(2)(i). Therefore, we believe this operational change mitigates adverse impacts on the continuity of care and the risk of coverage gaps because enrollees can choose to opt out and remain in QHP coverage without APTC, pursuant to the current regulation.

We also appreciate the concerns raised that non-Medicare family members could potentially lose coverage. We note that a special enrollment period will be available for family members of dual enrollees when such family members lose their coverage or their financial subsidies as a result of the PDM process described here.

Additionally, we continue to prioritize consumer and stakeholder education regarding dual enrollment and transitioning between coverage, and to engage in various outreach activities including distributing webinar, newsletter, and fact sheet content for assisters, agents, brokers, and issuers, as well as direct consumer notification and application help text. We also are working to develop educational materials to ensure that all Medicare beneficiaries understand the consequences of dual enrollment and
associated penalties for not enrolling in Medicare Part B when first eligible. We believe this will help reduce consumer confusion over their coverage options and the appropriate time to sign up for Medicare. We appreciate the comments and ideas for future education efforts for this population and will consider these suggestions as part of our Medicare PDM stakeholder outreach moving forward.

3. Eligibility redetermination during a benefit year (§155.330)

We proposed to add a new paragraph (d)(3) to §155.330, under which Exchanges would be required to conduct PDM at least twice each calendar year beginning with calendar year 2020. We are finalizing this proposal. However, we have changed the implementation date to the 2021 calendar year, and added clarifying language regarding State Exchanges that have fully integrated eligibility systems with their respective Medicaid agencies.

In accordance with §155.330(d), Exchanges must periodically examine available data sources to determine whether enrollees in a QHP through an Exchange with APTC or CSRs have been determined eligible for or enrolled in other qualifying coverage through Medicare, Medicaid, CHIP, or the BHP, if applicable. HHS has not previously defined “periodically.” Currently, Exchanges using the federal platform conduct Medicare PDM and Medicaid/CHIP PDM twice a year. To ensure that all Exchanges are taking adequate steps to identify enrollees who have become eligible for or enrolled in these other forms of MEC, and to terminate APTC and CSRs for those identified, we proposed to add paragraph (d)(3) to specify that Exchanges would be required to conduct Medicare, Medicaid/CHIP, and, if applicable, BHP PDM at least twice a calendar year, beginning with the 2020 calendar year. We indicated that this timeframe would likely
give Exchanges that are not already performing these PDM checks twice a year sufficient
time to implement any business, operational, and information technology changes needed
to comply with the proposed new requirement.

We explained our belief that this policy would reduce QHP premiums, since
Medicare and Medicaid/CHIP beneficiaries tend to have a higher risk profile than a
typical Exchange enrollee and, therefore, may have negative impacts on the risk pool.
Because this population includes significant numbers of older and disabled beneficiaries,
or persons that may have poorer health outcomes generally associated with lower income
 statuses, we expect that these populations typically will utilize health care services at a
greater rate as compared to other populations.\(^2\) So that the Exchanges could prioritize the
implementation of the proposed requirement to conduct PDM for Medicare, Medicaid,
CHIP, and, if applicable, BHP eligibility or enrollment at least twice yearly, we did not
also propose requiring Exchanges to perform PDM for death at least twice in a calendar
year, and will consider this as part of future rulemaking.

Since most State Exchanges that operate their own eligibility and enrollment
platform have a single shared, integrated eligibility system with their respective Medicaid
programs, the Medicaid/CHIP PDM requirements may be met differently by State
Exchanges. State Exchanges that have fully integrated eligibility systems generally have
controls in place to prevent concurrent or dual enrollment of an individual in both a QHP
through the Exchange with APTC/CSRs, and Modified Adjusted Gross Income
(MAGI)-based Medicaid/CHIP coverage, at any given time. We proposed at paragraph

\(^2\) For example, see Urban Institute and Center on Society and Health, How Are Income and Wealth Linked
to Health and Longevity? (April 2015), available at
(d)(3) that we will deem these State Exchanges to be in compliance with the requirement to perform Medicaid/CHIP PDM or, if applicable, BHP PDM. Thus, these State Exchanges would not need to perform additional Medicaid/CHIP PDM outside of the controls that are currently in place to prevent dual enrollment in their integrated eligibility system. State Exchanges that operate their own eligibility and enrollment platform and do not have fully integrated eligibility systems for APTC/CSRs and Medicaid/CHIP or BHP, if applicable, would be required to perform Medicaid/CHIP PDM at least twice a year.

We anticipate many State Exchanges will meet or exceed the proposed requirements for Medicare PDM, Medicaid/CHIP PDM and, if applicable, BHP PDM, based on operations reported to us through the State-based Marketplace Annual Reporting Tool (SMART). This view is also supported by information we have learned through technical assistance engagements. Furthermore, the new Medicaid/CHIP PDM requirement would not result in a significant administrative burden for State Exchanges because we believe most State Exchanges currently operate an integrated eligibility system and could be deemed to be in compliance with the proposed Medicaid/CHIP PDM requirements.

We did not propose specific penalties if State Exchanges do not comply with the proposed PDM requirements. However, we noted that, under current authority, HHS requires a State Exchange to take corrective action if it is not complying with applicable federal requirements. We utilize specific oversight tools (SMART, programmatic audits, etc., as described in the preamble to §155.1200) to identify issues with, and place corrective actions on, the Exchanges, and to provide technical assistance and ongoing
monitoring to track those actions until the Exchange comes into compliance.

Additionally, under section 1313(a)(4) of the PPACA, if HHS determines that an Exchange has engaged in serious misconduct with respect to compliance with Exchange requirements, it has the option to rescind up to 1 percent of payments due to a state under any program administered by HHS until it is resolved. These existing authorities would apply to the proposed periodic data matching requirements in §155.330(d). If HHS were to determine that it is necessary to apply this authority due to non-compliance by an Exchange with §155.330(d), HHS would also determine the HHS-administered program from which it would rescind payments that are due to that state.

Lastly, we proposed to make a technical correction in §155.330(d)(1) by adding an additional reference to the process and authority in §155.320(b). This reference was omitted previously, but the requirements in §155.320(b), specifying that Exchanges must verify whether an applicant is eligible for MEC other than through an eligible employer-sponsored plan using information obtained by transmitting identifying information specified by HHS to HHS for verification purposes, apply to the PDM process in §155.330.

We are finalizing this proposal to add paragraph (d)(3) as proposed, but have changed the implementation date to the 2021 calendar year, and have added some clarifying language with regard to fully integrated eligibility systems, as described below. A summary of comments received and our responses to those comments appear below.

Comment: We received multiple comments in support of PDM as an effort to improve Exchange program integrity. These commenters agreed that the process helps inform consumers of their enrollment in potentially duplicative other MEC such as
certain Medicare and Medicaid coverage, CHIP, or, if applicable, the BHP, and to help consumers avoid a tax liability for having to repay APTC received during months of overlapping coverage when reconciling at the time of annual federal income tax filing. Many commenters suggested improvements that could be made to current PDM processes.

Some commenters suggested that consumers, especially Medicare beneficiaries, could benefit from additional education or outreach from assisters, Navigators, or call center representatives to help these dually enrolled consumers make informed choices about their coverage options. Another commenter recommended that HHS work closely with SSA to identify which Medicare beneficiaries are approaching Medicare eligibility so that notices can be sent during the beneficiary’s initial enrollment period. Another commenter recommended that, in addition to periodic checks for other qualifying coverage, HHS should implement periodic checks for deceased enrollees and that these checks should occur before auto re-enrollment.

Response: We agree that the PDM process is an important tool to ensure that Exchange enrollees are enrolled in the appropriate coverage that best meets their needs and budget while reducing the risk for potential tax liabilities for having to repay APTC received during months of overlapping coverage. We also agree that outreach and education is critical for dual enrollees and we continue to work with Exchange stakeholders on education and outreach strategies, especially for the Medicare beneficiary population to ensure that consumers can make well-informed choices and sign up for Medicare coverage during the appropriate timeframes. In 2018, we added additional resources to the Exchange application that provided information on the appropriate
timeframes to enroll in Medicare Parts A and B to help consumers avoid incurring any late enrollment penalties. We also believe that periodic checks for deceased enrollees are a critical aspect to ensuring Exchange program integrity. Beginning in late 2019, Exchanges using the federal platform will conduct periodic checks for deceased enrollees in single member applications and subsequently end deceased enrollees’ QHP coverage. As noted previously, to ensure State Exchanges have appropriate time to implement the technical and operational changes necessary to conduct Medicare, Medicaid/CHIP, and, if applicable, BHP, PDM, we are not requiring that State Exchanges perform checks for deceased enrollees twice yearly, and will be considering changes as part of future rulemaking.

**Comment:** We received mixed comments regarding our proposal to require Exchanges to conduct Medicare, Medicaid/CHIP and, if applicable, BHP PDM twice a year. Many commenters stated that increasing the frequency of PDM, particularly Medicare PDM, may be burdensome on both consumers and State Exchanges, and could lead to increased consumer confusion, diversion of resources from customer service and outreach efforts, and potential loss of APTC due to potentially outdated data sources for Medicare enrollment and Medicaid/CHIP eligibility and enrollment. One commenter recommended that additional verification checks be incorporated into the final rule to ensure consumers are not removed from coverage due to outdated data. Two commenters noted that the twice yearly frequency was too infrequent and would not provide timely notice for those consumers who are dually enrolled in Medicare and Exchange coverage. One commenter recommended requiring that Exchanges only perform PDM checks once yearly, which taken together with the annual renewal process, would allow a check every
6 months. Another commenter expressed concerns that our proposed language would allow State Exchanges to perform PDM more than twice a year, which could cause consumers to lose coverage erroneously.

**Response:** We continue to believe that conducting Medicare, Medicaid/CHIP and, if applicable, BHP PDM serves a critical role in ensuring that consumers are enrolled in the appropriate coverage and ensures that APTC is paid appropriately. We continue to work with our partners throughout HHS to ensure the accuracy of Medicare, Medicaid, and CHIP data, and will continue to provide guidance to State Exchanges on notice language, especially regarding the availability of special enrollment periods for consumers who erroneously lose APTC or QHP coverage, as well as the consumer’s right to appeal an Exchanges’ determination. We disagree that conducting PDM checks twice yearly would cause consumer confusion or divert resources away from customer service and outreach because PDM provides valuable information to consumers regarding their dual enrollment in Medicare and/or Medicaid/CHIP and serves an important program integrity function by ensuring that only consumers eligible for APTC/CSRs receive them. We continue to prioritize consumer and stakeholder education related to dual enrollment and transitioning between coverage, including webinar, newsletter, and fact sheet content for assisters, agents, brokers, and issuers, as well as direct consumer notification and application help text. We encourage State Exchanges to prioritize these education efforts as well.

We appreciate commenters’ suggestions regarding the frequency of PDM checks, but we believe that requiring these checks at least twice a year strikes the appropriate balance between providing timely notice for dually enrolled consumers and not
overburdening Exchanges with potentially costly system changes and notice requirements. With respect to the comment regarding Exchanges conducting a Medicaid/CHIP or Medicare PDM check during the annual renewal process, this rule specifies the frequency, and not the precise timing, for when Exchanges must conduct the Medicaid/CHIP and Medicare PDM checks. Exchanges have the flexibility to conduct one of the required PDM checks during the annual renewal process.

Finally, we disagree that the changes outlined to PDM would increase burden on all Exchanges. We will deem State Exchanges that have implemented fully integrated eligibility systems with their respective Medicaid programs to be in compliance with the proposed Medicaid/CHIP PDM requirement. Thus, we anticipate the change to the Medicaid/CHIP PDM requirement will not increase burden for those State Exchanges because they will not have to build new functionality to meet this requirement. However, we do agree that any significant burden on State Exchanges would likely be on those that currently do not perform any Medicare PDM, or those that currently do not operate integrated eligibility systems and do not perform any Medicaid/CHIP PDM and, therefore, are not already in compliance with §155.330(d). Those Exchanges would likely be required to engage in information technology (IT) system development activities in order to communicate with these programs and act on enrollment data in a new way.

Comment: We received multiple comments that the proposed date of January 1, 2020 for the implementation of twice yearly Medicare, Medicaid/CHIP, and, if applicable, BHP PDM provides insufficient time for State Exchanges to implement the required technical changes. Commenters noted that State Exchanges that do not currently
conduct Medicare PDM, or do not have integrated eligibility systems with their State Medicaid programs and do not currently conduct Medicaid/CHIP PDM, would have to make significant changes to their eligibility systems and processes to confirm enrollment in Medicare or to verify Medicaid or CHIP eligibility, respectively. One commenter suggested 2021 as an appropriate implementation date. Two commenters also requested that HHS finalize a clear and certain definition of a fully integrated eligibility system to mean eligibility systems that have one eligibility rules engine, shared between the State Exchange and its respective Medicaid program, for MAGI-based Medicaid, CHIP, APTC, and if applicable, BHP, eligibility determinations.

Response: We agree with commenters that requiring implementation by the 2020 calendar year may not provide State Exchanges with a sufficient timeframe to implement these changes, especially for Exchanges without integrated eligibility systems that do not currently perform Medicaid/CHIP PDM or those that currently do not perform Medicare PDM. These Exchanges would need to implement new interfaces with their respective Medicaid programs and/or a new connection to federal data to confirm Medicare enrollment. Therefore, we are finalizing the proposal in §155.330(d)(3) to take effect beginning with the 2021 calendar year. We also agree on the importance of providing a clear and specific definition of “fully integrated eligibility system.” As described in the preamble to the proposed rule, by “fully integrated eligibility system,” we mean one where a State Exchange and its respective Medicaid program shares a single eligibility rules engine for determining eligibility for MAGI-based Medicaid/CHIP, APTC, and if applicable, BHP. We are finalizing paragraph (d)(3) with some additional language to codify this meaning.
**Comment:** We received three comments that were opposed to the proposed requirement to conduct Medicare, Medicaid/CHIP and, if applicable, BHP PDM, cautioning us that defining the precise frequency and nature of PDM encroaches upon the sovereignty of the State Exchanges. Two commenters noted that HHS has not provided enough evidence that there is a significant problem with duplicative enrollment in other qualifying coverage such as Medicare, Medicaid/CHIP, and BHP. One commenter expressed concern that additional requirements on State Exchanges could discourage consumers from applying for coverage.

**Response:** Ensuring that consumers are enrolled in the appropriate coverage remains a top priority for HHS. Additionally, ensuring that APTC is paid appropriately is a requirement set forth in §155.330(d)(1)(ii). Several Government Accountability Office (GAO) reviews have underscored the importance of continually re-verifying enrollee eligibility for APTC through PDM with other government entities.\(^3\) As such, we believe PDM plays a vital role in ensuring the health and integrity of all Exchanges by ensuring consumers are enrolled in the appropriate coverage, and reduces the risk that consumers will have to pay back all or some of APTC paid on their behalf during months of overlapping coverage when they file their annual federal income taxes. We disagree that the twice yearly requirement to conduct Medicare, Medicaid/CHIP and, if applicable, BHP PDM would discourage consumers from applying for and enrolling in QHP coverage, as the majority of consumers become dually enrolled inadvertently, such as by aging into Medicare or experiencing fluctuations in household income.

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4. General program integrity and oversight requirements (§155.1200)

As the Exchange Establishment grant program established under section 1311 of the PPACA has come to a conclusion and State Exchanges have become financially self-sustaining, HHS continues to develop and refine its mechanisms and tools for overseeing the ongoing compliance of State Exchanges and SBE-FPs with federal requirements for Exchanges, including eligibility and enrollment requirements under 45 CFR part 155.

HHS approves or conditionally approves a state to establish a State Exchange based on an assessment of a state’s attested compliance with applicable statutory and regulatory rules. Once approved or conditionally approved, State Exchanges must meet specific program integrity and oversight requirements identified at section 1313(a) of the PPACA, and the implementing regulations at §§155.1200 and 155.1210. These requirements outline HHS’s authority to oversee the Exchanges after their establishment. Currently, annual reporting requirements for State Exchanges at §155.1200(b) include the annual submission of (1) a financial statement in accordance with generally accepted accounting principles (GAAP); (2) eligibility and enrollment reports; and (3) performance monitoring data.

Additionally, under §155.1200(c), each State Exchange is required to contract with an independent external auditing entity that follows generally accepted government auditing standards (GAGAS) to perform annual independent external financial and programmatic audits. State Exchanges are required to provide HHS with the results of the annual external audits, including corrective action plans to address any material
weaknesses or significant deficiencies identified by the auditor. All corrective action plans are monitored by HHS until closed. Currently, the audits must address compliance with all Exchange requirements under 45 CFR part 155.

HHS designed and developed the SMART in 2014 to assist State Exchanges in conducting a defined set of oversight activities. The SMART was designed to facilitate State Exchanges’ reporting to HHS on how they are meeting federal program and operational requirements, including State Exchanges reporting their compliance with federal eligibility and enrollment program requirements under 45 CFR part 155 subparts D and E. The SMART, thus, enables HHS to evaluate and monitor State Exchange progress in coming into compliance with federal requirements where needed. Since then, HHS has come to utilize the SMART, along with the annual programmatic and financial audit reports, as primary oversight tools for identifying and addressing State Exchange non-compliance issues. HHS requires State Exchanges to take corrective actions to address issues that are identified through the SMART and annual audits, and HHS monitors the implementation of the corrective actions.

In the proposed rule, we proposed to modify §155.1200(b)(2) to reflect that HHS requires State Exchanges to submit annual compliance reports (such as the SMART), that encompass eligibility and enrollment reporting by State Exchanges, and also include reporting on compliance across other Exchange program requirements under 45 CFR part 155. We also proposed to modify §155.1200(b)(1) to eliminate the April 1st date by which State Exchanges must provide a financial statement to HHS, to provide HHS the flexibility to align the financial statement deadline with the SMART deadline,

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4 45 CFR 155.1200(c)(1) and (2).
5 45 CFR 155.1200(d)(2).
which is set annually by HHS. Because we proposed to remove the April 1st date, but intend to maintain the requirement that State Exchanges submit the required reports by a deadline, we also proposed to modify the introductory text to §155.1200(b) to specify that State Exchanges must provide the required annual reporting by deadlines to be set by HHS.

We proposed to retain the requirement at §155.1200(c) that an annual programmatic audit be conducted by State Exchanges, but proposed a minor change from “state” to “State Exchanges” to be consistent and clear on the entities to which this rule applies. We also proposed to add specificity to the annual programmatic audit requirement by proposing a clarification of §155.1200(d)(2) to make clear that HHS may specify or target the scope of a programmatic audit to address compliance with particular Exchange program areas or requirements. We explained that this would provide HHS with the ability to specify those Exchange functions that are most pertinent to a particular State Exchange model (either a traditional State Exchange that operates its own eligibility and enrollment system or an SBE-FP) and need to be regularly included in the audit; target those Exchange functions most likely to impact program integrity, such as eligibility verifications; and reduce burden on State Exchanges where possible. In addition, we proposed to modify §155.1200(d) by replacing existing paragraph (d)(4) with new paragraphs (d)(4) and (5). These proposed new requirements specify that State Exchanges must ensure that the independent audits implement testing procedures or other auditing procedures that assess whether a State Exchange is conducting accurate eligibility determinations and enrollment transactions under 45 CFR part 155 subparts D and E. Such auditing procedures can include the use of statistically valid sampling
methods in the testing or auditing procedures.

We indicated that we believe these proposed changes would strengthen our programmatic oversight and the program integrity of State Exchanges, while providing flexibility for HHS in the collection of information. We further explained that, through the Paperwork Reduction Act (PRA) process, we are able to make updates and refinements to the SMART reporting tool to align with our program integrity priorities for Exchanges as they evolve. In addition, allowing HHS to specify the scope of the programmatic audit at §155.1200(d)(2) would provide us the ability to target our oversight to specific Exchange program requirements based on the particular State Exchange model, our program integrity priorities, and the goal of reducing burden on State Exchanges where possible. We explained our belief that this approach would provide HHS and states with greater insight into State Exchange compliance with federal standards in a more cost-effective manner.

We also noted our belief that this approach would allow HHS to identify State Exchange non-compliance issues with more precision and efficacy. It would allow HHS to provide more effective, targeted technical assistance to State Exchanges in developing corrective action plans to address issues that are identified. We discussed how this approach could reduce administrative burden on State Exchanges while maintaining the traditional role of State Exchanges in managing and operating their Exchanges, with HHS maintaining its role of overseeing State Exchange compliance with federal requirements through structured reporting processes. We sought comments on these proposals.
After consideration of comments received, we are finalizing the amendments to §155.1200 as proposed. A summary of comments received and our responses to those comments appear below:

Comment: Commenters generally expressed support for some of the proposed changes to the annual reporting and programmatic audit requirement. They expressed support for removal of the April 1st financial statement deadline as long as the new deadline accommodates the state budget cycles for all State Exchanges. Some commenters supported the proposal to provide flexibility to specify the scope of the programmatic audit, such as focusing on eligibility and enrollment requirements under 45 CFR part 155 subparts D and E, while two commenters asked HHS to refrain from expanding the scope of the programmatic audit as it can divert funding from other Exchange functions and create administrative burden. Some commenters expressed concern with the timing and potential funding to implement the changes. These commenters urged HHS to provide State Exchanges with over a year of advanced notice to implement the changes, to ensure proper planning and funding.

One commenter requested that HHS clarify the proposed requirement for the State Exchange’s independent external auditor to use statistically valid sampling in their review of the State Exchange eligibility and enrollment transactions, noting that statistically significant sampling in the programmatic audit can be larger in scope and more costly in comparison to random sampling which can also identify programmatic issues. Another commenter recommended that HHS consider changing the frequency of the programmatic audit to biennially unless the programmatic audit shows irregularities.
Another commenter urged HHS to clarify that the proposed changes to the programmatic audit specific to eligibility and enrollment activities do not pertain to SBE-FPs, since SBE-FPs rely on HHS and the federal platform to perform eligibility and enrollment functions.

Response: We believe these proposed changes will strengthen our programmatic oversight and the program integrity of State Exchanges and thus are finalizing these amendments as proposed. As detailed in the proposed rule, these amendments are intended to allow for more targeted audits that focus HHS and State Exchange resources on compliance with particular Exchange program areas that have higher program integrity risks in a more consistent manner, rather than covering all program areas. These amendments are also intended to address requirements that are applicable only to a particular State Exchange model, in a more standardized manner. We are removing the April 1st deadline from §155.1200(b)(1) to allow HHS to align the deadline for submission of the financial statement to HHS with the deadline for submission of SMART reports, currently June 1. Going forward, we anticipate establishing the deadline for submission of the financial statement and SMART report on an annual basis through guidance and would seek to accommodate state budget cycles to the maximum extent practical when setting these dates. The general scope of these audits remains the same, that is, under the new paragraph (d)(2), HHS may specify that an audit focus on compliance with subparts D and E of 45 CFR part 155, or other requirements under 45 CFR part 155, as specified by HHS. However, we appreciate and considered the

6 This is consistent with the scope for audits in the existing regulation at 45 CFR 155.1200(d)(2), which currently requires State Exchanges to ensure these audits address compliance with “the requirements under this part.”
comments received. We understand that most State Exchanges negotiate their contracts with external auditing entities a year or more in advance and would need sufficient time to update their contracts to reflect any changes in the scope of the external programmatic audits. We also recognize that State Exchanges that operate their own eligibility and enrollment platforms would also need time to work with their contracted auditors to implement new procedures for testing the accuracy of eligibility determinations if their auditors have not previously employed such procedures for this purpose. Thus, subsequent to this rule, we will provide State Exchanges with technical operational guidance that will specify the first plan year for which changes to the scope of the programmatic audit would apply, taking into account the need to allow for a period of time for State Exchanges to implement the changes finalized in this rule.

In response to the comments regarding use of a statistically-significant sampling methodology versus a random sampling methodology, we clarify that, in this rule, we are not specifying a particular sampling methodology that must be used by all State Exchanges for testing the accuracy of eligibility determinations in the annual programmatic audits. In addition to State Exchanges and their contracted auditors using the generally accepted government auditing standards, CMS’s technical operational guidance would also outline procedures the independent external auditor can chose to implement to assess whether a State Exchange is conducting accurate eligibility determinations and enrollment transactions under 45 CFR part 155 subparts D and E. Going forward we intend to provide State Exchanges with this technical operational guidance on an annual basis to outline the deadline for submission of the applicable year’s reports, the scope of the applicable year’s external programmatic audit, and the requirements under
45 CFR part 155 that are applicable to each State Exchange model. We intend to release this guidance around April each year, to align with our existing timeframe for providing guidance to State Exchanges on the annual SMART process, so that State Exchanges have sufficient time to prepare, and administrative burden is minimized to the extent practical. Lastly, we agree with the overall notion of taking a risk-based approach towards determining the frequency by which State Exchanges are required to conduct the external programmatic audit. Specifically, we considered the recommendation to change the frequency of State Exchange programmatic audits to biennially unless the audit shows irregularities. We decline to make this change at this time because some State Exchanges currently are addressing active findings or corrective actions as a result of past programmatic audits, which we believe annual re-evaluations are still appropriate. However, we will consider this recommendation going forward and may propose to decrease the frequency of State Exchange audits in future rule-making.

Comment: Some commenters requested that certain regulatory language remain unchanged or be modified. One commenter urged HHS to retain the language under §§155.1200(b)(2) and 155.1200(d)(2) because the proposed language is broader and targeted auditing can create administrative burden. Another commenter requested that HHS limit the scope of the programmatic audit under §155.1200(d)(2) to solely cover the eligibility and enrollment requirements under 45 CFR part 155 subparts D and E and remove the language that allows HHS to include other Exchange requirements under 45 CFR part 155 in the scope of the programmatic audit. Another commenter requested that §155.1200(d)(2) remain unchanged because the general reference to compliance with 45 CFR part 155 is consistent with the HHS’s stated intent to specify the scope for
programmatic audits, and recommended that HHS make clear that the proposed changes to the review of State Exchange eligibility determinations under §155.1200(d)(4) applies to eligibility determinations for QHP/APTC only, and not to Medicaid eligibility determinations.

Response: We believe the proposed changes under §155.1200(d) will strengthen our programmatic oversight and the program integrity of State Exchanges and provide appropriate flexibility to target oversight and enforcement activities, as well as HHS and State Exchange resources, which, in turn, will reduce burden. As State Exchanges continue to evolve and mature, HHS will be able to focus oversight efforts, including making refinements to annual compliance reporting tools (such as the SMART), in response to changes in federal policy, as well as federal program integrity priorities and processes. We further note that, while these amendments provide flexibility for HHS to target these audits, they also retain the authority for HHS to require the audits to address other requirements under 45 CFR part 155, as specified by HHS. As such, HHS can still require audits with a broader scope when deemed appropriate or necessary. While we generally intend to focus programmatic audits on those Exchange functions most likely to impact program integrity, such as eligibility verifications, we do not agree with commenters that these audits should only focus on eligibility and enrollment functions because there may be changes to federal policy, priorities, or processes that result in the need for HHS to focus our oversight on other Exchange functions besides eligibility and enrollment. Also, not all State Exchanges perform their own eligibility and enrollment functions. For instance, SBE-FPs rely on HHS and the federal platform to perform their eligibility and enrollment functions, and thus HHS’s oversight of SBE-FPs would need to
focus on other Exchange functions that are more relevant or critical to the SBE-FP model. That is why HHS retains the authority, and the flexibility, under the amended §155.1200(d)(2) to require the audits to address other requirements under 45 CFR part 155, as specified by HHS. In addition, the amendments to §155.1200(d)(2) finalized in this rule give HHS flexibility to specify the Exchange functions that are most pertinent to the State Exchange model and most likely to impact program integrity.

In response to comments, we clarify that the changes to subparagraph §155.1200(d)(4) apply to State Exchange eligibility determinations for QHP/APTC, and not to Medicaid eligibility determinations. We recognize that not all State Exchanges make Medicaid eligibility determinations, but also wish to clarify that in accordance with §155.302, State Exchanges must conduct a MAGI-based assessment or determination of eligibility for Medicaid as part of determining eligibility for APTC. HHS will provide further guidelines on the auditing of State Exchange eligibility and enrollment transactions, and any other audit requirements applicable in a given year, in the annual technical operational guidance. We further clarify that the amendments to §155.1200(b)(2) do not reflect an expansion of State Exchange reporting obligations and instead capture the existing annual compliance reports (such as the SMART), that encompass eligibility and enrollment reporting, as well as compliance across other Exchange program requirements under 45 CFR part 155, that State Exchanges currently submit to HHS.

Comment: One commenter requested transparency regarding HHS’s oversight of the Federally-facilitated Exchanges’ (FFE’s) compliance with oversight standards. The commenter recommended that HHS publish a comparison of compliance standards and
activities to ensure the FFEs and State Exchanges are held to the same oversight requirements. Another commenter generally supported the proposed changes as enhancing the oversight and transparency of the State Exchanges.

**Response:** We appreciate and strive for transparency in the oversight of all Exchanges and will consider these suggestions. However, we note that the oversight standards under §155.1200, including the proposed amendments, are specific to State Exchanges. Therefore, the comments related to FFE oversight standards are outside the scope of this rulemaking. We also note that the FFEs are overseen through the efforts of other federal entities such as the Government Accountability Office and the HHS Office of the Inspector General.

**Comment:** Several commenters opposed HHS’s proposed changes to the annual reporting and programmatic audit requirements for State Exchanges. They stated that the proposed language expands federal authority and can add administrative burden to State Exchanges. Some commenters disagreed that the Federalism implications are substantially mitigated since the proposed changes only add specificity to existing requirements, stating that the proposed changes are open-ended and remove specificity. Additionally, some of these commenters expressed concern that HHS is eliminating the requirement of eligibility and enrollment reports under §155.1200(b)(2). These commenters also raised concerns with the disclosure of consumer information, as well as negative consumer impacts, due to the additional oversight on eligibility determinations being proposed.

**Response:** We believe these changes will strengthen our programmatic oversight and the program integrity of State Exchanges. Further, as detailed above, the
amendments do not represent an expansion of HHS’s authority to oversee and monitor compliance of State Exchanges. Under the existing language at §155.1200(d)(2), State Exchanges are currently required to ensure their respective annual programmatic audits address compliance with “the requirements under this part.” The changes to this provision finalized in this rule provide HHS with the flexibility to target the scope of the audits to the requirements applicable to each State Exchange model under 45 CFR part 155 and that most impact program integrity, which should generally reduce the administrative burdens associated with these audits. For example, we anticipate tailoring the requirements regarding audit of eligibility and enrollment activities by State Exchange model. Since SBE-FPs rely on the federal platform for eligibility and enrollment functions, we believe that they should not be subject to the same audit requirements as State Exchanges that perform all eligibility and enrollment activities because they operate their own technology platform for such activities.

We also clarify that we are not eliminating eligibility and enrollment reporting under §155.1200(b)(2). The amendments finalized to that provision reflect that HHS already requires State Exchanges to submit annual reporting (such as the SMART) that encompass eligibility and enrollment reporting, along with other information about compliance with requirements in other subparts under 45 CFR part 155. These changes recognize that HHS has come to utilize the SMART along with the annual programmatic and financial audit reports as the primary oversight tools to oversee State Exchange compliance with the applicable requirements under 45 CFR part 155, which includes compliance with eligibility and enrollment requirements. We further clarify that if we need additional information about a State Exchange’s compliance with applicable
requirements beyond what is reported through SMART, we would leverage the new flexibility under the new §155.1200(d)(2) to conduct a targeted audit.

Finally, in response to the comments expressing concern about the increased risk of disclosure of consumer information as a result of the additional oversight and auditor review of individual eligibility determinations made by State Exchanges that is contemplated in this rule, we note that, as part of the responsibilities of State Exchanges and their contracted entities in handling individual consumer data associated with core Exchange functions such as eligibility, enrollment, and consumer assistance, State Exchanges and their contracted non-Exchange entities must always comply with the privacy and security requirements under §§155.260 and 155.280 with respect to the protection and disclosure of personally identifiable information. Additionally, under §155.285, State Exchanges and their contracted entities are subject to civil monetary penalties for improper use or disclosure of personally identifiable information. Finally, HHS has authority under §155.280 to conduct audits and investigations to ensure compliance with Exchange privacy and security standards, and may pursue civil, criminal or administrative proceedings or actions as determined necessary.

After considering the comments received in response to the proposed rule and for the reasons discussed above, we are finalizing the modifications to §155.1200.

B. Health Insurance Issuer Standards under the Affordable Care Act, Including Standards Related to Exchanges

1. Segregation of Funds for Abortion Services (§156.280)

We proposed an amendment at §156.280(e)(2) relating to billing and payment of the policy holder’s portion of the premium attributable to abortion services for which
appropriated funds may not be used. Since 1976, Congress has included language, commonly known as the Hyde Amendment, in the Labor, Health and Human Services, Education and Related Agencies appropriations legislation. The Hyde Amendment, as currently in effect, permits federal funds subject to its funding limitations to be used for abortion services only in the limited cases of rape, incest, or if a woman suffers from a physical disorder, physical injury, or physical illness, including a life-endangering physical condition caused by or arising from the pregnancy itself, that would, as certified by a physician, place the woman in danger of death unless an abortion is performed (Hyde abortion services). Generally, when appropriated funds are subject to the Hyde Amendment’s funding limitations, an agency is prohibited, among other things, from using those funds to pay for coverage of abortion beyond these specific limited exceptions (non-Hyde abortion services). Section 1303(b)(2) of the PPACA prohibits the issuer of a QHP offering coverage for abortion services that are not exempt from the Hyde Amendment’s ban on the use of federal funds to pay for certain abortions, from using any amount attributable to PTC (including APTC) or CSRs (including advance payments of those funds to an issuer, if any) for abortions for which federal funds are prohibited, “based on the law as in effect as of the date that is 6 months before the beginning of the plan year involved.”

Section 1303 of the PPACA outlines specific accounting and notice requirements that QHPs covering non-Hyde abortion services must follow to ensure that no federal funding is used to pay for services for which public funds are prohibited. Under sections

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7 Accordingly, the Hyde Amendment is not permanent Federal law, but applies only to the extent reenacted by Congress from time to time in appropriations legislation.
8 Section 1303(b)(1)(B)(i) of the PPACA.
1303(b)(2)(B) and (b)(2)(D) of the PPACA, as implemented in §156.280(e)(2)(i) and (e)(4), QHP issuers must collect a separate payment from each enrollee in such a plan without regard to the enrollee’s age, sex, or family status, for an amount equal to the greater of the actuarial value of coverage of abortion services for which public funding is prohibited, or $1 per enrollee per month.

Section 1303(b)(2)(D) of the PPACA establishes certain requirements with respect to a QHP issuer’s estimation of the actuarial value of non-Hyde abortion services. Under section 1303(b)(2)(D) of the PPACA, the QHP issuer “may take into account the impact on overall costs of the inclusion of such coverage, but may not take into account any cost reduction estimated to result from such services, including prenatal care, delivery, or postnatal care.” The QHP issuer is also required to estimate such costs as if such coverage were included for the entire population covered, and may not estimate such a cost at less than $1 per enrollee, per month. If an enrollee’s premium is paid through employee payroll processes, section 1303(b)(2)(B) of the PPACA requires that the separate payments “shall each be paid by a separate deposit.” Accordingly, issuers that offer QHPs that provide coverage of non-Hyde abortion services must collect a separate payment of no less than $1 per enrollee in the plan per month, regardless of the actuarial value of coverage of non-Hyde abortion services and regardless of whether premiums are paid directly by enrollees or through payroll deductions.

In certain rare scenarios, the FFEs’ system allocated an amount of APTC to a QHP such that the share of the aggregate premium attributable to coverage of non-Hyde abortion services is less than $1, which falls below the minimum requirement under section 1303 of the PPACA. We made system changes for the open enrollment period.
for plan year 2019 to ensure that the minimum premium amount of $1 per enrollee per month is assigned to all enrollments into plans offering coverage of non-Hyde abortion services, so that issuers can separately collect this amount directly from enrollees for the portion of the total premium attributable to coverage of non-Hyde abortion services.

Pursuant to section 1303(b)(2)(C) of the PPACA, as implemented at §156.280(e)(3), QHP issuers must segregate funds for coverage of non-Hyde abortion services collected from enrollees into a separate allocation account that is to be used to pay for non-Hyde abortion services. Thus, if a QHP issuer disburses funds for a non-Hyde abortion on behalf of an enrollee, it must draw those funds from the segregated allocation account. The account cannot be used for any other purpose.⁹

Section 1303 of the PPACA and current implementing regulations at §156.280 do not specify the method a QHP issuer must use to comply with the separate payment requirement under section 1303(b)(2)(B)(i) of the PPACA and §156.280(e)(2)(i). In the 2016 Payment Notice, we provided guidance with respect to acceptable methods that a QHP issuer offering coverage of non-Hyde abortion services on an individual market Exchange may use to comply with the separate payment requirement. We stated that the QHP issuer could satisfy the separate payment requirement in one of several ways, including by sending the enrollee a single monthly invoice or bill that separately itemizes the premium amount for coverage of non-Hyde abortion services; sending the enrollee a separate monthly bill for these services; or sending the enrollee a notice at or soon after

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⁹ This means that funds from the allocation account into which premium amounts attributable to the non-Hyde abortion service benefit must be deposited are the only funds that may be used to pay for non-Hyde abortion services. It should not be read to suggest that the funds in the separate allocation account may not be used to cover administrative costs associated with coverage of non-Hyde abortion services. See 42 U.S.C. § 18023(b)(2)(D)(ii)(I) (when estimating per member, per month cost of non-Hyde abortion services, issuers may take into account the impact on overall costs of the inclusion of such coverage).
the time of enrollment that the monthly invoice or bill will include a separate charge for such services and specify the charge. In the 2016 Payment Notice, we also stated that an enrollee may make the payment for coverage of non-Hyde abortion services and the separate payment for coverage of all other services in a single transaction. On October 6, 2017, we released a bulletin that discussed the statutory requirements for separate payment, as well as this previous guidance with respect to the separate payment requirement.\textsuperscript{10}

As explained in the proposed rule, HHS now believes that some of the methods for billing and collection of the separate payment for coverage of non-Hyde abortion services described as permissible in the preamble to the 2016 Payment Notice do not adequately reflect Congress’s intent. We believe Congress intended that QHP issuers collect two distinct (that is, “separate”) payments, one for the coverage of non-Hyde abortion services, and one for coverage of all other services covered under the policy, rather than simply itemizing these two components in a single bill, or notifying the enrollee that the monthly invoice or bill will include a separate charge for these services.

We proposed an amendment at §156.280(e)(2) relating to billing and payment of the policy holder’s portion of the premium attributable to coverage of non-Hyde abortion services to reflect this interpretation of the statute. Specifically, we proposed that, as of the effective date of this final rule, QHP issuers (1) send an entirely separate monthly bill to the policy holder, the individual who is the party legally responsible for the payment of premiums (which we refer to in this final rule as the “policy holder”) for only the portion

of premium attributable to coverage of non-Hyde abortion services, and (2) instruct the policy holder to pay the portion of their premium attributable to coverage of non-Hyde abortion services in a separate transaction from any payment the policy holder makes for the portion of their premium not attributable to coverage of non-Hyde abortion services. We also proposed that if a policy holder pays the entire premium in a single transaction (both the portion attributable to coverage of non-Hyde abortion services, as well as the portion attributable to coverage for other services), the QHP issuer would not be permitted to refuse to accept such a combined payment on the basis that the policy holder did not send payment in two separate transactions as requested by the QHP issuer, and to then terminate the policy, subject to any applicable grace period, for non-payment of premiums. We also stated that the QHP issuer would be expected to counsel enrollees to pay in two separate transactions in the future. Finally, we proposed a technical change to §156.280(e)(2)(iii), as redesignated, to insert an appropriate cross reference to the explanation of the separate payments.

We are finalizing these policies at §156.280(e)(2), but with several changes explained below. We are also finalizing the technical revision to §156.280(e)(2)(iii) as redesignated, on which we received no comments, and are revising the heading of §156.280 so that it accurately describes the new requirements we are finalizing in this final rule.

Comment: Most commenters objected to the proposed changes to issuer billing for the portion of the premium attributable to coverage of non-Hyde abortion services, asking that we withdraw the proposals altogether. A minority of commenters summarily supported the policy.
Nearly all commenters objecting to the proposals stated that separately billing for one specific service would be an unnecessary change that would not enhance program integrity with respect to enrollee transparency or appropriate use of federal funds. These commenters noted that current requirements already adequately comply with the statute and ensure appropriate segregation of funds, without imposing the operational and administrative burdens of the proposed approach. These commenters asserted that the current regulatory structure allows enrollees to make and issuers to accept a single transfer of funds for the full amount of an enrollee’s premium payment including the amount attributable to coverage of non-Hyde abortion services, while still ensuring that the funds are ultimately segregated appropriately. Many commenters noted that requiring a separate bill and instructing enrollees to pay in separate transactions would be against industry practice, which permits one single bill outlining charges and allows for enrollees to make payments using a single transfer of funds which can be administratively separated by the insurer after payment is received.

Some commenters who supported the proposed changes stated that section 1303 of the PPACA contains an unambiguous statutory command that issuers separately bill and collect payments for the portion of a policy holder’s premium attributable to coverage of non-Hyde abortion services. These commenters stated that the proposals are necessary to remedy incorrect methods for billing and payment and will help to ensure issuer compliance with the segregation of funds and the requirement to collect separate payments under section 1303 of the PPACA.

Nearly all objecting commenters stated that the proposals would cause considerable and unnecessary confusion and frustration for enrollees that may jeopardize
their health insurance coverage. Commenters expressed concern that these billing changes would make it more difficult for policy holders to pay their premium bills, and could result in coverage being terminated for unintentional non-payment. Commenters expressed concerns that, despite issuer notices and communications to explain the second bill and separate payment requirement, enrollees would likely not understand this change in billing.

Among the many scenarios that commenters asserted could result in enrollees failing to pay the separate bill, commenters noted that enrollees might not realize or understand that there is a separate bill covering different services under their plan; enrollees may not realize that such payment is mandatory in order to fully satisfy their premium liability each month and avoid termination of coverage; or enrollees may not notice a second bill since it would be delivered in a separate mailing with which they are unfamiliar. Commenters expressed concern that in any of these scenarios, the enrollee would enter a grace period and, in most cases, have 90 days from the date of the missed payment to reconcile their balance, resulting in enrollees who fail to do so losing their health insurance coverage. Commenters expressed concern that such slight enrollee confusion as a result of the proposal could lead to the complete loss of coverage.

Commenters also stated that the proposal to allow enrollees to “not be penalized” for sending back a combined payment, would only send conflicting messages to enrollees and add to their confusion. Commenters stated that our proposal that issuers could accept combined payments from enrollees, but would then be expected to counsel enrollees to pay in two separate payments in the future, requiring issuers to repeatedly instruct enrollees to pay in separate transactions for each bill despite not being able to penalize
enrollees if they continuously fail to do so, adds additional burden on issuers and will lead to increased calls from confused enrollees.

   Many commenters stated they appreciated the enrollee protections prohibiting QHP issuers from refusing to accept a combined payment or terminating an enrollee’s coverage on this basis. However, commenters expressed concerns that this protection alone would not be enough for enrollees who fail to pay the second bill entirely and asked that HHS add protections to the policy to avoid termination of coverage for enrollees who inadvertently fail to make the additional payment due to confusion about the separate bill.

   Response: We continue to believe that the statute contemplates issuers billing separately for coverage of non-Hyde abortion services, consistent with Congress’s intent that issuers collect separate payments for such services. Requiring one bill for the portion of the policy holder’s premium attributable to coverage of non-Hyde abortion services and a separate bill for the portion of the policy holder’s premium attributable to coverage of all other services covered under the QHP will better align with the intent of section 1303 of the PPACA.

   HHS intentionally sought comment on ways to mitigate possible enrollee confusion from these proposals. After considering these comments, we believe there may be less confusing and less burdensome ways to implement these billing changes while also fulfilling section 1303 of the PPACA’s statutory mandates.

   Therefore, we are finalizing, as proposed in a new paragraph at §156.280(e)(2)(ii)(A), the requirement that QHP issuers must send an entirely separate monthly bill to the policy holder for only the portion of the premium attributable to coverage of non-Hyde abortion services. However, in an effort to mitigate issuer burden
associated with added postage and mailing costs, we will not require separate mailings with separate postage, as proposed. Rather, we are codifying that the QHP issuer may include the separate bill for coverage of non-Hyde abortion services in the same envelope or mailing as the bill for the portion of the premium attributable to coverage of all other services. As a result of finalizing this proposal, and to more accurately reflect the contents of §156.280, we are making a technical change to revise the section heading of §156.280 to now read, “Separate billing and segregation of funds for abortion services.”

We note that when issuers send a separate paper bill for the portion of the premium attributable to coverage of non-Hyde abortion services in the same mailing as the bill for the other portion of the policy holder’s premium, the bills must remain distinct and separate, on separate pieces of paper with separate explanations of the charges to ensure the policy holder understands the distinction between the two bills and understands that they are expected to pay the separate bills in separate transactions.

We are also codifying that issuers transmitting bills through email or other electronic means will still be required to transmit the separate bill for coverage of non-Hyde abortion services in a separate email or electronic communication than for the bill for the portion of the premium attributable to coverage of all other services. We assume that bills sent electronically can be sent at minimal cost such that requiring separate electronic communications will not significantly increase the burden this requirement places on issuers. We also believe policy holders are more likely to make a separate payment for coverage of non-Hyde abortion services when they receive a separate bill for such amount, and that receiving the separate bill in a separate communication further bolsters that likelihood. In deciding to finalize that QHP issuers
may send the separate bill in a single mailing when sending paper bills, but must send the separate bill in a separate email or electronic communication when sending bills electronically, we weighed the goal of separate payment with the competing concern of issuer burden resulting from sending separate paper bills, and the comparatively low burden in sending separate electronic bills.

We are also finalizing, as proposed in a new paragraph at §156.280(e)(2)(ii)(B) the requirement that issuers must instruct policy holders to pay the separate bill in a separate transaction. QHP issuers should make reasonable efforts to collect the payment separately. However, we continue to believe that potential loss of coverage would be an unreasonable result of an enrollee paying in full, but failing to adhere to the QHP issuer’s requested payment procedure. Therefore, at §156.280(e)(2)(ii)(B) we are also codifying, with minor non-substantive revisions, that the QHP issuer would not be permitted to refuse a combined payment on the basis that the policy holder did not send two separate payments as requested by the QHP issuer, and to then terminate the policy for non-payment of premiums. QHP issuers that receive combined enrollee premiums in a single payment must treat the portion of the premium attributable to coverage of non-Hyde abortion services as a separate payment and must disaggregate the amounts into the separate allocation accounts, consistent with §156.280(e)(2)(iii).

To mitigate enrollee confusion and satisfy the requirement to instruct policy holders to pay the separate bill in a separate transaction, QHP issuers should consider including—in the email or electronic communication containing the bill for the portion of the policy holder’s premium not attributable to coverage of non-Hyde abortion services—language notifying policy holders that they will be receiving a second, separate email or
electronic communication containing a separate bill for the portion of their premium attributable to coverage of non-Hyde abortion services that they should pay in a separate transaction. Regardless of whether the QHP issuer sends the bills as paper copies in a mailing or sends the bills through electronic communications, the QHP issuer must instruct their enrollees to pay the separate bill in a separate transaction and must still produce an invoice or bill that is distinctly separate from the invoice or bill for the other portion of the policy holder’s premium that is not attributable to coverage of non-Hyde abortion coverage, whether in paper or electronic format. We also suggest that issuers state clearly for policy holders on both bills that the policy holder is receiving two bills to cover the total amount of premium due for the coverage period, that the policy holder’s total premium due is inclusive of the amount attributable to coverage of non-Hyde abortion services, and that the policy holder should make separate payments for each bill. We believe including these statements on each bill, will help policy holders to understand that they are receiving two bills for the premiums due for the payment period, the total amount of premium they owe, and the need to make a separate payment for each bill. We believe this will help to ensure that policy holders return the full monthly amount due, thus preventing policy holders from entering grace periods for non-payment of the premium amounts for the non-Hyde abortion coverage.

We believe these changes will assist in managing enrollee confusion. However, we also acknowledge that additional outreach and education may still be necessary on the part of issuers and states to explain to enrollees why they are receiving a separate bill for a relatively small amount for which they are expected to submit payment in a separate transaction. As indicated above, we believe that QHP issuers should explain to the policy
holder in layperson terms on the separate bill for coverage non-Hyde abortion services, or otherwise communicate to enrollees through enrollee outreach and education, that non-payment of any premium due (including non-payment of the portion of the policy holder’s premium attributable to coverage of non-Hyde abortion services) would continue to be subject to state and federal rules regarding grace periods (unless the QHP issuer elects to take advantage of the enforcement discretion we outline later in this section), clarifying for policy holders that failure to pay the portion of the premium attributable to coverage of non-Hyde abortion services could ultimately result in termination of coverage.

We believe that including explanatory language on the bills as well as additional outreach and education by QHP issuers will decrease the likelihood that policy holders would inadvertently fail to pay the separate bill for the portion of their premium attributable to coverage of non-Hyde abortion services. However, we acknowledge commenters’ concerns that, even with fulsome outreach and education efforts to explain the billing scheme to the policy holder, consumer confusion could still lead to inadvertent coverage losses. This risk may be especially acute for enrollees whose plan choices likely were not motivated by the plan’s coverage of non-Hyde abortion services, such as men purchasing a QHP solely for themselves, consumers buying coverage for babies or toddlers, and those who otherwise may be unaware that the plan covers non-Hyde abortion services. However, we note that this risk is mitigated by the steps we have taken to improve transparency regarding QHP offerings, to make it easier for consumers to select QHPs that they believe are best suited to their needs and preferences, such as
information to more readily identify QHPs that offer coverage of non-Hyde abortion services.  

To address the risk of terminations related to inadvertent failure to pay the separately billed amount for coverage of non-Hyde abortion services, we intend to propose further rulemaking to change our regulations including, for example, our regulations governing termination for non-payment of premiums.  Although QHP issuers can implement premium payment thresholds under §155.400(g), those thresholds may not be effective at preventing termination of coverage for policy holders receiving higher APTC amounts who would have greater difficulty meeting the issuer’s premium payment threshold pursuant to §155.400(g). Until we can finalize regulatory changes through a separate rulemaking, we will exercise enforcement discretion as an interim step. Specifically, HHS will not take enforcement action against a QHP issuer that adopts and implements a policy, applied uniformly to all its QHP enrollees, under which an issuer does not place an enrollee into a grace period and does not terminate QHP coverage based solely on the policy holder’s failure to pay the separate payment for coverage of non-Hyde abortion services. In accordance with non-discrimination rules applicable to QHP issuers, we would expect issuers to apply such a policy uniformly to all of their enrollees for the duration of the applicable plan year. We also note that if a


12 CMS has yet to make determinations regarding specific requirements or rule changes CMS will propose to address the risk of terminations related to inadvertent failures to pay the separately bill amounts for coverage of non-Hyde abortion services. Accordingly, although CMS will undertake the described rulemaking, nothing in this preamble discussion should be construed as a representation or guarantee that CMS will propose changes to any specific rule or requirement.
QHP issuer chooses to take this approach, the QHP issuer would still be prohibited from using any federal funds for coverage of non-Hyde abortion services. Moreover, the QHP issuer would still be required to collect the premium for the non-Hyde abortion coverage, which means that the QHP issuer cannot relieve the policy holder of the duty to pay the amount of the premium attributable to coverage for non-Hyde abortion services. This enforcement posture will take effect upon the effective date of the separate billing requirements under 45 CFR 156.280, which is 6 months after publication of this final rule in the Federal Register. We encourage states and State Exchanges to take a similar enforcement approach.

We acknowledge that the enforcement posture described above may not mitigate all concerns identified by commenters. Some commenters expressed concern that the lack of transparency under current section 1303 billing requirements has contributed to unknowing purchases of QHPs that include coverage of non-Hyde abortion services by consumers who object to purchasing such coverage. As noted above, this risk is mitigated by the steps the FFEs have taken to improve transparency of the coverage of non-Hyde abortion services under FFE QHPs. However, even where consumers who hold religious or moral objections to coverage of non-Hyde abortion services may more easily detect whether a QHP offers coverage to which they object, they may still be deciding between purchasing a QHP that covers non-Hyde abortion services, or else going without the coverage they need, because there may not be a QHP available on the Exchange that omits coverage for non-Hyde abortion services.

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Until we are able to address these concerns through future rulemaking or other appropriate action, we also will not take enforcement action against QHP issuers that modify the benefits of a plan either at the time of enrollment or during a plan year to effectively allow enrollees to opt out of coverage of non-Hyde abortion services by not paying the separate bill for such services. This would result in the enrollees having a modified plan that does not cover non-Hyde abortion services, meaning that they would no longer have an obligation to pay the required premium for such services. We recognize that a QHP issuer’s ability to make changes to its QHPs to implement a policy holder’s opt out would be subject to applicable state law. We encourage states and State Exchanges to take an enforcement approach that is consistent with the one we intend to take, as described in this section.

Where a QHP issuer allows an enrollee to opt out of coverage of non-Hyde abortion services by not paying the separate bill for such services, the user fee a QHP issuer in an FFE or SBE-FP would pay would continue to be based on the original premium, which includes the portion of the premium attributable to non-Hyde abortion coverage. This is being done for operational reasons and issuer convenience, as making changes to the user fee system for FFEs and SBE-FPs to reflect a reduction in premium would result in only a minimal reduction in user fees owed. We do not believe the minimal reduction justifies the additional expense to FFEs and SBE-FPs related to the development of systems to receive and process such reports (which could then result in higher user fees in the future) or the additional cost to QHP issuers related to reporting the minimal changes in premiums.
We expect QHP issuers taking this approach to take appropriate measures to distinguish between a policy holder’s inadvertent non-payment of the separate bill for coverage of non-Hyde abortion services and a policy holder’s intentional nonpayment of the separate bill. A policy holder who inadvertently fails to pay the separate bill may have failed to pay because of unfamiliarity with receiving a separate bill for this portion of their premium and may still wish to retain coverage for non-Hyde abortion services if provided the opportunity to rectify nonpayment of the separate bill. A policy holder who intentionally does not pay the separate bill is likely to have made the conscious choice to opt-out of such coverage. To help ensure any modifications made by a QHP issuer under this enforcement approach to a policy holder’s plan align with the policy holder’s intent, the QHP issuer could include on the separate bill for coverage of non-Hyde abortion services or separate electronic communication an option (such as a check box or option button) where the policy holder can affirmatively indicate their intent to opt-out of such coverage by not paying the separate bill. We also recommend including an explanation for the policy holder that by affirmatively opting out, the policy holder would no longer have coverage for non-Hyde abortion services and would no longer have an obligation to pay the required premium for such services.

To be clear, we intend that a policy holder’s opt-out would have to be applied to all persons in the enrollment group under the policy. For example, if the policy holder does not pay the separate bill for the portion of the premium attributable to non-Hyde abortion coverage and therefore opts out of coverage for non-Hyde abortion, this opt-out would be applicable to all persons in the policy holder’s enrollment group, such as the policy holder’s spouse and/or family if they are also covered under the policy holder’s
policy. Further, our exercise of enforcement discretion would only permit issuers to make one-time changes to remove coverage of non-Hyde abortion services from the QHP coverage.

Accordingly, once a policy holder opts out of coverage for non-Hyde abortion services, the policy holder would not be allowed to retract their opt-out decision and reinstate coverage of non-Hyde abortion services for that benefit year, by paying premiums that could cover a portion of premium attributable to coverage of non-Hyde abortion services. Thus, an opt-out would be effective for the remainder of the benefit year.

Unlike the enforcement discretion policy we announce above to mitigate risk of inadvertent terminations, this enforcement posture will become effective on the effective date of this final rule, which will be 60 days after its publication in the Federal Register. The separate billing requirements we finalize here under 45 CFR 156.280 will address, among other things, stakeholder comments that the lack of transparency under current section 1303 billing requirements has contributed to unknowing purchases of QHPs that include coverage of non-Hyde abortion services by consumers who object to purchasing such coverage. Because the new billing requirements under these final rules will not take effect upon finalization of these rules, we believe it is important to take this enforcement posture as soon as possible to provide relief for the lack of transparency under current QHP billing requirements.

We are taking this approach to maintain protections against adverse selection, while mitigating the serious negative risks of coverage loss by enrollees who might experience difficulties adjusting from the manner in which enrollees are accustomed to
paying for insurance coverage or services under a single plan or contract. These interim policies will also provide relief to persons who may unknowingly purchase coverage to which they object because of the lack of transparency under current QHP billing requirements that do not require separate bills for non-Hyde abortion coverage. We believe these interim enforcement policies strike an appropriate balance between honoring PPACA section 1303’s requirement for collection of separate payments, protecting enrollees against inadvertent losses of coverage, and ensuring all enrollees have access to coverage that meets their needs and that does not result in their supporting coverage for non-Hyde abortion services to which they object.

Comment: Commenters stated that HHS greatly underestimated the burden on issuers caused by these proposals. Commenters stated that the proposed rule’s analysis of the expected costs and benefits was incomplete, such that HHS cannot accurately determine whether the benefits outweigh the quantitative and qualitative costs to justify finalizing the proposals. Many commenters stated that the burden and costs far outweigh any benefit and, as such, the proposals should not be finalized.

Commenters also stated that requiring issuers to send the separate bill in a different envelope or separate email communication would cost QHPs significantly more resources than HHS estimated for the multiple mailings, email communications, and personnel hours spent managing enrollee confusion, termination notices, and multiple bills. For example, commenters noted requiring a separate mailing would double the mailing and postage costs associated with current issuer billing. Commenters also explained that the technical build issuers would need to implement to comply with these proposals would be both complex and time consuming, and would alone require
substantial new upfront and annual costs for issuers that HHS did not account for. In general, commenters expressed concerns that requiring separate billing and instructing enrollees to make separate payments for a single policy would create substantial new operational administrative costs for health insurance issuers and, subsequently, for the enrollees they serve.

Commenters also expressed concerns with the burdens these changes would impose on Exchanges. Commenters noted Exchanges would need to make time consuming and resource intensive changes to their websites, enrollment systems, and customer service and outreach efforts to align with the separate billing and payment requirements, which would be costly and disrupt Exchange efficiency.

Commenters also expressed concern that HHS failed to address the adverse impacts on enrollees resulting from how issuers would react to being forced to allocate additional significant operational and administrative resources towards issuing and processing multiple bills and monthly payments from each policy holder. Many commenters stated that issuers would be required to consider these new costs when setting actuarially sound rates, which would lead to higher premiums for enrollees. Many commenters stated that the costs and requirements on QHP issuers that cover non-Hyde abortion services will in many cases be so high that it will result in QHP issuers dropping coverage for non-Hyde abortion services altogether, even if their enrollees desire such coverage. Commenters expressed concern that, in such scenarios, this would transfer the costs and burdens of accessing non-Hyde abortion services to enrollees who must seek coverage for non-Hyde abortion services elsewhere or pay out-of-pocket. Other commenters noted that issuers are likely to drop coverage of non-Hyde abortion services
if the alternative is terminating coverage for a substantial number of its enrollees due to enrollee confusion resulting in non-payment of miniscule amounts.

Many commenters stated that the proposals would threaten the mental and physical health, well-being, and economic security of enrollees, especially women, across the country. Commenters stated that health insurance should provide coverage for the full range of reproductive health care, including abortion, and that this rule threatens to take such coverage away by imposing burdensome requirements on issuers. Commenters also expressed concern that, should these proposals result in issuers ceasing to provide coverage of non-Hyde abortion services, it could impede a patient's ability to make the best medical decision for herself and her family in consultation with her physician given that many women would be unable to pay privately for such services due to high costs without insurance. Commenters noted that barriers to accessing affordable non-Hyde abortion services could have long-term, devastating effects on a woman and her family’s economic future.

Commenters noted that the proposals would have a greater impact on subsidized enrollees and might have a discriminatory effect on enrollees receiving higher APTC amounts who would have greater difficulty meeting the issuer’s premium payment threshold pursuant to §155.400(g). Commenters also stated that it would have damaging consequences on enrollees with specific conditions (like patients with cancer or chronic conditions), as any gaps in coverage as a result of confusion over billing may interrupt disease treatment schedules and could jeopardize health outcomes. Commenters also stated that the proposals would threaten the coverage gains made by the PPACA and have a disproportionate impact on enrollees who already face barriers to care, such as low-
income individuals and marginalized communities. HHS received many comments expressing concern that when legal abortion becomes inaccessible, women who seek to end their pregnancy turn to unsafe and illegal methods, risking arrest, serious injury, or even death. Commenters also expressed concern that HHS did not propose any requirements or guidelines for how issuers should educate, inform, and conduct outreach to enrollees regarding these changes in billing and payment if the proposed regulation is implemented as proposed. Commenters also expressed concern that the proposals didn’t address how individuals with limited English proficiency (LEP) or individuals with disabilities may experience barriers in complying with the proposed changes which commenters found particularly concerning, since individuals with LEP and individuals with disabilities already experience hardships in navigating and accessing health care.

**Response:** As we acknowledged in the proposed rule, we recognize that QHP issuers that cover non-Hyde abortion services may experience an increase in burden as a result of the proposals. We have carefully considered the comments that shared information about how the proposals would likely impact markets, issuers, and enrollees.

We agree with commenters that separately mailing the separate bill with separate postage could cause unintended additional burden and cost for issuers. Therefore, we are not finalizing the requirement that the separate bills be mailed separately with separate postage. However, we also acknowledge that QHP issuers will nevertheless still incur significant burden and costs as a result of implementing this new separate billing policy. We agree with commenters that QHP issuers are likely to consider these new costs when setting actuarially sound rates and that this will likely lead to higher premiums for enrollees. The potential premiums increases are discussed in further detail in section III,
“Collection of Information Requirements,” and section IV, “Regulation Impact Analysis,” of this rule. However, in spite of the potential premium increases, we do not agree that requiring issuers to send separate bills, instruct policy holders to pay in two separate transactions, and make reasonable efforts to collect the payments separately would be an inefficient use of resources. Rather, this instruction is important to achieving better alignment of the regulatory requirements for QHP issuer billing of enrollee premiums with the separate payment requirement in section 1303 of the PPACA. We understand commenters’ concerns that the issuer burden associated with this policy may result in issuers withdrawing coverage of non-Hyde abortion services altogether, requiring some enrollees to pay for these services out-of-pocket.

Subject to applicable state law, it is ultimately at the issuer’s discretion whether to cover non-Hyde abortion services in their QHPs, and thus to incur any associated burden, and it is ultimately the states’ and HHS’s duty to enforce the statutory provisions of the PPACA as they are written. Although section 1303 permits issuer flexibility in abortion coverage choices, it also requires that QHP issuers electing to cover non-Hyde abortion services take certain steps to ensure that no APTC or CSR funds are used to pay for these services, such as requiring the QHP issuer to collect a separate payment for these services. The finalized changes at §156.280(e)(2)(ii) may add issuer burden with regard to their payment and billing operations. However, the statute contemplates such burden in section 1303(b)(2)(B)(i) of the PPACA when it requires that issuers collect a separate payment for the portion of the premium attributable to coverage of non-Hyde abortion services and in section 1303(b)(2)(D) of PPACA when it specifies how QHP issuers are to calculate the basic per enrollee, per month cost, determined on an average actuarial
basis, for including coverage of non-Hyde abortions in QHPs. We believe that finalizing the rule to allow issuers to send both bills in a single mailing will mitigate the issuer and state burden that would be imposed if we were finalizing the policy as originally proposed, as well as any initial confusion on the part of enrollees. We estimate that these changes would eliminate much of the additional mailing costs for the second bill since issuers would no longer need to pay for additional postage and envelopes. We believe the changes we are finalizing at §156.280(e)(2)(ii) strike a balance between requiring the separate bill that we believe is required for better alignment with section 1303 of the PPACA, while also avoiding unnecessary enrollee confusion, enrollee harm, and issuer burden.

We understand that non-Hyde abortion services are services for which some enrollees may desire coverage, as they may be costly when not covered by insurance. However, we believe that requiring separate billing for the portion of the premium attributable to coverage of non-Hyde abortion services is a necessary change to better align issuer billing with the statutory requirements specified in section 1303 of the PPACA, which requires non-Hyde abortion services be treated differently from other covered services. We believe the changes we are finalizing at §156.280(e)(2)(ii) will impose less burden on issuers to implement this policy than if we were finalizing as originally proposed, decreasing the likelihood that issuers will drop this coverage or significantly raise their premiums. Although we acknowledge the changes we are finalizing will increase the burden associated with personnel hours spent managing enrollee confusion, termination notices, and multiple bills, we also believe the changes we are finalizing at §156.280(e)(2)(ii) minimize enrollee confusion surrounding receiving
a separate bill, helping to prevent situations where enrollees enter grace periods and subsequently have their coverage terminated for failing to inadvertently pay the second bill. We also believe policy holder confusion regarding the separate bill may decrease in future plan years as policy holders acclimate to this billing structure and as consumer education continues. However, we acknowledge that a policy holder enrolling for the first time after this policy is finalized in a QHP covering non-Hyde abortion services may still experience confusion regarding the separate bill. As finalized, we believe the inclusion of a second separate bill for these services in the same mailing and requiring issuers to instruct enrollees to pay in a separate transaction for the separate bill (whether sent electronically or by mail), but allowing issuers to accept combined payments if the enrollee fails to pay separately, will allow QHP issuers to continue providing coverage for non-Hyde abortion services subject to state and federal law and allow policy holders to continue accessing such coverage when available through their QHPs.

We understand commenters’ concern about how these proposals will impact individuals with LEP and other policy holders, especially those with disabilities. We note that, under the policy being finalized, issuers must still comply with all applicable enrollee assistance requirements for QHPs on the Exchange, such as those requirements at §155.205. In particular, we believe that the requirements at §155.205(c) will help to ensure that issuers are providing information regarding the separate bill and payment options to individuals with LEP and policy holders with disabilities in plain language and in an accessible manner as specified in regulation. We also suggest that issuers consider the needs of these enrollee groups when conducting enrollee education or outreach about the finalized changes.
A more detailed summary of comments discussing the potential burden associated with the proposals can be found in the sections III “Collection of Information Requirements” and IV “Regulation Impact Analysis” of this rule. In section III “Collection of Information Requirements” of this final rule, a detailed breakdown of the estimated one-time burden per issuer and the estimated one-time burden for all issuers can be found in tables 2 and 3, and a detailed breakdown of the estimated annual burden per issuer and the estimated annual burden for all issuers can be found in tables 4 and 5.

Comment: Many commenters expressed concern that the proposed effective date would be administratively and operationally infeasible. As proposed, issuers would be required to implement these proposals beginning on the effective date of the final rule, which is 60 days after the final rule is published in the Federal Register. Commenters explained that issuer billing and payment requirements are typically included in plan documents that are approved by the state regulator and provided to the enrollee at the time of enrollment. Commenters noted that a change in payment policies would mean that issuers would need to re-file their applications for all affected plans for approval by state regulators and that such a change could not be implemented mid-plan year. Commenters also stated that, given the substantial investment required to operationalize the new proposals and the associated complexities, issuers would need a minimum of 12 to 18 months to implement these changes. Further, because implementation would need to coincide with the beginning of a new plan year, many commenters stated that plan year 2021 would be the earliest at which implementation could occur given the likely publication timeline for this final rule. Commenters also stated that enrollees can more easily adapt to new payment arrangements at the beginning of a plan year, when
they expect premiums to be different and other changes to their plan to occur. Commenters also emphasized that the earlier the effective date, the more burdensome these proposals become.

One commenter noted that although state regulators are able to accept the responsibility of primary enforcement of this rule given appropriate lead time, they will be ill-equipped to enforce it if it is made effective immediately, since regulators will need time to develop enforcement policies in consultation with state stakeholders. This commenter also noted that, due to the small amounts issuers would separately bill for coverage of non-Hyde abortion services, many issuers may choose to revise their premium payment threshold policies permitted under §155.400, but would not have time to do so if the rule were made effective immediately.

Response: In response to comments that implementation will take longer than the proposed effective date would allow, we are finalizing that QHP issuers must be in compliance with the policies being finalized at §156.280(e)(2) on or before the day that is 6 months after publication of the final rule. If the date that is 6 months after publication of the final rule falls in the middle of a QHP issuer’s billing cycle (in other words, after the QHP issuer has already sent out bills to policy holders for that month), the QHP issuer would be expected to comply beginning with the next billing cycle immediately following that date. We acknowledge that requiring QHP issuers to begin complying mid-plan year may pose implementation challenges for some states and issuers. For example, as discussed further later in this response, QHP issuers offering coverage of non-Hyde abortion services will already have filed rates for the 2020 plan year and would be unable to update those rates until the following plan year to reflect the added
administrative costs they may experience as a result of the finalized separate billing policy. We also acknowledge requiring QHP issuer compliance mid-plan year would not provide QHP issuers offering coverage of non-Hyde abortion services an opportunity, in their discretion, to revise their plan and benefit designs, such as to remove coverage of non-Hyde abortion services, in order to avoid requirements under the separate billing policy.

We anticipate that State Exchanges that perform premium billing and payment processing that have QHP issuers that offer coverage for non-Hyde abortion services will face similar challenges to comply with the separate billing requirements within 6 months after publication of this final rule as QHP issuers that offer coverage for non-Hyde abortion services. However, we believe 6 months is sufficient for State Exchanges performing premium billing and payment processing and QHP issuers to implement the administrative and operational changes to billing processes necessary to comply with this policy. We also believe a 6-month implementation timeline appropriately prioritizes the goals of improved statutory alignment with the additional time State Exchanges and issuers may need to implement this policy. For those State Exchanges and QHP issuers that may face uncommon or unexpected impediments to timely compliance, HHS will consider extending enforcement discretion to an Exchange or QHP issuer that fails to timely comply with the separate billing policy as required under this final rule, if we find that the Exchange or QHP issuer attempted in good faith to timely meet the requirements.

Although we do not believe that it is necessary for state enforcement policies to have been developed prior to the effective and/or compliance date for the separate billing requirements, we believe this will offer state regulators enough time to develop
enforcement policies in consultation with state stakeholders. We also believe this implementation timeline will provide sufficient time for enrollee outreach and education to help mitigate any enrollee confusion resulting from the finalized policies, and to explain to enrollees how the QHP issuer’s previous payment policies will be changing to comply with these new billing requirements.

We believe it is important that QHP issuers implement these policy changes at the earliest date feasible to improve statutory alignment with section 1303 of the PPACA. Similarly, we do not believe that potential implementation challenges in connection with a mid-year implementation date should outweigh numerous commenters’ concerns regarding the lack of transparency as to whether their QHP covers non-Hyde abortion services, transparency that would be delayed by approximately a year if compliance were required by the first day of the 2021 plan year. We believe that further delaying implementation would be imprudent given that we are now aware of these consumer concerns and given that we believe it is operationally and administratively feasible for State Exchanges and QHP issuers to comply with the policy within 6 months after publication of the final rule.

We acknowledge that if QHP issuers are not able to take these additional costs into consideration when setting rates for the 2020 plan year, it is possible that some issuers may seek to exit the individual market in a state or incur losses. We believe that any such risk is small. QHP issuers will have the opportunity to adjust their plan and benefits design and rates in response to the separate billing policy for their plan year 2021 plan offerings. Moreover, we are aware that the actuarial value of the non-Hyde abortion coverage under QHPs generally may be less than the minimum $1 per enrollee, per
month QHP issuers must charge for such services under section 1303 of the PPACA; and we are not aware of any reason QHP issuers could not use funds from the allocation account into which premium amounts attributable to the non-Hyde abortion service benefit must be deposited to cover administrative costs associated with coverage of non-Hyde abortion services. This should mitigate the financial consequences to issuers of their not being able to update individual market rates prior to the 2021 plan year to incorporate the costs of implementing the processes required by this rule. We therefore believe that finalizing a longer, 6-month implementation timeline sufficiently mitigates the risk that some issuers would seek to exit the individual market to avoid the separate billing requirements under this final rule.

We acknowledge that State Exchanges’ and QHP issuers’ ability to comply within 6 months may depend on the current status of their billing systems and operations, and that State Exchanges and QHP issuers may be confronted with unexpected impediments to timely compliance. For this reason, HHS will consider extending enforcement discretion to an Exchange or QHP issuer that fails to timely comply with the separate billing policy as required under this final rule, if HHS finds that the Exchange or QHP issuer attempted in good faith to timely meet the requirements. Evidence of such good faith efforts might include records showing that planning for compliance with this final rule’s requirements was begun within a reasonable time following the publication of the final rule, but events outside the Exchange’s or QHP issuer’s control caused implementation delays. HHS will consider exercising this enforcement discretion based on

14See 42 U.S.C. § 18023(b)(2)(D)(ii)(I) (when estimating per member, per month cost of non-Hyde abortion services, issuers may take into account the impact on overall costs of the inclusion of such coverage).
on the circumstances of the particular Exchange or QHP issuer. We do not anticipate that HHS would exercise such discretion for an Exchange or QHP issuer that fails to meet the separate billing requirements after more than 1 year following publication of this final rule.

Comment: Many commenters who supported the proposals stated that these proposals would increase issuer compliance with the segregation of funds and separate payment requirements under section 1303 of the PPACA, and that the proposals would clarify and correct the previous administration’s interpretation of the statute. Many supporting commenters noted their dissatisfaction that abortion coverage of any kind is offered at all in the individual market, but expressed support that the proposals would better protect enrollees who object, based on their religious or moral beliefs (collectively, “conscience”), to coverage of non-Hyde abortion services.

Many commenters stated that it is a direct violation of their conscience rights to have to pay for abortion in any form, including subsidizing it through insurance coverage. Commenters stated that these proposals would increase transparency for enrollees as to what their health insurance covers and would allow enrollees to use this information to seek a plan that does not cover non-Hyde abortion services, consistent with their conscience.

Although many commenters expressed support for the proposals, many also objected to being required to pay this separate bill at all if they object to coverage of non-Hyde abortion services. Many commenters asked that HHS accommodate individuals who have conscience objections to these services by allowing enrollees in plans covering
non-Hyde abortion to “opt out” of this coverage by not paying the separate bill attributable to coverage of non-Hyde abortion services.

Many commenters stated they were unconvinced by the stated justification for the proposals (to better align the regulatory requirements for QHP issuer billing of enrollee premiums with the separate payment requirement in section 1303 of the PPACA) and instead stated that the motivation was to appease religious or political special interests. Commenters stated that the proposals would value the needs of enrollees with conscience objections to coverage of non-Hyde abortion services more highly than the needs of enrollees with a health interest in receiving coverage for non-Hyde abortion services. These commenters stated that the proposals address conscience objections of the few at the cost of the many women who need and value coverage of non-Hyde abortion services.

Many commenters asked that these proposals be withdrawn because they impose a narrow religious belief opposing a legal medical service on enrollees who do not share this viewpoint and need or value this coverage. Commenters also objected to the proposal because it singles out coverage of non-Hyde abortion services as the only service for which separate billing and payment is required, questioning why other services are not similarly subject to separate payment and billing requirements based on conscience objections. For example, one commenter expressed that they object based on their conscience to supporting coverage of individuals who get sick after refusing vaccinations for that illness. Another commenter noted that they object to having to pay for coverage of services for tobacco-related illnesses as they believe persons who
voluntarily choose to use tobacco products should not be subsidized by other enrollees for their unhealthy behaviors.

Response: Although we understand objecting commenters’ concerns, the changes are primarily meant to better align the regulatory requirements for QHP issuer billing of enrollee premiums with the statutory separate payment requirement in section 1303 of the PPACA. We acknowledge that the finalized policy regarding separate billing may increase transparency for policy holders who object on the basis of conscience to coverage of non-Hyde abortion services in their QHPs. And while it is true that this final rule treats coverage of non-Hyde abortion services differently from other covered services for purposes of QHP billing and payment, this differential treatment is based on the statutory PPACA requirement that non-Hyde abortion services be treated differently for billing, collection, payment, and federal-subsidy purposes; we are obligated to enforce the statute. Section 1303 of the PPACA has always required QHP issuers to estimate the basic per enrollee per month cost based on the average actuarial basis of the QHP’s coverage of non-Hyde abortion services, and prohibited QHP issuers from estimating that cost to be less than $1 per enrollee per month. Under the statute, QHP issuers must also collect a separate payment for that portion of the enrollee’s QHP premium attributable to coverage of non-Hyde abortion services and must segregate these payments in a separate allocation account that is to be used to pay for non-Hyde abortion services. Furthermore, section 1303 of the PPACA bars the use of PTCs or CSRs for such coverage. The changes we are finalizing at §156.280(e)(2)(ii) would strengthen regulatory alignment with the existing statutory requirements for QHP issuer billing of enrollee premiums with the separate payment requirement in section 1303 of the PPACA.
We further understand that policy holders who object, based on their conscience, to non-Hyde abortion services may prefer to not pay the separate bill attributable to coverage of these services, and thereby opt out of such coverage. We also acknowledge there may be other services covered by a plan that consumers object to or do not intend to use. As previously stated, the primary motivation for this rule is to better align the regulatory requirements for QHP issuer billing of premiums with the statutory separate payment requirement in section 1303 of the PPACA.

However, we agree that consumers are best served by the Exchanges when they can enroll in a QHP that meets their needs, from a conscience, as well as a care, perspective. In the Exchanges that use the federal platform, we have taken steps to improve transparency regarding QHP offerings to make it easier for consumers to select plans that they believe are best suited to their needs, preferences, and conscience concerns, such as information to more readily identify QHPs that offer coverage of non-Hyde abortion services. 15 State Exchanges that operate their own technology platforms have taken similar steps. For example, State Exchanges display different plan attributes to enrollees to foster the decision-making process, and allow consumers to view plan offerings by selecting filters that show plans with their desired plan characteristics. In addition, Summary of Benefits and Coverage (SBC) requirements help ensure that consumers have access to easy-to-understand information about coverage. Further, with regard to commenters that stated their dissatisfaction that abortion coverage is offered at all in the individual market, we note that section 1303(a)(1) of the PPACA specifies that

states may enact laws prohibiting QHP issuer coverage of abortion services on the Exchange. We also note that section 1303(a)(2) of the PPACA provides that a state may repeal such a law and provide for the offering of abortion coverage through the Exchange, and section 1303(b)(1)(A)(ii) of the PPACA allows QHP issuers to decide whether or not to offer coverage for abortion services, consistent with applicable state law.

Comment: Some commenters objected to HHS stating that it would enforce the requirements of section 1303 of the PPACA as codified at §156.280 directly in the event that State Exchanges do not enforce these requirements, arguing that it would be inconsistent with other HHS efforts to ensure that states can operate their programs with limited federal interference. Commenters also expressed concern that the proposed enforcement structure overrides the authority delegated to states in section 1303 of the PPACA over issuers that operate in their states, and will disrupt the nature of collaboration and partnership that the PPACA meant to create between the states and the federal government. Commenters also stated that the addition of new compliance reviews are unnecessary, as HHS does not articulate any facts or data establishing the current landscape of compliance—or lack of compliance—with existing regulations.

Many commenters stated that the 2014 U.S. Government Accountability Office report,\(^\text{16}\) which the proposed rule cites as evidence of potential remaining issuer compliance concerns, predates the 2016 Payment Notice, which clarified for issuers how to comply with the separate payment requirement. These commenters assert that HHS

offers no evidence that any compliance problems remain over 4 years later. Commenters also stated that the research to inform that report was conducted between February 2014 and September 2014, less than 1-full year after the Exchanges began operating and, as such, issuers were less likely to have fully implemented the compliance standards required under the PPACA.

Other commenters stated that compliance with section 1303 of the PPACA has been inconsistent and were supportive that the proposals would require greater oversight and transparency from State Exchanges and require them to meet the standards of section 1303 of the PPACA. Some commenters cited to the 2014 U.S. Government Accountability Office report\textsuperscript{17} as evidence of this noncompliance, and others cited to a letter sent prior to publication of the proposed rule by 102 members of Congress to HHS Secretary Alex Azar, which requested that new regulations be implemented "to remedy the severe problems with the ACA in regard to abortion coverage."\textsuperscript{18}

Response: We agree that oversight of issuer compliance with section 1303 of the PPACA is important to achieving greater transparency for consumers. We acknowledge that section 1303(b)(2)(E)(i) of the PPACA, as implemented at §156.280(e)(5), designates the state insurance commissioners as responsible for monitoring, overseeing, and enforcing the provisions in section 1303 of the PPACA related to QHP segregation of funds for non-Hyde abortion services. That is different than assigning the exclusive enforcement authority, with respect to all provisions in section 1303, to the states or to


\textsuperscript{18} Letter from Chris Smith, Member of Congress, to Alex Azar, Secretary, U.S. Department of Health and Human Services (Aug. 6, 2018), available at https://chrissmith.house.gov/uploadedfiles/2018-08-06_-smith_letter_on_section_1303_-_abortion_funding_transparency.pdf.
State Exchanges. As is the case with many provisions in the PPACA, states are generally the entities primarily responsible for implementing and enforcing the provisions in section 1303 of the PPACA related to individual market QHP coverage of non-Hyde abortion services.

However, where we are charged with directly enforcing statutory requirements in the FFE, we intend to do so fully in instances of issuer non-compliance with the separate payment requirement under section 1303 of the PPACA. Moreover, to the extent a state operating its own Exchange fails to substantially enforce these requirements, HHS is authorized to enforce them directly. Pursuant to section 1321(c)(2) of the PPACA, after determining that a state (or State Exchange) has failed to substantially enforce a federal requirement related to Exchanges and the offering of QHPs through Exchanges, including section 1303 of the PPACA’s separate payments requirement (or other requirements), the Secretary may step in to enforce the requirement against the non-compliant issuer. This enforcement structure strikes an appropriate balance between federal oversight and state flexibility with regard to the requirements of section 1303. Accordingly, unless HHS determines a state (or State Exchange) has failed to substantially enforce section 1303 of the PPACA requirements, we intend to continue to defer to states (or State Exchanges) that enforce section 1303 of the PPACA requirements. HHS disagrees that this enforcement structure in a state operating its own Exchange would override the state’s exercise of authority expressly delegated to states in section 1303 of the PPACA.

The compliance reviews governing QHP issuers participating in the FFE include reviews of compliance with section 1303 of the PPACA and §156.280. The compliance reviews for future benefit years will include the new requirements finalized in this rule.
for separate billing of the portion of the policy holder’s premium attributable to coverage of non-Hyde abortion services, as finalized at §156.280(e)(2). We continue to believe such compliance reviews will help to address remaining issuer compliance issues, if any, previously identified by the 2014 U.S. GAO report. However, commenters also expressed concern that the 2014 U.S. GAO report is outdated and that there is no evidence of ongoing compliance issues to support the changes we are finalizing regarding separate billing. But regardless of whether there are ongoing compliance issues, the changes we are finalize are primarily meant to better align the regulatory requirements for QHP issuer billing of enrollee premiums with the statutory separate payment requirement in section 1303 of the PPACA. This goal is related to overall compliance with section 1303, but has a different compliance focus than the compliance issues cited in the 2014 U.S. GAO report. Additionally, because we are amending the acceptable methods for issuers to comply with the separate payment requirement, we believe additional oversight during this transition time will be necessary to ensure that issuers are modifying their billing procedures appropriately.

FFE issuers subject to compliance reviews under §156.715 must retain all documents and records of compliance with section 1303 of the PPACA and these requirements in accordance with §156.705, and should anticipate making available to HHS the types of records specified at §156.715(b) that would be necessary to establish their compliance with these requirements. For example, FFE issuers subject to compliance reviews for §156.280 should anticipate supplying HHS with documentation

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of their estimate of the basic per enrollee per month cost, determined on an average actuarial basis, for including coverage of non-Hyde abortion services; detailed invoice and billing records demonstrating they are separately billing for and instructing policy holders to pay for in a separate transaction the portion of the policy holder’s premium attributable to coverage of non-Hyde abortion services as specified in this rule, the actuarial value which must be estimated to be no less than $1 per enrollee, per month; and appropriately segregating the funds collected from enrollees into a separate allocation account that is used to pay for non-Hyde abortion services.

We remind issuers that pursuant to §156.280(e)(5)(ii), any issuer offering coverage of non-Hyde abortion services on the Exchange must submit a plan to the relevant state insurance regulator that details the issuer’s process and methodology for meeting the requirements of section 1303(b)(2)(C), (D), and (E) of the PPACA (hereinafter, “segregation plan”). The segregation plan should describe the QHP issuer's financial accounting systems, including appropriate accounting documentation and internal controls, that would ensure the segregation of funds required by section 1303(b)(2)(C), (D), and (E) of the PPACA. Issuers should refer to §156.280(e)(5)(ii) for more information on precisely what issuers should include in their segregation plans to demonstrate compliance with these requirements. We also remind QHP issuers that pursuant to §156.280(e)(5)(iii) each QHP issuer participating in the Exchange must

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20 While we included compliance with section 1303(b)(2)(D) in the segregation plan that QHP issuers are required to submit to state insurance commissioners under our regulations at 45 C.F.R. 156.280(e)(5), we did not mean to suggest by that inclusion that such provision is part of the segregation requirements in the statutory subsection that are subject to the jurisdiction of state health insurance commissioners under section 1303(b)(2)(E).
provide to the state insurance commissioner an annual assurance statement attesting that the plan has complied with section 1303 of the PPACA and applicable regulations.

We also remind issuers offering medical QHPs in the FFEs that they already must attest to adhering to all applicable requirements of 45 CFR part 156 as part of the QHP certification application, including those requirements related to the segregation of funds for abortion services implemented in §156.280.21 As finalized, issuers in the FFE completing this attestation would also attest to adhering to these new separate billing and collection requirements. As part of the QHP certification process, issuers in states with FFEs where the states perform plan management functions must also complete similar program attestations attesting to adherence with §156.280.22 Issuers in states with State Exchanges that offer QHPs that cover non-Hyde abortion services should contact their state regarding the QHP certification process.

Comment: HHS received comments expressing a variety of legal arguments against the proposals. Many commenters stated that the proposals violate the Administrative Procedure Act (APA) because the proposals advance an unreasonable interpretation of law, are arbitrary and capricious, fail to provide adequate reasons or satisfactory explanations why HHS seeks to adopt a newly preferred interpretation of the requirement, and fail to adequately assess the costs and harms. Commenters also stated the proposals raise Federalism concerns under the Tenth Amendment because the proposals allegedly are designed to penalize states that have laws requiring QHPs to provide coverage of non-Hyde abortion services by requiring states—through their

respective Exchanges and the Department of Insurances (DOIs)—to adopt new oversight responsibilities, and make systemic changes to fit the alterations the proposals require. For these states, commenters stated that this effectively requires states to either divert extensive resources to implement these changes or change their sovereign laws to no longer require coverage of non-Hyde abortion services. Commenters also stated that the proposals exceed the federal government’s spending power by implementing new reporting and oversight obligations in the Exchanges that impose post-acceptance or retroactive conditions on states that were not originally anticipated. Commenters also stated that the proposals serve as a tax penalty on issuers for doing business in states with non-Hyde abortion services coverage requirements. One commenter stated that HHS improperly excluded the proposed changes to §156.280 among the rule changes with Federalism implications.

Commenters also stated that requiring QHP issuers to send a separate bill to enrollees about the plan’s coverage of non-Hyde abortion services constitutes a second separate notice outside of the notice included in the SBC indicating whether the plan covers abortions services and that, as such, these proposals violate section 1303(b)(3)(A) of the PPACA, which specifies that QHP issuers covering these services “shall provide a notice to enrollees, only as part of the summary of benefits and coverage explanation, at the time of enrollment, of such coverage.” Commenters further assert that the proposals violate section 1303(b)(3)(B), which states that all advertising used by issuers, any information provided by the Exchange, and “any other information specified by the Secretary” shall only provide information with respect to the total amount of the combined payments for all services.
Commenters also stated that the proposals violate section 1554 of the PPACA because these proposals will limit access to health care services, conflict with section 1557 of the PPACA, violate the Equal Protection Clause because the proposals place a heavy burden on a unique health care service only applicable to women, constitute an undue burden on a woman’s right to procreative choice, violate the unconstitutional conditions doctrine by penalizing those who choose to exercise a constitutionally-protected right by imposing unreasonable payment protocols to access abortion services, and violate the establishment clause of the First Amendment.

HHS also received many comments stating that the proposed interpretation of section 1303 of the PPACA violates congressional intent. Commenters stated that section 1303 of the PPACA makes clear that absent a state law to the contrary, issuers offering Exchange coverage can decide whether to cover non-Hyde abortion services and that these requirements effectively take that decision away from issuers. Commenters also stated that Congress specifically enacted section 1303 of the PPACA’s provisions after rejecting more extreme and restrictive alternatives that would have eliminated abortion coverage in the Exchanges or prohibited enrollees from using federal financial assistance to purchase a plan including abortion coverage, and that HHS is ignoring that legislative history by proposing changes that would have a net effect of reducing abortion coverage where issuers decide to eliminate coverage due to the regulatory burden. Commenters also noted that, although Congress decided to treat abortion differently when passing section 1303 of the PPACA, it did so specifically to ensure that private insurance plans could continue to decide whether or not to cover abortion in states that did not ban such coverage, and that this rule threatens that right. One commenter also
stated that HHS violated generally accepted principles of statutory interpretation and should have construed “separate payment” in line with industry practice.

Many commenters also stated that these proposals conflict with the Administration’s stated goals of reducing economic and regulatory burden, in conflict with several recently issued Executive Orders. Specifically commenters stated that the proposals would undermine Executive Order 13765 because these proposals would increase the administrative and economic burden of the PPACA, Executive Order 13813 which called for rules and guidelines to improve access to and the quality of information that Americans need to make informed healthcare decisions, Executive Order 13777 which orders federal agencies to alleviate unnecessary regulatory burden placed on the American people, and Executive Order 12866 because HHS did not “assess both the costs and the benefits of the intended regulation and… propose or adopt a regulation only upon a reasoned determination that the benefits of the intended regulation justify the costs,” as the Executive Order directs. Commenters also stated that the proposals would undermine CMS’s “Patients Over Paperwork” initiative aimed at reducing administrative burden on health plans and providers.

HHS also received comments arguing that these changes advance the congressional intent for the separate payment requirement in section 1303 of the PPACA, arguing that both the congressional record and the statutory language clearly demonstrate that Congress intended that billing for coverage of non-Hyde abortion services be separate.

Response: HHS disagrees with comments questioning its legal authority to make these policy changes, and disagrees that interpreting section 1303 of the PPACA to
require issuers to send a separate bill for the portion of the premium attributable to coverage of non-Hyde abortion services violates the APA. Section 1303 of the PPACA and regulations at §156.280 do not specify the method a QHP issuer must use to comply with the separate payment requirement under section 1303(b)(2)(B)(i) of the PPACA and §156.280(e)(2)(i). Although we recognized in the preamble to the proposed rule that the previous methods of itemizing or providing advance notice about the amounts noted as permissible in the preamble of the 2016 Payment Notice arguably identifies two “separate” amounts for two separate purposes, we continue to believe that requiring issuers to bill for two separate “payments” of these two amounts better aligns with, and better enables compliance with, the separate payment requirement in section 1303 of the PPACA. We also believe that consumers are more likely to make a separate payment for the non-Hyde abortion coverage when they receive a separate bill for such amount.

In fact, among the previously acceptable methods for QHP issuers to comply with the separate payment requirement outlined in the preamble to the 2016 Payment Notice was sending a separate monthly bill for these services.\footnote{Patient Protection and Affordable Care Act; HHS Notice of Benefit and Payment Parameters for 2016 (80 FR 10750, 10840).} As such, amending the policy to only permit this method of complying with the separate payment requirement does not wholly depart from the previous interpretation, it merely refines it to better reflect the statute.

Additionally, we have carefully considered the comments we received estimating the burden the proposals would impose on issuers, states, enrollees, and other entities, and agree – without accepting the estimates provided by commenters – that, as originally proposed, the actual burden would have exceeded HHS’s estimates. As such, we are
finalizing several changes described in responses to comments earlier in this section of
the preamble with the specific intent of mitigating the burden that would have been
imposed if we were finalizing as originally proposed.

HHS disagrees that the policy as originally proposed or as revised in the final rule violates state sovereignty, exceeds the federal government’s spending power, or raises other Federalism concerns. Because states are the entities primarily responsible for implementing and enforcing the provisions in section 1303 of the PPACA related to individual market QHP coverage of non-Hyde abortion services, we acknowledge that requiring issuers to separately bill for the portion of the premium attributable to these services means that states will likely adjust how they ensure issuer compliance with these new requirements. We also remind states concerned about enforcement and oversight of these requirements that, under section 1321(c) of the PPACA, states may elect not to establish and operate an Exchange, thereby deferring those responsibilities to HHS.

We are clarifying the existing statutory requirement by adding specificity to the regulatory requirement, for issuers to collect a separate payment for these services. As such, these changes do not directly impose new requirements on states other than to adjust how they check for compliance. We believe that any state oversight responsibility modified through these changes was already contemplated by section 1303 of the PPACA in identifying states as the entities primarily, but not exclusively, responsible for enforcing the provisions in section 1303. Further, as noted above, among the previously acceptable methods for QHP issuers to comply with the separate payment requirement was sending a separate monthly bill for coverage of non-Hyde abortion services. Therefore, states should already have developed mechanisms to confirm compliance with
separate monthly billing and payment for these services for any issuers that previously elected this option.

Setting aside the question of whether state laws requiring coverage of non-Hyde abortion services on the Exchange are consistent with statutory conditions on federal funding from the Department to the States, we acknowledge that some states have such laws. However, the changes we are finalizing do not preempt state law regarding coverage of non-Hyde abortion services or otherwise attempt to coerce states into changing these laws or to deny QHP issuers the ability to offer plans on the Exchanges that provide coverage of non-Hyde abortion services. HHS is simply refining the method by which issuers comply with the separate payment requirement.

HHS does not agree with commenters’ concerns that the proposals would inhibit enrollee access to appropriate and timely medical care in violation of section 1554 of the PPACA. We acknowledge that, as originally proposed, the combination of issuer burden and enrollee confusion could have potentially led to a reduction in the availability of coverage of non-Hyde abortion services (either by issuers choosing to drop this coverage to avoid the additional costs or by enrollees having their coverage terminated for failure to pay the second bill), thereby potentially increasing out-of-pocket costs for some women seeking those services. But such an effect of a separate billing requirement would not constitute a violation of section 1554. Moreover, we believe the changes we are finalizing will decrease the likelihood of these outcomes. Importantly, subject to state law, section 1303(b)(1)(A) of the PPACA makes it clear that it is ultimately at the issuer’s discretion whether to cover non-Hyde abortion services in their QHP; requiring a separate bill for these services does not limit that right.
HHS also disagrees that the policy in the proposed rule, as revised in this final rule, is inconsistent with sections 1303(b)(3)(A) or 1303(b)(3)(B) of the PPACA. Reading section 1303(b)(3) alongside section 1303(b)(2), which requires collection of separate payments, suggests that section 1303(b)(3) pertaining to notices should be read harmoniously with the separate payment requirement, rather than in conflict with those requirements, as commenters suggest. For example, the separate bill for the portion of the policy holder’s premium attributable to coverage of non-Hyde abortion services is primarily a means of ensuring separate QHP issuer collection of that portion of the policy holder’s premium, as required under section 1303(b)(2). This separate bill does not circumvent or conflict with the independent requirement in section 1303(b)(3) pertaining to notices. Further, any insight the policy holder gains from the separate bill for coverage of non-Hyde abortion services about the QHP’s coverage of non-Hyde abortion services is incidental to the primary purpose of the bill, which is to help ensure separate payment by the policy holder, and separate QHP issuer collection on this portion of the policy holder’s premium. We also note that requiring a separate bill for coverage of non-Hyde abortion services is not a violation of section 1303(b)(3), just as the separate itemization of the premium amount for such coverage on a single bill (as was previously one of the acceptable billing and premium collection methods for this amount) was not a violation of that section. Therefore, we believe it is a more reasonable interpretation of section 1303 of the PPACA that section 1303(b)(2) and 1303(b)(3) of the PPACA need not conflict when read in context with one another.

Section 1557 of PPACA prohibits discrimination on the basis of race, color, national origin, sex, age, or disability in certain health programs or activities. HHS
disagrees that the policy in the proposed rule and as revised in this final rule
discriminates against women or constitutes gender discrimination in violation of section
1557 of the PPACA or of the Equal Protection Clause. Although only women access
non-Hyde abortion services, the separate bill for the portion of the premium attributable
to coverage of these services, and any enrollee burden associated with that bill, is broadly
applicable to any policy holder in a plan that covers non-Hyde abortion services. In other
words, both men and women in plans covering non-Hyde abortion services will receive a
separate bill for the portion of the premium attributable to coverage of these services, not
just the women who may ultimately access such services.

Similarly, HHS disagrees that the proposals violate the unconstitutional
conditions doctrine, given that QHP issuers offering these services will be required to
send the separate bill to all policy holders in their plan, not just those who choose to
access non-Hyde abortion services. As such, although it may be true that enrollees who
would be most likely to need access to coverage of non-Hyde abortion services would be
most likely to intentionally enroll in a QHP with such coverage, any additional burden these
enrollees experience related to understanding and paying the second bill is unrelated to
whether enrollees actually do access coverage of non-Hyde abortion services. Therefore,
the finalized policy does not penalize enrollees for accessing their constitutionally
protected right to abortion. All policy holders would receive the separate bill for the
portion of their premium attributable to coverage of non-Hyde abortion services,
regardless of whether they could, intend to, or do, access the coverage for these services.

HHS also disagrees that the policy in the proposed rule, or as revised in this final
rule, violates the Establishment Clause or otherwise impedes the free exercise of religion.
Although it may be a secondary impact that the billing changes serve the interests of enrollees who object to coverage of non-Hyde abortion services based on their conscience, the objective for this policy change continues to be achieving better alignment with the statutory requirement for issuers to collect a separate payment for coverage of non-Hyde abortion services, as specified in section 1303 of the PPACA. As such, we reject commenter’s arguments that these proposals are religiously motivated.

We also disagree with commenters that this interpretation of section 1303 of the PPACA violates congressional intent. We acknowledge that, in drafting section 1303 of the PPACA, Congress rejected language that would have imposed more restrictive requirements on QHP issuers offering coverage of non-Hyde abortion services.24 However, although the language in section 1303 of the PPACA that Congress ultimately enacted into law permits issuers to offer coverage for non-Hyde abortion services subject to state law, this flexibility is not without limitations. As enacted, section 1303 of the PPACA requires that QHP issuers offering non-Hyde abortion coverage on the Exchanges follow specific actuarial, accounting, and notice requirements to ensure that federal funds are not used to pay for the costs of including coverage of these services under the QHP. We believe that by requiring issuers to collect separate payments, section 1303 of the PPACA contemplates sending to enrollees separate bills for these services to help ensure appropriate segregation of these funds. Furthermore, HHS previously listed “sending a separate monthly bill for these services” as one of the permissible methods for issuers to comply with the separate payment requirement in the 2016 Payment Notice.

HHS also disagrees with claims that the proposals impermissibly undermine the Executive Orders mentioned in comments. We interpret the proposals and the policy as finalized in this rule as consistent with Executive Order 13765 because the law is being “efficiently implemented” through better aligning the issuer requirements related to fulfilling section 1303 of the PPACA’s separate payment requirements with the statute. We also believe Executive Order 13813 supports the changes to the policy as finalized in this rule, since providing a separate bill to policy holders for the portion of the premium attributable to coverage of non-Hyde abortion services will “improve access to and the quality of information that Americans need to make informed healthcare decisions.”

We note that we also believe Executive Order 13877 supports the policy changes by enhancing the ability of enrollees “to choose the healthcare that is best for them” and to make “fully informed decisions about their healthcare.” Indeed, many commenters highlighted that this would be one of the positive impacts of the proposal—that the separate bill would serve to clarify for enrollees that their plan covers non-Hyde abortion services and at what cost, information which many commenters would use to decide whether to remain enrolled in that QHP or seek a QHP without such coverage. We also believe Executive Order 13777 supports the proposals and changes being finalized in this rule, since requiring a separate bill for coverage of these services helps to ensure that HHS is “prudent and financially responsible in the expenditure of funds,” by better aligning the requirements with the statute in a manner that will help to ensure that QHP issuers that offer coverage for non-Hyde abortion services collect a separate payment

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from policy holders for the portion of their premium attributable to non-Hyde abortion coverage which also helps to ensure that APTC or CSR funds are not used pay for such services.

Additionally, HHS did “assess both the costs and the benefits” of the proposed rule. However, we note that Executive Order 12866’s directive to only issue net-beneficial regulations applies only “to the extent permitted by law.” Although we have since adjusted the policy as well as the estimated burden to reflect a larger burden estimate, we continue to believe that requiring QHP issuers to separately bill the portion of the policy holder’s premium attributable to coverage of non-Hyde abortion services is a better interpretation of the statutory requirement for QHP issuers to collect a separate payment for coverage of these services, and, thus, justifies the costs.  

Lastly, although CMS’s “Patients Over Paperwork” initiative does include the goal of reducing unnecessary burden, HHS believes these changes and the added burdens associated with the changes are necessary, as the changes will better align issuer billing with the statutory requirements of the PPACA. Moreover, in line with this initiative, we believe enrollees will benefit from the additional clarity that the separate bill provides about their plan’s coverage of non-Hyde abortion services.

III. Collection of Information Requirements

This final rule contains information collection requirements as defined under the Paperwork Reduction Act of 1995 (PRA). We proposed and solicited comments on these information collection requirements (ICRs) in the notice of proposed rulemaking that

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26 This rule has been subject to interagency (including OMB) review under Executive Order 12866 and cleared by OMB for issuance and publication, indicating that the rule is consistent with Executive Orders.
published on November 9, 2018 (84 FR 56015). The information collection requirements and the reconciliation of any comment received on the requirements are discussed below.

In order to fairly evaluate whether an information collection should be approved by the Office of Management and Budget (OMB), section 3506(c)(2)(A) of the PRA requires that we solicit comment on the following issues:

- The need for the information collection and its usefulness in carrying out the proper functions of our agency.
- The accuracy of our estimate of the information collection burden.
- The quality, utility, and clarity of the information to be collected.
- Recommendations to minimize the information collection burden on the affected public, including automated collection techniques.

In our November 9, 2018 (83 FR 56015) proposed rule, we solicited public comment on each of the required issues under section 3506(c)(2)(A) of the PRA for the following ICRs.

A. Wage Estimates

To derive average costs, we generally used data from the Bureau of Labor Statistics to determine average labor costs (including a 100 percent increase for fringe benefits and overhead) for estimating the burden associated with the ICRs.\textsuperscript{27} Table 1 in this final rule presents the mean hourly wage (calculated at 100 percent of salary), the cost of fringe benefits and overhead, and the adjusted hourly wage.

As indicated, employee hourly wage estimates have been adjusted by a factor of 100 percent. This is necessarily a rough adjustment, both because fringe benefits and overhead costs vary significantly across employers, and because methods of estimating these costs vary widely across studies. However, we believe that doubling the hourly wage to estimate total cost is a reasonably accurate estimation method.

**TABLE 1: Adjusted Hourly Wages Used in Burden Estimates**

<table>
<thead>
<tr>
<th>Occupation Title</th>
<th>Occupational Code</th>
<th>Mean Hourly Wage ($/hour)</th>
<th>Fringe Benefits and Overhead ($/hour)</th>
<th>Adjusted Hourly Wage ($/hour)</th>
</tr>
</thead>
<tbody>
<tr>
<td>General and Operations Manager</td>
<td>11-1021</td>
<td>$59.56</td>
<td>$59.56</td>
<td>$119.12</td>
</tr>
<tr>
<td>Computer and Information Systems Manager</td>
<td>11-3021</td>
<td>$73.49</td>
<td>$73.49</td>
<td>$146.98</td>
</tr>
<tr>
<td>Computer Programmer</td>
<td>15-1131</td>
<td>$43.07</td>
<td>$43.07</td>
<td>$86.14</td>
</tr>
<tr>
<td>Computer System Analyst</td>
<td>15-1121</td>
<td>$45.01</td>
<td>$45.01</td>
<td>$90.02</td>
</tr>
<tr>
<td>Business Operations Specialist</td>
<td>13-1199</td>
<td>$37.00</td>
<td>$37.00</td>
<td>$74.00</td>
</tr>
<tr>
<td>Secretaries and Administrative Assistants</td>
<td>43-6014</td>
<td>$18.28</td>
<td>$18.28</td>
<td>$36.56</td>
</tr>
</tbody>
</table>

**B. Information Collection Requirements (ICRs)**

1. **ICRs Regarding General Program Integrity and Oversight Requirements (§155.1200)**

The burden associated with State Exchanges meeting the program integrity reporting requirements in §155.1200 have already been assessed and encompassed through SMART currently approved under OMB control number: 0938-1244 (CMS-10507). While we are finalizing proposals in this rule that would provide HHS the ability to focus State Exchange oversight and audit activities towards particular Exchange functions that have higher program integrity risks in a more consistent manner, and require State Exchanges and their auditors to employ auditing techniques or procedures in a more consistent manner, we do not envision these changes to have a material impact on the burden for State Exchanges. As detailed in the proposed rule and in the preamble of
this rule, these amendments are intended to allow for more targeted oversight and audits of State Exchanges that focus and direct existing HHS and State Exchange resources towards particular Exchange program areas that have higher program integrity risks, rather than having those Federal and State Exchange resources covering all program areas or covering program areas that have lower program integrity risks. Because existing resources would be directed away from certain program areas and towards program areas with higher program integrity impact across all State Exchanges, we believe the overall burden on State Exchanges would not change. Further, we are not specifying a particular sampling methodology that must be used by all State Exchanges for testing the accuracy of eligibility determinations in annual programmatic audits. This final rule therefore does not impose any new burden or revised information collection requirements pertaining to §155.1200.

2. ICRs Regarding Rules Relating to Segregation of Funds for Abortion Services (§156.280)

In §156.280(e)(2), we are finalizing that QHP issuers must send an entirely separate monthly bill to the policy holder covering only the portion of premium attributable to coverage of non-Hyde abortion, and instruct the policy holder to pay the portion of their premium attributable to coverage of non-Hyde abortion services in a separate transaction from any payment the policy holder makes for the portion of their premium not attributable to coverage of non-Hyde abortion services. Based on 2020 QHP certification data in the FFEs and SBE-FPs, we estimate that 23 QHP issuers will offer a total of 338 plans with coverage of non-Hyde abortion services in 9 FFE and SBE-FP states. For the 12 State Exchanges that will operate their own technology platforms in
2020 and have QHPs that offer coverage of non-Hyde abortion services, we have updated our methodology for identifying issuers with QHPs that offer coverage of non-Hyde abortion services, and now estimate that 71 QHP issuers will offer a total of approximately 1,129 plans that include coverage for non-Hyde abortions services. Three of those State Exchanges perform premium billing and payment processing, while the other 9 have their issuers perform premium billing and payment processing. In total, we now estimate that will be 94 QHP issuers offering a total of 1,467 plans (representing approximately 32 percent of individual market, on-Exchange plans) covering non-Hyde abortion services across 21 states in plan year 2020. As such, the ICRs associated with these proposals create a new burden on QHP issuers and State Exchanges that perform premium billing and payment processing, and thus will be submitted to OMB for final approval (OMB control number: 0938-1358 (Billing and Collection of the Separate Payment for Certain Abortion Services (CMS-10681)).

**Comment:** We used the estimated numbers of impacted issuers and plans to estimate the costs associated with the proposals regarding separate billing and payment for coverage of non-Hyde abortion services.

We received many comments from issuers, issuer associations, states, State Exchanges, state regulators, and other organizations arguing that we greatly underestimated the burden on issuers to implement the original proposals. For example, commenters stated that actual one-time costs for issuers to implement these proposals would be anywhere from $50,000 to $7,500,000 per issuer. Commenters also stated that annual costs per issuer would be anywhere from $40,000 to $10,800,000 annually. One commenter stated that the operational burden of a mid-size issuer (serving approximately
70,000 Exchange enrollees) would exceed HHS’s estimate by approximately 2,666 times for the first year alone. Commenters explained that the proposals would require changes to nearly every aspect of the enrollment and billing processes to identify impacted enrollees, generate and send multiple accurate invoices, collect multiple payments, and reconcile payment amounts.

Some commenters noted that many issuers do not have the ability to generate two separate bills for one policy and that, as such, the proposals would require them to issue two policies per policy holder (and enroll every policy holder into two separate policies to be able to bill them in the required way). Commenters stated that the proposals would consequently require that many issuers create separate member IDs in order to facilitate every enrollee receiving two bills and making two payments. Commenters stated that this would be an extraordinarily costly and difficult change for such issuers to make.

Commenters also expressed concern that requiring issuers to send the separate bill in a separate mailing would double an issuer’s postage and associated mailing costs, costing issuers an additional $15.6 to $31.2 million nationally per year, and expressed further concern that this cost was not accounted for in the proposed rule’s impact estimates. Many commenters explained that it is unrealistic to assume that issuers can save costs by enrollees switching to electronic billing, since many enrollees still elect to receive and pay their health coverage bills through the mail. Other commenters explained that many enrollees have no choice but to receive paper bills and send paper checks, as many enrollees in rural areas and many low-income individuals still do not have access to the internet.

Response: We appreciate these comments and after consideration, have adjusted
the estimated burden below. In response to these comments, we have updated the associated ICRs to reflect an increase in burden and costs for issuers. We believe that the original burden estimate in the proposed rule would not accurately reflect the actual costs issuers would have incurred if we finalized the provisions as proposed.

We estimate that allowing issuers to send the separate bill in the same mailing (though not in the same email or electronic communication) as the bill for other services would eliminate much of the commenter estimated $15.6 to $31.2 million that the second bill would have cost annually if we had finalized as proposed. By finalizing this policy to allow for combined mailings when sending paper bills, we ensure that issuers will not be required to incur the costs associated with additional postage and envelopes.

Issuers will incur burden to complete the one-time technical build to implement the necessary changes, which will involve activities such as planning, assessment, budgeting, contracting, building and testing their systems; as well as one-time changes such as billing-related outreach and call center training. We assume that this one-time burden will be incurred primarily in 2020. We estimate that, for each issuer, on average, it will take business operations specialists 2,500 hours (at $74 per hour), computer system analysts 6,500 hours (at $90.02 per hour), computer programmers 22,000 hours (at $86.14 per hour), computer and information systems managers 200 hours (at $146.98 per hour) and operations managers 300 hours (at $119.12 per hour) to complete this task. The total burden for an issuer will be approximately 31,500 hours on average, with an equivalent cost of approximately $2.7 million. We anticipate that implementing these changes within 6 months would result in issuers incurring additional costs such as higher contracting costs and overtime payments, which will increase the total cost for each
issuer by 50 percent, to approximately $4.1 million. For all 94 issuers, the total one-time burden will be 2,961,000 hours for a total cost of approximately $385 million.

We anticipate that the burden incurred by State Exchanges that perform premium billing and payment processing and have QHP issuers that offer coverage for non-Hyde abortion services will be similar to the burden incurred by QHP issuers offering coverage for non-Hyde abortion services. Therefore the total burden for a State Exchange that performs premium billing and payment processing will be approximately 31,500 hours on average, with a total cost of approximately $4.1 million. For all 3 State Exchanges that perform premium billing and payment processing, the total one-time burden will be 94,500 hours for a total cost of approximately $12.3 million.

**TABLE 2. Estimated One-time Burden per Issuer or State Exchange performing premium billing and payment processing**

<table>
<thead>
<tr>
<th>Occupation</th>
<th>Burden Hours per Respondent</th>
<th>Labor Cost per Hour</th>
<th>Total Cost per Respondent</th>
</tr>
</thead>
<tbody>
<tr>
<td>General and Operations Manager</td>
<td>300</td>
<td>$119.12</td>
<td>$35,736</td>
</tr>
<tr>
<td>Computer and Information Systems Manager</td>
<td>200</td>
<td>$146.98</td>
<td>$29,396</td>
</tr>
<tr>
<td>Computer Programmer</td>
<td>22,000</td>
<td>$86.14</td>
<td>$1,895,080</td>
</tr>
<tr>
<td>Computer System Analyst</td>
<td>6,500</td>
<td>$90.02</td>
<td>$585,130</td>
</tr>
<tr>
<td>Business Operations Specialist</td>
<td>2,500</td>
<td>$74.00</td>
<td>$185,000</td>
</tr>
<tr>
<td>Total Burden and Labor Cost per respondent</td>
<td>31,500</td>
<td></td>
<td>$2,730,342</td>
</tr>
<tr>
<td>Additional Costs due to Expedited Implementation</td>
<td>--</td>
<td></td>
<td>$1,365,171</td>
</tr>
<tr>
<td>Total per respondent</td>
<td>31,500</td>
<td></td>
<td>$4,095,513</td>
</tr>
</tbody>
</table>

**TABLE 3. Estimated One-time Burden for All Issuers and State Exchanges performing premium billing and payment processing**
<table>
<thead>
<tr>
<th>Type of Respondent</th>
<th>Number of Respondents</th>
<th>Number of Responses</th>
<th>Burden Hours Per Respondent</th>
<th>Total Burden Hours</th>
<th>Total Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Issuer</td>
<td>94</td>
<td>94</td>
<td>31,500</td>
<td>2,961,000</td>
<td>$384,978,222</td>
</tr>
<tr>
<td>State Exchange</td>
<td>3</td>
<td>3</td>
<td>31,500</td>
<td>94,500</td>
<td>$12,286,539</td>
</tr>
<tr>
<td>Total</td>
<td>97</td>
<td>97</td>
<td>31,500</td>
<td>3,055,500</td>
<td>$397,264,761</td>
</tr>
</tbody>
</table>

In addition to the one-time costs estimated, issuers will incur ongoing annual costs, such as those related to identifying impacted enrollees, ensuring billing accuracy, reconciliation, quality assurance, printing, record keeping, and document retention. We estimate that for each issuer, on average, it will take administrative assistants 20,000 hours (at $36.56 per hour), business operations specialists 2,000 hours (at $74 per hour), computer programmers 2,000 hours (at $86.14 per hour), and operations managers 120 hours (at $119.12 per hour) each year to perform these tasks. The total annual burden for each issuer will be 24,120 hours, with an equivalent cost of approximately $1.07 million. Assuming that issuers will start sending separate bills in July, 2020, the total burden for all 94 issuers for the 6 months in 2020 is estimated to be 1,133,640 hours with an equivalent cost of approximately $50.1 million. From 2021 onwards, we estimate the total annual burden for all 94 issuers will be approximately 2,267,280 hours with an associated cost of approximately $100.2 million.

We anticipate that State Exchanges performing premium billing and payment processing and which have QHP issuers that offer coverage for non-Hyde abortion services will incur costs similar to QHP issuers offering coverage of non-Hyde abortion services. Therefore, we estimate that for all 3 State Exchanges performing premium billing and payment processing, the total annual burden will be approximately 36,180 hours with an equivalent cost of approximately $1.6 million in 2020 and 72,360 hours
with an associated cost of approximately $3.2 million starting in 2021.

**TABLE 4. Estimated Annual Burden per Issuer or State Exchange performing premium billing and payment processing**

<table>
<thead>
<tr>
<th>Occupation</th>
<th>Burden Hours per Respondent</th>
<th>Labor Cost per Hour</th>
<th>Total Cost per Respondent</th>
</tr>
</thead>
<tbody>
<tr>
<td>Secretaries and Administrative Assistants</td>
<td>20,000</td>
<td>$36.56</td>
<td>$731,200</td>
</tr>
<tr>
<td>General and Operations Manager</td>
<td>120</td>
<td>$119.12</td>
<td>$14,294</td>
</tr>
<tr>
<td>Business Operations Specialist</td>
<td>2,000</td>
<td>$74.00</td>
<td>$148,000</td>
</tr>
<tr>
<td>Computer Programmer</td>
<td>2,000</td>
<td>$86.14</td>
<td>$172,280</td>
</tr>
<tr>
<td>Total per Respondent</td>
<td>24,120</td>
<td></td>
<td>$1,065,774</td>
</tr>
</tbody>
</table>

**TABLE 5. Estimated Annual Burden for All Issuers and State Exchanges performing premium billing and payment processing for 2020, 2021 and 2022**

<table>
<thead>
<tr>
<th>Type of Respondent</th>
<th>Year</th>
<th>Number of Respondents</th>
<th>Number of Responses</th>
<th>Burden Hours Per Respondent</th>
<th>Total Burden Hours Per Year</th>
<th>Total Labor Cost Per Year</th>
</tr>
</thead>
<tbody>
<tr>
<td>Issuer</td>
<td>2020</td>
<td>94</td>
<td>94</td>
<td>12,060</td>
<td>1,133,640</td>
<td>$50,091,397</td>
</tr>
<tr>
<td>State Exchange</td>
<td>2020</td>
<td>3</td>
<td>3</td>
<td>12,060</td>
<td>36,180</td>
<td>$1,598,662</td>
</tr>
<tr>
<td>Total</td>
<td>2020</td>
<td>97</td>
<td>97</td>
<td>12,060</td>
<td>1,169,820</td>
<td>$51,690,058</td>
</tr>
<tr>
<td>Issuer</td>
<td>2021, 2022</td>
<td>94</td>
<td>94</td>
<td>24,120</td>
<td>2,267,280</td>
<td>$100,182,794</td>
</tr>
<tr>
<td>State Exchange</td>
<td>2021, 2022</td>
<td>3</td>
<td>3</td>
<td>24,120</td>
<td>72,360</td>
<td>$3,197,323</td>
</tr>
<tr>
<td>Total</td>
<td>2021, 2022</td>
<td>97</td>
<td>97</td>
<td>24,120</td>
<td>2,339,640</td>
<td>$103,380,117</td>
</tr>
</tbody>
</table>

In response to comments, we reviewed our original enrollee estimates and have updated our estimates for accuracy. Based on 2019 QHP Certification Data in the FFES and SBE-FPs, we now estimate that there are approximately 442,400 enrollees in QHPs covering non-Hyde abortion services. In the 11 State Exchanges that operated their own technology platform and had issuers that offered coverage of non-Hyde abortion services in 2019, we estimate that there are approximately 2,597,700 enrollees in QHPs covering
non-Hyde abortion services. The total number of enrollees in QHPs covering non-Hyde abortion services is approximately 3.04 million in 2019. The number of QHPs covering non-Hyde abortion services will be higher in 2020 compared to 2019. Therefore, we are using the number of enrollees in such QHPs in 2019 as a lower bound for the number of enrollees who will experience an increase in burden as a result of the finalized policies.

Assuming 1.5 enrollees per policy, issuers and State Exchanges performing premium billing and payment processing will be required to send a separate bill to approximately 2 million policy holders. We understand that, although enrollees can often choose to pay electronically or by phone, choose to utilize automatic payment deductions, and often opt out of receiving paper bills, many enrollees still opt to receive physical mail detailing their coverage. We also understand that many enrollees face barriers to accessing the internet and have little choice but to receive paper bills. Because enrollees typically receive paper bills and because many enrollees already face barriers to accessing the internet, issuers are likely to experience an increased administrative cost in having to print an additional monthly bill for the majority of their policy holders. According to one commenter, issuers send paper bills to 92 percent of Exchange customers. We anticipate that the number of consumers opting for electronic bills will increase over time. Therefore, we assume that approximately 90 percent of policy holders will receive paper bills in 2020 and issuers and State Exchanges performing premium billing and payment processing will need to print and send approximately 1.82 million separate paper bills per month. Assuming materials and printing cost of $0.05 per page, issuers will incur additional monthly costs of approximately $91,200 to print separate bills for impacted policy holders in 2020. Assuming that issuers start sending
separate bills in July 2020, for the 6 months in 2020, total cost for all issuers is estimated to be approximately $547,225. Assuming that more consumers will opt to receive electronic bills over time, we estimate that approximately 88 percent of policyholders will receive paper bills in 2021, and the annual cost for all issuers to send separate paper bills will be approximately $1,070,129. We assume that, in 2022, approximately 86 percent of policyholders will receive paper bills, and the annual cost for all issuers to send separate paper bills will be approximately $1,045,808. The average annual materials and printing cost over 3 years (2020 to 2022) will be approximately $887,721. Since issuers and State Exchanges performing premium billing and payment processing will be permitted to send both bills together when sending bills in a physical mailing, they will not incur any additional mailing costs. We assume that bills sent electronically can be sent at minimal cost and note that we have incorporated any associated IT changes to accommodate electronic billing changes based on this rule above, where we discussed premium billing and payment processing costs to issuers and State Exchanges.

FFE issuers are subject to future HHS compliance reviews, requiring issuers in the FFE to maintain and submit records to HHS showing compliance with separately billing for the portion of the policy holder’s premium attributable to non-Hyde abortion services as specified in this rule. Commenters stated that HHS excluded an evaluation of the burden and cost for FFE issuers to participate in the additional HHS compliance reviews, ignoring the potential for any new costs associated with this requirement, such as documenting all efforts for audit purposes. We have revised our burden estimates to account for additional recordkeeping costs not reflected in the proposed rule’s estimates but reiterate that the requirements associated with compliance reviews were already
assessed and subsumed within issuer burdens described in previously finalized rules, including the information collection currently approved under OMB control number: 0938-1277 (Program Integrity: Exchange, Premium Stabilization Programs, and Market Standards; Amendments to the HHS Notice of Benefit and Payment Parameters for 2014 (CMS-10516)).

To show compliance with FFE standards and program requirements, all issuers seeking QHP certification in FFES are required to submit responses to program attestations as part of their QHP application. This response already includes an attestation that the issuer agrees to adhere to the requirements related to the segregation of funds for abortion services implemented in §156.280. We have determined that the requirements associated with QHP certification have already been assessed and encompassed by the information collection currently approved under OMB control number: 0938-1187 (Establishment of Exchanges and Qualified Health Plans; Exchange Standard for Employers (CMS-10433)).

C. Summary of Annual Burden Estimates for Proposed Requirements

**TABLE 6: Annual Recordkeeping and Reporting Requirements**

<table>
<thead>
<tr>
<th>Regulation Section(s)</th>
<th>OMB control number</th>
<th>Number of Respondents</th>
<th>Number of Responses</th>
<th>Burden per Response (hours)</th>
<th>Total Annual Burden (hours)</th>
<th>Total Labor Cost of Reporting ($)</th>
<th>Capital Costs (Printing and Materials) ($)</th>
<th>Total Cost ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>§156.280</td>
<td>0938-NEW</td>
<td>97</td>
<td>97</td>
<td>30,600</td>
<td>2,968,200</td>
<td>$218,571,684</td>
<td>$887,721</td>
<td>$219,459,405</td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td>97</td>
<td>97</td>
<td>30,600</td>
<td>2,968,200</td>
<td>$218,571,684</td>
<td>$887,721</td>
<td>$219,459,405</td>
</tr>
</tbody>
</table>

D. Submission of PRA-Related Comments

We have submitted a copy of this final rule to OMB for its review of the rule’s information collection and recordkeeping requirements. The requirements are not
effective until they have been approved by OMB.

IV. Regulatory Impact Analysis

A. Statement of Need

This final rule implements standards to ensure enrollees receive the correct amount of APTC and CSRs at the time of enrollment or re-enrollment via periodic data matching requirements. In addition, the provisions in this rule strengthen the mechanisms and tools for overseeing ongoing compliance by State Exchanges with federal program requirements. Finally, the provisions in this rule refine some of the methods for billing of the separate payment for the portion of the policy holder’s premium attributable to non-Hyde abortion services to better align with congressional intent regarding the separate payments provision of section 1303 of the PPACA. The following summary focuses on the benefits and costs of the requirements in this final rule.

B. Overall Impact

We have examined the impact of this rule as required by Executive Order 12866 on Regulatory Planning and Review (September 30, 1993), Executive Order 13563 on Improving Regulation and Regulatory Review (January 18, 2011), the Regulatory Flexibility Act (RFA) (September 19, 1980, Pub. L. 96-354), section 1102(b) of the Act, section 202 of the Unfunded Mandates Reform Act of 1995 (March 22, 1995; Pub. L. 104-4), Executive Order 13132 on Federalism (August 4, 1999), the Congressional Review Act (5 U.S.C. 804(2)), and Executive Order 13771 on Reducing Regulation and Controlling Regulatory Costs (January 30, 2017).
Executive Orders 12866 and 13563 direct agencies to assess all costs and benefits of available regulatory alternatives and, if regulation is necessary, to select regulatory approaches that maximize net benefits (including potential economic, environmental, public health and safety effects, distributive impacts, and equity), to the extent permitted by law. Section 3(f) of Executive Order 12866 defines a “significant regulatory action” as an action that is likely to result in a rule: (1) having an annual effect on the economy of $100 million or more in any 1 year, or adversely and materially affecting a sector of the economy, productivity, competition, jobs, the environment, public health or safety, or state, local or tribal governments or communities (also referred to as “economically significant”); (2) creating a serious inconsistency or otherwise interfering with an action taken or planned by another agency; (3) materially altering the budgetary impacts of entitlement grants, user fees, or loan programs or the rights and obligations of recipients thereof; or (4) raising novel legal or policy issues arising out of legal mandates, the President’s priorities, or the principles set forth in the Executive Order.

A regulatory impact analysis (RIA) must be prepared for rules with economically significant effects ($100 million or more in at least 1 year). This final rule is economically significant within the meaning of section 3(f)(1) of the Executive Order. Therefore, OMB has reviewed these regulations and HHS has provided an assessment of the potential costs, benefits, and transfers associated with this rule. Accordingly, we have prepared an RIA that presents the costs and benefits of this final rule.

C. Impact Estimates of the Program Integrity Provisions and Accounting Table

In accordance with OMB Circular A-4, Table 7 depicts an accounting statement summarizing HHS’s assessment of the benefits, costs, and transfers associated with this
regulatory action. Table 8 includes a summary of annualized values of costs, over a perpetual time horizon at 7 percent discount rate for Executive Order 13771 (E.O. 13771). This final rule implements standards that will have numerous effects, including ensuring that eligible enrollees receive the correct amount of APTC and CSR (as applicable); improving alignment with the separate payment requirement in section 1303 of the PPACA by requiring QHP issuers to send separate bills to policy holders for the portion of their premium attributable to non-Hyde abortion services; conducting effective and efficient monitoring and oversight of State Exchanges to ensure that enrollees are receiving the correct amount of APTC and CSRs in State Exchanges, and that State Exchanges are meeting the standards of federal law in a transparent manner; and protecting the interests of taxpayers, and enrollees, and the financial integrity of Exchanges through oversight of health insurance issuers, including ensuring compliance with the requirements of section 1303 of the PPACA. We are unable to quantify certain benefits and costs of this final rule – such as benefits to enrollees for timely notification of their dual enrollment in other qualifying coverage such as Medicare, Medicaid/CHIP, and, if applicable, the BHP, potential increases in cost to states for increased oversight activities and to establish access to federal data systems to verify eligibility for or enrollment in Medicaid/CHIP or Medicare, and potential costs to enrollees such as increased out-of-pocket costs related to billing changes due to the separate payment requirements for non-Hyde abortion services. The effects in Table 7 reflect qualitatively assessed impacts and estimated direct monetary costs and transfers resulting from the provisions of this final rule for health insurance issuers. States impacted by PDM requirements will incur costs of up to $6.9 million in 2020. In addition, we estimate that
issuers, State Exchanges, FFEs, and consumers impacted by the separate billing and payment requirements will incur costs of approximately $546.1 million in 2020, $232.1 million in 2021, $230.7 million in 2022, and $229.3 million 2023 onwards (see Table 10 below). We also expect that transfers from the federal government to consumers in the form of premium tax credits will decrease as a result of Exchanges conducting Medicare, Medicaid/CHIP, and, if applicable, BHP PDM, and increase as a result of separate billing and payment requirements. The net increase in premium tax credits is estimated to be approximately $106 million in 2021 and $96 million in 2022 onwards.

**TABLE 7: Accounting Table**

| Benefits: |
| Qualitative: |
| • Better alignment of the regulatory requirements for QHP issuer billing of premiums with the separate payment requirement in section 1303 of the PPACA. |
| • Clearer regulatory requirements for how frequently Exchanges should be conducting periodic checks for dual enrollment in other qualifying coverage. |
| • Clearer regulatory requirements for State Exchanges around CMS’s oversight and reporting process that allows for more effective oversight of State Exchanges. |

<table>
<thead>
<tr>
<th>Costs:</th>
<th>Estimate</th>
<th>Year</th>
<th>Discount Rate</th>
<th>Period Covered</th>
</tr>
</thead>
<tbody>
<tr>
<td>Annualized Monetized ($/year)</td>
<td>$304.09 million</td>
<td>2019</td>
<td>7 percent</td>
<td>2020-2024</td>
</tr>
<tr>
<td></td>
<td>$298.92 million</td>
<td>2019</td>
<td>3 percent</td>
<td>2020-2024</td>
</tr>
</tbody>
</table>

| Quantitative: |
| • Burden incurred by issuers, states, federal government and enrollees to comply with provisions related to coverage of non-Hyde abortion services and the segregation of premiums for such services. |
| • Costs for State Exchanges not in compliance with regulatory requirements to conduct Medicare, Medicaid/CHIP, and, if applicable, BHP PDM. |

| Qualitative: |
| • Potential increase in costs to states for increased oversight of separate payment requirements. |
| • Potential increased costs incurred by enrollees who choose to make separate payments for coverage of non-Hyde abortion services. |
| • Potential increased burden and costs for State Exchanges to authorize access to federal data sources to verify Medicare and Medicaid/CHIP eligibility and/or enrollment, notifying enrollees when dual enrollment is detected, and process QHP coverage terminations. |
| • Potential increased burden for assisters, agents and brokers to explain new billing process. |
| • Potential increase in public spending and out-of-pocket costs to enrollees if there is an increase in unplanned pregnancies due to loss of abortion coverage and, with respect to public spending, if those unplanned pregnancies are experienced by individuals who would be eligible for public benefit programs. |
| • Potential decrease in broker and issuer revenue due to decrease in QHP enrollment. |
Transfers:

<table>
<thead>
<tr>
<th>Description</th>
<th>Estimate</th>
<th>Year</th>
<th>Discount Rate</th>
<th>Period Covered</th>
</tr>
</thead>
<tbody>
<tr>
<td>Federal Annualized Monetized ($)</td>
<td>$76.2 million</td>
<td>2019</td>
<td>7%</td>
<td>2020-2024</td>
</tr>
<tr>
<td></td>
<td>$77.7 million</td>
<td>2019</td>
<td>3%</td>
<td>2020-2024</td>
</tr>
</tbody>
</table>

Quantitative:
- Total transfers from the federal government to enrollees due to an increase in premium tax credit payments.

Qualitative:
- Increase in premiums beginning in plan year 2021.
- Potential increase in out-of-pocket costs for enrollees who experience lapse in coverage for failing to make payments for coverage of non-Hyde abortion services due to confusion with new billing system.
- Potential increase in out-of-pocket costs for individuals who lose health insurance coverage due to increase in premiums.
- Potential increase in uncompensated care costs for people who lose health insurance coverage.

**TABLE 8: EO 13771 Summary Table (in $ Millions 2016 dollars, over a perpetual time horizon)**

<table>
<thead>
<tr>
<th>Description</th>
<th>Estimate (7% Discount rate)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Annualized Costs</td>
<td>$182.98</td>
</tr>
<tr>
<td>Annualized Cost Savings</td>
<td>$0</td>
</tr>
<tr>
<td>Annualized Net Costs</td>
<td>$182.98</td>
</tr>
</tbody>
</table>

1. Functions of an Exchange (§155.200)

   Our revisions to §155.200(c) specifying that Exchanges must perform oversight functions or cooperate with activities related to oversight and financial integrity requirements are a clarification and not a new function. Therefore, they will not impose additional burdens on State Exchanges.

2. Eligibility Redetermination During a Benefit Year (§155.330)

   Our requirement that Exchanges conduct Medicare PDM, Medicaid/CHIP PDM, and, if applicable, BHP PDM at least twice a year beginning with the 2021 calendar year, adds specificity to the existing requirement that Exchanges must periodically examine
available data sources to determine whether Exchange enrollees have been determined eligible for or enrolled in other qualifying coverage such as Medicare, Medicaid, CHIP, or, if applicable, the BHP. Therefore, we expect the costs associated with this requirement to be minimal. However, State Exchanges that are not already conducting PDM with the required frequency, or deemed in compliance with the Medicaid, CHIP, and, if applicable, BHP PDM requirements, will be required to engage in IT system development activity in order to communicate with these programs and act on enrollment data either in a new way, or in the same way more frequently. Thus, there may be additional associated administrative cost for these State Exchanges to implement the proposed PDM requirements. We anticipate a majority (up to eight) of the twelve State Exchanges that operate their own technology platforms would be exempt from the requirement to perform Medicaid/CHIP, and, if applicable, BHP PDM because they have shared, integrated eligibility systems with their respective Medicaid programs, as such they would be deemed in compliance with this requirement. However, we are not able to confirm the exact number because we have not yet set specific criteria and process to assess and confirm which State Exchanges would be exempt, and would need additional operational information from State Exchanges to confirm our assessment. We will establish and engage in that process after finalization of the rule. For a State Exchange not already conducting Medicare, Medicaid/CHIP, and, if applicable, BHP PDM at least twice a year, and that does not already have a shared, integrated eligibility system with its respective Medicaid/CHIP, and, if applicable, BHP programs, we estimate that it will cost approximately $1,740,000 per State Exchange (a total of $6,960,000 for all 4 nonexempt State Exchanges) to build such capabilities in their system. We assume that this cost will
be incurred primarily in 2020. These costs would be incurred by the State Exchange as they are required to be financially self-sustaining and do not receive federal funding for their establishment or operations.

We believe these changes will support HHS’s program integrity efforts regarding the Exchanges by helping promote a balanced risk pool for the individual market as Medicare and Medicaid/CHIP beneficiaries tend to be higher utilizers of medical services, ensuring that consumers are accurately determined eligible for APTC and income-based CSRs, and safeguarding consumers against enrollment in unnecessary or duplicative coverage. Such unnecessary or duplicative coverage, coupled with typically higher utilization, generally results in higher premiums across the individual market, leading to unnecessarily inflated expenditures of federal funds on PTC for taxpayers eligible for PTC in the individual market. We estimate that requiring State Exchanges to perform Medicare PDM twice a year will result in a reduction in PTC payments of approximately $500 million over a 9-year period (Table 9). We believe this will not have any discernable impact on premiums.

**TABLE 9: Medicare PDM effect on Premium Tax Credit Outlays**

<table>
<thead>
<tr>
<th>Fiscal Year</th>
<th>2021</th>
<th>2022</th>
<th>2023</th>
<th>2024</th>
<th>2025</th>
<th>2026</th>
<th>2027</th>
<th>2028</th>
<th>2029</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>PTC ($ millions)</td>
<td>-40</td>
<td>-50</td>
<td>-50</td>
<td>-50</td>
<td>-60</td>
<td>-60</td>
<td>-60</td>
<td>-60</td>
<td>-70</td>
<td>-500</td>
</tr>
</tbody>
</table>

3. General Program Integrity Oversight Requirements (§155.1200)

We do not anticipate the changes to §155.1200(b)(2) will result in any additional cost for State Exchanges because the changes leverage an existing reporting mechanism
currently used by all State Exchanges, the annual SMART, for meeting eligibility and enrollment reporting requirements. Additionally, State Exchanges are already required to annually contract with, and budget accordingly for, an external independent audit entity to perform an annual financial and programmatic audit as required under §155.1200(c). We believe the flexibility under the new §155.1200(d)(2) to permit HHS to target the scope of annual programmatic audits to focus on the program areas that are most pertinent to a State Exchange model (including SBE-FPs), or have the greatest program integrity implications, would allow State Exchanges to utilize the funds that they already allocate to contracting with an external independent audit entity in the most cost-effective manner. We also believe the flexibility we are providing to State Exchanges in the sampling method employed by their external independent audit entities for testing the accuracy of eligibility determinations in the annual programmatic audits, along with the flexibility for HHS to set the reporting deadlines for State Exchanges under §155.1200 on an annual basis, will also allow State Exchanges to utilize the funds that they have already allocated to these activities in the most cost-effective manner.

4. Segregation of Funds for Abortion Services (§156.280)

In §156.280, we proposed to amend billing and premium collection requirements related to the separate payment requirement for coverage of abortions for which public funding is prohibited pursuant to section 1303 of the PPACA, as implemented at §156.280. We originally proposed that QHP issuers send an entirely separate monthly bill in a separate envelope to the policy holder for only the portion of premium attributable to coverage of non-Hyde abortion services, and instruct the policy holder to pay the portion of their premium attributable to coverage of non-Hyde abortion services
in a separate transaction from any payment the policy holder makes for the portion of their premium not attributable to coverage of non-Hyde abortion services. We are also finalizing that QHP issuers must begin complying with these billing changes on or before the date that is 6 months after publication of the final rule. If the date that is 6 months after publication of the final rule falls in the middle of the QHP issuer’s billing cycle (in other words, after the QHP issuer has already sent out bills to policy holders for that month), QHP issuers would be expected to begin complying the next billing cycle immediately following that date. We will consider extending enforcement discretion to an Exchange or QHP that fails to timely comply with the separate billing policy as required under this final rule, if we find that the Exchange or QHP issuers attempted in good faith to timely meet the requirements. We believe these changes to the proposed policy will advance HHS’s goal of more closely aligning the regulatory requirements for QHP issuer billing of premiums with the separate payment requirement in section 1303 of the PPACA, while also mitigating the overall burden to affected issuers, states, and enrollees.

HHS received many comments stating that we greatly underestimated the burden caused by these proposals. Although we recognized in the proposed rule that QHP issuers that cover non-Hyde abortion services would experience an increase in burden as a result of finalizing these changes, we are committed to mitigating issuer burden where possible and, as such, are finalizing changes to §156.280(e)(2) that we believe will result in a lower overall regulatory burden than what issuers would have incurred if the provisions were finalized as originally proposed. Specifically, we are amending the proposals at §156.280(e)(2) to finalize in a new paragraph at §156.280(e)(2)(ii)(A) that
QHP issuers offering coverage of non-Hyde abortion services through an Exchange must send an entirely separate monthly bill to the policy holder for the portion of premium attributable to coverage of non-Hyde abortion services, but they will be permitted to send this separate bill in the same mailing (although not in the same email or electronic communication) as the bill for the portion of the policy holder’s premium not attributable to coverage of non-Hyde abortion services when sending paper copies of bills to policy holders. We are finalizing that, when issuers sending or issuing bills electronically, the issuer must send or issue a separate bill for the portion of the premium attributable to coverage of non-Hyde abortion services in a separate email or electronic communication from the bill for the rest of the policy holder’s premium. We are also finalizing at a new paragraph §156.280(e)(2)(ii)(B) the requirement that, although the QHP issuer would not be permitted to refuse a combined payment on the basis that the policy holder did not send two separate payments as requested by the QHP issuer, and to then terminate the policy, subject to any applicable grace period, for non-payment of premiums, the QHP issuer must continue to instruct the policy holder to pay the portion of their premium attributable to coverage of non-Hyde abortion services in a separate transaction from any payment the policy holder makes for the portion of their premium not attributable to coverage of non-Hyde abortion services. We are also finalizing that QHP issuers must begin complying with these billing changes on or before the date that is 6 months after publication of the final rule. We believe these changes to the proposed policy will advance HHS’s goal of more closely aligning the regulatory requirements for QHP issuer billing of premiums with the separate payment requirement in section 1303 of the PPACA, while also mitigating the overall burden to affected issuers, states, and enrollees.
However, we acknowledge that the changes we are finalizing will still result in additional burden for issuers. HHS received many comments on the original proposals arguing that the burden imposed on issuers would significantly exceed the estimated burden included in the proposed rule. Some commenters from the issuer community conducted internal surveys, providing detailed accounts to HHS of the various ways in which they believe HHS underestimated the burden and detailing the various issuer and Exchange activities that would be necessary for implementation that HHS failed to account for in estimating the burden.

The following one-time changes are issuer activities that commenters stated HHS should account for in response to the proposed policy, and that we expect may still be necessary for issuers under the amendments we are finalizing: planning, assessment, budgeting, funding approval, and allocating funds and resources for the actual technical build (a process of 6 to 9 months); changes to system architecture to allow multiple billing statements per policy holder; changes to enrollment systems to identify enrollees subject to separate billing and payment requirements; automating the processes to send separate invoices (mail or electronic communication); adding electronic communications and payment links (for example, to issuer’s online payment portal) for enrollees to pay separately for the separate bill; changes to call center training/scripting, response processes, billing-related outreach, and interactive voice response (IVR) technology; changes to enrollee notifications related to non-payment and the 3-month grace period; updating Health Insurance Casework System (HICS) and DOI complaint processes, changes to grievance/appeals processes; and testing to ensure accuracy of separate billing processes. Commenters also stated that HHS should have accounted for the development
of new training materials. Commenters explained that issuers would need to develop additional materials and training modules for customer service representatives, brokers, and agents, so that they could address member questions and educate them, particularly on the risk of losing coverage should members fail to pay the multiple bills.

We expect the following one-time activities to add burden for issuers as issuers must still make system changes to accommodate policy holders paying separately, potential changes to binder payment processing to collect two separate payments to effectuate enrollment; changes to processes to intake payments, including automating ability to match identity and match multiple payments from a policy holder; changes to pay-by-phone and online payment portal to support dual invoices and separate payments, while also supporting combined payments for enrollees who do not make separate payments; changes to processes for enrollment and payment reconciliation, including 834 matching to effectuate enrollments; and adding new processes to address scenarios where an enrollee’s payment is not processed because the bank flags payment as potentially fraudulent (expected to occur for multiple payments in the same day or $1 payments).

Commenters also noted several activities issuers would have to complete annually to effectively implement these proposals would also significantly raise the annual burden for issuers. The following annual changes are activities raised by commenters in response to the proposed policy, but that we expect will still be relevant under the amendments we are finalizing: generating separate billing statements (paper or electronic) and additional member education materials to explain separate billing; administrative expenses in generating twice as many bills; quality assurance to ensure accuracy of separate billing statements; additional customer service resources, including
additional staffing and training, to address enrollee questions, confusion, frustration, etc.;
increased resources for HICS/DOI case resolution; system testing for billing accuracy;
identifying enrollees who did not meet an issuer’s premium payment threshold and enter
a grace period for non-payment of premium if they fail to pay the second bill; managing
the grace period process for a higher volume of enrollees who enter a non-payment grace
period (notices, termination, appeals process, reinstatement), and verification and
reconciliation of the two separate bills. Commenters also stated that issuer costs should
account for additional staffing since issuers would need to hire additional FTEs for
reconciliation and auditing of the enrollment, billing, delinquency and payment processes
and to manage the added complexity for the Exchange back-end processes.

Because the policy as finalized will require QHP issuers to instruct the policy
holder to pay the portion of their premium attributable to coverage of non-Hyde abortion
services in a separate transaction from any payment the policy holder makes for the
portion of their premium not attributable to coverage of non-Hyde abortion services, we
anticipate that the burden associated with the following annual activities raised by
commenters will still be relevant: budgeting for fees for collecting and processing
multiple payments, such as bank processing fees; processing and reconciling separate
payments (paper and electronic) sent by enrollees; additional resources for manual review
where automated processes are not able to reconcile enrollments and payments; and
managing the grace period process for a higher volume of enrollees who enter a
non-payment grace period (notices, termination, appeals process, reinstatement).

Comment: Many commenters expressed concerns that these burdens would fall
hardest on those issuers in states that require QHPs to cover non-Hyde abortion services,
and that if issuers in these states find the requirements overly burdensome they would not have an option to eliminate coverage of non-Hyde abortion services and would thus have to absorb all associated costs or pass those costs onto enrollees. One commenter stated that the proposals are also likely to have an impact off-Exchange, as issuers offering plans on the Exchange are also generally required under guaranteed availability to offer the plans off the Exchange, and that because these administrative processes are fixed investments across all plans, it is likely that many plans would simply change their systems to apply to all plans even though the proposals would only require QHPs to comply.

Response: Setting aside the question of whether state laws requiring coverage of non-Hyde abortion services on the Exchange are consistent with statutory conditions on federal funding from the Department to the States, we acknowledge that some states have such laws. The changes we are finalizing do not preempt state law regarding coverage of non-Hyde abortion services or otherwise attempt to coerce states into changing these laws. Although we acknowledge that issuers in these states would incur additional costs if they choose to continue offering individual market plans, HHS is refining the method issuers use to comply with the separate payment requirement, changes that we believe are necessary to align issuer billing with the separate payment requirement in section 1303 of the PPACA.

The burden and costs related to the one-time technical changes have been previously estimated in section III “Collection of Information Requirements” of this final rule. We have also updated HHS’s estimates in the Collection of Information Requirements section to reflect some of the increased annual burden to be incurred by
issuers. Additionally, based on comments we received, we estimate that issuers will incur ongoing annual costs associated with activities such as processing and reconciling separate payments, support for enrollees who enter grace period for non-payments, customer service, outreach and compliance. We estimate that each issuer will incur additional annual costs of approximately $1 million for these activities. Assuming that issuers will start sending separate bills in July 2020, the total annual cost of for all 94 issuers will be approximately $47 million for the 6 months in 2020 and $94 million for 2021 onwards. Since issuers will not be able to take the costs incurred in 2020 into consideration when setting rates for the 2020 plan year, it is possible that some issuers will exit the individual market or incur losses. We acknowledge that QHP issuers may choose to make similar billing changes off-Exchange to maximize their investment in making system changes to comply with the separate billing policy required for on-Exchange QHPs. However, we note that the separate billing policy we are finalizing only requires QHP issuers to implement the required changes for their on-Exchange QHPs offering non-Hyde abortion coverage.

Comment: Commenters also stated that issuers would be required to consider the added operational and administrative costs when setting actuarially sound rates, which would lead to higher premiums for enrollees. Commenters also expressed concern that the additional administrative costs would be so high that they would place issuers at risk of not meeting the required Medical Loss Ratio (MLR) limits.

Response: We believe that the changes we are finalizing to §156.280(e)(2) will result in a lower burden than the provisions as originally proposed and as such will lessen the degree to which issuers have to raise enrollee premiums. However, we acknowledge
that issuers will still incur significant burden and costs as estimated above. Based on the total premiums in the 21 states that have QHP issuers offering non-Hyde abortion coverage, we estimate that there will be no premium impact in 2020 (as plan year 2020 premium rates will already be finalized), and an approximate premium impact of up to 1.0 percent in plan year 2021 and each year thereafter.

We also estimate that enrollment will be reduced in the impacted states very slightly as a result of the increase to premiums. In plan year 2021 and each year after, we estimate that APTC amounts will be increased by up to $146 million when premium rates reflect the projected additional administrative and operational expense burdens. We do not anticipate that the policies finalized at §156.280(e)(2) will measurably increase MLR rebates as we believe that QHP issuers would either cease offering coverage of non-Hyde abortion services (unless state law requires QHP issuers to offer coverage of non-Hyde abortion services) in the plan year following the effective date to avoid issuing additional MLR rebates or would pay for the increased administrative costs from a different revenue source. Further, as noted elsewhere in this rule, among the previously acceptable methods for QHP issuers to comply with the separate payment requirement was sending a separate monthly bill for these services. Therefore, if any issuers already elected this option, there should be no change or impact on MLR rebates as a result of the policies finalized at §156.280(e)(2). We believe these additional costs are necessary to achieve better alignment of issuer billing with the statute, and strikes a better balance between burden and benefit than if HHS were to require issuers to send the separate bill in a separate mailing.
**Comment**: Commenters also expressed concerns with the burdens these changes would impose on Exchanges, which commenters noted would need to make time consuming and resource intensive changes to their websites, enrollment systems, and customer service and outreach efforts (including the reallocation of marketing funds that currently provide critical enrollee outreach which drives Exchange success) to align with the separate billing and payment requirements, which would be costly and disrupt states’ Exchange efficiency. Commenters noted a variety of changes Exchanges would be required to make, including communicating the new separate billing and payment requirement to enrollees during the enrollment process; updating the online payment portal (the “Pay Now” button on HealthCare.gov) to collect the binder payment through two separate transactions; updating the enrollment materials and notices that reference binder payment requirements to effectuate coverage, updating call center scripting and customer service to address questions related to separate billing and payment (since questions related to payments should be referred to the issuer, but that the call center should be prepared to answer questions about why enrollees are required to make multiple payments); and update complaint processes to address complaints and questions related to separate bills and payments.

One commenter estimated that the proposed changes would cost $250,000 annually for its State Exchange customer service center, $152,000 annually for customer outreach, and $19,000 annually to resolve customer complaints and appeals. Another commenter estimated that the proposals would cost its state Exchange an additional $2.9 million annually in customer service costs, $2.25 - $2.75 million for IT system changes, and $3.6 million annually for outreach and education, which reflects one-quarter of that
state Exchange’s annual advertising and outreach budget. Commenters also stated that, because the proposed changes would lead to decreased QHP enrollment, the proposed rule would cause a corresponding loss of revenue to the Exchange. Commenters also highlighted how any lapse or loss of enrollee coverage due to these proposals would result in more individuals turning to state-funded programs or emergency care for their treatment needs and that any loss of coverage would decrease the size of the risk pool and increase the cost of uncompensated care, driving medical costs and health insurance rates higher generally. For example, one commenter estimated that each one percentage point decline in the uninsured rate is associated with a $167 million drop in uncompensated care.

**Response:** We acknowledge that these provisions will impact Exchange operations. Exchanges perform important enrollee-facing functions that could be integral to issuer and enrollee compliance with the new requirements. Ultimately, we believe the changes we are finalizing will mitigate some of the burden on Exchanges that would have been incurred if we were finalizing as proposed by decreasing potential enrollee confusion and lessening potential issuer burden.

We anticipate that State Exchanges will incur additional one-time costs associated with technical changes such as updating online payment portals to accept separate payments and updating enrollment materials and notices that reference binder payments. In addition, State Exchanges will incur ongoing annual costs associated with increased customer service, outreach, and compliance. Based on comments, we estimate that each State Exchange will incur, on average, one-time costs of $750,000 in 2020, and ongoing annual costs of approximately $200,000 for the 6 months in 2020 and $400,000 in 2021.
We anticipate that ongoing annual costs will decrease over time as consumers become used to receiving and paying separate bills. We estimate that ongoing annual costs will be approximately $300,000 for each State Exchange in 2022 and $200,00 in 2023 and after. The total one-time cost for all 12 State Exchanges affected by these requirements will be approximately $9 million in 2020. Total ongoing costs for all 12 State Exchanges is estimated to be approximately $2.4 million in 2020, $4.8 million in 2021, $3.6 million in 2022 and $2.4 million 2023 onwards. In addition, we anticipate that the 3 State Exchanges that perform premium billing and payment processing will incur annual ongoing costs similar to QHP issuers that offer coverage of non-Hyde abortion services, as discussed above. We estimate that each State Exchange that performs premium billing and payment processing will incur additional annual costs of approximately $1 million. The total annual cost for all 3 State Exchanges performing premium billing and payment processing will be approximately $1.5 million in 2020 and $3 million for 2021 onwards.

Comment: One commenter also stated that the federal government will incur additional expenses due to additional personnel time and other resources needed to ensure that QHPs on the FFEs comply with the proposed rule’s requirements and to ensure compliance if a State Exchange is unable to do so, costs that will be passed on to consumers in the form of taxes.

Response: We acknowledge that the FFEs will experience added burden as a result of the final policy. However, because federal government compliance efforts will be covered primarily by FFEs user fees, we disagree that the added costs on the FFEs will be passed on to consumers in the form of taxes (though any increase in user fees may be passed on to enrollees in the form of increased premiums). We do, however, anticipate
that the FFES will incur additional costs due to one-time technical changes and increased call volumes and additional customer services efforts. We do not anticipate that the FFES will need to make any operational changes to comply with these final policies. We estimate that the FFES will incur a one-time cost of $750,000 in 2020 and ongoing annual cost of approximately $400,000 in 2020 and $800,000 in 2021 to implement these provisions. As consumers become used to receiving and paying separate bills, the ongoing costs should decrease. We estimate that ongoing costs will be approximately $600,000 in 2022 and $400,000 in 2023 onwards.

**Comment:** Commenters stated that Navigators and in-person assisters will also need to invest time and training resources necessary to ensure that they can provide support to enrollees (especially populations who would be disproportionately impacted by these proposals, including the most financially vulnerable and those with limited English proficiency) as they become acquainted with additional steps needed to maintain coverage as a result of the proposed changes. Commenters also noted that any level of QHP disenrollment resulting from the proposed changes will result in decreased broker revenue and potential loss of broker participation in the market.

**Response:** Although there also may be an impact on Navigators, brokers, and other assisters, we believe these entities receive training and generally keep abreast of policy changes as part of their normal duties. As such, we believe these requirements will not amount to any additional burden above that already experienced by Navigators, brokers, and other assisters as a result of providing support to enrollees who are navigating these new billing requirements.
Comment: Many commenters also stated that enrollees would incur ancillary costs that would further drive up administrative costs and burden for enrollees, including postage costs, money order fees, or other banking fees for the second bill and cautioned that these costs will be felt most strongly by low income enrollees.

Many commenters stated that these proposals would transfer the costs and burdens of accessing non-Hyde abortion services to enrollees who must seek coverage for abortion elsewhere or pay out-of-pocket. Commenters estimated that non-Hyde abortions can cost between $400 and $1900. Commenters noted that low-income women who lack insurance coverage for abortion often struggle to pay for the procedure out-of-pocket, causing financial hardship that can drive families further into poverty. Commenters also expressed concern that when legal abortion is inaccessible, people who seek to end their pregnancy turn to unsafe and illegal methods, risking arrest, serious injury, or even death. Commenters also suggested that the changes would have a disproportionate effect on enrollee groups who already face barriers to care at higher rates such as low-income individuals, young people, people of color, individuals with LEP, lesbian, gay, bisexual, transgender and queer enrollees, the Latinx community, people with disabilities, rural residents, individuals without access to the internet, and American Indian/Alaskan Native populations.

Response: We acknowledge that as originally proposed, the combination of issuer burden and enrollee confusion could have potentially led to a reduction in the availability of coverage of non-Hyde abortion services in insurance (either by issuers choosing to drop this coverage to avoid the additional costs or by enrollees having their
coverage terminated for failure to pay the second bill), thereby increasing out-of-pocket costs for those seeking those services.

We understand that, even with the changes we are finalizing, the increased burden associated with issuers complying with the separate billing policy, could influence whether a QHP issuer continues offering coverage of non-Hyde abortion services in states that do not require it. However, we believe allowing the separate bill to be included in the same mailing (although not in the same email or other electronic communication), and allowing issuers to accept combined payments when policy holders fail to pay separately for the separate bill will mitigate some of the potential issuer and Exchange burden and consumer confusion associated with the proposed policy, thereby decreasing the likelihood that issuers will drop coverage of non-Hyde abortion services solely to avoid the burden associated with these changes or solely to avoid having to terminate enrollees coverage for non-payment of miniscule amounts.

We are also finalizing an enforcement posture that will further mitigate the risk of potential coverage loss. We intend to propose further rulemaking to change our regulations to mitigate this risk. Until we can effectuate such changes, we will exercise enforcement discretion as an interim step. Specifically, HHS will not take an enforcement action against a QHP issuer that adopts and implements a policy, beginning on or after the effective date for the separate billing policies, applied uniformly to all its QHP enrollees, under which an issuer does not place an enrollee into a grace period and does not terminate QHP coverage based solely on the policy holder’s failure to pay the separate payment for coverage of non-Hyde abortion services. We note that the QHP issuer would still be required to collect the premium for the non-Hyde abortion coverage.
We also will not take enforcement action against QHP issuers that, beginning upon the effective date of the final rule, modify the benefits of a plan either at the time of enrollment or during a plan year to effectively allow enrollees to opt out of coverage of non-Hyde abortion services by not paying the separate bill for such services, resulting in the enrollee having a modified plan that does not cover non-Hyde abortion services and that no longer obligates the enrollee to pay the required premium for such services. QHP issuers taking this approach should implement appropriate measures to distinguish between a policy holder’s inadvertent non-payment of the separate bill for non-Hyde abortion services and a policy holder’s intentional nonpayment of the separate bill. Although both of these approaches would be entirely optional for a QHP issuer, we believe that offering this enforcement discretion strikes an appropriate balance between honoring section 1303’s requirement for issuers to calculate the actuarial cost of non-Hyde abortion coverage and bill and collect premiums for such coverage in separate transactions, protecting enrollees against inadvertent losses of coverage, and ensuring all enrollees have access to coverage that meets their needs and that does not result in their supporting coverage for non-Hyde abortion services to which they object. We acknowledge that QHP issuers that do not utilize this available enforcement discretion may subsequently experience a higher number of enrollee terminations as a result of delinquent premium payments, which could influence whether a QHP issuer continues offering coverage of non-Hyde abortion services in states that do not require it.

Because enrollees will be instructed to make separate payments, those that follow the instructions may need to pay for additional postage, money order fees, credit card fees, or other banking fees for the second bill depending on how the QHP issuer
implements this policy. For example, policy holders who have funds automatically withdrawn from their bank accounts may need to arrange for a second withdrawal and may encounter additional fees. Additionally, because QHP issuers often incur fees for credit card transactions and these fees would double when a policy holder is paying in two separate transactions, QHP issuers may decide to transfer the cost of those credit card transaction fees onto policy holders choosing to pay via credit card rather than covering the cost of those transactions themselves. Policy holders that pay their premium bills via money order may need to pay an additional fee for the additional money order they submit for payment of the separate bill.

Comment: Many commenters stated that the proposals would cause considerable and unnecessary confusion and frustration for enrollees that may jeopardize their health insurance coverage by making it more difficult for policy holders to pay their premium bills, which could potentially result in their coverage being terminated for unintentional non-payment. Commenters also expressed concerns that despite consumer education and outreach, enrollees would likely not understand this change in billing.

Many commenters also stated that we underestimated the number of enrollees who would be impacted by these proposals. One commenter stated that there are 2 million enrollees alone in states where non-Hyde abortion coverage is required in all plans. Another commenter conducted an internal member survey, to which ten issuers responded, indicating that 2.4 million enrollees would be impacted across these ten issuers. This commenter noted that these ten issuers do not represent all health insurance issuers who would be required to comply with the proposals and that, thus, the number of affected enrollees would be greater than 2.4 million. Another commenter stated that the
rule would impact 3 million enrollees. As such, commenters stated that we underestimated how much it would cost enrollees annually to comply with the proposals. Commenters also objected that we excluded the cost of enrollees learning in our estimate.

**Response:** We based our initial estimates on 2018 QHP Certification data, and we acknowledge that the estimates may not have captured the exact number of enrollees that may be impacted by this final rule. In response to comments, we have reviewed our methodology and have updated our enrollee estimates accordingly. We also acknowledge that enrollees may initially be confused by receiving a separate bill for the portion of their premium attributable to coverage of non-Hyde abortion services in the same envelope as the bill for the rest of their premium. We believe that the provisions as finalized will minimize enrollee confusion surrounding the second bill for those receiving paper bills and will help to ensure that policy holders pay the entire premium due including the portion attributable to non-Hyde abortion services. There is still potential for confusion and loss of coverage for enrollees who receive electronic bills, due to failure to pay the second bill sent through a separate electronic communication, but the mechanisms by which electronic bills are paid may mitigate or lessen the potential for confusion over separate bills. We believe enrollee outreach and education will assist in further mitigating this risk.

Based on 2019 QHP certification data for the FFEs and SBE-FPs, we now estimate that there are approximately 442,400 enrollees in QHPs covering non-Hyde abortion services. In the 11 State Exchanges that operated their own technology platforms and had issuers that offered coverage of non-Hyde abortion services in 2019, we estimate that there are approximately 2,597,700 million enrollees enrolled in QHPs
offering coverage for non-Hyde abortion. As noted previously in section III “Collection of Information Requirements” of this final rule, we estimate that there are approximately 3.04 million enrollees impacted by these provisions. Assuming 1.5 enrollees per policy, issuers will be required to send a separate bill to approximately 2 million policy holders. We believe that finalizing the policies to allow for the separate bill to be sent in the same mailing with the bill for the rest of the policy holder’s premium will minimize enrollee confusion and burden.

We acknowledge that some policy holders will fail to pay in a separate transaction for both bills, and acknowledge that the burden may be moderately higher for those policy holders who follow instructions to pay in separate transactions. We also acknowledge that enrollees may experience burden in receiving a separate bill to which they are not yet accustomed in the same mailing as for the other portions of their premium or in a separate electronic communication. As such, using the May 2018 National Occupational Employment and Wage Estimates United States, Department of Labor’s Bureau of Labor Statistics (BLS) (https://www.bls.gov/oes/current/oes_stru.htm), listed national mean hourly wage for the 25th percentile, we estimate that for the 2020 plan year each policy holder will incur a burden of approximately 1 hour (at a cost of $12.37 per hour) to read and understand the separate bills received the first time and seek help from customer service if necessary, and approximately 5 minutes for each of the subsequent 5 months, resulting in a total estimated annual burden of 1.42 hours with an associated annual cost of approximately $18. For all policy holders we estimate that the

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28 The 25th percentile mean hourly wage most closely resembles the group of enrollees likely to be affected by this change as most enrollees enrolled in QHPs on the Exchange are between 100 percent and 400 percent of the federal poverty level.
initial 2020 burden will be approximately 2.9 million hours with and associated annual cost of $35.5 million. For subsequent years we estimate that enrollees will require approximately 5 minutes per month to read and understand their statements, resulting in an estimated annual burden of 1 hour with an associated annual cost of approximately $12. For all policy holders, we estimate that the annual enrollee burden will be approximately 2 million hours with an associated annual cost of approximately $25.1 million.

We also note that, although policy holders may experience burden related to reading and understanding the separate bills, there are non-quantifiable benefits to policy holders in QHPs covering non-Hyde abortion who hold conscience objections to such coverage or policy holders who seek a better understanding of what their health care dollars are purchasing.

HHS continues to believe that, although these changes will increase enrollee burden, this burden is reasonable and justified because it will achieve better alignment of the regulatory requirements for QHP issuer billing of premiums with the separate payment collection requirement in section 1303 of the PPACA.

**Table 10: Summary of Costs Related to Separate Billing and Payment**

<table>
<thead>
<tr>
<th>Requirements</th>
<th>2020</th>
<th>2021</th>
<th>2022</th>
<th>2023</th>
<th>2024</th>
</tr>
</thead>
<tbody>
<tr>
<td>Issuers</td>
<td>$482,616,844</td>
<td>$195,252,923</td>
<td>$195,228,601</td>
<td>$195,216,441</td>
<td>$195,216,441</td>
</tr>
<tr>
<td>States</td>
<td>$11,400,000</td>
<td>$4,800,000</td>
<td>$3,600,000</td>
<td>$2,400,000</td>
<td>$2,400,000</td>
</tr>
<tr>
<td>State Exchanges with payment portals</td>
<td>$15,385,201</td>
<td>$6,197,323</td>
<td>$6,197,323</td>
<td>$6,197,323</td>
<td>$6,197,323</td>
</tr>
<tr>
<td>Consumers</td>
<td>$35,517,268</td>
<td>$25,071,013</td>
<td>$25,071,013</td>
<td>$25,071,013</td>
<td>$25,071,013</td>
</tr>
<tr>
<td>Federal Government</td>
<td>$1,150,000</td>
<td>$800,000</td>
<td>$600,000</td>
<td>$400,000</td>
<td>$400,000</td>
</tr>
<tr>
<td>Total</td>
<td>$546,069,313</td>
<td>$232,121,259</td>
<td>$230,696,938</td>
<td>$229,284,777</td>
<td>$229,284,777</td>
</tr>
</tbody>
</table>
D. Regulatory Review Costs

If regulations impose administrative costs on private entities, such as the time needed to read and interpret this final rule, we estimate the cost associated with regulatory review. Due to the uncertainty involved with accurately quantifying the number of entities that will review the rule, we assume that the total number of unique reviewers on similar Exchange-related CMS rules will be the number of reviewers of this final rule. We acknowledge this assumption may understate or overstate the costs of reviewing this rule. It is possible that not all reviewers will review the rule in detail. For these reasons, we consider the number of past reviewers on similar CMS rules will be a fair estimate of the number of reviewers of this rule.

We recognize that different types of entities may be affected by only certain provisions of this final rule, and therefore, for the purposes of our estimate, we assume that each reviewer reads approximately 50 percent of the rule.

Using the wage information from the BLS for medical and health service managers (Code 11-9111), we estimate that the cost of reviewing this rule is $109.36 per hour, including overhead and fringe benefits.\(^{29}\) We estimate that it would take approximately 1 hour for each reviewer to review the relevant portions of this final rule. We received 75,439 comments, including 70,396 comments that were substantially similar to one of 13 different form letters, resulting in 5,043 unique comments on the proposed rule. We further assume that for the form letters received, only the staff at the organization that arranged for those letters will review the final rule. Therefore, we estimate that there will be 5,056 individuals that review the final rule resulting in an

\(^{29}\) [https://www.bls.gov/oes/current/oes_nat.htm](https://www.bls.gov/oes/current/oes_nat.htm)
estimated total cost of review of approximately $552,924 ($109.36 \times 5,056 \text{ reviewers}).

E. Regulatory Alternatives Considered

In developing the policies contained in this final rule, we considered numerous alternatives. Below we discuss the key regulatory alternatives that we considered.

For the eligibility determination during a benefit year, we considered not defining “periodically” for the frequency of Medicare, Medicaid/CHIP, or BHP, if applicable, PDM as twice a year in lieu of further outreach, education, and coordination with State Exchanges to identify and notice consumers who may also be enrolled in other qualifying coverage with APTC/CSRs. However, we believe it is critical that consumers receive timely notification of their potential dual enrollment in other qualifying coverage to ensure that consumers are accurately determined eligible for APTC and income-based CSRs, and to ensure that consumers are not enrolling in unnecessary or duplicative coverage. As previously discussed in the preamble of the proposed rule, such unnecessary or duplicative coverage, coupled with typically higher utilization generally results in higher premiums across the individual market leading to unnecessary expenditures of federal funds on PTC for taxpayers eligible for PTC in the individual market.

In finalizing the proposed changes to the general program integrity and oversight requirements in §155.1200, we considered not taking any action. However, because the existing requirements under §155.1200(b) did not accurately reflect the current structure of CMS’s oversight approach and reporting requirements for State Exchanges, not taking any action could have prevented HHS from being able to accurately describe our reporting requirements and strengthen our oversight processes for State Exchanges. In
particular, we needed to clarify that the eligibility and enrollment reports required under §155.1200(b)(2) were part of the annual compliance reports that State Exchanges were submitting to us, and did not require submission of a separate report. Thus, the amendments to §155.1200(b) do not reflect an expansion of State Exchange reporting obligations but instead were intended to capture the existing annual compliance reports (such as the SMART) that encompass eligibility and enrollment reporting, as well as compliance across other Exchange program requirements under 45 CFR part 155, that State Exchanges currently submit to HHS. Also, because the existing external programmatic audit requirements under §155.1200(d) did not specify how the audits needed to verify the accuracy of eligibility determinations made by State Exchanges, not taking any action would have prevented CMS from strengthening oversight processes by identifying a consistent procedure for these State Exchanges and their auditors to implement in order to ensure accurate eligibility determinations.

In finalizing the proposed changes to §155.1200(c) and (d), we also considered the alternative of narrowing the focus of the external programmatic audits to only 45 CFR part 155 subparts D and E, which cover Exchange eligibility and enrollment requirements. This approach would have focused the State Exchange’s auditing resources to the areas with highest program integrity impact. However, this approach would essentially exclude SBE-FPs from the external programmatic audit requirements altogether because SBE-FPs utilize the federal platform to carry out their eligibility and enrollment functions. Additionally, this approach would have limited our oversight in other program integrity areas that are important for all State Exchanges, such as consumer outreach and assistance. Because the external audit requirements under
§155.1200 is one of the only oversight tools we have for State Exchanges, we did not want to limit the scope of the Exchange functions that the external programmatic audits must cover. Instead, the approach finalized in this rulemaking allows us to specify the Exchange functions that are applicable to each State Exchange model through annual technical operational guidance. As State Exchanges continue to evolve and mature, this approach also provides HHS with the flexibility to focus the audits on emerging issues that raise program integrity concerns, while minimizing burden on State Exchanges to the extent possible.

In finalizing the requirement that issuers separately bill for the portion of the policy holder’s premium attributable to the cost of including coverage of non-Hyde abortion services in the QHP, and permit policy holders to pay for these amounts in a separate transaction if they so choose, as described at §156.280(e)(2), we considered maintaining the current methods of billing and collection without modification. We acknowledge that maintaining the current policy would promote stability for issuers and conserve administrative and operational resources by allowing QHP issuers to maintain their current process for billing for and collecting these separate payments. However, by requiring QHP issuers to separately bill for the portion of the policy holder’s premium attributable to coverage of non-Hyde abortion services, we believe we are strengthening alignment of issuer billing with the statutory requirements for collecting a separate payment for these services required under section 1303 of the PPACA.

We also considered finalizing the changes as originally proposed. However, we believe the changes we are finalizing will help to maximize the net benefit of achieving better statutory alignment while also mitigating burden where possible. For example, we
considered finalizing the proposed requirement that issuers would be required to send the separate bill in a separate mailing or electronic communication. This would have resulted in additional mailing costs of approximately $11 million in 2021 for all issuers. However, we believe allowing issuers to send the separate bill in the same mailing (although not in the same electronic communication) and allowing issuers to accept combined payments if a policy holder fails to pay the separate bill in a separate transaction will assist in mitigating the burden associated with this policy change by preventing unnecessary postage and mailing related costs and will mitigate issuer and Exchange burden and enrollee confusion generally associated with the proposed policy. We also believe the separate bill could assist in clarifying for enrollees that their plan covers non-Hyde abortion services and at what cost, increasing overall QHP transparency. Furthermore, we believe these changes will still better align issuer billing with section 1303 of the PPACA.

We also considered finalizing the rule without a requirement that issuers instruct policy holders to pay in a separate transaction. We understand that requiring issuers make this instruction and make reasonable efforts to collect the payment separately carries up-front and annual costs for issuers. However, we believe that instructing policy holders to pay the separate bill in a separate transaction is important to achieving better alignment of the regulatory requirements for QHP issuer billing of enrollee premiums with the separate payment requirement in section 1303 of the PPACA.

In addition, we considered requiring issuers to comply with the separate billing requirements within 3 months after the publication date of this final rule. We rejected this option because we estimated that one-time costs would have increased by 100
percent due to the shortened implementation period and estimated that total costs for issuers, State Exchanges, FFEs, and consumers would have been approximately $740 million in 2020. We opted to finalize a later effective date to avoid such a burden increase.

F. Regulatory Flexibility Act

The RFA requires agencies to prepare an initial RFA to describe the impact of the final rule on small entities, unless the head of the agency can certify that the rule will not have a significant economic impact on a substantial number of small entities. The RFA generally defines a “small entity” as (1) a proprietary firm meeting the size standards of the Small Business Administration (SBA), (2) a not-for-profit organization that is not dominant in its field, or (3) a small government jurisdiction with a population of less than 50,000. States and individuals are not included in the definition of “small entity.” HHS uses a change in revenue of more than 3 to 5 percent as its measure of significant economic impact on a substantial number of small entities.

In this final rule, we set standards for certain issuers related to the collection of a separate payment for the premium portion attributable to coverage for certain abortion services. Because we believe that insurance firms offering comprehensive health insurance policies generally exceed the size thresholds for “small entities” established by the SBA, we do not believe that an initial regulatory flexibility analysis is required for such firms.

For the purposes of the RFA, we expect health insurance issuers to be affected by this final rule. We believe that health insurance issuers would be classified under the North American Industry Classification System code 524114 (Direct Health and Medical
Insurance Carriers). According to SBA size standards, entities with average annual receipts of $38.5 million or less would be considered small entities for these North American Industry Classification System codes. Issuers could possibly be classified in 621491 (HMO Medical Centers) and, if this is the case, the SBA size standard would be $32.5 million or less.\footnote{https://www.sba.gov/document/support--table-size-standards} We believe that few, if any, insurance companies underwriting comprehensive health insurance policies (in contrast, for example, to travel insurance policies or dental discount policies) fall below these size thresholds.

Therefore, we are not preparing an analysis for the RFA because we have determined, and the Secretary certifies, that this final rule will not have a significant economic impact on a substantial number of small entities.

In addition, section 1102(b) of the Social Security Act requires us to prepare a regulatory impact analysis if a rule may have a significant impact on the operations of a substantial number of small rural hospitals. This analysis must conform to the provisions of section 604 of the RFA. For purposes of section 1102(b) of the Act, we define a small rural hospital as a hospital that is located outside of a metropolitan statistical area and has fewer than 100 beds. This final rule will not have a significant impact on small rural hospitals. Therefore, the Secretary has determined that this final rule will not have a significant impact on the operations of a substantial number of small rural hospitals.

G. Unfunded Mandates

Section 202 of the Unfunded Mandates Reform Act of 1995 (UMRA) requires that agencies assess anticipated costs and benefits and take certain other actions before issuing any rule that includes any federal mandate that may result in expenditures in any
1 year by a state, local, or Tribal government, in the aggregate, or by the private sector, of 
$100 million in 1995 dollars, updated annually for inflation. In 2019, that threshold is 
approximately $154 million. We anticipate that costs incurred by state, local, or tribal 
governments and the private sector will cross this threshold. States impacted by the 
separate billing and payment requirements at §156.280 may incur costs of approximately 
$26.8 million in 2020, 11 million in 2021, $9.8 million in 2022 and $8.6 million in 2023 
and each year after. In addition, states impacted by PDM requirements will incur costs of 
up to $6.9 million in 2020. Issuers impacted by the separate billing and payment 
requirements will incur costs of approximately $482.6 million in 2020 and approximately 
$195.3 million each year after.

H. Federalism

Executive Order 13132 establishes certain requirements that an agency must meet 
when it promulgates a proposed rule (and subsequent final rule) that imposes substantial 
direct requirement costs on state and local governments, preempts state law, or otherwise 
has Federalism implications. This final rule does not impose substantial direct costs on 
state and local governments or preempt state law. However, we believe the rule has 
Federalism implications.

In HHS’s view, this regulation has Federalism implications due to our 
requirements that Exchanges conduct Medicare, Medicaid/CHIP, and, if applicable, BHP 
PDM at least twice a year, beginning with the 2021 calendar year. As discussed earlier in 
this final rule, we received three comments that were opposed to the requirement to 
conduct Medicare, Medicaid/CHIP and, if applicable, BHP PDM at least twice yearly, 
cautioning us that defining the exact precise frequency and nature of PDM encroached
upon the sovereignty of the State Exchanges. However, HHS believes that the Federalism implications are substantially mitigated because the requirement sets only a minimum frequency with which Exchanges must conduct Medicare, Medicaid/CHIP, and, BHP, if applicable, PDM, which is already required to be conducted periodically; State Exchanges continue to have the flexibility to conduct PDM with greater frequency and the best way they see fit to implement the requirements set forth in §155.330(d). Additionally, as discussed earlier in this final rule, ensuring consumers are enrolled in the appropriate coverage remains a top priority for HHS and ensuring that APTC is paid appropriately is a requirement set forth in §155.330(d)(1)(ii) to mitigate the risk of federal dollars incorrectly leaving the federal Treasury in the form of APTC during the year. HHS believes that PDM plays a vital role in ensuring the health of all Exchanges, ensuring all consumers are enrolled in the appropriate coverage and in the case of Medicare enrollment, signing up at the appropriate time to avoid late enrollment penalties, and finally reduces the risk that consumers have to pay back all or some of APTC paid on their behalf during months of overlapping coverage when they file their federal income taxes.

Additionally, the changes to State Exchange oversight and reporting requirements in §155.1200 have Federalism implications since those rules require State Exchanges to submit certain reports to HHS and require them to enter into contracts with an external independent audit entity to perform audits, and incur the associated costs. However, HHS believes that the Federalism implications are substantially mitigated because the changes do not impose new requirements on State Exchanges, but rather add specificity and flexibility with respect to the existing requirements. Therefore, HHS believes it has
balanced states’ interests in operating State Exchanges with the need to ensure proper federal oversight. By doing so, it is HHS’s view that we have complied with the requirements of Executive Order 13132.

As discussed earlier in this final rule, commenters stated that the separate billing and payment proposals at §156.280 raise Federalism concerns under the Tenth Amendment because the proposals are designed to penalize states that have laws requiring QHPs to provide coverage of non-Hyde abortion services by requiring states—through their respective Exchanges and DOIs—to adopt new oversight responsibilities, and make systemic changes to fit the alterations the proposals require. As explained previously, we disagree that this policy raises Federalism concerns. Setting aside the question of whether state laws requiring coverage of non-Hyde abortion services on the Exchange are consistent with statutory conditions on federal funding from the Department to the States, we acknowledge that some states have such laws. However, the changes we are finalizing do not preempt state law regarding coverage of non-Hyde abortion services or otherwise attempt to coerce states into changing these laws. HHS is simply refining the method with which issuers use to comply with the separate payment requirement. We refer readers to section II.B of this final rule regarding the discussion of §156.280 for further information.

I. Reducing Regulation and Controlling Regulatory Costs

Executive Order 13771, entitled “Reducing Regulation and Controlling Regulatory Costs,” was issued on January 30, 2017 and requires that the costs associated with significant new regulations “shall, to the extent permitted by law, be offset by the elimination of existing costs associated with at least two prior regulations.” This final
rule is expected to be an Executive Order 13771 regulatory action. We estimate that this rule generates $182.98 million in annualized costs, discounted at 7 percent relative to year 2016, over a perpetual time horizon. Details on the estimated costs of this rule can be found in the preceding analyses.\textsuperscript{31}

**J. Congressional Review Act**

This final rule is subject to the Congressional Review Act provisions of the Small Business Regulatory Enforcement Fairness Act of 1996 (5 U.S.C. 801, \textit{et seq.}), which specifies that before a rule can take effect, the federal agency promulgating the rule shall submit to each House of the Congress and to the Comptroller General a report containing a copy of the rule along with other specified information, and has been transmitted to the Congress and the Comptroller General for review.

In accordance with the provisions of Executive Order 12866, this regulation was reviewed by the Office of Management and Budget.

\textsuperscript{31} We estimate costs of approximately $553.6 million in 2020, approximately $232.1 million in 2021, approximately $230.7 million in 2022, and annual costs of approximately $229.3 million thereafter. Thus the annualized value of costs, as of 2016 and calculated over a perpetual time horizon with a 7 percent discount rate, is $182.98 million.
List of Subjects

45 CFR Part 155

Administrative practice and procedure, Advertising, Brokers, Conflict of interests, Consumer protection, Grants administration, Grant programs-health, Health care, Health insurance, Health maintenance organizations (HMO), Health records, Hospitals, Indians, Individuals with disabilities, Intergovernmental relations, Loan programs-health, Medicaid, Organization and functions (Government agencies), Public assistance programs, Reporting and recordkeeping requirements, Technical assistance, Women and youth.

45 CFR Part 156

Administrative practice and procedure, Advertising, Advisory committees, Brokers, Conflict of interests, Consumer protection, Grant programs-health, Grants administration, Health care, Health insurance, Health maintenance organization (HMO), Health records, Hospitals, Indians, Individuals with disabilities, Loan programs-health, Medicaid, Organization and functions (Government agencies), Public assistance programs, Reporting and recordkeeping requirements, State and local governments, Sunshine Act, Technical assistance, Women, Youth.
For the reasons set forth in the preamble, the Department of Health and Human Services amends 45 CFR parts 155 and 156 as set forth below:

PART 155—EXCHANGE ESTABLISHMENT STANDARDS AND OTHER RELATED STANDARDS UNDER THE AFFORDABLE CARE ACT

1. The authority citation for part 155 continues to read as follows:


2. Section 155.200 is amended by revising paragraph (c) to read as follows:

§155.200 Functions of an Exchange.

(c) Oversight and financial integrity. The Exchange must perform required functions and cooperate with activities related to oversight and financial integrity requirements in accordance with section 1313 of the Affordable Care Act and as required under this part, including overseeing its Exchange programs and non-Exchange entities as defined in §155.260(b)(1).

3. Section 155.330 is amended by revising paragraph (d)(1) introductory text and adding paragraph (d)(3) to read as follows:

§155.330 Eligibility redetermination during a benefit year.

(d) (1) General requirement. Subject to paragraph (d)(3) of this section, the Exchange must periodically examine available data sources described in §§155.315(b)(1)
and 155.320(b) to identify the following changes:

* * * * *

(3) *Definition of periodically.* Beginning with the 2021 calendar year, the Exchange must perform the periodic examination of data sources described in paragraph (d)(1)(ii) of this section at least twice in a calendar year. State Exchanges that have implemented a fully integrated eligibility system with their respective State Medicaid programs, that have a single eligibility rules engine that uses MAGI to determine eligibility for advance payments of the premium tax credit, cost-sharing reductions, Medicaid, CHIP, and the BHP, if a BHP is operating in the service area of the Exchange, will be deemed in compliance with the Medicaid/CHIP PDM requirements and, if applicable, BHP PDM requirements, in paragraphs (d)(1)(ii) and (d)(3) of this section.

* * * * *

4. Section 155.1200 is amended by --

a. Revising paragraphs (b) introductory text, (b)(1) and (2), and (c) introductory text;

b. Revising paragraphs (d)(2) and (3);

c. Redesignating (d)(4) as paragraph (d)(5);

d. Adding a new paragraph (d)(4); and

e. Revising newly redesignated paragraph (d)(5).

The revisions and addition read as follows:

§155.1200  General program integrity and oversight requirements.

* * * * *

(b) *Reporting.* The State Exchange must, at least annually, provide to HHS, in a
manner specified by HHS and by applicable deadlines specified by HHS, the following
data and information:

(1) A financial statement presented in accordance with GAAP,

(2) Information showing compliance with Exchange requirements under this part 155 through submission of annual reports,

(c) External audits. The State Exchange must engage an independent qualified auditing entity which follows generally accepted government auditing standards (GAGAS) to perform an annual independent external financial and programmatic audit and must make such information available to HHS for review. The State Exchange must:

(d) Compliance with subparts D and E of this part 155, or other requirements under this part 155 as specified by HHS;

(3) Processes and procedures designed to prevent improper eligibility determinations and enrollment transactions, as applicable;

(4) Compliance with eligibility and enrollment standards through sampling, testing, or other equivalent auditing procedures that demonstrate the accuracy of eligibility determinations and enrollment transactions; and

(5) Identification of errors that have resulted in incorrect eligibility determinations, as applicable.
PART 156 – HEALTH INSURANCE ISSUER STANDARDS UNDER THE
AFFORDABLE CARE ACT, INCLUDING STANDARDS RELATED TO
EXCHANGES

5. The authority citation for part 156 is revised to read as follows:

Authority: 42 U.S.C. 18021-18024, 18031-18032, 18041-18042, 18044, 18054,

6. Section 156.280 is amended by --

a. Revising the section heading;

b. Redesignating paragraph (e)(2)(ii) as paragraph (e)(2)(iii);

c. Adding a new paragraph (e)(2)(ii); and

d. Revising newly redesignated paragraph (e)(2)(iii).

The addition and revision read as follows:

§156.280 Separate billing and segregation of funds for abortion services.

*   *   *   *   *
(e) *   *   *
(2) *   *   *

(ii) Beginning on or before the first billing cycle following June 27, 2020, to
satisfy the obligation in paragraph (e)(2)(i) of this section—

(A) Send to each policy holder of a QHP monthly bills for each of the amounts
specified in paragraphs (e)(2)(i)(A) and (B) of this section, either by sending separate
paper bills which may be in the same envelope or mailing, or by sending separate bills
electronically, which must be in separate emails or electronic communications; and
(B) Instruct the policy holder to pay each of the amounts specified in paragraphs (e)(2)(i)(A) and (B) of this section through separate transactions. Notwithstanding this instruction, if the policy holder fails to pay each of these amounts in a separate transaction as instructed by the issuer, the issuer may not refuse the payment and initiate a grace period or terminate the policy holder’s QHP coverage on this basis.

(iii) Deposit all such separate payments into separate allocation accounts as provided in paragraph (e)(3) of this section. In the case of an enrollee whose premium for coverage under the QHP is paid through employee payroll deposit, the separate payments required under paragraph (e)(2)(i) of this section shall each be paid by a separate deposit.

*   *   *   *   *

*   *   *   *   *

Seema Verma,
Administrator,
Centers for Medicare & Medicaid Services.

Dated: December 18, 2019.

Alex M. Azar II,
Secretary,
Department of Health and Human Services.

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