Privacy Protections for Virtual Clinical Trials

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Privacy Protections Matter

• Help assure people will seek care for sensitive health conditions
• 1/6 withhold information or decline to seek treatment due to concerns about confidentiality
• Of particular concern for sensitive health information - for example, as many as 1/4 adults in a given year is suffering from a diagnosable mental disorder, and nearly 2/3 do not seek treatment due in part to fear of disclosure, potential rejection from friends, and discrimination
Key to Privacy Protections: Building Trust in Appropriate Data Uses

• Aim of protections is to enable appropriate use
• Fair information practice principles (FIPPs) built on concept of responsible data stewardship
• Informed consent/autonomy is one principle – but it is not absolute
  – Others: transparency; data minimization; safeguards; accountability
Governing “research” uses of health data

- “Clinical Trial” and ”Virtual Clinical Trial” are not defined terms in HIPAA.
- Research = any systematic investigation that has as its primary purpose the development of, or contribution to, generalizable knowledge (same definition in Common Rule).
Governing “research” uses of health data

• If conducted by a HIPAA covered entity, HIPAA rules apply
  – Even if some of the data submitted to the covered entity for the research comes from entities not covered by HIPAA (e.g., devices)

• HIPAA also will govern the extent to which covered entities and business associates can contribute protected health information (PHI) to other entities for research
Does HIPAA cover data in consumer devices?

- It depends.
- Is a device/app a business associate?
  - In general, a business associate is a person [or entity] who creates, receives, maintains or transmits protected health information (PHI) on behalf of a covered entity or another business associate.
- FTC has authority to police unfair and deceptive practices of commercial entities.
Governing research uses of health data in US

- Heavy reliance on de-identification (or anonymization) of data
  - Consistent with data minimization principles
  - Historically has resulted in data “free” from U.S. regulation (if done appropriately)
- Otherwise, consent (or waiver) required – but recently U.S. regulators have allowed for more general consents for uses of identifiable data for research purposes (HIPAA guidance & Common Rule changes)
- There are paths forward – but risk aversion can create barriers.
Recent privacy law trends

• More attention focused on this issue – GDPR & passage of new, sweeping privacy legislation in California last summer
• Trend is toward requiring more explicit consent
• Also setting a higher bar for data to be considered “de-identified”
GDPR

• Went into effect on May 25, 2018
• Applies to data “controllers” and “processors” in the EU
  ○ Also applies to entities not located in the EU but who offer goods and services to EU residents or collect information from, or monitor the behavior of, EU data subjects within the EU
• Applies to “personal data”
  ○ Data still regulated but some relaxation of rules (individual rights provisions) regarding “pseudonymized” or “coded” data
  ○ Allows for scientific research, subject to safeguards
  ○ Doesn’t apply to data made public by the data subject
  ○ Doesn’t apply to data from individuals no longer living
“[P]ersonal data’ means any information relating to an identified or identifiable natural person (‘data subject’); an identifiable natural person is one who can be identified, directly or indirectly, in particular by reference to an identifier such as a name, an identification number, location data, an online identifier or to one or more factors specific to the physical, physiological, genetic, mental, economic, cultural or social identity of that natural person;”
CCPA (goes into effect 1/1/2020)

• Applies to a “Business” that:
  • Has annual gross revenues in excess of twenty-five million dollars ($25,000,000) (California revenue?);
  • Alone or in combination, annually buys, receives for the business’ commercial purposes, sells, or shares for commercial purposes, alone or in combination, the personal information of 50,000 or more CA consumers, households, or devices; or
  • Derives 50 percent or more of its annual revenues from selling CA consumers’ personal information. (Emphasis Added)
CCPA covers “personal information”

“Personal Information”

- SB 1121 1798.140(o)
- (1) “Personal Information” means information that identifies, relates to, describes, is capable of being associated with, or could reasonably be linked, directly or indirectly, with a particular consumer or household.
- Personal information includes, but is not limited to, the following if it identifies, relates to, describes, is capable of being associated with, or could be reasonably linked, directly or indirectly, with a particular consumer or household:
  - (A) Identifiers such as a real name, alias, postal address, unique personal identifier, online identifier, Internet Protocol address, email address, account name, social security number, driver’s license number, passport number, or other similar identifiers.
  - (B) Any categories of personal information described in subdivision (e) of Section 1798.80.
  - (C) Characteristics of protected classifications under California or federal law.
  - (D) Commercial information, including records of personal property, products or services purchased, obtained, or considered, or other purchasing or consuming histories or tendencies.
  - (E) Biometric information.
CCPA “personal information”

- (F) Internet or other electronic network activity information, including, but not limited to, browsing history, search history, and information regarding a consumer’s interaction with an Internet Web site, application, or advertisement.
- (G) Geolocation data.
- (H) Audio, electronic, visual, thermal, olfactory, or similar information.
- (I) Professional or employment-related information.
- (J) Education information, defined as information that is not publicly available personally identifiable information as defined in the Family Educational Rights and Privacy Act (20 U.S.C. section 1232g, 34 C.F.R. Part 99).
- (K) Inferences drawn from any of the information identified in this subdivision to create a profile about a consumer reflecting the consumer’s preferences, characteristics, psychological trends, predispositions, behavior, attitudes, intelligence, abilities, and aptitudes. (Emphasis added)
CCPA – definition of “deidentified”

SB 1121 1798.140

• (h) “Deidentified” means information that cannot reasonably identify, relate to, describe, be capable of being associated with, or be linked, directly or indirectly, to a particular consumer, provided that a business that uses deidentified information:
  – (1) Has implemented technical safeguards that prohibit reidentification of the consumer to whom the information may pertain.
  – (2) Has implemented business processes that specifically prohibit reidentification of the information.
  – (3) Has implemented business processes to prevent inadvertent release of deidentified information.
  – (4) Makes no attempt to reidentify the information.
• Believed to be stricter than HIPAA & Common Rule
CCPA Exemptions for Health Entities

1798.145
(c)(1) This title shall not apply to any of the following:

(A) Medical information governed by the Confidentiality of Medical Information Act (Part 2.6 (commencing with Section 56) of Division 1) or protected health information that is collected by a covered entity or business associate governed by the privacy, security, and breach notification rules issued by the U.S. Department of Health and Human Services, Parts 160 and 164 of Title 45 of the Code of Federal Regulations, established pursuant to the Health Insurance Portability and Accountability Act of 1996 (Public Law 104-191) and the Health Information Technology for Economic and Clinical Health Act (Public Law 111-5).

(B) A provider of health care governed by the Confidentiality of Medical Information Act (Part 2.6 (commencing with Section 56) of Division 1) or a covered entity governed by the privacy, security, and breach notification rules issued by the United States Department of Health and Human Services, Parts 160 and 164 of Title 45 of the Code of Federal Regulations, established pursuant to the Health Insurance Portability and Accountability Act of 1996 (Public Law 104-191), to the extent the provider or covered entity maintains patient information in the same manner as medical information or protected health information as described in subparagraph (A) of this section.

(C) Information collected as part of a clinical trial subject to the Federal Policy for the Protection of Human Subjects, also known as the Common Rule, pursuant to good clinical practice guidelines issued by the International Council for Harmonisation, or pursuant to the human subject protection requirements of the United States Food and Drug Administration.

(2) For purposes of this subdivision, the definitions of “medical information” and “provider of health care” in Section 56.05 shall apply and the definitions of business associate,” covered entity,” and “protected health information” in Section 160.103 of title 45 of the Code of Federal Regulations shall apply.
Impact of GDPR (& CCPA)

• Global companies have taken steps to implement
  o Raising the bar even for companies not required to comply?
• Will US lawmakers feel compelled to fill gaps in US law or to provide a federal standard that preempts the new CA law?
  o Will these efforts complement or contradict GDPR?
• Will increased attention to privacy result in greater — or less - data sharing for good?
• Impact of services empowering individuals to gather and share their data, including health data?
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