21st Century Cures Act: Interoperability, Information Blocking, and the ONC Health IT Certification Program Proposed Rule

Overview

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• ONC cannot address any comment suggestion or statement made by anyone attending the presentation or consider any such comment or suggestion in the rule writing process.

• Please submit comments through the formal process outlined in the Federal Register.
Agenda

• Purpose of the Rule
• Updates to the 2015 Edition Certification Criteria
• Conditions and Maintenance of Certification
• Enforcement of the Conditions and Maintenance of Certification Requirements

• Information Blocking
• Health IT for Pediatric Care and Practice Settings
• Additional Requests for Information
• Timeline
### Implementation of the 21st Century Cures Act

<table>
<thead>
<tr>
<th>KEY PROVISIONS IN TITLE IV OF THE CURES ACT</th>
<th>ONC’S WORK IN SUPPORT OF THE CURES ACT</th>
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</table>
| **Sec. 4001 Pediatrics**                   | • ONC engaged with stakeholders in the public and private sector.  
• ONC developed ten recommendations for the voluntary certification of health IT for pediatric care in response to the requirement set forth by Congress in Section 4001 of the Cures Act.  
• ONC proposes to adopt new and revised certification criteria to support the voluntary certification of health IT for use by pediatric health providers to support the health care of children.  
• ONC is also focused on non-regulatory initiatives that are nimble and responsive to stakeholders, including development of informational resources to support setting-specific implementation that aligns with the ONC Health IT Certification Program. |
| **Sec. 4002 Conditions of Certification**  | • ONC proposes an approach whereby the Conditions and Maintenance of Certification express initial and ongoing requirements for health IT developers and their certified Health IT Modules.  
• The Conditions of Certification with accompanying Maintenance of Certification requirements, consistent with the Cures Act, would focus on: (a) information blocking; (b) assurances; (c) communications; (d) application programming interfaces (APIs); (e) real world testing of certified health IT; and (f) attestations.  
• ONC proposes an enforcement approach to encourage consistent compliance with the requirements. The proposed rule outlines a corrective action process for ONC to review and take action for potential or known instances where a Condition or Maintenance of Certification requirement is not being met by a health IT developer under the ONC Health IT Certification Program. |
| **Sec. 4003 Interoperability Definition**  | • ONC proposes that interoperability means, with respect to health IT, such health IT that: (1) enables the secure exchange of electronic health information (EHI) with, and use of EHI from, other health IT without special effort on the part of the user; (2) allows for complete access, exchange, and use of all electronically accessible health information for authorized use under applicable state or federal law; and (3) does not constitute information blocking  
• The proposed definition is consistent with the Cures Act interoperability definition. |
## Implementation of the 21st Century Cures Act

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| Sec. 4004 Information Blocking            | - ONC proposes seven categories of practices that would be considered reasonable and necessary that, provided certain conditions are met, would not constitute information blocking. These categories were developed based on feedback from stakeholders and consultation with appropriate federal agencies.  
  - If the actions of a regulated actor (health care provider, health IT developer, or health information exchange or network) satisfy an exception, the actions would not be treated as information blocking and the actor would not be, as applicable, subject to civil penalties or other disincentives under the law. |
| Sec. 4005 Exchange with Registries        | - ONC’s proposed rule includes a Request for Information (RFI) on how a standards-based API might support improved information exchange between a health care provider and a registry in support of public health reporting, quality reporting, and care quality improvement.  
  - Public input on this RFI may be considered for future HHS rulemaking to support the bidirectional exchange of clinical data between health care providers and registries for a wide range of use cases. |
| Sec. 4006 Patient Access                  | - ONC proposes to promote policies that would ensure a patient’s EHI is accessible to that patient and the patient’s designees, in a manner that facilitates communication with the patient’s health care providers and other individuals, including researchers, consistent with such patient’s consent through the following proposals: United States Core Data for Interoperability (USCDI) standard; “EHI export” criterion; “standardized API for patient and population services” criterion, “data segmentation for privacy (DS4P)” criteria, “consent management for APIs” criterion; API Condition of Certification; and information blocking requirements, which include providing patients access to their EHI at no cost to them.  
  - Patient access to their EHI would be improved through the adoption of the following proposed 2015 Edition standard and certification criteria: USCDI standard; standardized APIs for patient and population services; and EHI export. |
# Implementation of Executive Orders

<table>
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<tr>
<th>EXECUTIVE ORDERS</th>
<th>ONC’S WORK IN SUPPORT OF EXECUTIVE ORDERS</th>
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| **Executive Order 13813**  
Promoting Healthcare Choice and Competition Across the United States | - ONC's proposed rule would contribute to fulfilling Executive Order 13813 by furthering patient (and health care provider) access to EHI and supporting competition in health care markets through new tools to access EHI and policies to address the hoarding of EHI.  
- ONC’s proposed rule calls on the health care industry to adopt standardized APIs, which would allow individuals to securely and easily access structured EHI using new and innovative applications for smartphones and other mobile devices.  
- The proposed rule would establish information blocking provisions, focusing on improving patient and health care provider access, exchange, and use of EHI. |
| **Executive Orders 13771 & 13777**  
Reducing Regulation and Controlling Regulatory Costs, and Enforcing the Regulatory Reform Agenda | - ONC reviewed and evaluated existing regulations to identify ways to reduce burden and implement deregulatory actions.  
- ONC proposes potential deregulatory actions that will reduce burden for health IT developers, providers, and other stakeholders. These six deregulatory actions are: (1) removal of a threshold requirement related to randomized surveillance; (2) removal of the 2014 Edition from the Code of Federal Regulations (CFR); (3) removal of the ONC-Approved Accrualor (ONC-AA) from the Certification Program; (4) removal of certain 2015 Edition certification criteria; (5) removal of certain Certification Program requirements; and (6) recognition of relevant Food and Drug Administration (FDA) certification processes with a request for information on the potential development of new processes for the ONC Health IT Certification Program. |
Purpose

Increase Innovation and Competition
by giving patients and their health care providers safe and secure access to health information and to new tools, allowing for more choice in care and treatment.

Reduce Burden and Advance Interoperability
through the use of United States Core Data for Interoperability (USCDI) standard, new API requirements, and EHI export capabilities for the purposes of switching health IT or to provide patients their electronic health information.

Promote Patient Access
through a provision requiring that patients can electronically access all of their electronic health information (structured and/or unstructured) at no cost.

https://www.healthit.gov/NPRM
Updates to the 2015 Edition Certification Criteria
This rule proposes to update the 2015 Edition by not only proposing criteria for removal, but by proposing to revise and add new certification criteria that would establish the capabilities and related standards and implementation specifications for the certification of health IT.

https://www.healthit.gov/sites/default/files/understanding-certified-health-it-2.pdf

*Note: this is a link to the current certification criteria but it will need to be updated after the Cures Act final rule is released.
Proposed Changes to the 2015 Edition Certification Criteria

## Removed Criteria

<table>
<thead>
<tr>
<th>2015 Base EHR Definition Criteria</th>
<th>Other Criteria</th>
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<tbody>
<tr>
<td>× Problem list (§ 170.315(a)(6))</td>
<td>× Drug formulary and preferred drug list checks (§170.315(a)(10))</td>
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<tr>
<td>× Medication list (§ 170.315(a)(7))</td>
<td>× Patient-specific education resource (§ 170.315(a)(13))</td>
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<tr>
<td>× Medication allergy list (§ 170.315(a)(8))</td>
<td>× Common Clinical Data Set summary record – create (§ 170.315(b)(4))</td>
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<tr>
<td>× Smoking status (§ 170.315(a)(11))</td>
<td>× Common Clinical Data Set summary record – receive (§ 170.315(b)(5))</td>
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<td></td>
<td>× Secure messaging (§ 170.315(e)(2))</td>
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## Updated Criteria

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<thead>
<tr>
<th>Remove</th>
<th>Update with</th>
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<tr>
<td>× Electronic prescribing (§ 170.315(b)(3))</td>
<td>✓ Electronic prescribing (§ 170.315(b)(11))</td>
</tr>
<tr>
<td>× Data export (§ 170.315(b)(6))</td>
<td>✓ Electronic health information (EHI) export (§ 170.315(b)(10))</td>
</tr>
<tr>
<td>× Data segmentation for privacy – send (§ 170.315(b)(7))</td>
<td>✓ Data segmentation for privacy – send (§ 170.315(b)(12))</td>
</tr>
<tr>
<td>× Data segmentation for privacy – receive (§ 170.315(b)(8))</td>
<td>✓ Data segmentation for privacy – receive (§ 170.315(b)(13))</td>
</tr>
<tr>
<td>× Application access – data category request (§ 170.315(g)(8))</td>
<td>✓ Standardized API for patient and population services (§ 170.315(g)(10))</td>
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## Revised Criteria

✓ Clinical Quality Measures (CQMs) – report criterion (§ 170.315(c)(3))

## New Criteria

+ Consent management for application programming interfaces § 170.315(g)(11)
+ Encrypt authentication credentials certification criterion § 170.315(d)(12)
+ Multi-factor authentication (MFA) criterion § 170.315(d)(13)
The United States Core Data for Interoperability Standard

We propose to remove the “Common Clinical Data Set” (CCDS) definition and its references from the 2015 Edition and replace it with the “United States Core Data for Interoperability” (USCDI) standard. This will increase the minimum baseline of data classes that must be commonly available for interoperable exchange.

USCDI reflects the same data classes referenced by the CCDS definition and includes the following new required data classes and data elements:

- Provenance
- Clinical Notes
- Pediatric Vital Signs
- Address & Phone Number

If adopted, health IT developers will need to update their certified health IT to support the USCDI for all certification criteria affected by this change.

USCDI Standard Annual Update Schedule
ONC intends to establish and follow a predictable, transparent, and collaborative process to expand the USCDI, including providing stakeholders with the opportunity to comment on the USCDI’s expansion.

https://www.healthit.gov/NPRM
Electronic Health Information (EHI) Export Criterion

- Our new proposal would require health IT developers to provide the capability to electronically export all EHI that they produce and electronically manage in a computable format.
- We propose to make this criterion part of the 2015 Edition Base EHR definition, and for providers and developers to implement this within 24 months of the final rule's effective date.

HOW WILL IT WORK?

The proposed EHI Export certification requirement requires that:

1. All EHI produced and electronically managed by a developer's health IT must be readily available to export for:
   A. a single patient upon request for their health data, and
   B. all patients when a provider seeks to change Health IT systems.

2. The export file must:
   A. be computable, and
   B. include documentation to allow for interpretation and use of EHI. The documentation must be made publicly available via a hyperlink.

Note: Health IT developers would have the flexibility to determine their products' export standards.

https://www.healthit.gov/NPRM
We propose to adopt a new API criterion in § 170.315(g)(10), which would replace the “application access – data category request” certification criterion (§ 170.315(g)(8)) and become part of the 2015 Edition Base EHR definition. This new certification criterion would require the use of Health Level 7 (HL7®) Fast Healthcare Interoperability Resources (FHIR®) standards and several implementation specifications.

https://www.hl7.org/fhir/overview.html

Supports two types of API-enabled services:

» Services for which a single patient’s data is the focus

» Services for which multiple patients’ data are the focus

https://www.healthit.gov/NPRM
Conditions and Maintenance of Certification
The 21st Century Cures Act (Section 4002) requires the Secretary of HHS to establish Conditions and Maintenance of Certification requirements for the ONC Health IT Certification Program. ONC proposes an approach whereby the Conditions and Maintenance of Certification express initial requirements and ongoing requirements for health IT developers and their certified Health IT Module(s). Any noncompliance with the proposed Conditions and Maintenance of Certification requirements would be subject to ONC direct review, corrective action, and enforcement procedures under the ONC Health IT Certification Program.

There are seven Conditions of Certification with accompanying Maintenance of Certification Requirements. They are:

1. Information Blocking
2. Assurances
3. Communications
4. Application Programming Interfaces (APIs)
5. Real World Testing
6. Attestations
7. (Future) Electronic Health Record (EHR) Reporting Criteria Submission

https://www.healthit.gov/NPRM
CONDITIONS OF CERTIFICATION

• As a Condition of Certification and to maintain such certification, a health IT developer must not take any action that constitutes information blocking as defined in section 4004 of the Cures Act.

• This provision is subject to the seven proposed exceptions to the information blocking definition, which define certain reasonable and necessary activities that would not constitute information blocking (e.g., exceptions for preventing harm, promoting the privacy of EHI, promoting the security of EHI, etc.).

MAINTENANCE OF CERTIFICATION

• No accompanying Maintenance of Certification requirements beyond ongoing compliance with the Condition.
**CONDITION OF CERTIFICATION**

A health IT developer must provide assurances to the Secretary (unless for reasonable and necessary activities identified by the Secretary) that it will not take any action that constitutes information blocking or any other action that may inhibit the appropriate exchange, access, and use of electronic health information (EHI).

<table>
<thead>
<tr>
<th>a.</th>
<th>Full Compliance and Unrestricted Implementation of Certification Criteria Capabilities</th>
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<tbody>
<tr>
<td></td>
<td>• A health IT developer must ensure that its certified health IT conforms to the full scope of the applicable certification criteria</td>
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<td></td>
<td>• Developers of certified health IT must provide assurance that they have made certified capabilities available in ways that enable them to be implemented and used in production environments for their intended purposes</td>
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<tr>
<th>b.</th>
<th>Certification to the “Electronic Health Information (EHI) Export” Criterion</th>
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<tr>
<td></td>
<td>• A health IT developer that produces and electronically manages EHI must certify health IT to the 2015 Edition “electronic health information export” certification criterion in § 170.315(b)(10)</td>
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<th>c.</th>
<th>Records and Information Retention</th>
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<td><strong>MAINTENANCE OF CERTIFICATION</strong></td>
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<tr>
<td></td>
<td>Such developer must provide all of its customers with certified health IT with the functionality included in § 170.315(b)(10) within 24 months of the final rule’s effective date or within 12 months of certification for a health IT developer that never previously certified health IT to the 2015 Edition, whichever is longer</td>
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<th>d.</th>
<th>Trusted Exchange Framework and the Common Agreement - Request for Information</th>
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<td>• ONC is requesting comment as to whether certain health IT developers should be required to participate in the Trusted Exchange Framework and adhere to the Common Agreement</td>
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<td></td>
<td>• This requirement would apply to health IT developers that certify to capabilities used for interoperability (i.e., §§ 170.315(b)(1), (c)(1) and (c)(2), (e)(1), (f), and (g)(9) through (11))</td>
</tr>
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</table>
CONDITIONS OF CERTIFICATION

• Requires that a health IT developer does not prohibit or restrict communication regarding the following subjects for the health IT:
  - Usability
  - Interoperability
  - Security
  - User experiences
  - Business practices
  - The manner in which a user of health IT has used such technology

• The Condition includes very limited exceptions to the prohibition (e.g., certain prohibitions and restrictions on communications by the health IT developer could be permissible when it would infringe the intellectual property rights existing in the developer's health IT)

MAINTENANCE OF CERTIFICATION

• A health IT developer must notify all customers within six months of the effective date of a subsequent final rule, that any communication or contract/agreement provision that violates the Communications Condition of Certification will not be enforced by the health IT developer

• Notice to customers would need to be provided annually up to and until the health IT developer amends the contract or agreement to remove or void any contractual provisions that violate this Condition of Certification
  - Developer must amend their contracts or agreements within a reasonable period of time, but not to exceed 2 years
## API TECHNOLOGY ROLES

<table>
<thead>
<tr>
<th>Role</th>
<th>Description</th>
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<tbody>
<tr>
<td><strong>API Technology Supplier</strong></td>
<td>Health IT developer that creates API technology presented for certification in the ONC Health IT Certification Program</td>
</tr>
<tr>
<td><strong>API Data Provider</strong></td>
<td>Health care organization that deploys the API technology</td>
</tr>
<tr>
<td><strong>API User</strong></td>
<td>Persons and entities that use or create software applications that interact with API technology</td>
</tr>
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</table>

ONC has designed API Conditions of Certification that will complement the technical capabilities described in our other proposals, while addressing the broader technology and business landscape in which these API capabilities will be deployed and used.

**Note:** The API Conditions of Certification only apply to API Technology Suppliers with health IT certified to any API-focused certification criteria.

### TRANSPARENCY

ONC has proposed that API Technology Suppliers make business and technical documentation necessary to interact with their APIs in production freely and publicly accessible.

### PERMITTED FEES

ONC has proposed to adopt specific conditions that would set boundaries for the fees API Technology Suppliers would be permitted to charge and to whom those permitted fees could be charged.

### PRO-COMPETITIVENESS

ONC has proposed that API Technology Suppliers would have to comply with certain requirements to promote an open and competitive marketplace.

[https://www.healthit.gov/NPRM](https://www.healthit.gov/NPRM)
CONDITIONS OF CERTIFICATION

- Requires health IT developers to publish APIs that allow health information from such technology to be accessed, exchanged, and used without special effort through the use of APIs or successor technology or standards, as provided for under applicable law
  - Through the APIs, a developer must also provide access to all data elements (i.e., the USCDI) of a patient’s EHR to the extent permissible under applicable privacy laws
- An API Technology Supplier must make business and technical documentation necessary to interact with their APIs in production freely and publicly accessible
- All fees related to API technology, not otherwise permitted by this section, are prohibited from being imposed by an API technology Supplier
- API Technology Suppliers must grant API Data Providers (i.e., health care providers who purchase or license API technology) the sole authority and autonomy to permit API Users to interact with the API technology

MAINTENANCE OF CERTIFICATION

- An API Technology Supplier must register and enable all applications for production use within one business day of completing its verification of an applications developer’s authenticity
- An API Technology Supplier must Support the publication of "Service Base URLs" (i.e., FHIR® server endpoints) for all of its customers, regardless of those that are centrally managed by the API Technology Supplier or locally deployed by an API Data Provider, and make such information publicly available at no charge
- An API Technology Supplier with API technology certified to § 170.315(g)(8) must provide all API Data Providers with a (g)(10)-certified API within 24 months of this final rule’s effective date
CONDITIONS OF CERTIFICATION

• Requires that health IT developers have successfully tested the real-world use of the technology for interoperability in the type of setting in which such technology would be marketed.

• This Condition of Certification applies to health IT developers with Health IT Module(s) certified to certain certification criteria focused on interoperability and data exchange (i.e., §§ 170.315(b), (c)(1) through (c)(3), (e)(1), (f), (g)(7) through (g)(11), and (h)).

MAINTENANCE OF CERTIFICATION

• Health IT developers must submit publicly available prospective annual real world testing plans and retrospective annual real world testing results for certified health IT products focused on interoperability.

• Standards Version Advancement Process: permits health IT developers to voluntarily use newer versions of adopted standards. Please also see the Standards Version Advancement Process fact sheet for more details.
Standards Version Advancement Process

• The Standards Version Advancement Process would allow developers to choose among the versions of standards and implementation specifications listed in regulation or National Coordinator (NC)-approved newer version updates for any or all standards applicable to criteria subject to real world testing requirements.

• This flexibility to choose among NC-approved versions of standards and implementation specifications would be available both when developers seek initial certification or to maintain certification of a Health IT Module.

HOW WILL IT WORK?

To take advantage of the flexibility to update to NC-approved versions, a developer will need to:

• Notify its ONC- Authorized Certification Body (ONC-ACB) and its customers of its intent to move to newer NC-approved standards version updates as well as whether, and for how long, they might plan to continue supporting prior versions of the standards.

• Ensure that the standards version updates are effectively implemented.

• Address standards version updates in the required annual real world testing plans and results.

• Demonstrate conformance of its health IT to all applicable criteria including, but not limited to, any NC-approved standards versions it has chosen to incorporate into its certified Health IT Module.

https://www.healthit.gov/NPRM
Attestations - § 170.406

CONDITION OF CERTIFICATION

• A health IT developer must provide an attestation, as applicable, to compliance with the Conditions and Maintenance of Certification, except for the "EHR reporting criteria submission" Condition of Certification

MAINTENANCE OF CERTIFICATION

• Health IT developers must submit their attestations every six months
Enforcement of the Conditions and Maintenance of Certification Requirements
Enforcement Approach

ONC DIRECT REVIEW OF THE CONDITIONS AND MAINTENANCE OF CERTIFICATION

• ONC would be the sole party responsible for enforcing compliance. ONC may, however, coordinate its review with the HHS Office of Inspector General (OIG) or defer to the OIG to lead review of a claim of information blocking.

• ONC will utilize the processes established for ONC direct review of certified health IT in the Enhanced Oversight and Accountability (EOA) final rule for enforcement. Using the established processes delivers multiple benefits:

  • EOA processes address non-conformities with Program requirements. Any noncompliance with the Conditions and Maintenance of Certification would constitute a Program non-conformity
  • Health IT developers are familiar with the ONC direct review provisions
  • The process established for working with health IT developers is thorough and transparent, with clear corrective action procedures and remedies for Program non-conformities
  • Direct review provides equitable opportunities for health IT developers to respond to ONC actions and appeal determinations
Enforcement Approach

**STEP 1**  Initiating Review and Health IT Developer Notice

**STEP 2**  Records Access

**STEP 3**  Corrective Action Plan

**STEP 4**  Certification Ban and/or Termination

**STEP 5**  Appeal

**STEP 6**  Public Listing of Certification Ban and/or Terminations
Information Blocking
Information Blocking

OVERVIEW

Section 4004 of the Cures Act authorizes the Secretary to identify reasonable and necessary activities that do not constitute information blocking.

In consultation with stakeholders, we have identified seven categories of practices that would be reasonable and necessary, provided certain conditions are met.

The seven categories of reasonable and necessary practices, and their corresponding conditions, are defined through the exceptions proposed at 45 CFR 171.201–207.

If the actions of a regulated actor (health care provider, health IT developer, or health information exchange or network) satisfy one or more exception, the actions would not be treated as information blocking and the actor would not be subject to civil penalties and other disincentives under the law.

"Actors" regulated by the information blocking provision:
- Health Care Providers
- Health IT Developers of Certified Health IT
- Health Information Exchanges
- Health Information Networks

https://www.healthit.gov/NPRM
What is information blocking?

A practice by a health care provider, health IT developer, health information exchange, or health information network that, except as required by law or specified by the Secretary as a reasonable and necessary activity, is likely to interfere with, prevent, or materially discourage access, exchange, or use of electronic health information.
We propose to define EHI to mean electronic protected health information (as defined in HIPAA), and any other information that:

- is transmitted by or maintained in electronic media (as defined in 45 CFR 160.103);
- identifies the individual, or with respect to which there is a reasonable basis to believe the information can be used to identify the individual;
- relates to the past, present, or future health or condition of an individual; the provision of health care to an individual; or the past, present, or future payment for the provision of health care to an individual.

Not limited to information that is created or received by a health care provider.

Does not include health information that is de-identified consistent with the requirements of 45 CFR 164.514(b).
Price Information – Request for Information

• The fragmented and complex nature of pricing within the health care system has decreased the efficiency of the health care system and has had negative impacts on patients, health care providers, health systems, plans, plan sponsors and other key health care stakeholders.

• Consistent with its statutory authority, the Department is considering subsequent rulemaking to expand access to price information for the public, prospective patients, plan sponsors, and health care providers.

• ONC has a unique role in setting the stage for such future actions by establishing the framework to prevent the blocking of price information.
  » We seek comment on the parameters and implications of including price information within the scope of EHI for purposes of information blocking.
  » The overall Department seeks comment on the technical, operational, legal, cultural, environmental and other challenges to creating price transparency within health care.
Information Blocking Exceptions

• **§ 171.201 Exception | Preventing Harm**
  » An actor may engage in practices that are reasonable and necessary to prevent physical harm to a patient or another person.
  » The actor must have a reasonable belief that the practice will directly and substantially reduce the likelihood of physical harm to a patient or another person.
  » The practice must implement an organizational policy that meets certain requirements or must be based on an individualized assessment of the risk in each case.

This proposed exception acknowledges that the public interest in protecting patients and other persons against unreasonable risks of harm can justify practices that are likely to interfere with access, exchange, or use of electronic health information (EHI).

• **§ 171.202 Exception | Promoting the Privacy of Electronic Health Information**
  » An actor may engage in practices that protect the privacy of EHI.
  » An actor must satisfy at least one of four discrete sub-exceptions that address scenarios that recognize existing privacy laws and privacy-protective practices:
    (1) practices that satisfy preconditions prescribed by privacy laws; (2) certain practices not regulated by HIPAA but which implement documented and transparent privacy policies; (3) denial of access practices that are specifically permitted under HIPAA; (4) practices that give effect to an individual's privacy preferences.
  » The information blocking provision will not require that actors provide access, exchange, or use of EHI in a manner that is not permitted under the HIPAA Privacy Rule.
  » General conditions apply to ensure that practices are tailored to the specific privacy risk or interest being addressed and implemented in a consistent and non-discriminatory manner.

This proposed exception would advance the goal of preventing information blocking for improper or self-interested purposes while maintaining and upholding the privacy rights that patients now have.
Information Blocking Exceptions

• **§ 171.203 Exception | Promoting the Security of Electronic Health Information**
  » An actor may implement measures to promote the security of EHI.
  » The practice must be directly related to safeguarding the confidentiality, integrity, and availability of EHI.
  » The practice must be tailored to specific security risks and must be implemented in a consistent and non-discriminatory manner.
  » The practice must implement an organizational security policy that meets certain requirements or must be based on an individualized determination regarding the risk and response in each case.

• **§ 171.204 Exception | Recovering Costs Reasonably Incurred**
  » An actor may recover costs that it reasonably incurs, in providing access, exchange, or use of EHI.
  » Fees must be:
    (1) charged on the basis of objective and verifiable criteria uniformly applied to all similarly situated persons and requests; (2) related to the costs of providing access, exchange, or use; and (3) reasonably allocated among all customers that use the product/service.
  » Fees must not be based on anti-competitive or other impermissible criteria.
  » Certain costs would be specifically excluded from coverage under this exception, such as costs that are speculative or subjective or costs associated with electronic access by an individual to their EHI.

This proposed exception acknowledges that actors should be able to recover costs that they reasonably incur to develop technologies and provide services that enhance interoperability and promote innovation, competition, and consumer welfare.
Information Blocking Exceptions

• **§ 171.205 Exception | Responding to Requests that are Infeasible**
  
  » An actor may decline to provide access, exchange, or use of EHI in a manner that is infeasible.
  
  » Complying with the request must impose a substantial burden on the actor that is unreasonable under the circumstances (taking into account the cost to the actor, actor's resources, etc.).
  
  » The actor must timely respond to infeasible requests and work with requestors to provide a reasonable alternative means of accessing the EHI.

  This proposed exception acknowledges that there may be legitimate practical challenges beyond an actor’s control that may limit its ability to comply with requests for access, exchange, or use of EHI.

• **§ 171.206 Exception | Licensing of Interoperability Elements on Reasonable and Non-discriminatory Terms**

  » An actor that controls technologies or other interoperability elements that are necessary to enable access to EHI will not be information blocking so long as it licenses such elements on reasonable and non-discriminatory terms.
  
  » The license can impose a reasonable royalty but must include appropriate rights so that the licensee can develop, market, and/or enable the use of interoperable products and services.
  
  » The terms of the license must be based on objective and verifiable criteria that are uniformly applied and must not be based on impermissible criteria, such as whether the requestor is a potential competitor.

  This proposed exception would allow actors to protect the value of their innovations and earn returns on the investments they have made to develop, maintain, and update those innovations.
Information Blocking Exceptions

• § 171.207 Exception | Maintaining and Improving Health IT Performance

» An actor may make health IT under its control temporarily unavailable in order to perform maintenance or improvements to the health IT.

» An actor must ensure that the health IT is unavailable for no longer than necessary to achieve the maintenance or improvements.

» The practice must be implemented in a consistent and non-discriminatory manner.

» In circumstances when health IT is supplied to an individual or entity, the individual or entity (e.g., customer) must agree to the unavailability of health IT.

The proposed exception recognizes that it may be reasonable and necessary for actors to make health IT, and in turn EHI, temporarily unavailable for the benefit of the overall performance of health IT.
• **Complaint Process**
  » Section 3022(d)(3)(A) of the PHSA directs the National Coordinator to implement a standardized process for the public to submit reports on claims of health information blocking.
  » We intend to implement and evolve this complaint process by building on existing mechanisms, including the complaint process currently available at [https://www.healthit.gov/healthit-feedback](https://www.healthit.gov/healthit-feedback).
  » We request comment on this approach and any alternative approaches that would best effectuate this aspect of the Cures Act.

• **Additional Exceptions**
  » We are considering whether we should propose, in a future rulemaking, a narrow exception to the information blocking provision for practices that are necessary to comply with the requirements of the Common Agreement.
  » We welcome comment on any potential new exceptions we should consider for future rulemaking.

• **Disincentives for Health Care Providers**
  » We request information on disincentives or if modifying disincentives already available under existing HHS programs and regulations would provide for more effective deterrents.
Health IT for Pediatric Care and Practice Settings
In response to the requirements set forth in section 4001 of the Cures Act, ONC has:

1. Developed ten recommendations for the voluntary certification of health IT for pediatric care that does NOT include a separate certification program for pediatric care and practice settings.


3. Focused on non-regulatory initiatives that are nimble and responsive to stakeholders, including development of informational resources to support setting-specific implementation that aligns with the ONC Health IT Certification Program.

https://www.healthit.gov/NPRM
ONC DEVELOPED RECOMMENDATIONS BASED ON STAKEHOLDER-IDENTIFIED CLINICAL PRIORITIES AND THE CHILDREN'S EHR FORMAT

Pediatric stakeholders identified clinical priorities and evaluated them with ONC.

ONC RECOMMENDATIONS FOR PEDIATRIC HEALTH IT VOLUNTARY CERTIFICATION CRITERIA
1. Use biometric-specific norms for growth curves and support growth charts for children
2. Compute weight-based drug dosage
3. Ability to document all guardians and caregivers
4. Segmented access to information
5. Synchronize immunization histories with registries
6. Age- and weight-specific single dose range checking
7. Transferrable access authority
8. Associate mother's demographics with newborn
9. Track incomplete preventative care opportunities
10. Flag special health care needs

ONC CERTIFICATION CRITERIA TO SUPPORT PEDIATRIC CARE AND PRACTICE SETTINGS
CURRENT 2015 EDITION CRITERIA:
- Transitions of Care
- Care Plan
- View, Download, Transmit
- Application Programming Interface (API)
- Data Segmentation for Privacy
- Problem List
- Electronic Prescribing
- Common Clinical Data Set (CCDS)
- Social, Psychological, and Behavioral Data
- Clinical Quality Measure (CQM)
- Clinical Decision Support
- Immunizations
- Demographic data capture
- Family health history
- Patient health data capture
- Privacy and security

PROPOSED NEW 2015 EDITION CRITERIA:
- United States Core Data Set for Interoperability (USCDI)
- Electronic prescribing
- FHIR-based API
- Data segmentation for privacy

https://www.healthit.gov/NPRM
Additional Requests for Information (RFIs)
ONC recognizes that health IT offers promising strategies to help medical specialties and sites of service as they combat opioid use disorder (OUD).

- We request public comment on how our existing Program requirements and the proposals in this rulemaking may support use cases related to opioid use disorder (OUD) prevention and treatment and if there are additional areas that ONC should consider for effective implementation of health IT to help address OUD prevention and treatment.
Patient matching is a critical component to interoperability and the nation’s health information technology infrastructure. Accurate patient matching helps health care providers access and share the right information on the right patient when and where it is needed.

- Section 4007 of the 21st Century Cures Act directed the Government Accountability Office (GAO) to conduct a study on patient matching.
  

- We seek comment on additional opportunities that may exist in the patient matching space and ways that ONC can lead and contribute to coordination efforts with respect to patient matching. ONC and CMS collaborated to jointly issue complementary requests for information regarding patient matching.
Section 4005 (a) and (b) of the Cures Act focuses on interoperability and bidirectional exchange between EHRs and registries, including clinician-led clinical data registries. ONC is approaching these provisions from several angles to address the technical capability of EHRs to exchange data with registries in accordance with applicable recognized standards.

- We include an RFI in the proposed rule on how a standards-based API might support improved information exchange between a health care provider and a registry to support public health reporting, quality reporting, and care quality improvement. Public input on this RFI may be considered for future HHS rulemaking to support the bidirectional exchange of clinical data between health care providers and registries for a wide range of use cases.
21st Century Cures Act NPRM –
Regulatory Implementation Milestones

21ST CENTURY CURES ACT NPRM – REGULATORY IMPLEMENTATION MILESTONES

*Last day for health IT developers to implement for customers (health care providers)
**Last day to remove “gag clauses” from health IT contracts
Questions

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