

Cabinet for Health and Family Services

Division of Health Information

Policies – Introduction

The following policies apply to the access, use and disclosure of protected health information by Participants of the Kentucky Health Information Exchange (KHIE). These are the policies adopted by the Office of Health and Data Analytics and Division of Health Information (DHI) according to the language found in the Participation Agreement signed by participants in the KHIE.

According to the Participation Agreement, the DHI may change or amend the Policies in a manner consistent with the Participation Agreement. The Participant shall be given notice of any proposed and final changes and Participants shall be given an opportunity to comment on such proposed and final changes. Any change to a policy is effective thirty days after the change unless an earlier effective date is required to address a legal requirement, a concern relating to privacy or security of Data, or an emergency. The DHI also postpone the effective date of a policy, if additional implementation time is needed.

The DHI believes the nine principles defined in “The Architecture for Privacy in a Networked Health Information Environment”¹ are essential for protecting privacy and developing a comprehensive privacy-protective architecture in a networked environment. The principles are as follows:

1. Openness and Transparency


There should be a broad and universal practice of transparency in the way data is handled. Individuals should be able to establish what information exists about them in the data and in databases.

2. Purpose Specification and Minimization

Data should never be collected without the subject of the data knowing why it is being collected and how it will be used. Data should only be used for the purpose for which it was collected.

¹ ©2006, Markle Foundation

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3. Collection Limitation

The collection of data should be obtained by lawful and fair means and with the knowledge and consent of persons.

4. Use Limitation

A minimization requirement would strictly limit whether data collected for one purpose could be reused in another context. Such use should not be permissible without the explicit consent of individuals.

5. Individual Participation and Control

Consistent with the scope of individual rights in HIPAA, an individual has a vital stake in, and needs to be, a participant in determining how his or her information is used.

Individuals should be seen as key participants in the process of information collection and dissemination, and not as mere subjects or passive spectators.

6. Data Integrity and Quality

Mechanisms need to be developed to address data corruption and for establishing accountability among those who maintain records. Individuals should have clear avenues to view all information that has been collected about them and to ensure that the information is accurate, complete and timely

7. Security Safeguards and Controls

Reasonable security safeguards must be built against loss, unauthorized access, destruction, use, modification, or disclosure of personal information. In addition, all data collectors and disseminators should be mandated to immediately disclose any security breach through a direct communication to those consumers affected.

8. Accountability and Oversight

There must be mechanisms to ensure that the responsibility for privacy and privacy violations is identifiable and that remedial action can be taken.

9. Remedies


There should be legal and financial means to remedy any privacy or security breaches.

² These principles are based upon the Organization for Economic Co-operation and Development's (OECD) Guidelines for the Protection of Privacy and Transborder Flows of Personal Data

http://www.oecd.org/document/18/0,3343,en_2649_34255_1815186_1_1_1_1,00.html#guidelines

The mission of OECD is to promote policies that will improve the economic and social well-being of people around the world.

These principles guide the DHI policies to the extent HIPAA, HITECH, state and federal laws allow. In some instances, current technology will not allow full implementation of the principles; however, these principles guide the policies. The following terms are used in the policies:





Definitions

American Recovery and Reinvestment Act means the appropriations bill signed into law on February 17, 2009.

Authorized User means an individual authorized by a Participant under a DHI Participation Agreement to use the Kentucky Health Information Exchange to access or provide Data for a Permitted Use.

Business Associate shall have the definition assigned to Business Associate under HIPAA by 45 C.F.R. § 160.103.

Cabinet for Health and Family Services or CHFS means the program cabinet of the Commonwealth of Kentucky established by K.R.S. § 12.250.

Covered Entity shall have the definition assigned to Covered Entity under HIPAA by 45 C.F.R. § 160.103.

Data means patient health information provided to KHIE by a Participant.

Division of Health Information was established by Executive Order dated May 14, 2018 to replace DHI with the same responsibilities and mission.

Exchange means the Kentucky Health Information Exchange (KHIE), the health information Exchange provided by DHI. The Exchange provides connection options for the capability to exchange key clinical information among Participants.


Help Desk means the service provided to Participants by KHIE to assist with questions concerning the functions and operation of the exchange.

HITECH means the Health Information Technology for Economic and Clinical Health Act, enacted as part of the American Recovery and Reinvestment Act of 2009.

HIPAA means the Health Insurance Portability and Accountability Act of 2013.

IP address means internet protocol address.

ePartnerViewer means the web-based portal that enables an Authorized User to see a patient's health information aggregated from multiple sources presented in a customizable, functional and easy-to-use format.





National Institute of Standards and Technology means the non-regulatory federal agency within the U.S. Department of Commerce.


Operations shall have the definition assigned to Health Care Operations under HIPAA as limited by 45 C.F.R. §164.506(c)(iv).


Participant means a Health Care Provider, as defined in 45 C.F.R. § 160.103, who is also a Covered Entity as defined by HIPAA, the Kentucky Department for Medicaid Services or the Kentucky State Laboratory, Division of Laboratory Services. The Participant must have entered into a DHI Participation Agreement and have not terminated the Participation Agreement.

Participation Agreement means the agreement between the DHI and Participants to contribute Data to the Kentucky Health Information Exchange.

Payment shall have the meaning set forth at 45 C.F.R. § 164.501 of the HIPAA regulations.

Permitted Use is defined in relationship to the use made by the type of Participant. Currently there are three types of participants providing data; health care providers, the Department of Medicaid Services and the Kentucky State Laboratory, Division of Laboratory Services. Permitted Use is defined as follows:

1. By Health Care Providers:
 - a. For Treatment of the individual who is the subject of the Data,
 - b. For the Payment activities of the Participant for the individual who is the subject of the Data which includes, but is not limited to, exchanging data in response to or to support a claim for reimbursement by Participant to a Health Plan.
 - c. For the Health Care Operations of either The provider of the Data if the provider is a Participant,
 - ii. A covered entity if the provider of the Data is exchanging Data on behalf of the covered entity, or
 - iii. The receiver of the data, if the receiver of the Data is a Participant who has an established Treatment relationship with the individual who the subject of the Data or the receiver of the Data is exchanging Data on behalf of a Participant, and: the purpose of the exchange is for those health care operations listed in paragraphs (1) or (2) of the definition of Health Care Operations in 45 CFR § 164.501 or health care fraud and abuse detection or compliance of such Health Care Provider. All Permitted Use by a Participant is such that patient authorization is not required under HIPAA; and
 - iv. To facilitate the implementation of “meaningful use” criteria as required under the American under the American Recovery and Reinvestment Act of 2009 and its related federal regulations, as permitted by HIPAA; and
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2. By the Department for Medicaid Services:
 - a. For Treatment and Payment for Medicaid patients and/or Operations such that patient authorization is not required under HIPAA, limited to functions related to case management, care coordination, and quality improvement activities; and
 - b. To facilitate the implementation of “meaningful use” criteria as required under the American Recovery and Reinvestment Act of 2009 and its related federal regulations, as permitted by HIPAA.
 3. By the Kentucky State Laboratory, Division of Laboratory Services:
 - a. For Treatment and Payment for patients and/or Operations such that patient authorization is not required under HIPAA, limited to functions related to case management, care coordination, and quality improvement activities for the Kentucky newborn screening program as authorized in KRS 214.155 and cited as the James William Lazzaro and Madison Leigh Heflin Newborn Screening Act; and
 - b. To facilitate the implementation of “meaningful use” criteria as required under the American Recovery and Reinvestment Act of 2013 and its related federal regulations, as permitted by HIPAA.
 4. Public health activities and reporting as permitted by applicable law, including the HIPAA regulation at 45 CFR § 164.512(b) or 164.514(e); and
 5. Uses and disclosures pursuant to an Authorization provided by the individual who is the subject of the Data exchanged or such individual’s personal representative as described in 42 CFR § 164.502(g) of HIPAA.

Privacy Officer shall have the definition assigned to it by the HIPAA regulations at 45 CFR § 164.530(a)(1).

Privacy Rule means those provisions of 45 C.F.R. Part 160 and Subparts A and E of 45 C.F.R. Part 164 of the HIPAA regulations that regulate the privacy of individually identifiable health information.

Protected Health Information shall have the definition assigned to Protected Health Information in 45 C.F.R. § 160.103 of the HIPAA regulations.

Security Rule means those provisions of 45 C.F.R. Part 160 and Subparts A and C of Part 164 of the HIPAA regulations that regulate the security of individually identifiable health information.

Treatment shall have the meaning set forth at 45 C.F.R. § 164.501 of the HIPAA regulations.

Review Date: 01-23-2020

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