

FEDERAL TRADE COMMISSION**16 CFR Part 318**

RIN 3084-AB56

Health Breach Notification Rule**AGENCY:** Federal Trade Commission.**ACTION:** Final rule.

SUMMARY: The Federal Trade Commission (“FTC” or “Commission”) is amending the Commission’s Health Breach Notification Rule (the “HBN Rule” or the “Rule”). The HBN Rule requires vendors of personal health records (“PHRs”) and related entities that are not covered by the Health Insurance Portability and Accountability Act (“HIPAA”) to notify individuals, the FTC, and, in some cases, the media of a breach of unsecured personally identifiable health data.

DATES: The amendments are effective July 29, 2024.

ADDRESSES: Relevant portions of the record of this proceeding, including this document, are available at <https://www.ftc.gov> and <https://www.regulations.gov>.

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SUPPLEMENTARY INFORMATION: The amendments: (1) clarify the Rule’s scope, including its coverage of developers of many health applications (“apps”); (2) clarify what it means for a vendor of personal health records to draw PHR identifiable health information from multiple sources; (3) revise the definition of breach of security to clarify that a breach of security includes data security breaches and unauthorized disclosures; (4) revise the definition of PHR related entity; (5) modernize the method of notice; (6) expand the content of the notice; (7) alter the Rule’s timing requirement for notifying the FTC of a breach of security; and (8) improve the Rule’s readability by clarifying cross-references and adding statutory citations, consolidating notice and timing requirements, articulating the penalties for non-compliance, and incorporating a small number of non-substantive changes.

I. Background

Congress enacted the American Recovery and Reinvestment Act of 2009

(“Recovery Act” or “the Act”),¹ in part to advance the use of health information technology and, at the same time, strengthen privacy and security protections for health information. Recognizing that certain entities that hold or interact with consumers’ personal health records were not subject to the privacy and security requirements of HIPAA,² Congress created requirements for such entities to notify individuals, the Commission, and, in some cases, the media of the breach of unsecured identifiable health information from those records.

Specifically, section 13407 of the Recovery Act created certain protections for “personal health records” or “PHRs,”³ electronic records of PHR identifiable health information on an individual that can be drawn from multiple sources and that are managed, shared, and controlled by or primarily for the individual.⁴ Congress recognized that vendors of personal health records and PHR related entities (*i.e.*, companies that offer products and services through PHR websites or access information in or send information to personal health records) were collecting consumers’ health information but were not subject to the privacy and security requirements of HIPAA. Accordingly, the Recovery Act directed the FTC to issue a rule requiring these non-HIPAA covered entities, and their third party service providers, to provide notification of any breach of unsecured PHR identifiable health information. The Commission issued its Rule implementing these provisions in 2009.⁵ FTC enforcement of the Rule began on February 22, 2010.

The Rule the Commission issued in 2009 (“2009 Rule”) requires vendors of personal health records and PHR related entities to provide: (1) notice to consumers whose unsecured PHR identifiable health information has been breached; (2) notice to the Commission; and (3) notice to prominent media outlets⁶ serving a State or jurisdiction, in cases where 500 or more residents are

confirmed or reasonably believed to have been affected by a breach.⁷ The Rule also requires third party service providers (*i.e.*, those companies that provide services such as billing, data storage, attribution, or analytics) to vendors of personal health records and PHR related entities to provide notification to such vendors and entities following the discovery of a breach.⁸

The 2009 Rule requires notice to individuals “without unreasonable delay and in no case later than 60 calendar days” after discovery of a data breach.⁹ If the breach affects 500 or more individuals, notice to the FTC must be provided “as soon as possible and in no case later than ten business days” after discovery of the breach.¹⁰ The FTC makes available a standard form for companies to use to notify the Commission of a breach,¹¹ and posts a list of breaches involving 500 or more individuals on its website.¹²

The 2009 Rule applies only to breaches of “unsecured” health information, which the Rule defines as health information that is not secured through technologies or methodologies specified by the Department of Health and Human Services (“HHS”). The Rule does not apply to businesses or organizations covered by HIPAA,¹³ HIPAA-covered entities and their “business associates” must instead comply with HHS’s breach notification rule.¹⁴

⁷ 16 CFR 318.3, 318.5.

⁸ *Id.* § 318.3(b).

⁹ *Id.* § 318.4(a).

¹⁰ *Id.* § 318.5(c).

¹¹ Fed. Trade Comm’n, Notice of Breach of Health Information, https://www.ftc.gov/system/files/documents/rules/health-breach-notification-rule/health_breach_form.pdf.

¹² Fed. Trade Comm’n, Notices Received by the FTC Pursuant to the Health Breach Notification Rule, https://www.ftc.gov/system/files/ftc_gov/pdf/Health%20Breach%20Notices%20Received%20by%20the%20FTC.pdf (last visited Dec. 2, 2022).

¹³ Per HHS guidance, electronic health information is “secured” if it has been encrypted according to certain specifications set forth by HHS, or if the media on which electronic health information has been stored or recorded is destroyed according to HHS specifications. *See* 74 FR 19006; *see also* U.S. Dep’t of Health & Human Servs., *Guidance to Render Unsecured Protected Health Information Unusable, Unreadable, or Indecipherable to Unauthorized Individuals* (July 26, 2013), <https://www.hhs.gov/hipaa/for-professionals/breach-notification/guidance/index.html>. PHR identifiable health information would be considered “secured” if such information is disclosed by, for example, a vendor of personal health records, to a PHR related entity or a third party service provider, in an encrypted format meeting HHS specifications, and the PHR related entity or third party service provider stores the data in an encrypted format that meets HHS specifications and also stores the encryption and/or decryption tools on a device or at a location separate from the data.

¹⁴ 45 CFR 164.400 through 164.414.

¹ Am. Recovery and Reinvestment Act of 2009, Public Law 111-5, 123 Stat. 115 (2009).

² Health Ins. Portability and Accountability Act, Public Law 104-191, 110 Stat. 1936 (1996).

³ 42 U.S.C. 17937.

⁴ 42 U.S.C. 17921(11).

⁵ 74 FR 42962 (Aug. 25, 2009) (“2009 Final Rule”).

⁶ The Recovery Act does not limit this notice to particular types of media. Thus, an entity can satisfy the requirement to notify “prominent media outlets” by, for example, disseminating press releases to a number of media outlets, including internet media in appropriate circumstances, where most of the residents of the relevant State or jurisdiction get their news. This will be a fact-specific inquiry that will depend on what media outlets are “prominent” in the relevant jurisdiction. 74 FR 42974.

Since the Rule's issuance, apps and other direct-to-consumer health technologies, such as fitness trackers and wearable blood pressure monitors, have become commonplace.¹⁵ Further, as an outgrowth of the COVID-19 pandemic, consumer use of such health-related technologies has increased significantly.¹⁶

In May 2020, the Commission announced its regular, ten-year review of the Rule and requested public comment about potential Rule changes.¹⁷ The Commission requested comment on, among other things, whether changes should be made to the Rule in light of technological changes, such as the proliferation of apps and similar technologies. The Commission received 26 public comments.¹⁸

Many of the commenters in 2020 encouraged the Commission to clarify that the Rule applies to apps and similar technologies.¹⁹ In fact, no commenter opposed this type of clarification regarding the Rule's coverage of health apps. Several commenters pointed out examples of health apps that have abused users' privacy, such as by

disclosing sensitive health information without consent.²⁰ Several commenters noted the urgency of this issue, as consumers have further embraced digital health technologies during the COVID-19 pandemic.²¹ Commenters argued the Commission should take additional steps to protect unsecured PHR identifiable health information that is not covered by HIPAA, both to prevent harm to consumers²² and to level the competitive playing field among companies dealing with the same health information.²³ To that end, commenters not only urged the Commission to revise the Rule, but also to increase its enforcement efforts.²⁴

A. The Commission's 2021 Policy Statement

On September 15, 2021, the Commission issued a Policy Statement providing guidance on the scope of the Rule. The Policy Statement clarified that the Rule covers most health apps and similar technologies that are not covered

by HIPAA.²⁵ The Rule defines a "personal health record" as "an electronic record of PHR identifiable health information on an individual that can be drawn from multiple sources and that is managed, shared, and controlled by or primarily for the individual."²⁶ As the Commission explained in the Policy Statement, many makers and purveyors of health apps and other connected devices are vendors of personal health records covered by the Rule because their products are electronic records of PHR identifiable health information.

The Commission explained that PHR identifiable health information includes individually identifiable health information created or received by a health care provider,²⁷ and that "health care providers" include any entities that "furnish[] health care services or supplies."²⁸ Because these health app purveyors furnish health care services to their users through the mobile applications they provide, the information held in the app is PHR identifiable health information, and therefore many health app purveyors likely qualify as vendors of personal health records.²⁹

The Policy Statement further explained that the statute directing the FTC to promulgate the Rule requires that a "personal health record" be an electronic record that can be drawn from multiple sources.³⁰ Accordingly, health apps and similar technologies likely qualify as personal health records covered by the Rule if they are capable of drawing information from multiple sources. The Commission further clarified that health apps and other products experience a "breach of security" under the Rule when they disclose users' sensitive health information without authorization;³¹ a breach is "not limited to cybersecurity intrusions or nefarious behavior."³²

²⁵ Statement of the Commission on Breaches by Health Apps and Other Connected Devices, Fed. Trade Comm'n (Sept. 15, 2021), https://www.ftc.gov/system/files/documents/public_statements/1596364/statement_of_the_commission_on_breaches_by_health_apps_and_other_connected_devices.pdf ("Policy Statement").

²⁶ 16 CFR 318.2.

²⁷ *Id.* § 318.2, incorporating in part the definition from section 1171(6) of the Social Security Act (42 U.S.C. 1320d(6)).

²⁸ *Id.* § 318.2; 42 U.S.C. 1320d(6), d(3).

²⁹ See Policy Statement at 1.

³⁰ The Policy Statement provided this example: "[I]f a blood sugar monitoring app draws health information only from one source (e.g., a consumer's inputted blood sugar levels), but also takes non-health information from another source (e.g., dates from your phone's calendar), it is covered under the Rule." *Id.* at 2.

³¹ 16 CFR 318.2.

³² Policy Statement at 2. In the Statement of Basis and Purpose to the 2009 Final Rule published in the

¹⁵ See, e.g., Kokou Adzo, *App Development in Healthcare: 12 Exciting Facts*, TechnoChops (Jan. 3, 2023), <https://www.technochops.com/programming/4329/app-development-in-healthcare/>; Emily Olsen, *Digital health apps balloon to more than 350,000 available on the market, according to IQVIA report*, MobiHealthNews (Aug. 4, 2021), <https://www.mobihealthnews.com/news/digital-health-apps-balloon-more-350000-available-market-according-iqvia-report>; Elad Natanson, *Healthcare Apps: A Boon, Today and Tomorrow*, Forbes (July 21, 2020), <https://www.forbes.com/sites/eladnatanson/2020/07/21/healthcare-apps-a-boon-today-and-tomorrow/?sh=21df01ac1bb9>.

¹⁶ See *id.* See also Lis Evenstad, *Covid-19 has led to a 25% increase in health app downloads, research shows*, ComputerWeekly.com (Jan. 12, 2021), <https://www.computerweekly.com/news/252494669/Covid-19-has-led-to-a-25-increase-in-health-app-downloads-research-shows> (finding that COVID-19 has led to a 25% increase in health app downloads); Jasmine Pennic, *U.S. Telemedicine App Downloads Spikes During COVID-19 Pandemic*, HIT Consultant (Sept. 8, 2020), <https://hitconsultant.net/2020/09/08/u-s-telemedicine-app-downloads-spikes-during-covid-19-pandemic/> ("US telemedicine app downloads see dramatic increases during the COVID-19 pandemic, with some seeing an 8,270% rise YoY.").

¹⁷ 85 FR 31085 (May 22, 2020).

¹⁸ Comments are available at <https://www.regulations.gov/docket/FTC-2020-0045/comments>.

¹⁹ E.g., Am. Health Info. Mgmt. Ass'n ("AHIMA") at 2; Kaiser Permanente at 3; Allscripts at 3; Am. Acad. of Ophthalmology at 2; All. for Nursing Informatics ("ANI") at 2; Am. Med. Ass'n ("AMA") at 4; Am. Coll. of Surgeons at 6; Physicians' Elec. Health Rec. Coal. ("PEHRC") at 4 ("Apps that collect health information, regardless of whether or not they connect to an EHR, must be regulated by the FTC Health Breach Notification Rule to ensure the safety and security of personal health information."); Am.'s Health Ins. Plans ("AHIP") and Blue Cross Blue Shield Ass'n ("BCBS") at 2; The App Ass'n's Connected Health Initiative ("CHI") at 3.

²⁰ Kaiser Permanente at 7; The Light Collective at 2; Am. Acad. of Ophthalmology at 2; PEHRC at 2-3.

²¹ Lisa McKeen at 2-3; Kaiser Permanente at 7-8; AMA at 3; Off. of the Att'y Gen. for the State of Cal. ("OAG-CA") at 3-4; Healthcare Info. and Mgmt. Sys. Soc'y ("HIMSS") and Personal Connected Health All. ("PCH Alliance") at 4-5.

²² Georgia Morgan; Am. Acad. of Ophthalmology at 2-3 (arguing that consumers do not know all the ways their data is being used by third parties, and the downstream consequences of data being used in this way may ultimately erode a patient's privacy and willingness to disclose information to his or her physician); Coll. of Healthcare Info. Mgmt. Exec.'s ("CHIME") at 3 (arguing that apps' privacy practices impact the patient-provider relationship because providers do not know what technologies are sufficiently trustworthy for their patients); AMA at 2-3 (expressing concern that patients share less health data with health care providers, perhaps because of "spillover from privacy and security breaches").

²³ Kaiser Permanente at 2, 4; Workgroup for Elec. Data Interchange ("WEDI") at 2; AHIP and BCBS at 3 ("[HIPAA] covered entities, such as health plans, that use or disclose protected health information should not be subject to stricter notification requirements than those imposed on vendors of personal health records or other such entities. Otherwise, the Federal government will be providing market advantages to particular industry segments with the effect of dampening competition and harming consumers.").

²⁴ Kaiser Permanente at 4; Fred Trotter at 1; Casey Quinlan at 1; CARIN Alliance at 2. At the time of this document's publication, the Commission has brought two enforcement actions under the Rule; the first against digital health company GoodRx Holdings, Inc., and the second against an ovulation-tracking mobile app marketed under the name "Premom" and developed by Easy Healthcare, Inc. *United States v. GoodRx Holdings, Inc.*, No. 23-cv-460 (N.D. Cal. Feb. 17, 2023), <https://www.ftc.gov/legal-library/browse/cases-proceedings/2023090-goodrx-holdings-inc>; *United States v. Easy Healthcare Corp.*, No. 1:23-cv-3107 (N.D. Ill. June 22, 2023), <https://www.ftc.gov/legal-library/browse/cases-proceedings/202-3186-easy-healthcare-corporation-us-v>.

B. Enforcement History

In 2023, the Commission brought its first enforcement actions under the Rule against vendors of personal health records. In February 2023, the Commission brought an enforcement action alleging a violation of the Rule against GoodRx Holdings, Inc. (“GoodRx”), a digital health company that sells health-related products and services directly to consumers, including prescription medication discount products and telehealth services through its website and mobile applications.³³

In its complaint, the Commission alleged that between 2017 and 2020, GoodRx, as a vendor of personal health records, disclosed more than 500 consumers’ unsecured PHR identifiable health information to third party advertising platforms like Facebook and Google, without the authorization of those consumers. As charged in the complaint, these disclosures violated explicit privacy promises the company made to its users about its data sharing practices (including about its sharing of PHR identifiable health information). The Commission alleged GoodRx broke these promises and disclosed its users’ prescription medications and personal health conditions, personal contact information, and unique advertising and persistent identifiers. The Commission charged GoodRx with violating the Rule by failing to provide the required notifications, as prescribed by the Rule, to (1) individuals whose unsecured PHR identifiable health information was acquired by an unauthorized person, (2) the Federal Trade Commission, and (3) media outlets. 16 CFR 318.3 through 318.6. The Commission entered into a settlement that imposed injunctive relief and required GoodRx to pay a \$1.5 million civil penalty for its alleged violation of the Rule.³⁴

Similarly, on May 17, 2023, the Commission brought its second

Federal Register (“2009 Rule Commentary”), the Commission, in addressing questions about how the extent of individual authorization should be determined, stated data sharing to enhance consumers’ experience with a PHR is authorized only if such use is consistent with the entity’s disclosures and individuals’ reasonable expectations. For anything beyond such uses, the Commission expects vendors of personal health records and PHR related entities to limit the sharing of consumers’ information, unless the consumers exercise “meaningful choice” in allowing sharing. The Commission believes burying disclosures in lengthy privacy policies does not satisfy the standard of “meaningful choice.” 74 FR 42967.

³³ *United States v. GoodRx Holdings, Inc.*, No. 23-cv-460 (N.D. Cal. Feb. 17, 2023), <https://www.ftc.gov/legal-library/browse/cases-proceedings/2023090-goodrx-holdings-inc>.

³⁴ In addition, the Commission alleged GoodRx’s data sharing practices were deceptive and unfair, in violation of section 5 of the FTC Act.

enforcement action under the Rule against Easy Healthcare Corporation (“Easy Healthcare”), a company that publishes an ovulation and period tracking mobile application called Premom, which allows its users to input and track various types of health and other sensitive data. Similar to the conduct alleged against GoodRx, Easy Healthcare disclosed PHR identifiable health information to third party companies such as Google and AppsFlyer, contrary to its privacy promises, and did not comply with the Rule’s notification requirements. The Commission entered into a settlement that imposed injunctive relief and required Easy Healthcare to pay a \$100,000 civil penalty for its alleged violation of the Rule.³⁵

C. Notice of Proposed Rulemaking

Having considered the public comments on the regulatory review notification and its Policy Statement, on June 9, 2023, the Commission issued a notice of proposed rulemaking (“NPRM”) ³⁶ proposing to revise the Rule, 16 CFR part 318, in seven ways:

- First, the Commission proposed to revise several definitions in order to clarify the Rule and better explain its application to health apps and similar technologies not covered by HIPAA. Consistent with this objective, the NPRM modified the definition of “PHR identifiable health information” and added two new definitions (“health care provider” and “health care services or supplies”). These proposed changes were consistent with a number of public comments supporting the Rule’s coverage of these technologies.

- Second, the Commission proposed to revise the definition of “breach of security” to clarify that a breach of security includes an unauthorized acquisition of PHR identifiable health information in a personal health record that occurs as a result of a data security breach or an unauthorized disclosure.

- Third, the Commission proposed to revise the definition of “PHR related entity” in two ways. Consistent with its proposal to clarify that the Rule applies to health apps, the Commission first proposed clarifying the definition of “PHR related entity” to make clear that the Rule covers entities that offer products and services through the online services, including mobile applications, of vendors of personal health records. In addition, the

Commission proposed revising the definition of “PHR related entity” to provide that entities that access or send unsecured PHR identifiable health information to a personal health record—rather than entities that access or send *any* information to a personal health record—are PHR related entities.

- Fourth, the Commission proposed to clarify what it means for a personal health record to draw PHR identifiable health information from multiple sources.

- Fifth, in response to public comments expressing concern that mailed notice is costly and not consistent with how consumers interact with online technologies like health apps, the Commission proposed to revise the Rule to authorize electronic notice in additional circumstances. Specifically, the proposed Rule adjusted the language in the “method of notice section” and added a new definition of the term “electronic mail.” The proposed Rule also required that any notice delivered by electronic mail be “clear and conspicuous,” a newly defined term, which aligns closely with the definition of “clear and conspicuous” codified in the FTC’s Financial Privacy Rule.³⁷

- Sixth, the Commission proposed to expand the required content of the notice to individuals, to require that consumers whose unsecured PHR identifiable health information has been breached receive additional important information, including information regarding the potential for harm from the breach and protections that the notifying entity is making available to affected consumers. In addition, the proposed Rule included exemplar notices, which entities subject to the Rule could use to notify consumers in terms that are easy to understand.

- Seventh, in response to public comments, the Commission proposed to make a number of changes to improve the Rule’s readability. Specifically, the Commission proposed to include explanatory parentheticals for internal cross-references, add statutory citations in relevant places, consolidate notice and timing requirements in single sections, respectively, of the Rule, and add a new section that plainly states the penalties for non-compliance.

The NPRM also included a section discussing several alternatives the

³⁷ 16 CFR 313.3(b). The FTC’s Financial Privacy Rule requires financial institutions to provide particular notices and to comply with certain limitations on disclosure of nonpublic personal information. Using a comprehensive definition of “clear and conspicuous” based on the Financial Privacy Rule definition aims to ensure consistency across the Commission’s privacy-related rules.

³⁵ *United States v. Easy Healthcare Corporation*, No. 1:23-cv-3107 (N.D. Ill. June 22, 2023), <https://www.ftc.gov/legal-library/browse/cases-proceedings/2023186-easy-healthcare-corporation-us-v>.

³⁶ 88 FR 37819 (“2023 NPRM”).

Commission considered but did not propose. Although the Commission did not put forth any proposed modifications on those issues, the Commission nonetheless sought public comment on them.

The Commission received approximately 120 comments in response to the NPRM from a wide spectrum of stakeholders, including consumers, consumer groups, trade associations, think tanks, policy organizations, private sector entities, and members of Congress.³⁸ As discussed in detail below, commenters addressed the seven topics on which the Commission proposed changes, responded to particular points on which the Commission requested comment, offered additional comment on alternatives that the Commission considered but did not propose, and provided comment on other topics. The majority of commenters expressed support for the Commission's proposed changes.

The Commission believes the amendments are consistent with the language and intent of the Recovery Act, address the concerns raised by the public comments in response to the NPRM, and will ensure the Rule remains current in the face of changing business practices and technological developments.

II. Analysis of the Final Rule

The following discussion analyzes the amendments to the Rule.

A. Clarification of Entities Covered

1. The Commission's Proposal To Clarify the Entities Covered

The Commission proposed changes to several definitions in § 318.2 to clarify the Rule's application to health apps and similar technologies not covered by HIPAA. First, the proposed Rule revised the definition of "PHR identifiable health information" to remove a cross-reference and instead import language from section 1171(6) of the Social Security Act, 42 U.S.C. 1320d(6), which is also referenced directly in section 13407 of the Recovery Act. The proposed Rule defined "PHR identifiable health information" as information (1) that is provided by or on behalf of the individual; (2) that identifies the individual or with respect to which there is a reasonable basis to believe that the information can be used to identify the individual; (3) relates to the past, present, or future physical or mental 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; and (4) is created or received by a health care provider, health plan (as defined in 42 U.S.C. 1320d(5)), employer, or health care clearinghouse (as defined in 42 U.S.C. 1320d(2)).

The Commission explained that this proposed definition covers traditional health information (such as diagnoses or medications), health information derived from consumers' interactions with apps and other online services (such as health information generated from tracking technologies employed on websites or mobile applications or from customized records of website or mobile application interactions), as well as emergent health data (such as health information inferred from non-health-related data points, such as location and recent purchases). The Commission sought comment as to whether any further amendment of the definition was needed to clarify the scope of data covered.

Second, the NPRM proposed to define the term "health care provider" that appears in the proposed definition of "PHR identifiable health information" ("is created or received by a health care provider"). The Commission proposed to define this term in a manner similar to the definition of "health care provider" found in 42 U.S.C. 1320d(3) (and referenced in 42 U.S.C. 1320d(6), which is directly referenced in section 13407 of the Recovery Act), to mean a provider of services (as defined in 42 U.S.C. 1395x(u)), a provider of medical or other health services (as defined in 42 U.S.C. 1395x(s)), or any other entity furnishing health care services or supplies. The Commission observed that this proposed definition, which is consistent with the statutory scheme, differs from, but does not contradict, the definitions or interpretations adopted by HHS. The Commission sought comment on defining this term more broadly than the term is used in other contexts.

Third, the NPRM proposed to define "health care services or supplies" (the final term in the definition of "health care provider") to include any online service, such as a website, mobile application, or internet-connected device that provides mechanisms to track diseases, health conditions, diagnoses or diagnostic testing, treatment, medications, vital signs, symptoms, bodily functions, fitness, fertility, sexual health, sleep, mental health, genetic information, diet, or that provides other health-related services or tools. The Commission explained that this change clarified that the Rule

applies generally to online services, including websites, apps, and internet-connected devices that provide health care services or supplies, and clarified that the Rule covers online services related not only to medical issues (by including in the definition terms such as "diseases, diagnoses, treatment, medications") but also wellness issues (by including in the definition terms such as "fitness, sleep, and diet").

The Commission explained that these proposed changes to the definitions clarified that developers of health apps and similar technologies providing "health care services or supplies" qualify as "health care providers," such that any individually identifiable health information these products collect or use would constitute "PHR identifiable health information" covered by the Rule. The Commission explained that these proposed changes further clarified that a mobile health application can be a "personal health record" covered by the Rule and the developers of such applications can be "vendors of personal health records."

2. Public Comments Regarding the Commission's Proposal To Clarify the Entities Covered

The Commission received numerous comments on the application of the Rule to health apps and similar technologies. A substantial number of commenters supported the Rule's application to health apps and similar technologies not covered by HIPAA as necessary in light of the explosion of health apps and the associated dangers to the privacy and security of consumers' health information.³⁹ Notably, support for the

³⁹ See generally, Am. Acad. of Fam. Physicians ("AAFP"); AHIP; AHIMA; Ass'n of Health Info. Outsourcing Serv.'s ("AHIOS"); AMA; Am. Med. Informatics Ass'n ("AMIA"); ANI; Anonymous 1; Anonymous 2; Anonymous 3; Anonymous 4; Anonymous 9; Anonymous 10; Anonymous 11; Anonymous 14; Am. Osteopathic Ass'n ("AOA"); Ella Balasa; Beth Barnett; Lauren Batchelor; Bipartisan Pol'y Ctr. ("BPC"); Alan Brewington; Ctr. for Democracy & Tech. ("CDT"); Ctr. for Digit. Democracy ("CDD"); Confidentiality Coal.; Consumer Rep.'s; Elec. Frontier Found. ("EFF"); Elec. Priv. Info. Ctr. ("EPIC"); Dave K.; Members of the House of Representatives; MRO Corp. ("MRO"); Omada Health; Planned Out; Planned Parenthood Federation of Amer. ("Planned Parenthood"); CB Sanders; Robb Streicher; SYNGAP1 Foundation and SYNGAP1 Foundation 2; Devin Thompson; Janice Tufte; Michael Turner; U.S. Public Interest Research Group ("U.S. PIRG"); UL Sol.'s; Grace Vinton; WEDI; Anli Zhou. Some commenters elaborated on the nature of the risks to consumers' health data and on the importance to consumers. Two commenters, for example, described research they had performed regarding mental health and/or reproductive health apps' disclosure of consumers' health data to third parties. Mozilla at 3-4; Consumer Reports at 2. Another commenter, a public interest group and advocacy organization, attached a petition containing 9,659 signatures

³⁸ Comments are available at <https://www.regulations.gov/document/FTC-2023-0037-0001/comment>.

Commission's proposals came from a variety of commenters—industry associations,⁴⁰ businesses,⁴¹ members of Congress,⁴² consumer or patient advocacy groups,⁴³ individual consumers,⁴⁴ and anonymous sources.⁴⁵ Many commenters argued that safeguards for non-HIPAA covered health data are essential,⁴⁶ particularly because consumers generally are not aware of varying legal protections for health data.⁴⁷ Indeed, according to some commenters, requiring notification to consumers of the breach of health information not protected by HIPAA is precisely what Congress intended by authorizing the FTC to issue this Rule; the Commission's proposed changes are, therefore, consistent with the goals of the Recovery Act.⁴⁸ Some commenters argued that Federal privacy legislation is needed to protect non-HIPAA covered health data, but, in the interim, the Commission should strengthen its Rule to protect consumer health data to the extent possible.⁴⁹ Other commenters

asking for strong rules to protect digital health privacy. US PIRG at 5–230.

⁴⁰ E.g., AAFP, AHIMA, AHIOS, AMA, AMIA, AOA; Network Advert. Initiative (“NAI”).

⁴¹ E.g., Mozilla; MRO; Omada Health; UL Sol.’s.

⁴² See Members of the House of Representatives (six members of Congress expressing support for the proposed changes).

⁴³ E.g., CDD; CDT; EFF; U.S. PIRG.

⁴⁴ Ella Balasa; Beth Barnett; Lauren Batchelor; Alan Brewington; Sean Castillo; Dave K.; CB Sanders; Robb Streicher; Devin Thompson; Janice Tuft; Michael Turner; Grace Vinton; Anli Zhou.

⁴⁵ Anonymous 1; Anonymous 2; Anonymous 3; Anonymous 4; Anonymous 5; Anonymous 6; Anonymous 9; Anonymous 10; Anonymous 11; Anonymous 14.

⁴⁶ See, e.g., AAFP at 1–2; AHIMA at 2; AHIOS at 2; Anonymous 5 at 1; AOA at 1; Am. Speech-Language-Hearing Ass’n (“ASHA”) at 1; Am. Psychiatric Ass’n (“APA”) at 1; CDT at 3–4; CHIME at 2; EFF at 1; Generation Patient at 1; HIMSS at 2; HIMSS Elec. Health Rec. Ass’n (“HIMSS EHR Ass’n”) at 1; MRO at 1–2; Omada Health at 2; PharmedOut at 1; Planned Parenthood at 2–3; Michael Turner at 1; WEDI at 1–4.

⁴⁷ AHIMA at 2; Anonymous 5 at 1; ASHA at 1; EFF at 1; WEDI at 2. One commenter, a software company that assists digital health companies with legal compliance, argued that three factors, in particular, support greater protection for digital health data: (1) consumers mistakenly believe HIPAA covers all health data; (2) there is a culture within some digital health companies that favors rapid adoption of products to secure venture capital even when compliance infrastructure is lacking; and (3) digital health products deal with sensitive data and inherently present a greater privacy risk given their heavy reliance on data and data exchange compared to traditional medicine. Tranquil Data at 1.

⁴⁸ Confidentiality Coal. at 2; Consumer Rep.’s at 4.

⁴⁹ See, e.g., AAFP at 2. One commenter, an industry coalition focused on health IT and health care information exchange, emphasized a significant privacy problem adjacent to the Rule: whether HIPAA covered entities should warn patients about the privacy risks associated with health apps and what the Federal government can do to apply equal privacy protections to health data,

urged the Commission to take even broader measures in this Rule, such as imposing breach prevention measures,⁵⁰ banning health-based surveillance technologies or targeted advertising,⁵¹ banning selling or sharing of health data not necessary to provide patient care or mandating data retention limits and deletion,⁵² or requiring adherence to standardized terms of service with strong privacy protections.⁵³

Although many commenters expressed support for the proposed changes, several business coalitions, industry associations and individual firms opposed the changes, which, they argued, are inconsistent with Congress’s intent in the Recovery Act to address a narrow subset of “personal health records” and therefore exceed the FTC’s statutory authority.⁵⁴ According to some comments, Congress should address any privacy issues that exceed the narrow scope of the Recovery Act. These commenters also contend that if the Commission believes there has been a violation of section 5, then the Commission needs to engage in an FTC Act section 18 rulemaking.⁵⁵ One commenter argued further that consumers have different privacy expectations for an electronic health record offered by their physician versus a fitness app (for example) that they download themselves, and the Commission’s Rule should respect those differing expectations.⁵⁶

Some commenters opposed to the changes also argued that the revised definitions would reduce choice and access in the marketplace,⁵⁷ stifle innovation,⁵⁸ or create disincentives for advertising⁵⁹ because (1) firms would risk initiating breaches by sharing user data with their partners and (2) in

notwithstanding HIPAA’s limitations. See WEDI at 3. One commenter supported the proposed changes but argued the Commission should work with Congress to update antiquated terms like “personal health record.” HIMSS at 3.

⁵⁰ Ella Balasa at 2; PharmedOut at 1.

⁵¹ Light Collective at 5.

⁵² EFF at 2.

⁵³ Texas Med. Ass’n (“TMA”) at 1–2.

⁵⁴ See, e.g., Ass’n of Nat’l Advertisers, Inc. (“ANA”) at 4–5; Comput. & Comm’n’s Indus. Ass’n (“CCIA”) at 2–3; Chamber of Com. (“Chamber”) at 1–3; CHI at 2; Consumer Tech. Ass’n (“CTA”) at 2; Lab’y Access and Benefits Coal. (“LAB”) at 1; Priv. for Am. at 1–2; TechNet at 2.

⁵⁵ Priv. for Am. at 2–3; Chamber at 6–7; Health Innovation All. (“HIA”) at 1. See also Advanced Med. Tech. Ass’n (“AdvaMed”) at 1 (recommending the Commission adopt a privacy framework pursuant to the advanced notice of proposed rulemaking (R111004) regarding commercial surveillance and data security (87 FR 51273, Aug. 22, 2022)).

⁵⁶ CCIA at 4.

⁵⁷ Am. Telemedicine Ass’n (“ATA Action”) at 1.

⁵⁸ TechNet at 1–2; CTA at 5.

⁵⁹ ANA at 3.

accepting data from health apps, partners such as advertising and analytics firms would risk being covered by the Rule.⁶⁰ According to some commenters, placing such strictures on the advertising and service provider ecosystem would raise prices (by, for example, undermining ad-supported services) and thereby harm competition.⁶¹ One commenter argued that while robust protections for consumer health data are needed, the Rule should not be a vehicle for such protections, because it will result in over-notification of consumers (who have largely learned to disregard breach notices) and be a barrier to legislative change on privacy and data security issues more generally.⁶² Another commenter argued against a breach notification rule altogether, asserting that the Commission should instead focus on requiring robust data security practices to prevent breaches in the first instance.⁶³

Some commenters specifically addressed the proposed changes to the definitions of “PHR identifiable health information” and the new definitions of “health care provider” and “health care services or supplies.” First, a number of comments addressed the scope of “PHR identifiable health information.” Some commenters urged greater breadth, arguing, for example, that the definition of “PHR identifiable health information” should be expanded to include other types of data, such as data *about* an individual—not just data provided by or on behalf of an individual.⁶⁴ Other commenters urged the Commission to state expressly that its definition encompasses particular types of information, such as unique persistent identifiers⁶⁵ or information about sexual health⁶⁶ or substance use or treatment.⁶⁷ By contrast, some commenters urged the Commission to narrow the definition or otherwise clarify its limits, by, for example, exempting data relating to clinical research or trials⁶⁸ or data that has been de-identified.⁶⁹

Relatedly, some commenters urged the Commission to create a definition of or standard for “identifiable data,” “de-identification” or “de-identified

⁶⁰ Priv. for Am. at 3.

⁶¹ E.g., ANA at 3; Priv. for Am. at 1, 3–4.

⁶² World Priv F. (“WPF”) at 4.

⁶³ HIA at 2.

⁶⁴ Consumer Rep.’s at 3.

⁶⁵ *Id.*

⁶⁶ BPC at 1–2; Planned Parenthood at 5.

⁶⁷ Legal Action Ctr. & Opioid Pol’y Inst. at 1–2.

⁶⁸ Soc’y for Clinical Rsch. Sites (“SCRS”) at 1.

⁶⁹ Future of Priv. F. (“FPF”) at 3.

data,”⁷⁰ such as by adopting HHS’s de-identification standard,⁷¹ or by stating that information is identifiable if it is “reasonably linkable to an identified or identifiable individual.”⁷² Commenters argued that clarifying what constitutes “identifiable” data is necessary both because of the increasing ability for de-identified data to be re-identified⁷³ and because the market needs clarity to enable uninhibited flow of de-identified health data for research, public health, and commercial activities.⁷⁴ Indeed, according to one commenter, failure to clarify the standard could complicate or chill public health research and other innovation.⁷⁵ One commenter argued that an objective standard of “reasonable linkability” is better than what the commenter described as the Rule’s knowledge-based standard (*i.e.*, whether the company has a reasonable basis to believe it can be used to identify an individual).⁷⁶ One commenter urged the Commission to issue a new notice of proposed rulemaking on the issue of de-identification alone.⁷⁷

Second, many commenters specifically addressed the Commission’s proposed new definition of “health care provider.” One commenter applauded the Commission’s revised definition of “health care provider,” arguing that taking a crabbed view of that or related terms would lead to further fragmentation of health data, which is already fragmented by HIPAA’s limited purview.⁷⁸ Another commenter noted the Commission’s definition of “health care provider” is simply a logical outgrowth of how consumers interact with health apps: consumers look to health apps to provide health-related services—the quintessential function of a health care provider.⁷⁹

Other commenters, however, raised concerns that the proposed definition of “health care provider” is confusing in its departure from HIPAA’s terminology or is otherwise overbroad.⁸⁰ Some commenters argued this departure from the traditional meaning of the term is

not what Congress intended.⁸¹ A few commenters suggested reducing the confusion with the traditional term by re-naming the definition. These commenters suggested the Commission instead use one of the following terms: “non-HIPAA-regulated health care provider,”⁸² “PHR provider,”⁸³ “Health-related vendor,”⁸⁴ “HIPAA covered entity,”⁸⁵ or “health-related service provider.”⁸⁶ Another commenter recommended eliminating the confusion by stating within the definition that it excludes HIPAA-covered entities and their business associates.⁸⁷ Another commenter urged the Commission to affirm that its definition would have no impact on the term “health care provider” as used in other regulations.⁸⁸

Several comments also expressed concern with the final phrase of the definition of “health care provider” (“any other entity furnishing health care services or supplies”), as overly broad and confusing. Commenters argued its breadth (and the breadth of the accompanying definition of “health care services or supplies”) would have perverse results, turning retailers of tennis shoes, shampoo, or vitamins into entities covered by the Rule, which is not what Congress intended.⁸⁹ Moreover, it would result not only in compliance burdens for companies (with the downstream effect of raising prices for consumers) but also in massive over-notification of consumers, who will become desensitized to the onslaught of notices.⁹⁰

Several commenters urged the Commission to address this problem by dropping the phrase “any other entity furnishing health care services or supplies” entirely—or at least excising the word “supplies”—from the definition of “health care provider.”⁹¹ One commenter recommended replacing the phrase with a different phrase: “any other person or organization who furnishes, bills, or is paid for health care in the normal course of business.”⁹² Another commenter recommended expressly

excluding retailers.⁹³ Commenters requested further clarification of certain terms within the definition of “health care provider,” including the terms “furnishing”⁹⁴ and “health care.”⁹⁵ And another commenter argued a better approach would be to jettison the definitions of “health care provider” and “health care services and supplies” entirely and instead apply the Rule to any entity that “promotes its offering as addressing, improving, tracking or informing matters about a consumer’s health.”⁹⁶

Third, some commenters addressed the proposed definition of “health care services or supplies.”⁹⁷ Several commenters requested more clarity as to what constitutes an “online service,”⁹⁸ as nearly all commercial activities have some online presence.⁹⁹ Several commenters recommended deleting the final phrase of the definition (“or that provides other health-related services or tools”) to limit the definition’s breadth.¹⁰⁰ Conversely, some commenters urged the Commission to reinforce its breadth, by expressly stating that “health care services or supplies” include services related to “wellness”¹⁰¹ or to specific health conditions, such as substance abuse disorder diagnosis, treatment, medication, recurrence of use (“relapse”) and recovery.¹⁰²

3. The Commission Adopts the Proposed Changes To Clarify the Entities Covered

After considering the comments received, the Commission adopts the proposed changes to the Rule (with only non-substantive, organizational improvements noted below) to clarify that the Rule applies to mobile health applications and similar technologies. The Commission agrees with the substantial number of comments, from many different types of entities and individuals, who argued that such clarification is necessary in light of changing technology (*i.e.*, the mass adoption of health apps) and the privacy and data security risks to consumer health data collected by that technology. The Commission also agrees with

⁷⁰ SCRS at 2; Chamber at 7; EPIC at 7–9; FPF at 3–4; LAB at 2; MRO at 4; Network for Pub. Health L. and Texas A&M Univ. (“Network”) at 3.

⁷¹ LAB at 2; Network at 3; SCRS at 2.

⁷² FPF at 3.

⁷³ SCRS at 2.

⁷⁴ FPF at 3; Network at 3–4.

⁷⁵ Network at 3.

⁷⁶ FPF at 3.

⁷⁷ Chamber at 7.

⁷⁸ CDT at 11.

⁷⁹ Confidentiality Coal. at 3–4.

⁸⁰ AAFP at 2–3; AdvaMed at 3–4; AHIP at 2; AMA at 2–3; ATA Action at 1; CARIN Alliance at 2–3; CCA at 3; CTA at 4, 6–9; Datavant at 2; Invitae Corp. (“Invitae”) at 4; NAI at 3–4; Software & Info. Indus. Ass’n (“SIIA”) at 1–2; TechNet at 2; TMA at 2–3; WPF at 7.

⁸¹ ANA at 5; ATA Action at 1; Invitae at 4–5; Priv. for Am. at 4.

⁸² Planned Parenthood at 6.

⁸³ WPF at 7.

⁸⁴ AHIP at 2.

⁸⁵ AMA at 3.

⁸⁶ AHIP at 2.

⁸⁷ Datavant at 2.

⁸⁸ AAFP at 2–3.

⁸⁹ ANA at 7–8; CCIA at 4; CHI at 3–4; CTA at 7–8; SIIA at 2.

⁹⁰ ANA at 3; SIIA at 1.

⁹¹ AdvaMed at 4; CHI at 4; CTA at 9; TechNet at 2.

⁹² AdvaMed at 4.

⁹³ CTA at 8–9.

⁹⁴ EPIC at 2.

⁹⁵ AdvaMed at 3 (urging the Commission to define “health care” and “health care provider” as in 45 CFR 160.103).

⁹⁶ WPF at 10.

⁹⁷ AdvaMed at 3; AAFP at 3; AHIP at 3; Priv. for Am. at 6–7.

⁹⁸ MRO at 2; WPF at 7–8.

⁹⁹ WPF at 8.

¹⁰⁰ NAI at 4.

¹⁰¹ EPIC at 4.

¹⁰² Legal Action Ctr. & Opioid Pol’y Inst. at 3.

commenters who argued that the proposed changes to the Rule are consistent with the Recovery Act, which was intended to bolster breach notifications for consumer health data that falls outside HIPAA. Although the Commission agrees with commenters who argue that consumer health data should enjoy substantial and unfragmented privacy protections, this Rule addresses breach notification, not omnibus privacy protections. While this rulemaking does not address omnibus privacy protections, the Commission observes that companies collecting or holding consumers' sensitive health data should engage in many of the practices commenters described, such as imposing data retention limits, enabling deletion options, and preventing breaches through robust privacy and data security practices.¹⁰³

The Commission is not persuaded that applying the Rule to health apps and similar technologies will have deleterious consequences for individual firms or competition or result in over-notification of consumers. Importantly, the only obligation the Rule imposes is to notify the Commission, consumers, and, in some cases, the media of a breach of unsecured PHR identifiable health information. As noted in the NPRM, many State laws already impose similar, or significantly broader, data breach obligations.¹⁰⁴ Moreover, firms can avoid notification costs entirely by avoiding breaches—by reducing the amount of unsecured PHR identifiable health information they access and maintain (which can be achieved by securing PHR identifiable health information), by de-identifying health information, and by implementing other privacy and data security measures appropriate to the sensitivity of the data. Congress intended for consumers to learn of breaches of their unsecured PHR identifiable health information that fall outside HIPAA; the changes to the Rule help ensure consumers will receive the notification Congress intended.

The Commission carefully considered the arguments commenters raised that the definitional changes depart from the language or spirit of the Recovery Act. The Commission does not agree. The definitions hew closely to the language of the Recovery Act and to the

definitions directly referenced by the Recovery Act in section 1171(6) of the Social Security Act, 42 U.S.C. 1320d(6). As many commenters noted, while health apps did not exist when Congress passed the Recovery Act, they function in a similar manner to the personal health records that existed at the time.

For these reasons, the Commission is adopting the proposed definitions, with minor clarifications. First, the Commission has retained the definition of “PHR identifiable health information” as set out in the NPRM, with non-substantive organizational changes noted below. In response to comments that the definition of “PHR identifiable health information” should be broader, the Commission notes the definition, which closely follows the statutory language, already encompasses most of the categories of data that commenters identified. For example, unique, persistent identifiers (such as unique device and mobile advertising identifiers), when combined with health information, constitute “PHR identifiable health information,” if these identifiers can be used to identify or re-identify an individual. Moreover, “PHR identifiable health information” encompasses information about sexual health and substance abuse disorders, because the information “relates to the past, present, or future physical or mental 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.” The Recovery Act states PHR identifiable health information is information provided “by or on behalf of the individual,” so the Commission declines to change this phrase to “about,” as one commenter suggested.¹⁰⁵ The Commission notes, however, that information provided “by or on behalf of the individual” will encompass much information “about” an individual, as the consumer is the original source of most data; many inferences “about” the individual originate from information provided “by or on behalf of the individual.”

The Commission does not agree with commenters who sought to narrow the definition of PHR identifiable health information out of concern for the Rule's overall breadth. The Commission notes that liability under the Rule does not arise from a single definition. While data used for public health research, for example, may, in some instances, meet the definition of “PHR identifiable health information,” the firm using that data is subject to the Rule only if other

conditions are met (*i.e.*, the firm is an entity covered by the Rule).

The Commission declines to create a new definition of “de-identified data” or another similar term, because the definition of de-identification is already embedded in the second part of the definition of PHR identifiable health information (“that identifies the individual or with respect to which there is a reasonable basis to believe that the information can be used to identify the individual”). Where there is no “reasonable basis to believe that the information can be used to identify the individual,” the information is not identifiable; rather, it is de-identified. If data has been de-identified according to standards set forth by HHS, then there is not a “reasonable basis to believe that the information can be used to identify the individual,” as the definition of PHR identifiable health information requires. Because the Commission's standard is consistent with HHS's, the Commission's Rule poses no impediment to health-related research or other flows of de-identified data. The Commission does not view the existing language as a subjective standard that turns on a company's knowledge, as one commenter suggested; by requiring a “reasonable basis to believe” that the information is not identifiable, the Rule creates an objective standard. Whether such reasonable basis exists will depend on whether the data can reasonably be linked to an individual consumer. There is no need for a supplemental notice of proposed rulemaking on this issue, as the Commission is not changing this aspect of the Rule, which closely follows the statute.¹⁰⁶

Second, the Commission is modifying the proposed definition of “health care provider” to “covered health care provider” to distinguish that term from interpretations of the term “health care provider” in other contexts, which may be more limited in scope. As commenters requested, the Commission affirms its definition of “covered health care provider” is unique to the Rule; it does not bear on the meaning of “health care provider” as used in other regulations enforced by other government agencies. The Commission adopts this change merely to dispel confusion in terminology; the Commission is not making any substantive change from the definition as proposed. The Commission does not need to state expressly, either in this definition or elsewhere, that the Rule's notification requirements do not apply to HIPAA-covered entities and their business associates, as § 318.1 of the

¹⁰³ In the 2009 Final Rule, the Commission similarly underscored the importance of maintaining protections for health information, stating: “In addition, as noted in the NPRM, the Commission expects entities that collect and store unsecured PHR identifiable health information to maintain reasonable security measures, including breach detection measures, which should assist them in discovering breaches in a timely manner.” 74 FR 42971 n.93 (2009).

¹⁰⁴ 88 FR 37832 n.103.

¹⁰⁵ Consumer Rep.'s at 4.

¹⁰⁶ 42 U.S.C. 17937(f)(2).

Rule already includes this proviso. The Commission declines to remove the phrase “any other entity furnishing health care services or supplies” from the definition of “health care provider,” because this phrase is nearly identical to the language that appears in 42 U.S.C. 1320d(3), which is referenced in the definition of individually identifiable health information in 42 U.S.C. 1320d(6), which is in turn referenced in the definition of PHR identifiable health information in section 13407(f)(2) of the Recovery Act, 42 U.S.C. 17937.¹⁰⁷ The Commission declines to define the terms “furnish” and “health care” as the Commission believes the plain meaning of the term “furnish” (to supply someone with something) is already clear and adding a definition of “health care” is unnecessary in light of the definition of “covered health care provider” and “health care services and supplies.” Differences from HHS’s regulations pursuant to HIPAA are appropriate, as the Recovery Act differs from HIPAA, and the Recovery Act’s mandate is specifically to cover entities *not* covered by HIPAA.

Third, the Commission is adopting the proposed definition of “health care services or supplies,” with one minor modification: the Commission has substituted the word “means” for “includes” to avoid implying greater breadth than the Commission intends. The Commission adopts this change merely to dispel confusion about undue breadth; the Commission does not intend any substantive change from the definition proposed. The Commission otherwise affirms the proposed definition without change. The Commission believes the term “online service” in the definition of “health care services or supplies” is sufficiently clear because of the examples of “online services” given within the definition itself: website, mobile application, or internet-connected device. Providing an exhaustive list of what constitutes an online service would prevent the definition from being sufficiently flexible to account for future innovation in types of online services. The Commission also retains the catch-all “or that provides other health-related services or tools” for the same reason: to ensure the Rule’s language can accommodate future changes in technology. There is no undue breadth, because that phrase’s meaning is in the

context of the preceding phrase (“provides mechanisms to track diseases, health conditions, diagnoses or diagnostic testing, treatment, medications, vital signs, symptoms, bodily functions, fitness, fertility, sexual health, sleep, mental health, genetic information, diet”).

In response to some commenters’ concerns that the proposed Rule’s definition of “health care provider” and “health care services or supplies” would impermissibly cause the Rule to cover retailers of general-purpose items like tennis shoes, shampoo, or vitamins, the Commission disagrees this would necessarily be the case. A threshold inquiry under the Rule is whether an entity is a “vendor of personal health records,” which the Recovery Act defines as “an entity . . . that offers or maintains a personal health record.”¹⁰⁸ The Recovery Act usage of the term “vendor of” in connection with “personal health records” underscores that entities that are not in the business of offering or maintaining (*e.g.*, selling, marketing, providing, or promoting) a health-related product or service are not covered—in other words, they are not “vendors” of personal health records. Thus, to be a vendor of personal health records under the Rule, an app, website, or online service must provide an offering that relates more than tangentially to health.¹⁰⁹

The Commission notes a general retailer (one that sells food products, children’s toys, garden supplies, healthcare products (such as pregnancy tests), or apparel (such as maternity clothes)) offering consumers an app to purchase and access purchases of these products—by itself—would not make the retailer a vendor of personal health records. In this scenario, purchase information relating to certain items—such as a pregnancy test or maternity clothes from a retailer—may reveal information about that person’s health. While this purchase information may be PHR identifiable health information, the retailer in this scenario is not a vendor of personal health records because the app is only tangentially related to

health. The Commission notes, however, there may be scenarios where a general-purpose retailer described above may become a vendor of personal health records under the Rule, such as where the retailer offers an app with features or functionalities that are sold, marketed, or promoted as more than tangentially relating to health.

In addition, the Commission reiterates a personal health record must be an electronic record of PHR identifiable health information on an individual, must have the technical capacity to draw information from multiple sources, and must be managed, shared, and controlled by or primarily for the individual. The Commission also notes that purchases of items at a brick and mortar retailer where there is no app, website, or online service to access or track that purchase information electronically is not a personal health record, because there is no electronic record at issue. Contrary to the assertions of some commenters, these definitions do not result in undue breadth, because they do not function in isolation. The Commission provides the following examples to illustrate the interplay of these definitions with the definition of “personal health record”:

- *Example 1:* Health advice app or website A, which is not covered by HIPAA, provides information to consumers about various medical conditions. Its function is purely informational; it does not provide any mechanism through which the consumer may track or record information. Health advice app or website A is not a personal health record, because it is not an electronic record of PHR identifiable health information on an individual.

- *Example 2:* Health advice app or website B, which is not covered by HIPAA, provides information to consumers about various medical conditions and provides a symptom tracker, available to consumers who log into the site with a username and password, in which consumers may input symptoms and receive potential diagnoses. Health advice app or website B is an electronic record of PHR identifiable health information on an individual, because its information is provided by the individual, it identifies the individual (via username and password), it relates to the individual’s health conditions (the symptoms), and is received by a health care provider (*i.e.*, the entity providing the site itself, as that entity is furnishing the health care service of an online service that provides mechanisms to track symptoms). However, health advice app or website B is not a personal health

¹⁰⁸ 42 U.S.C. 17921(18); *see also* 42 U.S.C. 17937.

¹⁰⁹ At least one commenter urged a somewhat similar interpretation, contending that a relevant inquiry in determining whether a service offers a personal health record is “the terms under which a product or service is offered to consumers. If an entity promotes its offering as addressing, improving, tracking, or informing matters about a consumer’s health, then that entity’s offering would be subject to the rule. Thus, any product or services that tracks or addresses physical activity, blood pressure, heart rate, digestion, strength, genetics, sleep, weight, allergies, pain, and similar characteristics would be subject to a PHR rule.” *See* WPF at 10.

¹⁰⁷ The definition of “covered health care provider” in § 318.2 substitutes “entity” for “person”—*i.e.*, “any other *entity* furnishing health care services or supplies”—because the rest of the Rule speaks in terms of “entities,” but the definition in § 318.2 is otherwise identical to the statutory definition in 42 U.S.C. 1320d(3).

record to the extent the site does not have the technical capacity to draw information from multiple sources (*i.e.*, if the consumer is its only source of information).

- *Example 3:* Health advice website C, which is not covered by HIPAA, functions in the same way as health advice app or website B, except that it collects geolocation data via an application programming interface (“API”). For the reasons stated in Example 2, it is an electronic record of PHR identifiable health information on an individual. It also has the technical capacity to draw information from multiple sources (consumer inputs and collection of geolocation data through the API. It is managed primarily for the individual (*i.e.*, to provide the individual health advice). Therefore, health advice app or website C is a personal health record.

- *Example 4:* Health advice app or website D, which is not covered by HIPAA, functions in the same way as health advice app or website B, except that it also draws information from a data broker and connects that information to some of its individual users to provide them with more accurate diagnostic suggestions. For the reasons stated in Example 2, it is an electronic record of PHR identifiable health information on an individual. It also has the technical capacity to draw information from multiple sources (the consumer and the data broker) and is managed by or primarily for the individual. Therefore, health advice app or website D is a personal health record.

Whether a health app or other electronic record constitutes a personal health record (and is therefore subject to the Rule) is a fact-intensive inquiry whose outcome depends not only on the nature of the information contained in that record, but also on numerous other factors, such as its “technical capacity,” its source(s) of information, and its relationship to the individual.

Finally, the Commission notes a non-substantive, organizational change relating to the definition of “PHR identifiable health information.” In the 2023 NPRM, the Commission proposed revising “PHR identifiable health information” by importing language from section 1171(6) of the Social Security Act, 42 U.S.C. 1320d(6), which is referenced directly in section 13407 of the Recovery Act. To hew more closely to the organization of the Recovery Act, and to preserve the word “includes” in the phrase “includes information that is provided by or on behalf of the individual,” the Commission revised slightly the order of

the elements in the definition of “PHR identifiable health information.”

B. Clarification of What It Means for a Personal Health Record To Draw Information From Multiple Sources

1. The Commission’s Proposal Regarding What It Means for a Personal Health Record To Draw Information From Multiple Sources

The Commission proposed amending the definition of the term “personal health record” to clarify what it means for a personal health record to draw information from multiple sources. Under the 2009 Rule, a personal health record is defined as an electronic record of PHR identifiable health information that can be drawn from multiple sources and that is managed, shared, and controlled by or primarily for the individual. Under the Commission’s proposed definition, a “personal health record” would be defined as an electronic record of PHR identifiable health information on an individual that has the technical capacity to draw information from multiple sources and that is managed, shared, and controlled by or primarily for the individual.

Changing the phrase “that can be drawn from multiple sources” to “has the technical capacity to draw information from multiple sources” serves several purposes. First, it clarifies a product is a personal health record if it can draw information from multiple sources, even if the consumer elects to limit information to a single source only, in a particular instance. For example, a depression management app that accepts consumer inputs of mental health states and has the technical capacity to sync with a wearable sleep monitor is a personal health record, even if some customers choose not to sync a sleep monitor with the app. Thus, whether an app qualifies as a personal health record would not depend on the prevalence of consumers’ use of a particular app feature, like sleep monitor-syncing. Instead, the analysis of the Rule’s application would be straightforward: either the app has the technical means (*e.g.*, the application programming interface or API) to draw information from multiple sources, or it does not. Next, adding the phrase “technical capacity to draw information” clarifies a product is a personal health record if it can draw any information from multiple sources, even if it only draws health information from one source. This change further clarifies the Commission’s interpretation of the

Recovery Act, as explained in the Policy Statement.¹¹⁰

The Commission sought public comment as to whether this revised language sufficiently clarifies the Rule’s application to developers and purveyors of products that have the technical capacity to draw information from more than one source. The Commission invited comment on its interpretation that an app is a personal health record because it has the technical capacity to draw information from multiple sources, even if particular users of the app choose not to enable the syncing features. The Commission also requested comment about whether an app (or other product) should be considered a personal health record even if it only draws health information from one place (in addition to non-health information drawn elsewhere); or only draws identifiable health information from one place (in addition to non-identifiable health information drawn elsewhere). The Commission further requested comment about whether the Commission’s bright-line rule (apps with the “technical capacity to draw information” are covered) should be adjusted to take into account consumer use, such as where no consumers (or only a de minimis number) use a feature, and about the likelihood of such scenarios. For example, the Commission offered an example of an app that might have the technical capacity to draw information from multiple sources, but its API is entirely or mostly unused, either because it remains a Beta feature, has not been publicized, or is not popular.

2. Public Comments Regarding What It Means for a Personal Health Record To Draw Information From Multiple Sources

Many commenters supported the Commission’s proposal amending the definition of a “personal health record.”¹¹¹ Commenters noted, for instance, this change would help to ensure that many services that collect PHR identifiable health information are covered by the Commission’s Rule,¹¹² and would help to promote greater privacy and security for health information,¹¹³ while still “hewing to

¹¹⁰ Policy Statement at 2.

¹¹¹ Ella Balasa at 1; TMA at 4 (arguing that “PHRs include applications with the technical capacity to draw information from multiple sources, regardless of the patient’s preference to activate the technical capability.”); Consumer Rep.’s at 6; AAFP at 3; AHIMA at 4–5; AMA at 4; CHIME at 4; CDT at 13; AOA at 3.

¹¹² AHIMA at 4–5.

¹¹³ AAFP at 3.

the limitations of the statute.”¹¹⁴ Some commenters noted without this change, developers of personal health records (such as app developers) might have incentives to design their products in ways that would intentionally skirt the Rule’s requirements (such as by restricting a consumer’s ability to import data from other sources).¹¹⁵ Others noted the importance of the Rule covering apps with the technical capacity to draw information from multiple sources even where such capacity is not used by the consumer.¹¹⁶

Other commenters opposed this proposal.¹¹⁷ Some argued the proposed clarification regarding what drawing information from multiple sources means runs counter to Congress’s statutory intent,¹¹⁸ because virtually every app has some sort of integration (e.g., for analytics) through which it draws information other than from the consumer.¹¹⁹ One commenter asserted the change would broaden the scope of the Rule to the point that it would sweep in online services that should not be thought of as a personal health record (such as email apps),¹²⁰ or otherwise create confusing standards for app developers or reduce innovation.¹²¹ In addition, commenters expressed concern this change would sweep in apps or online services that have the technical capacity to draw from multiple sources during the development or testing phase of the product, or would sweep in products with unused, unavailable, or unpublicized APIs or integrations that count as a source.¹²² One commenter

expressed concern about lack of clarity, such as in scenarios where a user is required to pay for an upgrade to access a feature or integration that draws information from another source.¹²³ Some commenters also expressed concern that apps and online services that are subject to HIPAA (i.e., HIPAA-covered entities or business associates) should be carved out of the definition of a personal health record.¹²⁴ Other commenters expressed broader concern with the definition of “personal health record,” urging the Commission to, for example, abandon the purportedly outdated term in favor of a more modern one.¹²⁵ For instance, some commenters urged that the Commission abandon or tweak the requirement that the personal health record be “managed, shared, and controlled by or primarily for the individual.”¹²⁶

Another commenter expressed concern the proposed change could sweep in services that draw any information from multiple sources, regardless of whether that information is identifiable health information.¹²⁷

3. The Commission Adopts the Proposed Changes Clarifying What It Means for a Personal Health Record To Draw Information From Multiple Sources

After considering the comments received, the Commission adopts the proposed amendment without change. This amendment will help clarify the types of entities covered by the Rule. The definition does not create undue breadth or deviate from Congressional intent; rather, the changes are consistent with the language of the Recovery Act, and only serve to give meaning to the phrase “can be drawn” in the Recovery Act in a way that is consistent with the current state of technology. They are also necessary to keep pace with technological change, which has enabled firms to offer consumers mobile electronic records of their health information that contain numerous integrations. To illustrate the intended meaning of the proposed revisions to

health record.” (emphasis in original); CTA at 11 (arguing Rule should instead have bright-line test that assesses whether the app actually draws health information from multiple sources); AdvaMed at 5 (arguing the Commission should decline to adopt multiple sources changes because it could cause confusion and potentially sweep in apps or services with features that have not been made available to consumers, such as APIs connected to the PHR that have not been publicized).

¹²³ WPF at 9.

¹²⁴ Omada at 5; Datavant at 3.

¹²⁵ HIMSS at 3 (urging the Commission to work with Congress to craft a definition more consonant with technological realities).

¹²⁶ AHIOS at 4; MRO at 4.

¹²⁷ NAI at 6.

the term “personal health record,” the Commission reiterates examples from the 2023 NPRM of two non-HIPAA covered diet and fitness apps available for consumer download in an app store. Under the amended Rule, each is a personal health record.

- *Example 1:* Diet and fitness app Y allows users to sync their app with third-party wearable fitness trackers. Diet and fitness app Y has the technical capacity to draw identifiable health information both from the user (e.g., name, weight, height, age) and the fitness tracker (e.g., user’s name, miles run, heart rate), even if some users elect not to connect the fitness tracker.

- *Example 2:* Diet and fitness app Y has the ability to pull information from the user’s phone calendar via the calendar API to suggest personalized healthy eating options. Diet and fitness app Y has the technical capacity to draw identifiable health information from the user (e.g., name, weight, height, age) and non-health information (e.g., calendar entry info, location, and time zone) from the user’s calendar.

As these examples make clear, and in response to one commenter’s concern that the changes would sweep in services that do not draw any health information,¹²⁸ the Commission notes the Rule still requires drawing PHR identifiable health information from at least one source to count as a personal health record.

The Commission declines to make other requested changes to the definition of personal health record. First, the Commission declines to include an express exemption for HIPAA-covered entities within the definition of personal health record because § 318.1 of the Rule already specifically exempts businesses or organizations covered by HIPAA.¹²⁹ Second, the Commission declines to exempt apps and services where there are available but unused or unpublicized APIs or integrations. Similarly, the Commission declines to exempt apps and services from the definition just because they are drawing information from multiple sources while undergoing product or beta testing and are not yet in their final form.¹³⁰ The Commission notes a product feature or integration that exists

¹²⁸ NAI at 6.

¹²⁹ See, e.g., 16 CFR 318.1(a) (Rule “does not apply to HIPAA-covered entities, or to any other entity to the extent that it engages in activities as a business associate of a HIPAA-covered entity.”); see also 16 CFR 318.2 (exempting business associates and HIPAA-covered entities from the Rule’s definitions of “PHR related entity” and “vendor of personal health records.”).

¹³⁰ ACLA at 1–2; CTA at 11; AdvaMed at 5.

¹¹⁴ Consumer Reports at 5–6.

¹¹⁵ AHIP at 2–3; CDT at 13 (arguing that changes remove “incentives for companies to technically design products and services to not trigger the HBNR to avoid any need to provide consumer notice.”).

¹¹⁶ AHIOS at 4; CARIN Alliance at 4.

¹¹⁷ NAI at 6 (urging that the Commission make clear that a personal health record is one that “not only has the technical capacity to draw PHR identifiable health information from multiple sources, but that it also has the functionality and actually does incorporate data from multiple sources.”); ANA at 7; ACLA at 1–2.

¹¹⁸ NAI at 6.

¹¹⁹ Chamber at 4–5; Priv. for Am. at 5–6; NAI at 6.

¹²⁰ CCIA at 6.

¹²¹ CTA at 11; AdvaMed at 5; CHI at 5.

¹²² CHI at 5 (asking the Commission to clarify that an “app having the ability to draw from multiple sources with some changes to the app’s coding/APIs is not within this definition’s threshold.”); ACLA at 1 (arguing “[i]f a feature is unused by individuals ‘because it remains a Beta feature,’ then in fact it does not have the ‘technical capacity’ to draw an individual’s information from other sources, unless and until its functionality has been enabled by the vendor. The mere possibility that an application vendor *might* sometime in the future enable that functionality should not bring the electronic record within the scope of the definition of ‘personal

and that is able to draw PHR identifiable health information counts as a source under the Rule. Exempting such instances would be contrary to the purpose of the Rule and would impermissibly limit notification of breaches just because a product feature is not widely disseminated, used, or in its final form. The Commission notes under the Rule, a covered entity that experienced a breach of security of unsecured PHR identifiable health information triggering the Rule would not be exempt because the breach occurred in the context of such scenarios.

Further, and importantly, the Rule is triggered only by breaches of unsecured PHR identifiable health information and does not apply to information that is protected or “secured” through the use of a technology or methodology specified by the Secretary of Health and Human Services in the guidance issued under section 13402(h)(2) of the American Reinvestment and Recovery Act of 2009, 42 U.S.C. 17932(h)(2).¹³¹ The Rule, therefore, creates appropriate incentives for product testing with de-identified data or that secures information through certain specifications, such as through specified encryption methods.

Third, the Commission declines, as one commenter requested,¹³² to expressly exempt scenarios where a change is required to an app’s coding to draw information from another source. The Commission notes, however, it does not intend to cover instances where an app can draw from multiple sources only through changes to the design or underlying software code and where the app developer does not implement those changes.

¹³¹ Per HHS guidance, electronic health information is “secured” if it has been encrypted according to certain specifications set forth by HHS, or if the media on which electronic health information has been stored or recorded is destroyed according to HHS specifications. See 74 FR 19006; see also U.S. Dep’t of Health & Human Servs., *Guidance to Render Unsecured Protected Health Information Unusable, Unreadable, or Indecipherable to Unauthorized Individuals* (July 26, 2013), <https://www.hhs.gov/hipaa/for-professionals/breach-notification/guidance/index.html>. PHR identifiable health information would be considered “secured” if such information is disclosed by, for example, a vendor of personal health records, to a PHR related entity or a third party service provider, in an encrypted format meeting HHS specifications, and the PHR related entity or third party service provider stores the data in an encrypted format that meets HHS specifications and also stores the encryption and/or decryption tools on a device or at a location separate from the data.

¹³² CHI at 5 (asking the Commission to clarify that an “app having the ability to draw from multiple sources with some changes to the app’s coding/APIs is not within this definition’s threshold.”).

In addition, the Commission declines to remove from the definition of personal health record the requirement that it be “managed, shared, and controlled by or primarily for the individual.” This language mirrors the Recovery Act’s statutory definition of personal health record.¹³³ Further, this language provides a boundary to the definition. Even if a website or app has the technical capacity to draw information from multiple sources (for example, because it has integrations for advertising or analytics), it must still be “managed, shared, and controlled by or primarily for the individual” to be covered by the Rule.

Generally, a personal health record is an electronic record of an individual’s health information by which the individual maintains access to the information and may have, for example, the ability to manage, track, control, or participate in his or her own health care. If these elements are not present, the website or app may not be “managed, shared, and controlled by or primarily for the individual,” and would not, therefore, constitute a personal health record.

C. Clarification Regarding Types of Breaches Subject to the Rule

1. The Commission’s Proposals

a. The Commission’s Proposal Regarding “Breach of Security”

The Commission proposed a definitional change to clarify that a breach of security under the Rule encompasses unauthorized acquisitions that occur as a result of a data breach or an unauthorized disclosure. The Commission’s proposal underscores that a breach of security is not limited to data exfiltration, and includes unauthorized disclosures (such as, but not limited to, a company’s unauthorized sharing or selling of consumers’ information to third parties that is inconsistent with the company’s representations to consumers). The Rule previously defined “breach of security” as the acquisition of unsecured PHR identifiable health information of an individual in a personal health record without the authorization of the individual, which language mirrored the definition of “breach of security” in section 13407(f)(1) of the Recovery Act.

Accordingly, consistent with the Recovery Act definition, the Policy Statement, FTC enforcement actions under the Rule, and public comments received, the Commission proposed amending the definition of “breach of security” in § 318.2 by adding the

¹³³ 42 U.S.C. 17921(11).

following sentence to the end of the existing definition: “[a] breach of security includes an unauthorized acquisition of unsecured PHR identifiable health information in a personal health record that occurs as a result of a data breach or an unauthorized disclosure.” The change was intended to make clear to the marketplace that a breach includes an unauthorized acquisition of identifiable health information that occurs as a result of a data breach or an unauthorized disclosure, such as a voluntary disclosure made by the PHR vendor or PHR related entity where such disclosure was not authorized by the consumer.

The NPRM, like the 2009 Rule, continued to include a rebuttable presumption for unauthorized access to an individual’s data; it stated when there is unauthorized access to data, unauthorized acquisition will be presumed unless the entity that experienced the breach “has reliable evidence showing that there has not been, or could not reasonably have been, unauthorized acquisition of such information.”

b. The Commission’s Related Proposal To Not Define the Term “Authorization” in the Rule

In the 2023 NPRM, the Commission stated it had considered defining the term “authorization,” which appears in § 318.2’s definition of “breach of security,” but did not propose any such change in the NPRM.

The Commission considered defining “authorization” to mean the affirmative express consent of the individual and then defining “affirmative express consent” consistent with State laws that define consent, such as the California Consumer Privacy Rights Act, Cal. Civ. Code 1798.140(h).¹³⁴ Such changes would have ensured notification is required anytime there is acquisition of

¹³⁴ As noted in the 2023 NPRM, the Commission considered defining “affirmative express consent” as any freely given, specific, informed, and unambiguous indication of an individual’s wishes demonstrating agreement by the individual, such as by a clear affirmative action, following a clear and conspicuous disclosure to the individual, apart from any “privacy policy,” “terms of service,” “terms of use,” or other similar document, of all information material to the provision of consent. Acceptance of a general or broad terms of use or similar document that contains descriptions of agreement by the individual along with other, unrelated information, does not constitute affirmative express consent. Hovering over, muting, pausing, or closing a given piece of content does not constitute affirmative consent. Likewise, agreement obtained through use of user interface designed or manipulated with the substantial effect of subverting or impairing user autonomy, decision-making, or choice, does not constitute affirmative express consent. See 88 FR 37830 n.78.

unsecured PHR identifiable health information without the individual's affirmative express consent for that acquisition—such as when an app discloses unsecured PHR identifiable health information to another company, having obtained nominal “consent” from the individual by using a small, greyed-out, pre-selected checkbox following a page of dense legalese.

The Commission did not, however, propose to define “authorization” because (1) the 2009 Rule Commentary already provided guidance on the types of disclosures the Commission considers to be “unauthorized”;¹³⁵ (2) recent Commission orders, such as the Commission's enforcement actions against GoodRx and Easy Healthcare,¹³⁶ also make clear that the use of “dark patterns,” which have the effect of manipulating or deceiving consumers, including through use of user interfaces designed with the substantial effect of subverting or impairing user autonomy and decision-making, do not satisfy the standard of “meaningful choice”; and (3) Commission settlements establish important guidelines involving authorization (the Commission's recent settlement with GoodRx, alleging violations of the Rule, highlights that disclosures of PHR identifiable health information inconsistent with a company's privacy promises constitute an unauthorized disclosure).

The Commission sought public comment about:

- Whether the commentary above and FTC enforcement actions under the Rule provide sufficient guidance to put companies on notice about their obligations for obtaining consumer authorization for disclosures, or whether defining the term “authorization” would better inform companies of their compliance obligations.

- To the extent that including such definitions would be appropriate, the definitions of “authorization” and “affirmative express consent,” as described above, and the extent to which such definitions are consistent with the language and purpose of the Recovery Act.

- What constitutes an acceptable method of authorization, particularly

when unauthorized sharing is occurring.¹³⁷

- Whether there are certain types of sharing for which authorization by consumers is implied because such sharing is expected and/or necessary to provide a service to consumers.

2. Public Comments

a. Public Comments Regarding “Breach of Security”

Many commenters supported the Commission's proposed amendment to the definition of “breach of security.”¹³⁸ One commenter noted the change is consistent with the broad definition of “breach of security” in the Recovery Act, which refers explicitly to the acquisition of PHR identifiable health information without the authorization of an individual (rather than the authorization of an entity holding the data, as is the case where a breach involves data theft or exfiltration).¹³⁹ Commenters also noted the amendment would ensure notice, accountability, and regulatory oversight, regardless of the underlying cause of the unauthorized acquisition.¹⁴⁰ Commenters noted that breaches encompass more than just cybersecurity intrusions.¹⁴¹ Commenters also argued that a company's voluntary unauthorized disclosure can be just as damaging as data theft.¹⁴² For instance, a commenter noted that unauthorized disclosures of health information may cause embarrassment, perpetuate stigma about patients' conditions, deter patients from seeking care, interfere in the patient-physician relationship, or impact patients' employment.¹⁴³ Moreover, voluntary, unauthorized disclosures increase the risk of additional unauthorized acquisition and

sharing of this information among bad actors.¹⁴⁴

Some commenters supported expanding or changing the definition further. Specifically, some commenters urged the Commission to amend the definition to encompass (1) exceeding authorized access or use of PHR identifiable health information, such as where a company collects data for one purpose, but later uses or discloses that data for a second, undisclosed purpose;¹⁴⁵ or (2) the collection or retention of PHR identifiable health information beyond what is necessary to provide the associated service to an individual consumer.¹⁴⁶ One commenter asked the Commission to clarify that the Rule would be triggered by unauthorized use of or access to information derived from PHR identifiable health information, and to define the phrase acquisition.¹⁴⁷

Some commenters, however, urged the Commission to not amend the definition at all. These commenters expressed concern the amendment would cause the Rule to exceed what Congress intended in the Recovery Act and transform the Rule into an opt-in notice and consent privacy regime.¹⁴⁸ Commenters argued further the proposed changes would cause consumer notice fatigue,¹⁴⁹ consumer panic,¹⁵⁰ or over-reporting by companies.¹⁵¹ One commenter urged the Commission to limit the definition of “acquisition” to actual acquisition, and exclude instances of access or disclosure where the information was not actually acquired by a third party.¹⁵² Commenters argued the proposed definition would be burdensome and force companies to limit certain beneficial disclosures to certain third parties, such as disclosures to support internal operations, detect security vulnerabilities or fraud, for law enforcement, and other purposes.¹⁵³

Some commenters also urged that the Commission adopt carve-outs so that certain conduct would not be deemed breaches of security under the Rule. Commenters requested exemptions consistent with or found in HIPAA or

¹³⁷ For example, the Commission sought comment about when a vendor of personal health records or a PHR-related entity is sharing information covered by the Rule, is it acceptable for that entity to obtain the individual's authorization to share that information when an individual clicks “agree” or “accept” in connection with a pre-checked box disclosing such sharing? Is it sufficient if an individual agrees to terms and conditions disclosing such sharing but that individual is not required to review the terms and conditions? Or is it sufficient if an individual uses a health app that discloses in its privacy policy that such sharing occurs, but the app knows via technical means that the individual never interacts with the privacy policy? See 88 FR 37832.

¹³⁸ See, e.g., TMA at 3; U.S. PIRG at 2–3; AAFP at 3; AHIMA at 3; AMA at 3–4; AMIA at 3; AOA at 2–3; AHIOS at 3; CDT at 11–12; CHIME at 4; EPIC at 5–6.

¹³⁹ Consumer Rep.'s at 4.

¹⁴⁰ CDT at 11–12; U.S. PIRG at 2–3.

¹⁴¹ AMA at 4; CDT at 11–12; EPIC at 5.

¹⁴² AAFP at 3; CDT at 11–12.

¹⁴³ AOA at 2.

¹⁴⁴ AHIMA at 3.

¹⁴⁵ PPF at 12–15.

¹⁴⁶ EPIC at 5–7; U.S. PIRG at 2–3.

¹⁴⁷ Mozilla at 6–7.

¹⁴⁸ Chamber at 6; Priv. for Am. at 2–5; ANA at 6–7.

¹⁴⁹ SIIA at 3; CTA at 13–14.

¹⁵⁰ CCIA at 4–5, 7 (arguing that requiring notification for unauthorized disclosures could cause consumers to worry in the absence of harm, such as where it is “typical” to disclose such information.)

¹⁵¹ CTA at 13–14.

¹⁵² *Id.* at 14–16.

¹⁵³ TechNet at 3; Chamber at 7; CCIA at 5–6.

¹³⁵ See, e.g., 74 FR 42967.

¹³⁶ *United States v. GoodRx Holdings, Inc.*, No. 23–cv–460 (N.D. Cal. 2023), <https://www.ftc.gov/legal-library/browse/cases-proceedings/2023090-goodrx-holdings-inc>; *United States v. Easy Healthcare Corp.*, No. 1:23–cv–3107 (N.D. Ill. 2023), <https://www.ftc.gov/legal-library/browse/cases-proceedings/202-3186-easy-healthcare-corporation-us-v>.

under State breach notification laws, such as exemptions for disclosures to certain types of entities or for certain purposes, or where there is inadvertent or unintentional access, use, or disclosure.¹⁵⁴ Commenters also proposed safe harbors for companies that implement recognized security or privacy safeguards;¹⁵⁵ and one commenter proposed safe harbors that would apply where data is shared with “affiliated businesses,” where there is inadvertent but “good-faith” access by a company employee, where a company makes good faith efforts to inform consumers of disclosures to third parties, and where companies take steps to contractually limit downstream uses of the data.¹⁵⁶ Other commenters expressed support for exempting disclosures of PHR identifiable health information to public health authorities for public health purposes, noting the amended definition could discourage such disclosures.¹⁵⁷

b. Public Comments Regarding Defining “Authorization”

Commenters were divided as to whether the Commission should define “authorization.” Some commenters supported defining “authorization” to provide greater guidance to companies, to promote transparency, and to discourage buried or inconspicuous disclosures relating to health information, or approaches to consent that are not meaningful because they are

¹⁵⁴ CHI at 4 (stating the FTC “should explicitly except the same situations from disclosure that are excepted from HIPAA disclosures, and/or try to align exceptions with those found in State privacy statutes.”); CTA at 16; HIA at 2; TechNet at 3 (arguing the Rule should adopt exemptions that encompass “actions taken to prevent and detect security incidents, to comply with a civil, criminal, or regulatory inquiry or investigation, to cooperate with law enforcement agencies concerning conduct or activity that the data controller reasonably and in good faith believes may be illegal, to perform internal operations consistent with a consumer’s expectations, and to provide a product or service that a consumer requested.”); CCIA at 5–6 (arguing the Rule should exempt disclosures relating to a host of purposes, including: preventing and detecting security incidents and fraud, complying with legal process, cooperating with law enforcement, performing internal operations consistent with consumer expectations, providing a service requested by the consumer, protecting “the vital interests of the consumer,” or processing data relating to public health); Chamber at 7 (arguing if the Commission does amend the definition of breach of security, it “should provide exceptions for legitimate and societally beneficial uses of data that other privacy laws have for failure to honor opt-in including but not limited to network security, prevention and detection of fraud, protection of health, network maintenance, and service/product improvement.”); LAB at 2.

¹⁵⁵ DirectTrust at 1–2.

¹⁵⁶ ATA Action at 2.

¹⁵⁷ Network for Pub. Health L. and Texas A&M Univ. at 1–2.

confusing or coercive.¹⁵⁸ To further regulatory consistency, some commenters supported adding a definition of “authorization” that is consistent with how that term is defined in other health-related laws, such as under HIPAA¹⁵⁹ or State health privacy laws that define consent or authorization (such as the California Consumer Privacy Rights Act¹⁶⁰ or the Washington My Health, My Data Act).¹⁶¹

By contrast, some commenters opposed defining the term—or opposed a requirement under the Rule that entities be required to get authorization before disclosing PHR identifiable health information.¹⁶² Commenters argued that Congress had not granted the Commission the authority to define “authorization” in the Recovery Act,¹⁶³ or that doing so would import a substantive consent requirement that is outside the scope of the Rule, converting a breach notice Rule into an opt-in privacy regime.¹⁶⁴ Other commenters noted that requiring a specifically defined authorization would create an inflexible standard that would not evolve with changes in technology.¹⁶⁵ Other commenters opposed a requirement that consumers should be required to review terms before agreeing to use a service, contending that this would not increase consumer understanding of terms.¹⁶⁶

Some commenters endorsed other approaches that would exempt from any requirement of affirmative express consent certain types of disclosures of

¹⁵⁸ AHIP at 4; Light Collective at 4; MRO at 2–3; Mozilla at 4; CARIN Alliance at 10; Consumer Rep.’s at 9; *see also* PharmedOut at 3 (arguing that defining “authorization” is crucial but urging the Commission go further and place substantive restrictions on what companies can do with consumer health data.).

¹⁵⁹ AdvaMed at 7 (arguing that any definition of “authorization” or “affirmative express consent” should take into account the necessity for medical technologies and medical technology companies to be able to operate and communicate under standards consistent with those governing HIPAA covered entities and others in the health care ecosystem. These standards permit certain uses and disclosures of individually identifiable health information without express consent where necessary for the provision of timely and effective health care); MRO at 3; AHIMA at 7–8.

¹⁶⁰ AHIOS at 3.

¹⁶¹ Consumer Rep.’s at 9.

¹⁶² HIA at 2 (arguing that “[r]outine disclosures of data should be allowed in certain contexts without additional need for authorizations”); CTA at 16–17; AdvaMed at 7–8; ACLA at 6; Confidentiality Coal. at 4–5.

¹⁶³ Confidentiality Coal. at 4–5.

¹⁶⁴ CTA at 16–17 (arguing that the Rule does not allow the Commission to impose “substantive consent requirements” that would be burdensome and “likely not administrable for many companies.”).

¹⁶⁵ SIIA at 4.

¹⁶⁶ CHI at 7.

PHR identifiable health information, such as to service providers, data processors, and entities that assist with combatting fraud and promoting safety.¹⁶⁷ Some commenters urged a disclosure be deemed authorized if the disclosure is consistent with a company’s privacy notices or policies or where applicable State privacy laws require affirmative consent or provide for the right to opt-out, without the need to define affirmative express consent under the Rule.¹⁶⁸ One commenter argued that authorization should be met when a consumer agrees to opt-in to certain data sharing, such as by clicking a box proximate to a disclosure of material terms.¹⁶⁹

3. The Commission Adopts the Proposed Changes to the Definition of “Breach of Security”

After carefully considering the public comments, the Commission adopts the proposed amendment without change. The final rule definition is consistent with the statutory definition in the Recovery Act, the Policy Statement,¹⁷⁰ and recent Commission enforcement actions under the Rule. The Commission notes the statutory definition in the Recovery Act is sufficiently broad to cover both cybersecurity intrusions as well as a company’s intentional but unauthorized disclosures of consumers’ PHR identifiable health information to third party companies. In addition, the Commission finds persuasive the comment noting the Recovery Act’s definition of “breach of security” refers to the acquisition PHR identifiable health information without the authorization of an individual, rather than the authorization of the entity holding the data.¹⁷¹ The definition is

¹⁶⁷ PPF at 10 (arguing that “an organization may share information with a service provider operating on their behalf to provide storage; may share information to protect the safety or vital interests of an individual or react to a public health emergency; or to protect themselves against security incidents and fraud. In each of these situations, data protection laws typically invoke a variety of non-consent measures, including data minimization, transparency, notice to the end-user or the regulator, and opportunities to object.”); Chamber at 7.

¹⁶⁸ Confidentiality Coal. at 4–5; SIIA at 4; CHI at 7.

¹⁶⁹ CTA at 17.

¹⁷⁰ The Commission’s Policy Statement makes clear that “[i]ncidents of unauthorized access, including sharing of covered information without an individual’s authorization, triggers notification obligations under the Rule,” and that a breach “is not limited to cybersecurity intrusions or nefarious behavior.” Policy Statement at 2.

¹⁷¹ Consumer Rep.’s at 5 (noting “the Recovery Act frames breaches of security in relation to individuals, rather than to vendors of personal health records or PHR related entities,” and defines

also consistent with public comments received by the Commission in 2020 (when the Commission announced its regular, ten-year review of the Rule and requested public comments about potential Rule changes¹⁷²), which urged the Commission to clarify what constitutes an unauthorized acquisition under the Rule.¹⁷³ Importantly, the amendment to the definition of “breach of security” in § 318.2 does not depart from the 2009 Rule Commentary or the Commission’s enforcement policy under the Rule. Instead, it further underscores the 2009 Rule Commentary and subsequent Commission enforcement actions that unauthorized disclosures (*i.e.*, sharing inconsistent with consumer expectations) can be a “breach of security” that triggers the Rule.¹⁷⁴

The Commission declines to adopt any specific exemptions or safe harbors to the definition of breach of security. Unlike the section of the Recovery Act that governs breach notifications under HIPAA,¹⁷⁵ Congress did not provide for

breach of security as “acquisition of such information without the authorization of the individual.”)

¹⁷² 85 FR 31085 (May 22, 2020).

¹⁷³ See Public Comments in response to May 2020 Request for Public Comments in connection with regular, ten-year review of Rule: AMA at 5–6 (“The FTC should define ‘unauthorized access’ as presumed when entities fail to disclose to individuals how they access, use, process, and disclose their data and for how long data are retained. Specifically, an entity should disclose to individuals exactly what data elements it is collecting and the purpose for their collection”; “[T]he FTC should define ‘unauthorized access’ as presumed when an entity fails to disclose to an individual the specific secondary recipients of the individual’s data.”); AMIA at 2 (recommending the FTC “[e]xpand on the concept of ‘unauthorized access’ under the definition of ‘Breach of security,’ to be presumed when a PHR or PHR related entity fails to adequately disclose to individuals how user data is accessed, processed, used, reused, and disclosed.”); OAG–CA at 5–6 (urging the FTC to include “impermissible acquisition, access, use, disclosure” under the definition of breach.). These comments can be found at <https://www.regulations.gov/docket/FTC-2020-0045>.

¹⁷⁴ The 2009 Rule Commentary noted other examples illustrating that unauthorized sharing or transferring of information constitutes a breach of security, including that the unauthorized downloading or transfer of information by an employee can constitute a breach of security; that inadvertent access by an unauthorized employee reading or sharing information triggers the Rule’s notification obligations; and notes that given the highly personal nature of health information, “the Commission believes that consumers would want to know if such information was read or shared without authorization.” See 74 FR 42966–67.

¹⁷⁵ 42 U.S.C. 17921; see also U.S. Dep’t of Health & Human Servs., *Breach Notification* (July 26, 2013), <https://www.hhs.gov/hipaa/for-professionals/breach-notification/index.html>. Under the Recovery Act’s definition of “breach of security” for the Rule governing HIPAA-covered entities and business associates, the statute explicitly provides for three exceptions: (1) unintentional acquisition, access, or use of

any specific, enumerated exemptions for breaches under the Commission’s Rule. Moreover, the Commission’s Rule provides for a rebuttable presumption for certain types of access: when there is unauthorized access to data, unauthorized acquisition will be presumed unless the entity that experienced the breach “has reliable evidence showing that there has not been, or could not reasonably have been, unauthorized acquisition of such information.” That is, companies can rebut the presumption of acquisition in instances of unauthorized access by providing reliable evidence disproving acquisition. The Commission has previously offered guidance on what counts as unauthorized access and reiterates that guidance here.¹⁷⁶

protected health information by a workforce member or person acting under the authority of a covered entity or business associate, if such acquisition, access, or use was made in good faith and within the scope of authority; (2) the inadvertent disclosure of protected health information by a person authorized to access protected health information at a covered entity or business associate to another person authorized to access protected health information at the covered entity or business associate, or organized health care arrangement in which the covered entity participates; and (3) if the covered entity or business associate has a good faith belief that the unauthorized person to whom the impermissible disclosure was made, would not have been able to retain the information. See 45 CFR 164.400 through 164.414. In the first two cases, the information cannot be further used or disclosed in a manner not permitted by the Privacy Rule. These exceptions are not found in the provisions of the Recovery Act authorizing the FTC’s Health Breach Notification Rule; this makes sense, given there is no analogous Privacy Rule, Security Rule, or required Business Associate agreements outside the HIPAA sphere governing entities covered by the FTC’s Health Breach Notification Rule.

¹⁷⁶ The Rule continues to provide that, when there is unauthorized access to data, unauthorized acquisition will be presumed unless the entity that experienced the breach “has reliable evidence showing that there has not been, or could not reasonably have been, unauthorized acquisition of such information.” As noted in the 2009 Rule Commentary, the presumption was intended to address the difficulty of determining whether access to data (*i.e.*, the opportunity to view the data) did or did not lead to acquisition (*i.e.*, the actual viewing or reading of the data). In these situations, the Commission stated that the entity that experienced the breach is in the best position to determine whether unauthorized acquisition has taken place. In describing the rebuttable presumption, the Commission provided several examples. It noted that no breach of security has occurred if an unauthorized employee inadvertently accesses an individual’s PHR and logs off without reading, using, or disclosing anything. If the unauthorized employee read the data and/or shared it, however, he or she “acquired” the information, thus triggering the notification obligation in the Rule. Similarly, the Commission provided an example of a lost laptop: If an entity’s employee loses a laptop in a public place, the information would be accessible to unauthorized persons, giving rise to a presumption that unauthorized acquisition has occurred. The entity can rebut this presumption by showing, for example, that the laptop was recovered, and that forensic analysis revealed that

4. The Commission Affirms Its Proposal Not To Define “Authorization”

After carefully considering the public comments, the Commission declines to define “authorization,” as that term appears in § 318.2’s definition of “breach of security.” The Commission finds persuasive the public comments suggesting that imposing an affirmative express consent requirement would not be appropriate or warranted in all cases.

The Commission believes whether a disclosure is authorized under the Rule is a fact-specific inquiry that will depend on the context of the interactions between the consumer and the company; the nature, recipients, and purposes of those disclosures; the company’s representations to consumers; and other applicable laws. The Commission reiterates the 2009 Rule Commentary, which states a use of data is “authorized” only where it is consistent with a company’s disclosures and consumers’ reasonable expectations and where there is meaningful choice in consenting to sharing—buried disclosures do not suffice.¹⁷⁷

The Commission’s recent enforcement actions alleging violations of the Rule against GoodRx and Easy Healthcare further highlight that disclosures of PHR identifiable health information inconsistent with a company’s privacy promises constitute an unauthorized disclosure. These recent Commission orders also make clear that the use of “dark patterns,” which have the effect of manipulating or deceiving consumers, including through use of user interfaces designed with the substantial effect of subverting or impairing user autonomy and decision-making, undercut an entity’s assertion that consumers exercised “meaningful choice.”

In response to public comments seeking more guidance on what constitutes an unauthorized disclosure under the Rule,¹⁷⁸ the Commission

files were never opened, altered, transferred, or otherwise compromised. See 74 FR 42966.

¹⁷⁷ The 2009 Rule Commentary states: “[g]iven the highly personal nature of health information, the Commission believes that consumers would want to know if such information was read or shared without authorization.” It further states that data sharing to enhance consumers’ experience with a PHR is authorized only “as long as such use is consistent with the entity’s disclosures and individuals’ reasonable expectations” and that “[b]eyond such uses, the Commission expects that vendors of personal health records and PHR related entities would limit the sharing of consumers’ information, unless the consumers exercise meaningful choice in consenting to such sharing. Buried disclosures in lengthy privacy policies do not satisfy the standard of ‘meaningful choice.’” 74 FR 42967.

¹⁷⁸ TechNet at 4; Tranquil Data at 4.

offers the following, non-exhaustive examples relating to authorization:

- *Example 1—Unauthorized Disclosure (Affirmative Misrepresentation)*: A medication app offers a personal health record (not covered by HIPAA) which allows users to track information about their prescription medication history, such as prescription names, dosages, pharmacy and refill information, and the user's health conditions. The app voluntarily discloses PHR identifiable health information to third party companies for advertising and advertising-related analytics, in violation of the app's privacy representations to its users. The third parties that receive the PHR identifiable health information are able to use the information for their own business purposes, such as to improve the third party's own products and services, to infer information about consumers, or to compile profiles about consumers to use for targeted advertising. These disclosures are not authorized under the Rule because they are inconsistent with consumer expectations—the disclosures violate the app's privacy representations, and consumers would also not expect their PHR identifiable health information (which they input into the app to track their medications and health conditions) would be disclosed to, and used by, third party companies that use the data for their own economic benefit.

- By contrast, disclosures of PHR identifiable health information by the app in Example 1 would be authorized if made to service providers in the following circumstances: (1) the service providers assist with functions that are necessary to the operation and functioning of the medication app, or with services the consumer requested; (2) the service providers are contractually prohibited from using, sharing, or disclosing the PHR identifiable health information for any purpose beyond providing services to the medication app; and (3) the medication app's privacy notice clearly and conspicuously discloses the specific purposes for which it shares users' PHR identifiable health information with these service providers. Such authorized disclosures could include those to cloud storage providers that host user data in the health record in a secure fashion; payment processors who process user payments to the app; vendors that facilitate refill reminders or other communications from the app developer that directly relate to the provision of the personal health record or services the consumer requested; analytics providers that assist with tracking analytics relating to the app's

functionality;¹⁷⁹ or companies that help to detect, prevent, or mitigate fraud or security vulnerabilities. Such disclosures are authorized because they are consistent with consumer expectations. Importantly, this sharing is disclosed to consumers in a clear and conspicuous manner, and is essential, and limited to, sharing the PHR identifiable health information with service providers solely to provide users with a safe and reliable personal health record experience.

- *Example 2—Unauthorized Disclosure (Deceptive Omission)*. The medication app from Example 1 shares PHR identifiable health information with a third party for purposes of targeting consumers with ads. The app does not disclose the sharing and also fails to obtain affirmative express consent from users whose information it shares. The third party company can use the PHR identifiable health information to market and advertise—on behalf of the medication app, on behalf of other companies, or on behalf of itself. It can also use the information to improve its own products and services. Such disclosures are not authorized because they are not consistent with consumer expectations (*i.e.*, without disclosure and without affirmative express consent, consumers would not expect that their PHR identifiable health information would be shared, sold, or otherwise exploited for a purpose other than providing the user with a personal health record, and are neither essential nor limited to sharing the PHR identifiable health information solely to provide users with a safe and reliable personal health record experience). This conclusion is also consistent with Commission enforcement actions relating to the sharing of health information (*e.g.*, GoodRx and Easy Healthcare), and those relating to the sharing of other types of sensitive information.¹⁸⁰

- *Example 3—Authorized Disclosure (Public Health Reporting)*: A COVID-19 contact tracing app not covered by HIPAA allows users to self-report their COVID-19 diagnosis, and to notify the user's contacts of their diagnosis, or others with whom the individual may have come into physical contact. PHR identifiable health information about

the individual's COVID-19 diagnosis is transmitted to public health authorities for public health-related purposes, such as public health reporting and analysis or to track areas where the virus is spreading the most rapidly. The contact tracing app discloses to users clearly and conspicuously the specific purposes for which it shares their PHR identifiable health information with public health authorities. These disclosures are authorized, and consistent with consumer expectations, because they are consistent with the company's relationship with the consumer (a PHR that allows a user to report their COVID-19 diagnosis in order to notify others) and are also appropriately disclosed.

Examples 1 and 3 provide guidance about scenarios in which limited disclosures of PHR identifiable health information are permitted without opt-in consent because it is necessary to provide a personal health record to a consumer, is consistent with consumer expectations, the sharing is disclosed to consumers, and (in the case of Example 1) the sharing is subject to protections like service provider agreements that limit the use of the data only for the purpose of providing that service to the consumer. Examples 1 and 3 are also consistent with HIPAA and State health privacy laws.¹⁸¹ For instance, HIPAA permits disclosures for treatment, payment, and operations without patient authorization.

The Commission notes "breach of security" could cover more than just an unauthorized disclosure to a third party. For example, depending on the facts and scope of the authorizations, such as in the company's promises and disclosures to consumers, a "breach of security" could include unauthorized *uses*. There may be a "breach of security" where an entity exceeds authorized access to use PHR identifiable health information, such as where it obtains the data for one legitimate purpose, but later uses that data for a secondary purpose that was not originally authorized by the individual.

Finally, the Commission notes unauthorized access or use of derived PHR identifiable health information may also constitute a breach of security. The Commission noted in its 2023 NPRM that PHR identifiable health information includes "health

¹⁷⁹This would include an analytics provider whose services are essential to the proper functioning of the app and not tied to marketing or advertising—this includes analytics tools to assist with crash reporting or to assess usage patterns (such as the frequency of use of certain features).

¹⁸⁰*Fed. Trade Comm'n et al. v. Vizio, Inc. et al.*, No. 17-cv-00758 (D.N.J. 2017), <https://www.ftc.gov/legal-library/browse/cases-proceedings/162-3024-vizio-inc-vizio-inscape-services-llc>.

¹⁸¹For example, Washington State's My Health, My Data Act permits sharing consumer health data to the "extent necessary to provide a product or service that the consumer to whom such consumer health data relates has requested from such regulated entity or small business." See Revised Code of Washington (RCW) 19.373.030 (1)(b)(ii).

information derived from consumers' interactions with apps and other online services (such as health information generated from tracking technologies employed on websites or mobile applications or from customized records of website or mobile application interactions), as well as emergent health data (such as health information inferred from non-health-related data points, such as location and recent purchases).¹⁸²

D. Clarification of What Constitutes a "PHR Related Entity"

1. The Commission's Proposal Regarding "PHR Related Entity"

The NPRM proposed to revise the definition of "PHR related entity" in two ways. Consistent with its clarification that the Rule applies to health apps, the Commission proposed amending the definition of "PHR related entity" to make clear the Rule covers entities that offer products and services through the online services, including mobile applications, of vendors of personal health records. In addition, the Commission proposed revising the definition of "PHR related entity" to provide that entities that access or send unsecured PHR identifiable health information to a personal health record—rather than entities that access or send any information to a personal health record—are PHR related entities.

The Commission explained the first change (to cover online services) was necessary as websites are no longer the only means through which consumers access health information online. The Commission explained the second change—narrowing the scope of "PHR related entities" to entities that access or send unsecured PHR identifiable health information—was intended to eliminate potential confusion about the Rule's breadth and promote compliance by narrowing the scope of entities that qualify as PHR related entities.¹⁸³ The

Commission identified remote blood pressure cuffs, connected blood glucose monitors, and fitness trackers as examples of internet-connected devices that could qualify as a PHR related entity when individuals sync them with a personal health record (e.g., a health app).¹⁸⁴ The Commission explained, however, that a grocery delivery service that sends information about food purchases to a diet and fitness app would not be a PHR related entity if it does not access unsecured PHR identifiable health information in a personal health record or send unsecured PHR identifiable health information to a personal health record.

The proposed Rule also revised § 318.3(b) by adding language establishing that a third party service provider is not rendered a PHR related entity when it accesses unsecured PHR identifiable health information in the course of providing services. The Commission explained it did not intend for any entity (such as a firm performing attribution and analytics services for a health app) to be considered both a PHR related entity (to the extent it accesses unsecured PHR identifiable health information in a personal health record) and a third party service provider, which could create competing notice obligations and confuse consumers with notice from an unfamiliar company. The Commission explained it considers such firms to be third party service providers that must notify the health app developers for whom they provide services, who in turn would notify affected individuals.

The Commission explained that distinguishing between third party service providers and PHR related entities would create incentives for responsible data stewardship and for de-identification because a firm would only

identifiable health information to a personal health record—rather than entities that access or send any information to a personal health record—are PHR related entities. Otherwise, many entities could be a PHR related entity under the definition's third prong and such entities would then, in the event of a breach, need to analyze whether they experienced a reportable breach under the Rule. If an entity, per the proposed revision, does not qualify as a PHR related entity in the first place, there would be no need to consider whether it experienced a reportable breach. 88 FR 37825 n.54.

¹⁸⁴ The Commission explained, for example, the maker of a wearable fitness tracker may be both a vendor of personal health records (to the extent that its tracker interfaces with its own app, which also accepts consumer inputs) and a PHR related entity (to the extent that it sends information to another company's health app). The Commission noted that regardless of whether the maker of the fitness tracker is a vendor of personal health records or a PHR related entity, its notice obligations are the same: it must notify individuals, the FTC, and in some case, the media, of a breach. 16 CFR 318.3(a), 318.5(b). 88 FR 37825 n.55.

become an entity covered by the Rule in relation to unsecured PHR identifiable health information. To the extent that firms must deal with unsecured PHR identifiable health information, PHR vendors would have incentives to select and retain service providers capable of treating data responsibly (e.g., by not engaging in any onward disclosures of data that could result in a reportable breach) and incentives to oversee their service providers to ensure ongoing responsible data stewardship (which would avoid a breach).

The Commission observed in most cases, third party service providers are likely to be non-consumer facing. The Commission noted examples of PHR related entities would include, as noted above, makers of fitness trackers and health monitors when consumers sync their devices with a mobile health app. The Commission noted further examples of third party service providers would include entities that provide support or administrative functions to vendors of personal health records and PHR related entities.

2. Public Comments Regarding "PHR Related Entity"

The Commission received numerous public comments about the changes to the definition of PHR related entity. Most commenters supported the Commission's approach.¹⁸⁵ One commenter, an industry association for advertisers, noted that addition of the term "unsecured" in the definition of "PHR related entity" created a limitation on the definition's scope that counterbalances the breadth of including "any online service" in the definition.¹⁸⁶ Moreover, this commenter noted, the addition of "unsecured" creates appropriate incentives for firms to secure PHR identifiable health information and to choose partners who will be good data stewards.¹⁸⁷ This commenter noted that limiting the definition to "unsecured" PHR identifiable health information was consistent with the original intent of the Rule, to cover only the most sensitive types of data not covered by HIPAA.¹⁸⁸

A few commenters proposed changes to the definition of "third party service provider" to further distinguish the term from "PHR related entity." One commenter recommended defining "third party service provider" as an

¹⁸⁵ ANI at 1; AAFP at 3; AHIMA at 3; AHIOS at 4; AOA at 3; CARIN Alliance at 3; CDT at 12; CHIME at 3; Confidentiality Coal. at 6; Consumer Rep.'s at 6; CHI at 5; DirectTrust at 4; EFF at 2; EPIC at 7.

¹⁸⁶ NAI at 4–5.

¹⁸⁷ *Id.* at 5.

¹⁸⁸ *Id.* at 4.

¹⁸² 88 FR 37823.

¹⁸³ The proposed definition stated that a PHR related entity is an entity, other than a HIPAA-covered entity or an entity to the extent that it engages in activities as a business associate of a HIPAA-covered entity, that (1) offers products or services through the website, including any online service, of a vendor of personal health records; (2) offers products or services through the websites, including any online services, of HIPAA-covered entities that offer individuals personal health records; or (3) accesses unsecured PHR identifiable health information in a personal health record or sends unsecured PHR identifiable health information to a personal health record. Although the Rule is only triggered when there is a breach of security involving unsecured PHR identifiable health information, the Commission explained it believed there is a benefit to revising the third prong of PHR related entity to make clear that only entities that access or send unsecured PHR

entity that only processes data.¹⁸⁹ This commenter argued the Commission could then impose liability on service providers for further use, sale, disclosure for incompatible purposes.¹⁹⁰ Another commenter recommended aligning the definition of “third party service provider” with the definition of “business associate” under HIPAA.¹⁹¹

Some commenters raised concerns that the Commission’s approach did not provide sufficient clarity for companies trying to understand their obligations as either a third party service provider or PHR related entity.¹⁹² Some commenters requested more examples of types of firms falling within each definition (e.g., examples clearly establishing the status of health data brokers, health marketing firms, search engines, email providers, cloud storage providers)¹⁹³—to facilitate compliance,¹⁹⁴ avoid overlapping notice requirements¹⁹⁵ and to prevent a loophole through which firms may attempt to avoid obtaining consumers’ authorization for data disclosures and to avoid providing breach notifications.¹⁹⁶ One commenter urged the Commission to exempt from the definition of “PHR related entity” any firm that complies with the privacy and data security requirements of HIPAA.¹⁹⁷

In response to the Commission’s request for comment on whether an analytics firm would be a third party service provider, many commenters responded that an analytics firm should fall within that definition¹⁹⁸ for the reasons the Commission articulated: It would be confusing to consumers to receive a notice from a back-end service provider rather than the firm with whom the consumer has the relationship, and categorizing analytics firms (and firms that provide other services) as service providers will create incentives for PHR vendors and PHR related entities to choose their service providers with care. A few commenters, however, expressed concern about covering advertising, analytics, and cloud firms—and health information service providers (“HISPs”) more generally—as they are unable to determine whether the data they receive contains unsecured PHR identifiable health information; only the vendor of

the PHR knows what their data transmissions contain.¹⁹⁹ One commenter urged the Commission to address the data recipient’s unawareness of the content of the data by creating a safe harbor that exempts advertising, analytics and cloud providers that contractually limit their customers, vendors, or partners from sharing health information with them.²⁰⁰

3. The Commission Adopts the Proposed Changes to “PHR Related Entity”

After considering the comments received, the Commission adopts the proposed changes regarding “PHR related entity” without further change. The Commission affirms that (1) PHR related entities include entities offering products and services not only through the websites of vendors of personal health records, but also through any online service, including mobile applications; (2) PHR related entities encompass only entities that access or send unsecured PHR identifiable health information to a personal health record; and (3) while some third party service providers may access unsecured PHR identifiable health information in the course of providing services, this does not render the third party service provider a PHR related entity.

In response to commenters who expressed concern that certain data recipients will not be able to understand their obligations under the Rule because they are unaware of the content of the data transmissions they receive, the Commission highlights § 318.3(b), which states: “For purposes of ensuring implementation of this requirement, vendors of personal health records and PHR related entities shall notify third party service providers of their status as vendors of personal health records or PHR related entities subject to this Part.” This requirement puts data recipients on notice about the potential content of the data transmissions they receive.

Firms may also facilitate compliance by stipulating by contract whether transmissions of data will contain unsecured PHR identifiable health information. Both the sender and recipient of the data can monitor for compliance with those contractual agreements through the use of automated tools, internal auditing, external auditing, or other mechanisms, as appropriate to the size and sophistication of the firms and the

sensitivity of the data. For example, a large advertising platform that has routinely received unsecured PHR identifiable health information, notwithstanding partners’ promises not to send this information, may have different obligations to monitor the data it receives than small firms that do not engage in high-risk activities where the contract precludes sending such data and there is no history of such transmissions.

The Commission believes this approach—notice to service providers pursuant to § 318.3(b) coupled with contracts and oversight—is more appropriate than creating a safe harbor in the Rule that exempts firms that enter into contracts, as there is evidence from FTC cases that firms do not always abide by contractual obligations to safeguard data.²⁰¹

The Commission declines to change the definition of “third party service provider” to distinguish it further from a “PHR related entity,” for two reasons. First, the Commission notes the current definitions of “third party service provider” and “PHR related entity” align closely with the language prescribed by section 13407 and section 13424(b)(1)(A) of the Recovery Act. Jettisoning the current language entirely, as some commenters suggested, would not be consistent with the Recovery Act’s requirements. Second, the Commission believes the current language, in conjunction with the examples provided below, will provide sufficient guidance to the market as to which types of firms fit within each definition.

In response to comments that requested examples of the types of firms that fall into the category of “third party service provider” or “PHR related entity,” the Commission provides the following examples. The Commission believes these examples, in conjunction with the language in § 318.3(b), will provide sufficient clarity about the obligations of third party service providers and PHR related entities to promote compliance, avoid overlapping notice, and prevent loopholes.

²⁰¹ Compl. at ¶ 21, *In the Matter of Flo Health, Inc.*, FTC File No. 1923133 (Jan. 13, 2021), <https://www.ftc.gov/legal-library/browse/cases-proceedings/192-3133-flo-health-inc>; Compl. at ¶ 14(d), *In the Matter of UPromise, Inc.*, FTC File No. 1023116 (Mar. 27, 2012), <https://www.ftc.gov/legal-library/browse/cases-proceedings/102-3116-c-4351-upromise-inc>; Cf. Compl. at ¶ 40, *U.S. v. Easy Healthcare Corporation*, No. 1:23-cv-3107 (N.D. Ill. 2023), <https://www.ftc.gov/legal-library/browse/cases-proceedings/202-3186-easy-healthcare-corporation-us-v> (alleging that the defendant’s disclosures of consumers’ health information violated the policies of platforms to which it had agreed).

¹⁸⁹ FPF at 10.

¹⁹⁰ *Id.*

¹⁹¹ AdvaMed at 8.

¹⁹² SIIA at 3; CARIN Alliance at 4.

¹⁹³ AHIMA at 3–4; AMIA at 3–4; CHI at 5; Direct Trust at 1; Light Collective at 4–5.

¹⁹⁴ SCRS at 1.

¹⁹⁵ NAI at 5.

¹⁹⁶ MRO at 3.

¹⁹⁷ AdvaMed at 5.

¹⁹⁸ NAI at 5; TMA at 3; Consumer Rep.’s at 11.

¹⁹⁹ CCIA at 7–8; CTA at 9–10; SIIA at 3; Direct Trust at 5.

²⁰⁰ CTA at 13.

• *Example 1:* Four separate firms provide data security, cloud computing, advertising and analytics services to a health app (a personal health record), as specified by their service provider contracts, for the health app vendor's benefit. To perform the services specified in their respective contracts, the firms access unsecured PHR identifiable health information. The firms are "third party service providers" of the vendor of the personal health record (the maker of the health app) because they provide services to a vendor of a personal health record (the maker of the health app) in connection with the offering or maintenance of the app, and they access unsecured PHR identifiable health information as a result of these services. In the event of a breach, they should abide by their obligations as third party service providers.

• *Example 2:* An analytics firm provides analytics services to a health app (a personal health record). The analytics firm and health app vendor do not have a customized service provider contract, although the health app vendor agrees to the analytics firm's standard terms of service. The analytics firm accesses unsecured PHR identifiable health information (device identifier and whether the consumer has paid for therapy). The analytics firm uses that data both to provide analytics services to the health app and for its own benefit, for research and development and product improvement. The analytics firm is a third party service provider to the extent that it provides analytics services to the health app for the health app's benefit because it is then providing services to a vendor of a PHR in connection with the offering of the PHR and accessing unsecured PHR identifiable health information as a result of such services. However, the analytics firm is a PHR related entity, rather than a third party service provider, to the extent that it offers its services through the health app for its own purposes (*i.e.*, for research and development and product improvement) rather than to provide the services. In the event of a breach, the analytics firm must fulfill its notification obligations under the Rule according to which function it was performing in connection with the breach. If the functions are indistinguishable, then, pursuant to § 318.3(b), the Commission will consider the firm a third party service provider for policy reasons: a firm that functions, at least in part, as a service provider may not be consumer-facing, such that the consumer may be surprised by a breach notification from

that entity. As a policy matter, it is better for the consumer to receive notice from the health app with whom the consumer directly interacts.

• *Example 3:* A health tracking website (a personal health record) integrates a search bar branded with its maker's logo, which enables its maker (a search engine firm) to offer its services through the website. The search engine firm is a PHR related entity because it offers its services through the website, which is a personal health record. The search bar branded with its maker's logo is consumer-facing, so the consumer would not be surprised to receive a notice from that company if it experiences a reportable breach. By contrast, if the health tracking website had contracted with the search engine firm to provide back-end search services to the website (rather than offering its own branded product or service through the website), and the search engine firm had accessed unsecured PHR identifiable health information as a result of such services, it would be a third party service provider. In the event of a breach, it should abide by its obligations as a third party service provider.

• *Example 4:* Digital readings from a fitness tracker offered by Company A can be integrated into a sleep app offered by Company B (in which the consumer may input other health information). Company A is a PHR related entity to the extent that it offers its fitness tracker product through an online service (Company B's sleep app), and to the extent that it sends unsecured PHR identifiable health information (fitness tracker readings) to a personal health record (the sleep app).

E. Facilitating Greater Opportunity for Electronic Notice

1. The Commission's Proposal Regarding Electronic Notice

The Commission proposed to authorize expanded use of email and other electronic means of providing clear and effective notice of a breach to consumers. In furtherance of this objective, the Commission proposed to update § 318.5 to specify that vendors of personal health records or PHR related entities that discover a breach of security must provide written notice at the last known contact information of the individual. Such written notice may be sent by electronic mail, if an individual has specified electronic mail as the primary contact method, or by first-class mail. The Commission proposed defining "electronic mail" in § 318.2 to mean email in combination with one or more of the following: text

message, within-application messaging, or electronic banner. The Commission further specified that any notification delivered via electronic mail should be clear and conspicuous, and the proposed Rule defined "clear and conspicuous." To assist entities that are required to provide notice to individuals under the Rule, the Commission developed a model notice for entities to use to notify individuals.²⁰²

2. Public Comments Regarding Electronic Notice

Nearly every comment submitted on this proposed change supported the Commission's efforts to update the Rule to allow for greater electronic notice.²⁰³ One commenter noted electronic notices increase the likelihood that individuals will receive the notice, may reduce the time it takes for individuals to receive notice, and reduce the burden on entities providing notice.²⁰⁴ Many commenters also supported the Commission's efforts to provide notice via more than one channel through the new definition of "electronic mail."²⁰⁵

However, not all commenters agreed with the Commission's proposal and some commenters offered other suggestions. Some objected to defining "electronic mail" to mean anything more than "email," stating that electronic mail is commonly understood to mean email and nothing else.²⁰⁶ A few commenters noted that defining multiple forms of electronic notice could result in entities collecting more information than necessary (and consumers having to provide more information than needed) in order to comply with the Rule.²⁰⁷ Others preferred a single notice, arguing that multiple forms of notice is burdensome

²⁰² This model notice was attached as appendix A to the NPRM. 88 FR 37837.

²⁰³ AHIP at 5; AAFP at 3; AHIMA at 5; AHIOS at 3; Anonymous 3 at 1; Anonymous 10 at 1; Beth Barnett; CARIN Alliance at 7; CHI at 5–6; CHIME at 4; Consumer Reports at 8–9; CTA at 21; EPIC at 10; HIMSS at 4; George Mathew at 1; MRO at 3; NAI at 7; Dharini Padmanabhan at 1; Nancy Piwowar at 1. One commenter also stated while there are clear advantages to allowing increased use of electronic notification of data breaches, this notification method could also increase the likelihood that breaches escape public scrutiny. Identity Theft Res. Ctr. ("ITRC") at 2.

²⁰⁴ AdvaMed at 5.

²⁰⁵ AAFP at 3; AHIMA at 5; Anonymous 3 at 1; CARIN Alliance at 7; CHIME at 4; CCLA at 7; EPIC at 10; NAI at 7.

²⁰⁶ ACLA at 5; Mass. Health Data Forum ("MHDF") at 9.

²⁰⁷ Consumer Rep.'s at 7–8; CTA at 22. Consumer Reports further suggested the Commission clarify that substitute notice may be effectuated under the Rule via text message, in-app messaging, or electronic banners for consumers that do not wish to share a mailing or email address. Consumer Rep.'s at 8.

and could result in over-notification, confusion, and notice fatigue among consumers.²⁰⁸ One commenter stated the Commission should revise the definition of “electronic mail” to mean “one or more of the following that is reasonable and appropriate based on the relationship between the individual and the relevant vendor of personal health records or PHR related entity: email, text message, within-application messaging, or electronic banner.”²⁰⁹ Another commenter encouraged the FTC to clarify the in-app messaging method must include push notifications in the event of a breach so consumers are made aware of a breach as soon as possible.²¹⁰ One commenter urged the Commission to specify in § 318.5(i) that a banner notice in the affected app or a website home page notice must be posted for a period of 90 days.²¹¹ Another commenter noted that the different mechanisms listed in the proposed rule are not equivalent—this commenter noted that some are push notifications that a consumer is likely to see without directly interacting with the application, website, or device and some require consumer interaction with the application, website, or device in order to see the notification.²¹² This commenter recommended that the requirement be selection of one push notification but that additional options like in-app notifications and website banners be supported as additional, secondary notice options.²¹³ One commenter stated the FTC may want to consider adding a provision allowing an individual to request a copy of the notice in other accessible formats, such as for hearing- or vision-impaired people, or in a non-English language.²¹⁴ Another commenter argued the Commission should take into consideration TCPA and CAN-SPAM compliance regarding the delivery of electronic notification. Another commenter stated the Commission’s proposal to require two contact methods imposes a higher requirement than HIPAA and State breach notification laws.²¹⁵

Many commenters endorsed the Commission’s proposal that any notification delivered via electronic mail should be “clear and conspicuous,” a newly defined term in

the Rule.²¹⁶ One commenter stated that consistent with FTC’s desire for entities to provide a clear and conspicuous notice, the Commission should consider requiring an email subject line that starts with “Breach of Your Health Information” so that attention is appropriately drawn to the importance of the message content.²¹⁷ One commenter disagreed with the new definition, arguing that the definition is unnecessary and confusing, and urged the Commission to insert the “clear and conspicuous” definition directly into § 318.5 of the Rule.²¹⁸

Regarding the model notice, nearly all who commented on this topic urged the Commission to make the model notice voluntary.²¹⁹ One commenter suggested that using the model should be a safe harbor that shields entities from enforcement.²²⁰

3. The Commission Adopts the Proposed Changes Regarding Electronic Notice

The Commission adopts without change the modifications regarding § 318.5 involving electronic notice and adopts without change the definition of “electronic mail” in § 318.2. The Commission declines to make the other changes commenters requested. First, the Commission believes it is critical, especially given how consumers are accessing information today, to modernize the methods of notice to facilitate greater opportunities for electronic notice. The Commission believes the changes to § 318.5 and the new definition of “electronic mail”²²¹ in § 318.2 accomplish this objective.

²¹⁶ AMA at 5; CHIME at 5; EPIC at 9.

²¹⁷ TMA at 4.

²¹⁸ NAI at 7.

²¹⁹ AdvaMed at 6; AHIP at 6; AMA at 6; CCLIA at 7; CHI at 6; Consumer Rep.’s at 8–9; NAI at 7–8.

One commenter stated that making the model notice mandatory can lead to industry consistency and it may be easier for consumers to understand the message and the contents if they are familiar with a uniform, standardized notice. AHIMA at 5. While the Commission generally agrees that uniform, consistent notices assist with consumer comprehension, the Commission declines to make the model notice compulsory because the facts and circumstances of each breach will vary. Plus, § 318.6 sets forth certain required elements of the content of the notice, so the presence of these elements in all breach notices achieves some degree of consistency across notices.

²²⁰ AHIP at 6.

²²¹ The Commission disagrees with the commenters who urged the Commission to avoid defining “electronic mail” to mean anything more than “email.” ACLA at 5; MHDF at 9. The definition in § 318.2 is clear and unambiguous. Plus, section 13402(e)(1) of the Recovery Act requires that notification be provided via “written notification by first-class mail” or “electronic mail.” Accordingly, the Commission must use “electronic mail.”

In response to concerns raised about the two-part electronic notice, the Commission agrees with commenters who stated it increases the likelihood that individuals will encounter such notices.²²² The Commission does not agree that it is burdensome for entities to comply with this requirement. For example, an entity who complies with the notice requirement by notifying consumers via email plus posting a website notice likely would not need to expend significant additional time and resources by issuing the second part of the notice (*i.e.*, the website notice), and any “cost” of posting such a notice is outweighed by the benefit to consumers of learning of a breach involving their health information. The Commission also is not persuaded that consumers who, for example, receive an email about a breach coupled with an in-app notice about the same breach will be confused. The Commission believes consumers will understand that such notices relate to the same incident, especially given the Rule’s requirement that the notices be “clear and conspicuous.” The Commission also does not find it problematic that the Rule requires notice effectuated via “electronic mail” to occur via two methods while other breach notice laws require one method. The Commission also notes while these amendments are intended to facilitate greater electronic notice, the Rule still permits notice via first-class mail. Accordingly, the contention that this Rule requires two methods of electronic notice is incorrect.

The Commission also declines, in response to public comments,²²³ to mandate how notifications are effectuated when sent via “electronic mail,” as the Commission believes it is important to not be overly prescriptive given rapidly changing technologies.

²²² AAFP at 3–4 (noting AAFP appreciates “the proposed structure of providing notice in two different electronic formats to increase the likelihood individuals will see them”); CHIME at 5 (“CHIME is supportive of the FTC’s approach to revise the “method of notice section” and to structure the breach notification in two parts in order to increase the likelihood that consumers encounter the notice.”); EPIC at 10 (“By requiring email *and* an in-app or website notice option, the expanded definition enables entities to have the best chance at notifying consumers regardless of whether they reliably check their email or continue to use the entity’s app or website.”). The Commission also disagrees with the commenter who recommended that the Commission abandon the two-part notice and create a new definition of “electronic mail” where, for example, only a website notice alone would satisfy the notice requirement if such a notice was “reasonable and appropriate.” AdvaMed at 6. The Commission disagrees with this approach and declines to adopt it.

²²³ See *supra* notes 210–213.

²⁰⁸ AdvaMed at 6; ACLA at 5; AHIP at 5; CTA at 21–22;

²⁰⁹ AdvaMed at 6.

²¹⁰ AHIMA at 5.

²¹¹ TechNet at 5.

²¹² MHDF at 10.

²¹³ *Id.*

²¹⁴ AHIP at 5.

²¹⁵ CHI at 6.

The Commission emphasizes though, as described below, that the notice must satisfy the Rule's definition of "clear and conspicuous."

Nor does the Commission believe, as some commenters argued, the two-part electronic notification will result in additional collections of information by notifying entities. The Commission agrees with commenters who stated entities are generally already collecting the information needed for notice via "electronic mail" and a data minimization issue does not exist.²²⁴

In response to the commenter who suggested the FTC consider adding a provision allowing an individual to request a copy of the notice in other accessible formats, such as for hearing- or vision-impaired people, or in non-English languages,²²⁵ the Commission previously addressed a similar comment in the 2009 Rule Commentary. There, the Commission noted that section 13402(e)(l) of the Recovery Act requires that notification be provided via "written notification by first-class mail" or "electronic mail." The Commission emphasized then, as we do today, that the Rule does not preclude notifications in accessible formats. The Commission supports their use in appropriate circumstances, in addition to the forms of notice prescribed by the Rule.²²⁶

The Commission also adopts without modification the definition of "clear and conspicuous." The Commission agrees with the commenter who indicated it is imperative that a breach notice be reasonably understandable and call attention to the significance of the information that is included in the notice.²²⁷ The Commission believes its definition of "clear and conspicuous" will assist in achieving this objective. The Commission declines, however, to mandate specific language for the email subject line to satisfy the Rule's "clear and conspicuous" requirement, as one commenter had suggested.²²⁸ The Commission emphasizes, however, that the clear and conspicuous requirement would require a notifying entity to use an email subject line that draws the reader's attention to the email notice. The Commission also declines to adopt the suggestion that the definition of "clear and conspicuous" be incorporated directly into § 318.5. The Commission believes the entities seeking information on what "clear and

conspicuous" means will find it clearer to consult the definition in § 318.2.

Turning to the model notice,²²⁹ as the Commission noted in the NPRM, the model was intended for entities to use, in their discretion, to notify individuals, and the Commission adopts the same position here.²³⁰ The model is voluntary and while the Commission believes it represents a best practice, using the model is not required to achieve compliance with the Rule.

The Commission declines to adopt the position that use of the model notice provides a safe harbor, although the Commission would take into consideration in an enforcement action an entity who follows the model notice. Further, the Commission notes an entity who follows the model notice can nevertheless violate the Rule in other ways. For example, an entity could follow the model notice but fail to provide timely notice. In such instances, providing a safe harbor because the entity utilized the model notice would be inappropriate.

F. Revisions to the Required Content of Notice

1. The Commission's Proposal Regarding Content of Notice

The Commission proposed five changes to the content of the notice. First, in § 318.6(a), as part of relaying what happened regarding the breach, the Commission proposed the notice to individuals also include a brief description of the potential harm that may result from the breach, such as medical or other identity theft. Second, the Commission proposed to amend the requirements for the notice under § 318.6(a) to include the full name, website, and contact information (such as a public email address or phone number) of any third parties that acquired unsecured PHR identifiable health information as a result of a breach of security, if this information is known to the vendor of personal health records or PHR related entity (such as where the breach resulted from disclosures of users' sensitive health information without authorization). Third, the Commission proposed modifications to § 318.6(b), which requires that the notice include a description of the types of unsecured PHR identifiable health information that were involved in the breach. The Commission proposed this exemplar list be expanded to include additional types of PHR identifiable health information, such as health diagnosis or condition,

lab results, medications, other treatment information, the individual's use of a health-related mobile application, and device identifier. Fourth, the Commission proposed revising § 318.6(d) of the Rule to require the notice to individuals include additional information providing a brief description of what the entity that experienced the breach is doing to protect affected individuals, such as offering credit monitoring or other services. Fifth, the Commission proposed modifying § 318.6(e) so the contact procedures specified by the notifying entity must include two or more of the following: toll-free telephone number; email address; website; within-application; or postal address.

2. Public Comments Regarding Content of Notice

a. Proposal That Notice Include Description of Potential Harm That May Result From a Breach

The Commission's proposal to modify § 318.6(a) to include in the notice to individuals a brief description of the potential harm that may result from a breach drew a wide range of comments. On the one hand, many commenters supported the Commission's proposal.²³¹ For example, one commenter noted this proposal would help individuals better understand the connection between the information breached and the potential harm that could result from the breach of such information.²³² Other commenters stated that providing the potential harms from a breach better equips consumers to address injuries and mitigate harms from it.²³³ One commenter stated including some potential harms would be helpful, but notifying entities should also include language in the notice stating that other harms may occur.²³⁴ This same commenter suggested the Commission consider selecting the most common types of breaches and listing some but not all of the potential consequences from each.²³⁵

On the other hand, many commenters criticized this proposal.²³⁶ Some

²²⁴ CARIN Alliance at 6; EPIC at 10.

²²⁵ See *supra* note 214.

²²⁶ 74 FR 42972.

²²⁷ AMA at 5.

²²⁸ See *supra* note 217.

²²⁹ The model notice is found in appendix A.

²³⁰ 88 FR 37827.

²³¹ AAFP at 4; AMA at 6; AOA at 5; Anonymous 3; AHIOS at 3; CARIN Alliance at 7–8; CHIME at 3, 6; Consumer Reports at 9–10; EFF at 2; EPIC at 10–11; HIMSS at 3–4; ITRC at 2; Members of the House of Representatives at 1–2; Dharini Padmanabhan at 1.

²³² AMA at 6.

²³³ Consumer Rep.'s at 9–10; EPIC at 10–11.

²³⁴ MHDF at 10–11.

²³⁵ *Id.*

²³⁶ AdvaMed at 6–7; AHIP at 6; ACLA at 4–5; Confidentiality Coal. at 7; CTA at 23–24; MHDF at 10; NAI at 9.

commenters argued this proposal will result in notifying entities having to speculate about potential harms that may never occur or providing a list of harms that may be incomplete.²³⁷ Others pointed out that notifying individuals about potential harms could cause consumer anxiety, consumer confusion, and detract from actions the individuals should take.²³⁸ One commenter noted the Commission's proposal might lead consumers to believe the harms listed in the notice are the only possible harms from a breach, when in fact consumers may suffer other harms not disclosed in the notice.²³⁹ This same commenter also noted it is opposed to entities stating there are no known harms that may result from a breach solely because a notifying entity is unaware of any specific bad outcomes.²⁴⁰

b. Proposal That Notice Include Full Name, Website and Contact Information of Third Parties That Acquired Unsecured PHR Identifiable Health Information

Next, the Commission proposed to amend the requirements for the notice under § 318.6(a) to include the full name, website, and contact information (such as a public email address or phone number) of any third parties that acquired unsecured PHR identifiable health information as a result of a breach of security. Although several commenters supported this proposal,²⁴¹ many others pointed out it is problematic in certain circumstances.²⁴² A few commenters noted the proposal is ill-suited for security breaches, such as a hacking, where providing consumers with the name and contact information of an actor who committed a security breach (e.g., a hacker) could result in further malicious action against the target entity.²⁴³ One commenter noted for security breaches, the malicious actor or hacker would not be responsive to consumers.²⁴⁴ Further, one commenter noted this requirement could hamper law enforcement efforts.²⁴⁵ One commenter also indicated this requirement could

frustrate investigative efforts or have a chilling effect on an inadvertent recipient from reporting a wrongful disclosure.²⁴⁶

c. Proposal That Notice Include Description of Types of Unsecured PHR Identifiable Health Information Involved in a Breach

Third, the Commission proposed modifications to § 318.6(b), which requires the notice to individuals include a description of the types of unsecured PHR identifiable health information that were involved in the breach. The Commission proposed this exemplar list be expanded to include additional types of PHR identifiable health information, such as health diagnosis or condition, lab results, medications, other treatment information, the individual's use of a health-related mobile application, and device identifier. Several commenters supported this proposal.²⁴⁷ One commenter noted it is important for consumers to receive notice of the specific types of PHR identifiable health information involved in a breach, given that the exposure of health information can lead to a wide spectrum of harms.²⁴⁸ Another commenter stated providing individuals with a more expansive list of exposed data points will also give them a more complete picture of the risks they face.²⁴⁹

d. Proposal That Notice Include Description of What Entity Is Doing To Protect Affected Individuals

Fourth, the Commission proposed revising § 318.6(d) of the Rule to require that the notice to individuals include additional information providing a brief description of what the entity that experienced the breach is doing to protect affected individuals, such as offering credit monitoring or other services. This proposal attracted support from multiple commenters.²⁵⁰ One commenter stated that informing individuals about these steps is important so that they know what additional actions they should take to protect themselves from potential harm.²⁵¹ Another similarly stated that knowing what the notifying entity is doing to protect affected individuals can help consumers who are considering

making purchase decisions for fraud detection or credit monitoring.²⁵² One commenter stated that requiring notifying entities to share this information will incentivize them to take proactive measures to mitigate harms to consumers.²⁵³

Some commenters, however, raised concerns about this proposal. For instance, one commenter believed the Rule already encompasses this requirement and therefore the Commission's proposal could result in duplicative information being provided in the notice.²⁵⁴ Another commenter stated the FTC needs to go further in ensuring that notification requirements help consumers understand what remedies are available when their health information is breached.²⁵⁵

e. Proposal That Notice Include Two or More Contact Procedures

Fifth, the Commission proposed amendments to § 318.6(e) so the contact procedures specified by the notifying entity in its breach notification must include two or more of the following: toll-free telephone number; email address; website; within-application; or postal address. Many commenters expressed support for this proposal.²⁵⁶ One commenter noted multiple contact options ensures that victims of all backgrounds and technical capabilities are able to contact the notifying entity to learn more about how to protect themselves after a breach.²⁵⁷ Another commenter noted that providing multiple contact options encourages and facilitates communication between the individual and the notifying entity.²⁵⁸ One commenter, however, expressed concern the proposal is burdensome, the HIPAA breach notice rule requires only one method of contact, and HHS has not identified any concerns with individuals having difficulty obtaining information from covered entities using one contact method under HIPAA's breach notice rule.²⁵⁹

²³⁷ AdvaMed at 6–7; AHIP at 6; MHDF at 10; NAI at 9.

²³⁸ ACLA at 4–5; AMIA at 5; NAI at 9.

²³⁹ MHDF at 10.

²⁴⁰ *Id.* at 10–11.

²⁴¹ AAFP at 4; AHIMA at 5–6; AMA at 6; AMIA at 5; AOA at 5; CARIN Alliance at 7; Consumer Rep.'s at 9–10; EFF at 2; EPIC at 10–11; HIMSS at 3–4; ITRC at 2; Members of the House of Representatives at 1–2.

²⁴² ACLA at 4–5; AHIP at 6; CHI at 6; Confidentiality Coalition at 7; CTA at 24.

²⁴³ ACLA at 4–5; Confidentiality Coal. at 7.

²⁴⁴ Confidentiality Coal. at 7.

²⁴⁵ CTA at 24.

²⁴⁶ AHIP at 6.

²⁴⁷ AAFP at 4; AHIMA at 6; AMA at 6; AOA at 5; CARIN Alliance at 7; Consumer Rep.'s at 9–10; Ella Balasa at 2; HIMSS at 3–4; ITRC at 2; NAI at 9.

²⁴⁸ Light Collective at 2.

²⁴⁹ ITRC at 2.

²⁵⁰ AAFP at 4; AMA at 6; AOA at 4; CARIN Alliance at 7–8; HIMSS at 3–4; ITRC at 2.

²⁵¹ AMA at 6.

²⁵² AHIMA at 5–6.

²⁵³ Consumer Rep.'s at 9–10.

²⁵⁴ Confidentiality Coal. at 7.

²⁵⁵ Light Collective at 6–7.

²⁵⁶ AAFP at 4; AHIMA at 6; AHIP at 5; Anonymous 3 at 1; AOA at 5; CARIN Alliance at 8; Consumer Rep.'s at 9–10; EPIC at 9–10; HIMSS at 3–4; ITRC at 2; Dharini Padmanabhan at 1.

²⁵⁷ AHIMA at 6.

²⁵⁸ AMA at 6.

²⁵⁹ AdvaMed at 6–7.

3. The Commission Changes Regarding Content of Notice

a. The Commission Declines To Adopt Proposal That Notice Include Description of Potential Harm That May Result From a Breach

The Commission believes, in light of the public comments, that the downsides of requiring in the notice a description of the potential harms that may result from a breach outweigh the upsides. The Commission is concerned about requiring a consumer notice to include possible harms that may never materialize. In such cases, consumers may experience needless anxiety and take actions that are not necessary, leading to consumer frustration. The Commission also is concerned this proposal may result in entities describing potential harms so generically that the description provides minimal value to consumers, or, alternatively, that entities will provide a laundry list of potential harms, making such a list meaningless to consumers. The Commission also agrees with one commenter who noted this proposal might lead consumers to believe the harms listed in the notice are the only possible harms from a breach, when in fact consumers may suffer other harms not disclosed in the notice.²⁶⁰

Accordingly, the Commission declines to adopt this proposal.²⁶¹ The Commission believes the remaining elements of the content of the notice will supply individuals with sufficient information about a breach, especially given the other modifications to § 318.6. The Commission also emphasizes in certain cases where harms are concrete and known, notifying entities should as a best practice inform individuals about those harms in the notice.

b. The Commission Modifies Proposal That Notice Include Full Name, Website, and Contact Information of Third Parties That Acquired Unsecured PHR Identifiable Health Information

In light of the public comments, the Commission is modifying § 318.6(a) to require notifying entities to provide the full name or identity (or where providing name or identity would pose a risk to individuals or the entity providing notice, a description) of the third parties that acquired the PHR identifiable health information as a result of a breach of security.²⁶² The Commission believes it is important for consumers to know who acquired their

PHR identifiable health information as a result of a breach. At the same time, the Commission acknowledges in some scenarios it could be problematic to require notifying entities to provide the contact information of those who acquired PHR identifiable health information.

Accordingly, this revised provision is intended to still provide individuals with information about who acquired their health information. Under § 318.6(a), notifying entities are required to provide the full name or identity of the third parties that acquired the PHR identifiable health information as a result of a breach of security, except where providing the full name or identity of the third parties would pose a risk to affected individuals or the entity providing notice. In cases where providing the name or identity of the third parties that acquired the PHR identifiable health information as a result of a breach of security would pose a risk to affected individuals or the entity providing notice (e.g., providing the name of hacker could subject affected individuals or the entity providing notice to further harm), § 318.6(a) permits notifying entities to describe the type of third party (e.g., hacker) who acquired individuals' PHR identifiable health information.

c. The Commission Adopts Proposal That Notice Include Description of Types of Unsecured PHR Identifiable Health Information Involved in a Breach

The Commission agrees with the many public comments supporting this proposal.²⁶³ The Commission concurs with the commenter who noted it is important for consumers to receive notice of the specific types of PHR identifiable health information involved in a breach,²⁶⁴ and the commenter who stated that providing affected individuals with a more expansive list of health data points implicated in a breach will help them better understand the risks they face.²⁶⁵ The Commission adopts this proposal without modification.

d. The Commission Adopts Proposal That Notice Include Description of What Entity Is Doing To Protect Affected Individuals

Several commenters supported the Commission proposal that the notice to individuals include a description of what the notifying entity is doing to protect affected individuals.²⁶⁶ The

Commission concurs with the commenter who stated that informing affected individuals about the steps notifying entities are taking to protect them is important so that affected individuals know what additional actions they should take to protect themselves from potential harm.²⁶⁷ The Commission similarly agrees with the commenter who stated that knowing what the notifying entity is doing to protect affected individuals can help consumers who are considering making purchase decisions like fraud detection or credit monitoring.²⁶⁸ The Commission also agrees with the commenter who stated that requiring notifying entities to share information about what they are doing to protect affected individuals will incentivize notifying entities to take proactive measures to mitigate harms to consumers.²⁶⁹

In response to the one commenter who noted the 2009 Rule already includes this proposed requirement,²⁷⁰ the Commission notes § 318.6(d) from the 2009 Rule requires notifying entities to include in the notice to individuals what the entity is doing to investigate the breach, to mitigate any losses, and to protect against any further breaches. Accordingly, under the 2009 Rule, there is no explicit requirement for the notifying entity to state in the individual notice what the entity is doing to protect affected individuals. Given this, the Commission does not believe individuals will receive duplicative information.

In response to the commenter who argued the Commission needs to help consumers understand post-breach remedies,²⁷¹ the Commission believes this concern is addressed by the combination of § 318.6(c), which requires notifying entities to include in the notice steps individuals should take to protect themselves from potential harm resulting from the breach, and § 318.6(d), which requires notifying entities to include in the notice the steps the notifying entity is taking to protect affected individuals following the breach.

The Commission adopts proposed § 318.6(d) without modification.

e. The Commission Adopts Proposal That Notice Include Two or More Contact Procedures

In response to the comment that providing two or more contact

²⁶⁰ MHDF at 10.

²⁶¹ The Commission has updated the model notice in appendix A to reflect this change.

²⁶² The Commission has updated the model notice in appendix A to reflect this change.

²⁶³ See *supra* note 247.

²⁶⁴ See *supra* note 248.

²⁶⁵ See *supra* note 249.

²⁶⁶ See *supra* note 250.

²⁶⁷ See *supra* note 251.

²⁶⁸ See *supra* note 252.

²⁶⁹ See *supra* note 253.

²⁷⁰ See *supra* note 254.

²⁷¹ See *supra* note 255.

procedures in the notice is burdensome,²⁷² the Commission believes if this proposal results in any burden to notifying entities, such burden will be minimal given the ease with which compliance with this provision can be achieved, and outweighed by the benefits to consumers who will have increased options to communicate with notifying entities. Second, in response to the comment that the HIPAA Breach Notification Rule requires only one contact method,²⁷³ the Commission notes while there are many similarities between the FTC's and HHS's respective breach notification rules and the agencies have consulted to harmonize the two rules, there are differences between them, and the Commission believes it is important to update this provision to reflect new modes of communication and facilitate greater opportunities for communication between affected individuals and notifying entities.

The Commission notes multiple commenters supported this proposal.²⁷⁴ Specifically, the Commission agrees with the commenter who stated multiple contact procedures enables greater opportunities for affected individuals to communicate with notifying entities.²⁷⁵ The Commission also agrees with the commenter who noted multiple contact options ensures that affected individuals from all backgrounds and technical capabilities are able to contact the notifying entity following a breach.²⁷⁶ The Commission therefore adopts proposed § 318.6(e) without modification.

G. Timing of Notice to the FTC

1. The Commission's Proposal Regarding Timing of Notice

Although the Commission did not propose any timing changes in the NPRM, the Commission requested comments on several issues related to timing, including the timing of the notification to the FTC. Regarding the notification timeline to the FTC, the Commission sought comment on whether it should extend the timeline to give entities more time to investigate breaches and better ascertain the number of affected individuals or whether an extension would simply facilitate dilatory action and minimize the opportunity for an important dialogue with Commission staff during

the fact-gathering stage immediately following a breach.

2. Public Comments Regarding Timing of Notice

Several commenters expressed support for extending the notification timeline to the FTC.²⁷⁷ Commenters provided several reasons why the existing requirement of notice to the FTC "as soon as possible and in no case later than ten business days following the date of discovery of the breach" for breaches involving 500 or more individuals should be amended. For example, commenters noted that ten days does not provide entities with sufficient time to adequately investigate incidents and fully understand the facts, possibly leading to notices that may be incomplete and require amendment or correction.²⁷⁸ Others commented that the existing requirement diverts key resources from investigating potential breaches, indicating when a breach is suspected or has been discovered, the target entity's focus should be responding to the incident, conducting a thorough investigation of what may have occurred, and addressing and mitigating vulnerabilities to ensure additional information is not compromised.²⁷⁹

Several commenters urged the FTC to align the timeframe to notify the FTC with the timing requirement under HIPAA's Health Breach Notification Rule,²⁸⁰ which requires notification to the Secretary of HHS without unreasonable delay and in no case later than 60 calendar days following a breach.²⁸¹ One commenter, irrespective of HIPAA, suggested the Commission give entities up to 60 days to investigate a breach and provide notification to the Commission.²⁸² One commenter recommended the FTC adopt a "risk-based" notification approach whereby the agency could create a shorter notification timeline for high-risk incidents and a longer notification timeline or even no notification for low-risk incidents.²⁸³

3. The Commission Adopts Changes to the Timing of Notice

Having considered the public comments, the Commission agrees with

²⁷⁷ AdvaMed at 9; AHIP at 7; ACLA at 3–4; ATA Action at 2; CCIA at 8; CHI at 6; CTA at 20–21; TechNet at 5.

²⁷⁸ AdvaMed at 9; ACLA at 3–4; AHIP at 7; TechNet at 5–6.

²⁷⁹ ACLA at 3–4; CTA at 19–21.

²⁸⁰ 45 CFR 164.400 through 414.

²⁸¹ AdvaMed at 9; AHIP at 7; ACLA at 3; ATA Action at 2; TechNet at 5–6.

²⁸² ACLA at 3–4.

²⁸³ CTA at 19–21.

commenters who recommended that the notification timeline to the FTC for breaches of security involving 500 or more individuals should be adjusted. The Commission agrees that in certain incidents, especially large, complex breaches, it can be challenging for entities to fully understand the scope of a breach in ten business days, leading to the possibility of incomplete breach notices.

Accordingly, the Commission is revising § 318.4(b) to read: "All notifications required under § 318.5(c) involving the unsecured PHR identifiable health information of 500 or more individuals shall be provided contemporaneously with the notice required by paragraph (a) of this section." This change requires entities, for breaches involving 500 or more individuals, to notify the FTC consistent with the notice required by § 318.4(a)—*i.e.*, without unreasonable delay and in no case later than 60 calendar days after the discovery of a breach of security. This change also requires the notice to the FTC be sent at the same time as the notice to the individuals. This requirement thus ensures the notice to the FTC includes all of the information provided in the notice to the individual. It also avoids a scenario where individuals receive notice before the FTC receives notice and affected individuals contact the FTC about a breach for which the Commission has not been notified.

As a result of this change, the Commission anticipates entities will have sufficient time to provide complete and fulsome notifications to the Commission. The Commission emphasizes, however, that notice to the FTC should occur "without unreasonable delay," with 60 days serving as the outer limit.²⁸⁴ The Commission believes, consistent with public comments, this change effectively harmonizes the notification timeline to the FTC with the notification timeline to the Secretary of HHS under the HIPAA Breach Notification Rule.

²⁸⁴ As the Commission stated in the 2009 Rule Commentary, in some cases, it may be an "unreasonable delay" to wait until the 60th day to provide notification. For example, if a vendor of personal health records or PHR related entity learns of a breach, gathers all necessary information, and has systems in place to provide notification within 30 days, it would be unreasonable to wait until the 60th day to send the notice. Similarly, the Commission noted there may be circumstances where a vendor of personal health records discovers that its third party service provider has suffered a breach before the service provider notifies the vendor that the breach has occurred. In such circumstances, the vendor should begin taking steps to address the breach immediately, and should not wait until receiving notice from the service provider. 74 FR 42971 n.94 (2009).

²⁷² See *supra* note 259.

²⁷³ *Id.*

²⁷⁴ See *supra* note 256.

²⁷⁵ See *supra* note 258.

²⁷⁶ See *supra* note 257.

The Commission also believes this notification timeline satisfies the Recovery Act requirement that notice be provided “immediately.”²⁸⁵ The Commission also notes this change does not affect in any way the timing of the notice to the FTC for breaches involving less than 500 individuals.

Finally, a small number of commenters addressed other issues related to timing, such as the timeline for providing notice to consumers or the media. The Commission believes, for the reasons stated in the commentary accompanying the 2009 NPRM and the 2009 Rule Commentary, the current timelines are appropriate to give consumers and the media timely notice without overburdening notifying firms.²⁸⁶

H. Proposed Changes To Improve Rule’s Readability

1. The Commission Proposed Changes To Promote Readability

The Commission proposed several changes to improve the Rule’s readability. Specifically, the Commission proposed to include explanatory parentheticals for internal cross-references, add statutory citations in relevant places, consolidate notice and timing requirements in single sections, and revise the Enforcement section to state more plainly the penalties for non-compliance.

2. Public Comments Regarding Readability

Commenters supported the Commission’s proposed changes to improve the Rule’s readability and promote comprehension by including explanatory parentheticals and statutory citations.²⁸⁷ Commenters also expressed support for the proposed changes to improve the Rule’s readability and promote compliance by consolidating into single sections, respectively, the Rule’s breach notification and timing requirements.²⁸⁸ Commenters also favored the proposal to modify § 318.7 to make plain that a violation of the Rule constitutes a violation of a rule promulgated under section 18 of the FTC Act and is subject to civil penalties,

stating this clarification will decrease the burden on the FTC in enforcement actions and prevent unintended barriers to enforcement.²⁸⁹

3. The Commission Adopts Changes Regarding Readability

In light of support from commenters and the Commission’s belief that these proposed changes improve readability, the Commission adopts these changes without modification.²⁹⁰

²⁸⁹ AHIMA at 7; AMA at 6–7; AHIOS at 5; MRO at 4. As part of its comment, AMA recommended the FTC, as Rule violations are filed, use actual examples as case study models for future educational resources. The Commission notes that its existing enforcement actions under the Rule already provide guidance for the marketplace and the FTC also has issued business guidance regarding the Rule. E.g., Fed. Trade Comm’n, *Collecting, Using, or Sharing Consumer Health Information? Look to HIPAA, the FTC Act, and the Health Breach Notification Rule* (Sept. 2023), <https://www.ftc.gov/business-guidance/resources/collecting-using-or-sharing-consumer-health-information-look-hipaa-ftc-act-health-breach> (last visited Jan. 11, 2023); Fed. Trade Comm’n, *Health Breach Notification Rule: The Basics for Business* (Jan. 2022), <https://www.ftc.gov/business-guidance/resources/health-breach-notification-rule-basics-business> (last visited Jan. 11, 2024); Fed. Trade Comm’n, *Complying with FTC’s Health Breach Notification Rule* (Jan. 2022), <https://www.ftc.gov/business-guidance/resources/complying-ftcs-health-breach-notification-rule-0> (last visited Jan. 11, 2024). One commenter also asserted the Commission was seeking to apply the NPRM’s proposed changes retrospectively to breaches of security that were discovered on or after September 24, 2009. This commenter urged the Commission to modify § 318.8 so that the Rule would only apply to breaches of security discovered at least 30 days after the effective date of this final rule. TechNet at 5–6. The 2023 NPRM set out the entire part for the convenience of commenters but did not propose any changes to § 318.8. The Commission notes this effective date section was codified in 2009 when part 318 was added to the CFR and has been in effect since September 24, 2009. As explained in the 2009 Rule Commentary, “the Commission does not have discretion to change the effective date of the rule because the Recovery Act establishes the effective date.” See 74 FR 42976; see also 42 U.S.C. 17937(g)(1) (“The provisions of this section shall apply to breaches of security that are discovered on or after the date that is 30 days after the date of publication of such interim final regulations.”). The Commission emphasizes that this final rule does not apply retroactively.

²⁹⁰ Relatedly, the Commission also is making a non-substantive grammatical change to § 318.5(a)(2)(ii), which involves substitute notice. This provision currently states: “Such a notice in media or web posting shall include a toll-free phone number, which shall remain active for at least 90 days, where an individual can learn whether or not the individual’s unsecured PHR identifiable health information may be included in the breach.” The Commission is revising § 318.5(a)(2)(ii) so it reads: “Such a notice in media or web posting shall include a toll-free phone number, which shall remain active for at least 90 days, where an individual can learn if the individual’s unsecured PHR identifiable health information may have been included in the breach.” The Commission made this grammatical change to improve the rule’s readability; the change does not alter the provision’s substantive meaning.

III. Paperwork Reduction Act

The Paperwork Reduction Act (“PRA”), 44 U.S.C. chapter 35, requires Federal agencies to seek and obtain Office of Management and Budget (“OMB”) approval before undertaking a collection of information directed to ten or more persons.²⁹¹ This final rule is modifying an existing collection of information,²⁹² which OMB has approved through July 31, 2025 (OMB Control No. 3084–0150). As required by the PRA, the Commission sought OMB review of the modified information collection requirement at the time of the publication of the NPRM. OMB directed the Commission to resubmit its request at the time the final rule is published. Accordingly, simultaneously with the publication of this final rule, the Commission is resubmitting its clearance request to OMB. FTC staff has estimated the burdens associated with the amendments as set forth below.

FTC staff estimates the amendments to 16 CFR part 318 will likely result in more reportable breaches by covered entities to the FTC. In the event of a breach of security, the covered firms will be required to investigate and, if certain conditions are met, notify consumers, the Commission, and, in some cases, the media.²⁹³

Based on industry reports, FTC staff estimates the amendments will cover approximately 193,000 entities, which, in the event they experience a breach, may be required to notify consumers, the Commission, and, in some cases, the media. While there are approximately 1.8 million apps in the Apple App Store²⁹⁴ and 2.4 million apps in the Google Play Store,²⁹⁵ as of March 2024, it appears that roughly 193,000 of the apps offered in either store are categorized as “Health and Fitness.”²⁹⁶

²⁹¹ 44 U.S.C. 3502(3)(A)(i).

²⁹² See 44 U.S.C. 3502(3)(A)(i).

²⁹³ Third party service providers who experience a breach are required to notify the vendor of personal health records or PHR related entity, which in turn is then required to notify consumers. The Commission expects the cost of notification to third party service providers would be small, relative to the entities that have to notify consumers. As part of the NPRM, the Commission solicited public comment on this issue and data that may be used to quantify the costs to third party service providers. The Commission did not receive any responsive submissions pertaining to this issue.

²⁹⁴ See App Store—Apple, <https://www.apple.com/app-store/>.

²⁹⁵ See AppBrain: Number of Android Apps on Google Play (Mar 2024), <https://www.appbrain.com/stats/number-of-android-apps>.

²⁹⁶ See Business of Apps, “App Data Report: App Store Stats, Downloads, Revenues and App Rankings,” <https://www.businessofapps.com/data/report-app-data/> (reporting 90,913 apps in the Apple iOS App Store and 102,402 apps in the Google Play Store were categorized as “Health and

Continued

²⁸⁵ 42 U.S.C. 17932(e)(3). Like the Department of Health and Human Services previously concluded with respect to notification to the Secretary under the HIPAA Breach Notification Rule (74 FR 42753 (2009)), the Commission concludes this interpretation satisfies the statutory requirement that notifications of larger breaches be provided to the FTC immediately as compared to the notifications of smaller breaches (i.e., those involving less than 500 individuals), which the statute allows to be reported annually to the FTC.

²⁸⁶ 74 FR 17918 (2009); 74 FR 42971 (2009).

²⁸⁷ AMA at 6; CARIN Alliance at 9.

²⁸⁸ AHIMA at 7; AMA at 6–7.

The Commission received three comments in response to the NPRM arguing the Rule's scope is broader than apps categorized as "Health and Fitness" and the NPRM's PRA analysis therefore underestimated the number of covered entities and the resulting number of reportable breaches.²⁹⁷ As discussed above,²⁹⁸ the Commission is adopting these amendments to clarify that the Rule applies to mobile health applications and similar technologies. The Commission also highlighted several key limitations to the Rule's scope.²⁹⁹ Thus, the 193,000 covered entities is a rough proxy for all covered PHRs, because it encompasses mobile health applications categorized as "Health and Fitness." Similar health technologies are included in the roughly 193,000 covered entities because most websites and connected health devices that will be covered by the amendments act in conjunction with an app.³⁰⁰

FTC staff estimates these entities will, cumulatively, experience 82 breaches per year for which notification may be required. With the proviso that there is insufficient data at this time about the number and incidence rate of breaches at entities covered by the amendments (due to underreporting prior to issuance of the Policy Statement), FTC staff determined the number of estimated breaches by calculating the breach incidence rate for HIPAA-covered entities, and then applied this rate to the estimated total number of entities that will be subject to the amendments.³⁰¹

Fitness"). Together, this suggests there are approximately 193,000 Health and Fitness apps. This figure is likely both under- and over-inclusive as a proxy for covered entities. For example, this figure does not include apps categorized elsewhere (i.e., outside "Health and Fitness") that may be PHRs. However, at the same time, this figure also overestimates the number of covered entities, since many developers make more than one app and may specialize in the Health and Fitness category.

²⁹⁷ See Chamber at 2; CHI at 6–7; CCIA at 8–9.

²⁹⁸ See section II.1.c.

²⁹⁹ *Id.*

³⁰⁰ Indeed, one of the commenters who argued the Rule's coverage is broader than projected in the NPRM's PRA analysis acknowledged that there has been growth in the number of websites and apps since the 2009 PRA analysis estimated 700 covered entities to be covered by the Rule. Chamber at 2. Further, the approximately 193,000 covered entities may overestimate the number of covered entities, as some apps or websites may not qualify as a covered entity given the Rule's boundaries. For example, a website or app must have the technical capacity to draw information from multiple sources and that same website or app must still be "managed, shared, and controlled by or primarily for the individual" to be covered by the Rule.

³⁰¹ FTC staff used information publicly available from HHS on HIPAA related breaches because the HIPAA Breach Notification Rule is similarly constructed. However, while there are similarities between HIPAA-covered entities and HBNR-covered entities, it is not necessarily the case that rates of breaches would follow the same pattern.

Additionally, as the number of breaches per year has grown significantly in the recent years,³⁰² and FTC staff expects this trend to continue, FTC staff relied on the average number of breaches from 2021 through 2023 to estimate the annual breach incidence rate for HIPAA-covered entities.

Specifically, HHS's OCR reported 715 breaches in 2021, 719 breaches in 2022, and 733 breaches in 2023,³⁰³ which results in an average of 722 breaches between 2021 and 2023. Based on the 1.7 million entities that are covered by the HIPAA Breach Notification Rule³⁰⁴ and the average number of breaches for 2021–2023, FTC staff determined an annual breach incidence rate of 0.000425 (722/1.7 million). Accordingly, multiplying the breach incidence rate (0.000425) by the estimated number of entities covered by the amendments (193,000) results in an estimated 82 breaches per year.³⁰⁵

For instance, HIPAA-covered entities are generally subject to stronger data security requirements under HIPAA, but also may be more likely targets for security incidents (e.g., ransomware attacks on hospitals and other medical treatment centers covered by HIPAA have increased dramatically in recent years); thus, this number could be an under- or overestimate of the number of potential breaches per year.

³⁰² According to HHS's Office for Civil Rights ("OCR"), the number of breaches per year grew from 276 in 2013 to 739 breaches in 2023. See *Breach Portal*, U.S. Dep't of Health & Human Servs., Office for Civil Rights, https://ocrportal.hhs.gov/ocr/breach/breach_report.jsf (last visited March 1, 2024). The data was downloaded on March 1, 2024, resulting in limited data for 2024. Thus, breaches from 2024 were excluded from the calculations. However, breach investigations that remain open (under investigation) from years prior to 2024 are included in the count of yearly breaches.

³⁰³ See *Breach Portal*, U.S. Dep't of Health & Human Servs., Office for Civil Rights, https://ocrportal.hhs.gov/ocr/breach/breach_report.jsf (last visited March 1, 2024).

³⁰⁴ In a **Federal Register** publication titled "Proposed Modifications to the HIPAA Privacy Rule to Support, and Remove Barriers to, Coordinated Care and Individual Engagement", OCR proposes increasing the number of covered entities from 700,000 to 774,331. 86 FR 6446, 6497 (Jan. 21, 2021). For purposes of calculating the annual breach incidence rate, FTC staff utilized 700,000 covered entities because the proposed estimate of 774,331 covered entities represents a projected increase that has not been finalized by OCR. The OCR publication also lists the number of covered Business Associates as 1,000,000. 86 FR 6528. FTC staff arrived at 1.7 million entities subject to the HIPAA Breach Notification Rule by adding 700,000 covered entities and 1,000,000 Business Associates.

³⁰⁵ One commenter argued that basing the NPRM's projection of the annual number of breaches on the breach incidence rate for HIPAA-covered entities is problematic because the NPRM's proposed definition of a breach of security "goes far and beyond" the HIPAA definition of a breach. CCIA at 8–9. To the extent the commenter is referring to the fact that the Rule's definition of breach of security covers unauthorized disclosures, the Commission notes the HIPAA Breach Notification Rule similarly covers unauthorized disclosures. See *Breach Notification Rule*, U.S.

Costs

To determine the costs for purposes of this analysis, FTC staff has developed estimates for two categories of potential costs: (1) the estimated annual burden hours and labor cost of determining what information has been breached, identifying the affected customers, preparing the breach notice, and making the required report to the Commission; and (2) the estimated capital and other non-labor costs associated with notifying consumers.

Estimated Annual Burden Hours: 12,300.

Estimated Annual Labor Cost: \$883,140.

First, to determine what information has been breached, identify the affected customers, prepare the breach notice, and make the required report to the Commission, FTC staff estimates covered firms will require per breach, on average, 150 hours of employee labor at a cost of \$10,770.³⁰⁶ This estimate does not include the cost of equipment or other tangible assets of the breached firms because they likely will use the equipment and other assets they have for ordinary business purposes. Based on the estimate that there will be 82 breaches per year the annual hours of burden for affected entities will be 12,300 hours (150 hours × 82 breaches) with an associated labor cost of \$883,140 (82 breaches × \$10,770).

Estimated Capital and Other Non-Labor Costs: \$91,984,370.

The capital and non-labor costs associated with breach notifications depend upon the number of consumers contacted and whether covered firms are likely to retain the services of a forensic expert. For breaches affecting large numbers of consumers, covered firms are likely to retain the services of a forensic expert. FTC staff estimates, for each breach requiring the services of forensic experts, forensic experts will spend approximately 40 hours to assist in the response to the cybersecurity intrusion, at an estimated cost of \$20,000.³⁰⁷ FTC staff estimates the

Dep't of Health & Human Servs., Office for Civil Rights, <https://www.hhs.gov/hipaa/for-professionals/breach-notification/index.html> ("A breach is, generally, an impermissible use or disclosure under the Privacy Rule that compromises the security or privacy of the protected health information.").

³⁰⁶ This estimate is the sum of 40 hours of marketing managerial time (at an average wage of \$76.10), 40 hours of computer programmer time (\$49.42), 20 hours of legal staff (\$78.74), and 50 hours of computer and information systems managerial time (\$83.49). See *Occupational Employment and Wage Statistics*, U.S. Bureau of Labor Statistics (May 2022), https://www.bls.gov/oes/current/oes_nat.htm#00-0000.

³⁰⁷ This estimate is the sum of 40 hours of forensic expert time at a cost of \$500 per hour,

services of forensic experts will be required in 60% of the 82 breaches. Based on the estimate that there will be 49 breaches per year requiring forensic experts (60% × 82 breaches), the annual hours burden for affected entities will be 1,960 hours (49 breaches requiring forensic experts × 40 hours) with an associated cost of \$980,000 (49 breaches requiring forensic experts × \$20,000).

Using the data on HIPAA-covered breach notices available from HHS for the years 2018–2023, FTC staff estimates the average number of individuals affected per breach is 93,497.³⁰⁸ Given an estimated 82 breaches per year, FTC staff estimates an average of 7,666,754 consumers per year will receive a breach notification (82 breaches × 93,497 individuals per breach).

Based on a recent study of data breach costs, FTC staff estimates the cost of providing notice to consumers to be \$11.87 per breached record.³⁰⁹ This estimate includes the costs of electronic notice, letters, outbound calls or general notice to data subjects; and engagement of outside experts.³¹⁰ Applied to the above-stated estimate of 7,666,754 consumers per year receiving breach notification yields an estimated total annual cost for all forms of notice to consumers of \$91,004,370 (7,666,754 consumers × \$11.87 per record). Accordingly, the estimated capital and non-labor costs total \$91,984,370 (\$90,000 + \$91,004,370).

FTC staff notes these estimates likely overstate the costs imposed by the amendments because FTC staff made conservative assumptions in developing many of the underlying estimates. Moreover, many entities covered by the amendments already have similar notification obligations under State data

which yields a total cost of \$20,000 (40 hours × \$500/hour).

³⁰⁸ HHS Breach Data, *supra* note 303. This analysis uses the last six years of HHS breach data to generate the average, in order to account for the variation in number of individuals affected by breaches observed in the HHS data over time.

³⁰⁹ See IBM Security, Costs of a Data Breach Report 2023 (2023), <https://www.ibm.com/reports/data-breach> (“2023 IBM Security Report”). The research for the 2023 IBM Security Report is conducted independently by the Ponemon Institute, and the results are reported and published by IBM Security. Figure 2 of the 2023 IBM Security Report shows that cost per record of a breach was \$165 per record in 2023, \$164 in 2022, and \$161 in 2021, resulting in an average cost of \$163.33. Figure 5 of the 2023 IBM Security Report shows that 8.3% (\$0.37m/\$4.45m) of the average cost of a data breach are due to “Notification” costs. The fraction of average breach costs due to “Notification” were 7.1% in 2022 and 6.4% in 2021 (IBM Security, Costs of a Data Breach Reports 2022 and 2021). Using the average of these numbers (7.27%), FTC staff estimates that notification costs per record across the three years are 7.27% × \$163.33 = \$11.87 per record.

³¹⁰ See 2023 IBM Security Report at 72.

breach laws.³¹¹ In addition, the Commission has taken several steps designed to limit the potential burden on covered entities that are required to provide notice, including by providing exemplar notices that entities may choose to use if they are required to provide notifications and expanding the use of electronic notifications.

IV. Regulatory Flexibility Act

The Regulatory Flexibility Act (RFA)³¹² requires that the Commission provide an Initial Regulatory Flexibility Analysis (“IRFA”) with a proposed rule and a Final Regulatory Flexibility Analysis (“FRFA”) with a final rule, unless the Commission certifies that the rule will not have a significant economic impact on a substantial number of small entities. As discussed in the IRFA, the Commission believes the final rule will not have a significant economic impact upon small entities.

In this document, the Commission largely adopts the amendments proposed in its NPRM. The Commission believes the amendments will not have a significant economic impact upon small entities, although they may affect a substantial number of small businesses. Among other things, the amendments clarify certain definitions, revise the disclosures that must accompany notice of a breach under the Rule, and modernize the methods of notice to allow additional use of electronic notice such as email by entities affected by a breach. In addition, the amendments improve the Rule’s readability by clarifying cross-references and adding statutory citations. The Commission does not anticipate that these changes will add significant additional costs for entities covered by the Rule, and by authorizing electronic notice in additional circumstances, the amendments may reduce costs for many entities covered by the Rule. Therefore, the Commission certifies that the amendments will not

³¹¹ Many State data breach notification statutes require notification when a breach occurs involving certain health or medical information of individuals in that State. See, e.g., Ala. Code 8–38–1 *et seq.*; Alaska Stat. 45.48.010 *et seq.*; Ariz. Rev. Stat. 18–551 *et seq.*; Ark. Code 4–110–101 *et seq.*; Cal. Civ. Code 1798.80 *et seq.*; Cal. Health & Safety Code 1280.15; Colo. Rev. Stat. 6–1–716; Del. Code Ann. tit. 6 12B–101 *et seq.*; D.C. Code 28–3851 *et seq.*; Fla. Stat. 501.171; 815 Ill. Comp. Stat. 530/5 *et seq.*; Md. Code Com. Law 14–3501 *et seq.*; Mo. Rev. Stat. 407.1500; Nev. Rev. Stat. 603A.010 *et seq.*; N.H. Rev. Stat. 359–C:19–C:21; N.H. Rev. Stat. 332–I:5; N.D. Cent. Code 51–30–01–07; Or. Rev. Stat. 646A.600–646A.628; R.I. Gen. Laws 11–49.3–1–11–49.3–6; SDCL 22–40–19–22–40–26; Tex. Bus. & Com. Code 521.002, 521.053, 521.151–152; 9 V.S.A. 2430, 2435; Va. Code 18.2–186.6; Va. Code 32.1–127.1:05; Va. Code 58.1–341.2; Wash. Rev. Code 19.255.010 *et seq.*

³¹² 5 U.S.C. 601–612.

have a significant economic impact on a substantial number of small entities. Although the Commission certifies under the RFA that the Rule will not have a significant impact on a substantial number of small entities, and hereby provides notice of that certification to the Small Business Administration (“SBA”), the Commission has determined, nonetheless, that it is appropriate to publish an FRFA to inquire into the impact of the proposed amendments on small entities.

A. Need for and Objectives of the Amendments

The objective of the amendments is to clarify existing notice obligations for entities covered by the Rule. The legal basis for the amendments is section 13407 of the Recovery Act.

B. Significant Issues Raised in Public Comments

Although the Commission received several comments that argued that the amendments would be burdensome for businesses, none argued specifically that smaller businesses in particular would be subject to special burdens. The Commission did not receive any comments filed by the Chief Counsel for Advocacy of the SBA.

C. Small Entities to Which the Amendments Will Apply

The amendments, like the current Rule, will apply to vendors of personal health records, PHR related entities, and third party service providers, including developers and purveyors of health apps, connected health devices, and similar technologies. As discussed in the Commission’s PRA estimates above, FTC staff estimates the amendments will apply to approximately 193,000 covered entities. The Commission estimates that a substantial number of these entities likely qualify as small businesses. According to the Statistics on Small Businesses Census data, approximately 94% of “Software Publishers” (the category to which health and fitness apps belong) are small businesses.³¹³

³¹³ 2017 SUSB Annual Data Tables by Establishment Industry, U.S. Census Bureau (May 2021), <https://www.census.gov/data/tables/2017/econ/susb/2017-susb-annual.html>, using “Data by Enterprise Receipts Size.” The U.S. Small Business Administration (“SBA”) categorizes Software Publishers as a small business if the annual receipts are less than \$41.5 million; the 2017 data is the most recent data available reporting receipts size.

D. Projected Reporting, Recordkeeping, and Other Compliance Requirements, Including Classes of Covered Small Entities and Professional Skills Needed To Comply

The Recovery Act and the amendments contain certain reporting requirements. The amendments will clarify which entities are subject to those reporting requirements. Specifically, the Act and amendments require vendors of personal health records and PHR related entities to provide notice to consumers, the Commission, and in some cases the media in the event of a breach of unsecured PHR identifiable health information. The Act and amendments also require third party service providers to provide notice to vendors of personal health records and PHR related entities in the event of such a breach. If a breach occurs, each entity covered by the Act and amendments will expend costs to determine the extent of the breach and the individuals affected. If the entity is a vendor of personal health records or a PHR related entity, additional costs will include the costs of preparing a breach notice, notifying the Commission, compiling a list of consumers to whom a breach notice must be sent, and sending a breach notice. Such entities may incur additional costs in locating consumers who cannot be reached, and in certain cases, posting a breach notice on a website, notifying consumers through media advertisements, or sending breach notices through press releases to media outlets.

In-house costs may include technical costs to determine the extent of breaches; investigative costs of conducting interviews and gathering information; administrative costs of compiling address lists; professional/legal costs of drafting the notice; and potentially, costs for postage, web posting, and/or advertising. Costs may also include the purchase of services of a forensic expert. As discussed in the context of the PRA, FTC staff estimates that compliance with these requirements will likely result in \$883,148 in labor costs and \$91,984,370 in capital and other non-labor costs. The estimated cost per covered entity is \$481 (the total labor, capital, and non-labor costs of \$92,867,518 divided by 193,000 covered entities). The SBA categorizes Software Publishers with annual receipts under \$41.5 million as a small business; the per entity cost of \$481 represents 0.0001% of this annual receipts threshold.

E. Significant Alternatives to the Amendments

In drafting the Rule, the Commission has made every effort to avoid unduly burdensome requirements for entities. In particular, the Commission believes that the changes to facilitate electronic notice will assist small entities by significantly reducing the costs of sending breach notices. In addition, the Commission is making available exemplar notices that entities covered by the Rule may use, in their discretion, to notify individuals. The Commission anticipates these exemplar notices will further reduce the burden on entities that are required to provide notice under the Rule. The Commission is not aware of alternative methods of compliance that will reduce the impact of the amendments on small entities, while also comporting with the Recovery Act. The statutory requirements are specific as to the timing, method, and content of notice.

V. Other Matters

Pursuant to the Congressional Review Act (5 U.S.C. 801 *et seq.*), the Office of Information and Regulatory Affairs designated this rule as not a “major rule,” as defined by 5 U.S.C. 804(2).

List of Subjects in 16 CFR Part 318

Breach, Consumer protection, Health, Privacy, Reporting and recordkeeping requirements, Trade practices.

■ Accordingly, the Federal Trade Commission revises and republishes 16 CFR part 318 to read as follows:

PART 318—HEALTH BREACH NOTIFICATION RULE

Sec.	
318.1	Purpose and scope.
318.2	Definitions.
318.3	Breach notification requirement.
318.4	Timeliness of notification.
318.5	Methods of notice.
318.6	Content of notice.
318.7	Enforcement.
318.8	Applicability date.
318.9	Sunset.

Authority: 42 U.S.C. 17937 and 17953.

§ 318.1 Purpose and scope.

(a) This part, which shall be called the “Health Breach Notification Rule,” implements section 13407 of the American Recovery and Reinvestment Act of 2009, 42 U.S.C. 17937. This part applies to foreign and domestic vendors of personal health records, PHR related entities, and third party service providers, irrespective of any jurisdictional tests in the Federal Trade Commission (FTC) Act, that maintain information of U.S. citizens or residents.

This part does not apply to HIPAA-covered entities, or to any other entity to the extent that it engages in activities as a business associate of a HIPAA-covered entity.

(b) This part preempts State law as set forth in section 13421 of the American Recovery and Reinvestment Act of 2009, 42 U.S.C 17951.

§ 318.2 Definitions.

Breach of security means, with respect to unsecured PHR identifiable health information of an individual in a personal health record, acquisition of such information without the authorization of the individual. Unauthorized acquisition will be presumed to include unauthorized access to unsecured PHR identifiable health information unless the vendor of personal health records, PHR related entity, or third party service provider that experienced the breach has reliable evidence showing that there has not been, or could not reasonably have been, unauthorized acquisition of such information. A breach of security includes an unauthorized acquisition of unsecured PHR identifiable health information in a personal health record that occurs as a result of a data breach or an unauthorized disclosure.

Business associate means a business associate under the Health Insurance Portability and Accountability Act, Public Law 104–191, 110 Stat. 1936, as defined in 45 CFR 160.103.

Clear and conspicuous means that a notice is reasonably understandable and designed to call attention to the nature and significance of the information in the notice.

(1) *Reasonably understandable.* You make your notice reasonably understandable if you:

- (i) Present the information in the notice in clear, concise sentences, paragraphs, and sections;
- (ii) Use short explanatory sentences or bullet lists whenever possible;
- (iii) Use definite, concrete, everyday words and active voice whenever possible;

- (iv) Avoid multiple negatives;
- (v) Avoid legal and highly technical business terminology whenever possible; and

- (vi) Avoid explanations that are imprecise and readily subject to different interpretations.

(2) *Designed to call attention.* You design your notice to call attention to the nature and significance of the information in it if you:

- (i) Use a plain-language heading to call attention to the notice;
- (ii) Use a typeface and type size that are easy to read;

(iii) Provide wide margins and ample line spacing;

(iv) Use boldface or italics for key words; and

(v) In a form that combines your notice with other information, use distinctive type size, style, and graphic devices, such as shading or sidebars, when you combine your notice with other information. The notice should stand out from any accompanying text or other visual elements so that it is easily noticed, read, and understood.

(3) *Notices on websites or within-application messaging.* If you provide a notice on a web page or using within-application messaging, you design your notice to call attention to the nature and significance of the information in it if you use text or visual cues to encourage scrolling down the page if necessary to view the entire notice and ensure that other elements on the website or software application (such as text, graphics, hyperlinks, or sound) do not distract attention from the notice, and you either:

(i) Place the notice on a screen that consumers frequently access, such as a page on which transactions are conducted; or

(ii) Place a link on a screen that consumers frequently access, such as a page on which transactions are conducted, that connects directly to the notice and is labeled appropriately to convey the importance, nature and relevance of the notice.

Covered health care provider means a provider of services (as defined in 42 U.S.C. 1395x(u)), a provider of medical or other health services (as defined in 42 U.S.C. 1395x(s)), or any other entity furnishing health care services or supplies.

Electronic mail means email in combination with one or more of the following: text message, within-application messaging, or electronic banner.

Health care services or supplies means any online service such as a website, mobile application, or internet-connected device that provides mechanisms to track diseases, health conditions, diagnoses or diagnostic testing, treatment, medications, vital signs, symptoms, bodily functions, fitness, fertility, sexual health, sleep, mental health, genetic information, diet, or that provides other health-related services or tools.

HIPAA-covered entity means a covered entity under the Health Insurance Portability and Accountability Act (HIPAA), Public Law 104–191, 110 Stat. 1936, as defined in 45 CFR 160.103.

Personal health record (PHR) means an electronic record of PHR identifiable health information on an individual that has the technical capacity to draw information from multiple sources and that is managed, shared, and controlled by or primarily for the individual.

PHR identifiable health information means information that:

(1) Relates to the past, present, or future physical or mental 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; and

(i) Identifies the individual; or
(ii) With respect to which there is a reasonable basis to believe that the information can be used to identify the individual; and

(2) Is created or received by a:

(i) Covered health care provider;
(ii) Health plan (as defined in 42 U.S.C. 1320d(5));
(iii) Employer; or
(iv) Health care clearinghouse (as defined in 42 U.S.C. 1320d(2)); and
(3) With respect to an individual, includes information that is provided by or on behalf of the individual.

PHR related entity means an entity, other than a HIPAA-covered entity or an entity to the extent that it engages in activities as a business associate of a HIPAA-covered entity, that:

(1) Offers products or services through the website, including any online service, of a vendor of personal health records;

(2) Offers products or services through the websites, including any online service, of HIPAA-covered entities that offer individuals personal health records; or

(3) Accesses unsecured PHR identifiable health information in a personal health record or sends unsecured PHR identifiable health information to a personal health record.

State means any of the several States, the District of Columbia, Puerto Rico, the Virgin Islands, Guam, American Samoa, and the Northern Mariana Islands.

Third party service provider means an entity that:

(1) Provides services to a vendor of personal health records in connection with the offering or maintenance of a personal health record or to a PHR related entity in connection with a product or service offered by that entity; and

(2) Accesses, maintains, retains, modifies, records, stores, destroys, or otherwise holds, uses, or discloses unsecured PHR identifiable health information as a result of such services.

Unsecured means PHR identifiable information that is not protected through the use of a technology or methodology specified by the Secretary of Health and Human Services in the guidance issued under section 13402(h)(2) of the American Reinvestment and Recovery Act of 2009, 42 U.S.C. 17932(h)(2).

Vendor of personal health records means an entity, other than a HIPAA-covered entity or an entity to the extent that it engages in activities as a business associate of a HIPAA-covered entity, that offers or maintains a personal health record.

§ 318.3 Breach notification requirement.

(a) *In general.* In accordance with §§ 318.4 (regarding timeliness of notification), 318.5 (regarding methods of notice), and 318.6 (regarding content of notice), each vendor of personal health records, following the discovery of a breach of security of unsecured PHR identifiable health information that is in a personal health record maintained or offered by such vendor, and each PHR related entity, following the discovery of a breach of security of such information that is obtained through a product or service provided by such entity, shall:

(1) Notify each individual who is a citizen or resident of the United States whose unsecured PHR identifiable health information was acquired by an unauthorized person as a result of such breach of security;

(2) Notify the Federal Trade Commission; and

(3) Notify prominent media outlets serving a State or jurisdiction, following the discovery of a breach of security, if the unsecured PHR identifiable health information of 500 or more residents of such State or jurisdiction is, or is reasonably believed to have been, acquired during such breach.

(b) *Third party service providers.* A third party service provider shall, following the discovery of a breach of security, provide notice of the breach to an official designated in a written contract by the vendor of personal health records or the PHR related entity to receive such notices or, if such a designation is not made, to a senior official at the vendor of personal health records or PHR related entity to which it provides services, and obtain acknowledgment from such official that such notice was received. Such notification shall include the identification of each customer of the vendor of personal health records or PHR related entity whose unsecured PHR identifiable health information has been, or is reasonably believed to have been, acquired during such breach. For

purposes of ensuring implementation of this paragraph (b), vendors of personal health records and PHR related entities shall notify third party service providers of their status as vendors of personal health records or PHR related entities subject to this part. While some third party service providers may access unsecured PHR identifiable health information in the course of providing services, this does not render the third party service provider a PHR related entity.

(c) *Breaches treated as discovered.* A breach of security shall be treated as discovered as of the first day on which such breach is known or reasonably should have been known to the vendor of personal health records, PHR related entity, or third party service provider, respectively. Such vendor, entity, or third party service provider shall be deemed to have knowledge of a breach if such breach is known, or reasonably should have been known, to any person, other than the person committing the breach, who is an employee, officer, or other agent of such vendor of personal health records, PHR related entity, or third party service provider.

§ 318.4 Timeliness of notification.

(a) *In general.* Except as provided in paragraph (d) of this section (exception for law enforcement), all notifications required under § 318.3(a)(1) (required notice to individuals), (a)(3) (required notice to media), and (b) (required notice by third party service providers), shall be sent without unreasonable delay and in no case later than 60 calendar days after the discovery of a breach of security.

(b) *Timing of notice to FTC.* All notifications required under § 318.5(c) (regarding notice to FTC) involving the unsecured PHR identifiable health information of 500 or more individuals shall be provided contemporaneously with the notice required by paragraph (a) of this section. All logged notifications required under § 318.5(c) (regarding notice to FTC) involving the unsecured PHR identifiable health information of fewer than 500 individuals may be sent annually to the Federal Trade Commission no later than 60 calendar days following the end of the calendar year.

(c) *Burden of proof.* The vendor of personal health records, PHR related entity, and third party service provider involved shall have the burden of demonstrating that all notifications were made as required under this part, including evidence demonstrating the necessity of any delay.

(d) *Law enforcement exception.* If a law enforcement official determines that

a notification, notice, or posting required under this part would impede a criminal investigation or cause damage to national security, such notification, notice, or posting shall be delayed. This paragraph (d) shall be implemented in the same manner as provided under 45 CFR 164.528(a)(2), in the case of a disclosure covered under § 164.528(a)(2).

§ 318.5 Methods of notice.

(a) *Individual notice.* A vendor of personal health records or PHR related entity that discovers a breach of security shall provide notice of such breach to an individual promptly, as described in § 318.4 (regarding timeliness of notification), and in the following form:

(1) Written notice at the last known address of the individual. Written notice may be sent by electronic mail if the individual has specified electronic mail as the primary method of communication. Any written notice sent by electronic mail must be Clear and Conspicuous. Where notice via electronic mail is not available or the individual has not specified electronic mail as the primary method of communication, a vendor of personal health records or PHR related entity may provide notice by first-class mail at the last known address of the individual. If the individual is deceased, the vendor of personal health records or PHR related entity that discovered the breach must provide such notice to the next of kin of the individual if the individual had provided contact information for his or her next of kin, along with authorization to contact them. The notice may be provided in one or more mailings as information is available.

(2) If, after making reasonable efforts to contact all individuals to whom notice is required under § 318.3(a), through the means provided in paragraph (a)(1) of this section, the vendor of personal health records or PHR related entity finds that contact information for ten or more individuals is insufficient or out-of-date, the vendor of personal health records or PHR related entity shall provide substitute notice, which shall be reasonably calculated to reach the individuals affected by the breach, in the following form:

(i) Through a conspicuous posting for a period of 90 days on the home page of its website; or

(ii) In major print or broadcast media, including major media in geographic areas where the individuals affected by the breach likely reside. Such a notice in media or web posting shall include a toll-free phone number, which shall

remain active for at least 90 days, where an individual can learn if the individual's unsecured PHR identifiable health information may have been included in the breach.

(3) In any case deemed by the vendor of personal health records or PHR related entity to require urgency because of possible imminent misuse of unsecured PHR identifiable health information, that entity may provide information to individuals by telephone or other means, as appropriate, in addition to notice provided under paragraph (a)(1) of this section.

(b) *Notice to media.* As described in § 318.3(a)(3), a vendor of personal health records or PHR related entity shall provide notice to prominent media outlets serving a State or jurisdiction, following the discovery of a breach of security, if the unsecured PHR identifiable health information of 500 or more residents of such State or jurisdiction is, or is reasonably believed to have been, acquired during such breach.

(c) *Notice to FTC.* Vendors of personal health records and PHR related entities shall provide notice to the Federal Trade Commission following the discovery of a breach of security, as described in § 318.4(b) (regarding timing of notice to FTC). If the breach involves the unsecured PHR identifiable health information of fewer than 500 individuals, the vendor of personal health records or PHR related entity may maintain a log of any such breach and submit such a log annually to the Federal Trade Commission as described in § 318.4(b) (regarding timing of notice to FTC), documenting breaches from the preceding calendar year. All notices pursuant to this paragraph (c) shall be provided according to instructions at the Federal Trade Commission's website.

§ 318.6 Content of notice.

Regardless of the method by which notice is provided to individuals under § 318.5 (regarding methods of notice), notice of a breach of security shall be in plain language and include, to the extent possible, the following:

(a) A brief description of what happened, including: the date of the breach and the date of the discovery of the breach, if known; and the full name or identity (or, where providing the full name or identity would pose a risk to individuals or the entity providing notice, a description) of any third parties that acquired unsecured PHR identifiable health information as a result of a breach of security, if this information is known to the vendor of

personal health records or PHR related entity;

(b) A description of the types of unsecured PHR identifiable health information that were involved in the breach (such as but not limited to full name, Social Security number, date of birth, home address, account number, health diagnosis or condition, lab results, medications, other treatment information, the individual's use of a health-related mobile application, or device identifier (in combination with another data element));

(c) Steps individuals should take to protect themselves from potential harm resulting from the breach;

(d) A brief description of what the entity that experienced the breach is doing to investigate the breach, to mitigate harm, to protect against any further breaches, and to protect affected individuals, such as offering credit monitoring or other services; and

(e) Contact procedures for individuals to ask questions or learn additional information, which must include two or more of the following: toll-free telephone number; email address; website; within-application; or postal address.

§ 318.7 Enforcement.

Any violation of this part shall be treated as a violation of a rule promulgated under section 18 of the Federal Trade Commission Act, 15 U.S.C. 57a, regarding unfair or deceptive acts or practices, and thus subject to civil penalties (as adjusted for inflation pursuant to § 1.98 of this chapter), and the Commission will enforce this part in the same manner, by the same means, and with the same jurisdiction, powers, and duties as are available to it pursuant to the Federal Trade Commission Act, 15 U.S.C. 41 *et seq.*

§ 318.8 Applicability date.

This part shall apply to breaches of security that are discovered on or after September 24, 2009.

§ 318.9 Sunset.

If new legislation is enacted establishing requirements for notification in the case of a breach of security that apply to entities covered by this part, the provisions of this part shall not apply to breaches of security discovered on or after the effective date of regulations implementing such legislation.

By direction of the Commission, Commissioners Holyoak and Ferguson dissenting.

April J. Tabor,
Secretary.

Note: The following appendices will not appear in the Code of Federal Regulations.

Appendix A—Health Breach Notification Rule Exemplar Notices

The notices below are intended to be examples of notifications that entities may use, in their discretion, to notify individuals of a breach of security pursuant to the Health Breach Notification Rule. The examples below are for illustrative purposes only. You should tailor any notices to the particular facts and circumstances of your breach. While your notice must comply with the Health Breach Notification Rule, you are not required to use the notices below.

Mobile Text Message and In-App Message Exemplars

Text Message Notification Exemplar 1

Due to a security breach on our system, *the health information you shared with us through [name of product] is now in the hands of unknown attackers*. Visit [add non-clickable URL] to learn what happened, how it affects you, and what you can do to protect your information. We also sent you an email with additional information.

Text Message Notification Exemplar 2

You shared health information with us when you used [product name]. *We discovered that we shared your health information with third parties for [describe why the company shared the info] without your permission*. Visit [add non-clickable URL] to learn what happened, how it affects you, and what you can do to protect your information. We also sent you an email with more information.

In-App Message Notification Exemplar 1

Due to a security breach on our system, *the health information you shared with us through [name of product] is now in the hands of unknown attackers*. This could include your [Add specifics—for example, your name, email, address, blood pressure data]. Visit [URL] to learn what happened, how it affects you, and what you can do to protect your information. We also sent you an email with additional information.

In-App Message Notification Exemplar 2

You shared health information with us when you used [product name]. *We discovered that we shared your health information with third parties for [if known, describe why the company shared the info] without your permission*. This could include your [Add specifics—for example, your name, email, address, blood pressure data]. Visit [URL] to learn what happened, how it affects you, and what you can do to protect your information. We also sent you an email with additional information.

Web Banner Exemplars

Web Banner Notification Exemplar 1

Due to a security breach on our system, *the health information you shared with us through [name of product] is now in the hands of unknown attackers*. This could include your [Add specifics—for example, your name, email, address, blood pressure data]. Visit [URL] to learn what happened, how it affects you, and what you can do to protect your information.

- **Recommend:** Include clear “Take action” call to action button, such as the example below:

Due to a security breach on our system, **the health information you shared with us through [name of product] is now in the hands of unknown attackers**. This could include your [Add specifics – **for example, your name, email, address, blood pressure data**]. Visit [URL] to learn what happened, how it affects you, and what you can do to protect your information.

Take action

Web Banner Notification Exemplar 2

You shared health information with us when you used [product name]. *We discovered that we shared your health information with third parties for [if known,*

describe why the company shared the info] without your permission. This could include your [Add specifics—for example, your name, email, address, blood pressure data]. Visit [URL] to learn what happened, how it

affects you, and what you can do to protect your information.

- **Recommend:** Include clear “Take action” call to action button, such as the example below:

You shared health information with us when you used [product name]. **We discovered that we shared your health information with third parties for [if known, describe why the company shared the info] without your permission.** This could include your [Add specifics – for example, your name, email, address, blood pressure data]. Visit [URL] to learn what happened, how it affects you, and what you can do to protect your information.

Take action

Email Exemplars

Exemplar Email Notice 1

Email Sender: [Company] <company email>
Email Subject Line: [Company] Breach of Your Health Information
 Dear [Name],

We are contacting you because an attacker recently gained unauthorized access to our system and stole health information about our customers, including you.

What happened and what it means for you
 On [March 1, 2024], we learned that an attacker had accessed a file containing our customers' health information on [February 28, 2024]. The file included your name, the name of your health insurance company, your date of birth, and your group or policy number.

What you can do to protect yourself

You can take steps now to reduce the risk of identity theft.

1. *Review your medical records, statements, and bills for signs that someone is using your information.* Under the health privacy law known as HIPAA, you have the right to access your medical records. Get your records and review them for any treatments or doctor visits you don't recognize. If you find any, report them to your healthcare provider in writing. Then go to www.IdentityTheft.gov/steps to see what other steps you can take to limit the damage.

Also review the Explanation of Benefits statement your insurer sends you when it pays for medical care.

Some criminals wait before using stolen information so keep monitoring your benefits and bills.

2. *Review your credit reports for errors.* You can get your free credit reports from the three credit bureaus at www.annualcreditreport.com or call 1-877-322-8228. Look for medical billing errors, like medical debt collection notices that you don't recognize. Report any medical billing errors to all three credit bureaus by following the "What To Do Next" steps on www.IdentityTheft.gov.

3. *Sign up for free credit monitoring to detect suspicious activity.* Credit monitoring detects and alerts you about activity on your credit reports. Activity you don't recognize could be a sign that someone stole your identity. We're offering free credit monitoring for two years through [name of service]. Learn more and sign up at [URL].

4. *Consider freezing your credit report or placing a fraud alert on your credit report.* A credit report freeze means potential creditors can't get your credit report without your permission. That makes it less likely that an identity thief can open new accounts in your name. A freeze remains in place until you ask the credit bureau to temporarily lift it or remove it.

A fraud alert will make it harder for someone to open a new credit account in your name. It tells creditors to contact you before they open any new accounts in your name or change your accounts. A fraud alert lasts for one year. After a year, you can renew it.

To freeze your credit report, contact *each of the three credit bureaus*, Equifax, Experian, and TransUnion.

To place a fraud alert, contact *any one of the three credit bureaus*, Equifax, Experian, and TransUnion. As soon as one credit bureau confirms your fraud alert, the others are notified to place fraud alerts on your credit report.

Credit bureau contact information

Equifax, www.equifax.com/personal/credit-report-services, 1-800-685-1111

Experian, www.experian.com/help, 1-888-397-3742

TransUnion, www.transunion.com/credit-help, 1-888-909-8872

Learn more about how credit report freezes and fraud alerts can protect you from identity theft or prevent further misuse of your personal information at www.consumer.ftc.gov/articles/what-know-about-credit-freezes-and-fraud-alerts.

What we are doing in response

We hired security experts to secure our system. We are working with law enforcement to find the attacker. And we are investigating whether we made mistakes that made it possible for the attackers to get in.

Learn more about the breach.

Go to [URL] to learn more about what happened and what you can do to protect yourself. If we have any updates, we will post them there.

If you have questions or concerns, call us at [telephone number], email us at [address], or go to [URL].

Sincerely,
 First name Last Name
 [Role], [Company]

Exemplar Email Notice 2

Email Sender: [Company] <company email>
Email Subject Line: Unauthorized disclosure of your health information by [Company]
 Dear [Name],

We are contacting you because you use our company's app [name of app]. When you downloaded our app, we promised to keep your personal health information private. Instead, we disclosed health information about you without your approval.

What happened?

We told [insert Company name, identity, or, where providing full name or identity would pose a risk to individuals or the entity providing notice, a description of type of company] that you use our app, and between [January 10, 2024] and [March 1, 2024], we

gave them your name and your email address.

We gave [insert Company name, identity, or where providing full name or identity would pose a risk to individuals or the entity providing notice, a description of type of company] this information so they could use it for advertising and marketing purposes. For example, to target you for ads for cancer drugs.

What we are doing in response

We will stop selling or sharing your health information with other companies. We will stop using your health information for advertising or marketing purposes. We have asked Company XYZ to delete your health information, but it's possible they could continue to use it for advertising and marketing.

What you can do

We made important changes to our app to fix this problem. Download the latest updates to our app then review your privacy settings. You can also contact Company XYZ to request that it delete your data.

Learn more

Learn more about our privacy and security practices at [URL]. If we have any updates, we will post them there.

If you have any questions or concerns, call us at [telephone number] or email us at [address].

Sincerely,
 First name Last Name
 [Role], [Company]

Exemplar Email Notice 3

Email Sender: [Company] <company email>
Email Subject Line: [Company] Breach of Your Health Information
 Dear [Name],

We are contacting you about a breach of your health information collected through the [product], a device sold by our company, [Company].

What happened?

On [March 1, 2024], we discovered that our employee had accidentally posted a database online on [February 28, 2024]. That database included your name, your credit or debit card information, and your blood pressure readings. We don't know if anyone else found the database and saw your information. If someone found the database, they could use personal information to steal your identity or make unauthorized charges in your name.

What you can do to protect yourself

You can take steps now to reduce the risk of identity theft.

1. *Get your free credit report and review it for signs of identity theft.* Order your free credit report at www.annualcreditreport.com. Review it for accounts and activity you don't recognize. Recheck your credit reports periodically.

2. Consider freezing your credit report or placing a fraud alert on your credit report. A credit report freeze means potential creditors can't get your credit report without your permission. That makes it less likely that an identity thief can open new accounts in your name. A freeze remains in place until you ask the credit bureau to temporarily lift it or remove it.

A fraud alert will make it harder for someone to open a new credit account in your name. It tells creditors to contact you before they open any new accounts in your name or change your accounts. A fraud alert lasts for one year. After a year, you can renew it.

To freeze your credit report, contact *each of the three credit bureaus*, Equifax, Experian, and TransUnion.

To place a fraud alert, contact *any one of the three credit bureaus*, Equifax, Experian, and TransUnion. As soon as one credit bureau confirms your fraud alert, the others are notified to place fraud alerts on your credit report.

Credit bureau contact information

Equifax, www.equifax.com/personal/credit-report-services, 1-800-685-1111

Experian, www.experian.com/help, 1-888-397-3742

TransUnion, www.transunion.com/credit-help, 1-888-909-8872

Learn more about how credit report freezes and fraud alerts can protect you from identity theft or prevent further misuse of your personal information at www.consumer.ftc.gov/articles/what-know-about-credit-freezes-and-fraud-alerts.

3. Sign up for free credit monitoring to detect suspicious activity. Credit monitoring detects and alerts you about activity on your credit reports. Activity you don't recognize could be a sign that someone stole your identity. We're offering free credit monitoring for two years through [name of service]. Learn more and sign up at [URL].

What we are doing in response

We are investigating our mistakes. We know the database shouldn't have been online and it should have been encrypted. We are making changes to prevent this from happening again.

We are working with experts to secure our system. We are reviewing our databases to make sure we store health information securely.

Learn more about the breach.

Go to [URL] to learn more about what happened and what you can do to protect yourself. If we have any updates, we will post them there.

If you have questions or concerns, call us at [telephone number], email us at [address], or go to [URL].

Sincerely,

First name Last Name

[Role], [Company]

Appendix B—Joint Statement by FTC Chair and Commissioners

Joint Statement of Chair Lina M. Khan, Commissioner Rebecca Kelly Slaughter, and Commissioner Alvaro M. Bedoya

Today, the FTC finalizes an update to the Health Breach Notification Rule (“the Final

Rule”) that ensures its protections keep pace with the rapid proliferation of digital health records. We do so to fulfill a clear statutory directive given to us by Congress.

In 2009, as part of the American Recovery and Reinvestment Act (“ARRA”), Congress passed the Health Information Technology for Economic and Clinical Health Act (“HITECH Act”).³¹⁴ Among other things, the HITECH Act sought to fill the gaps left by the privacy and security protections created under the Health Insurance Portability and Accountability Act (“HIPAA”), which was passed more than a decade earlier.³¹⁵ Specifically, it expanded the kinds of entities subject to the privacy and security provisions of HIPAA,³¹⁶ gave state attorneys general enforcement powers,³¹⁷ and—most relevant here—directed the Commission to issue a rule requiring entities not covered by HIPAA to provide notification of any breach of unsecured health records.³¹⁸ The Commission issued the original rule in 2009.³¹⁹ In 2020, the Commission initiated its regular decennial rule review and, in 2021, the Commission issued a policy statement clarifying how the rule applies to health apps and other connected devices.³²⁰ In the years since, the Commission has brought enforcement actions against health apps alleging violations of the Health Breach Notification Rule.³²¹ Today's issuance of the Final Rule codifies this approach, honoring the statutory directive that people must be notified when their health records are breached.

The dissent argues that the Commission's action “exceeds the Commission's statutory authority.”³²² But its analysis contravenes a plain reading of the statute.

In the HITECH Act, Congress directed the FTC to issue rules requiring vendors of

³¹⁴ Am. Recovery and Reinvestment Act of 2009, Public Law 111–5, 123 Stat. 115 (2009) at Sec. 13400 *et seq.*

³¹⁵ Health Insurance Portability and Accountability Act, Public Law 104–191, 110 Stat. 1936, 2022 (1996) at Sec. 1171, codified at 42 U.S.C. 1320d.

³¹⁶ Health Information Technology for Economic and Clinical Health Act, Public Law 111–5, Div. A, Title XIII, Subtitle D, sections 13401 and 13404 (codified at 42 U.S.C. 17937(a))

³¹⁷ *Id.* 13410(e).

³¹⁸ *Id.* 13407(g)(1).

³¹⁹ 74 FR 42962 (Aug. 25, 2009).

³²⁰ Statement of the Commission on Breaches by Health Apps and Other Connected Devices (Sept. 15, 2021), https://www.ftc.gov/system/files/documents/public_statements/1596364/statement_of_the_commission_on_breaches_by_health_apps_and_other_connected_devices.pdf.

³²¹ See, e.g., Fed. Trade Comm'n, FTC Enforcement Action to Bar GoodRx from Sharing Consumers' Sensitive Health Info for Advertising (Feb. 1, 2023), <https://www.ftc.gov/news-events/news/press-releases/2023/02/ftc-enforcement-action-bar-goodrx-sharing-consumers-sensitive-health-info-advertising>; Fed. Trade Comm'n, Ovulation Tracking App Premom Will Be Barred from Sharing Health Data for Advertising Under Proposed FTC Order (May 17, 2023), <https://www.ftc.gov/news-events/news/press-releases/2023/05/ovulation-tracking-app-premom-will-be-barred-sharing-health-data-advertising-under-proposed-ftc>.

³²² Dissenting Statement of Comm'rs Melissa Holyoak and Andrew Ferguson at 1 (Apr. 25, 2024) (hereinafter “Dissent”).

personal health records (“PHR”) to notify consumers and the FTC following “a breach of security of unsecured PHR identifiable health information.”³²³ The statute defines the term “PHR identifiable health information” as “individually identifiable health information, as defined in section 1320d(6) of this title.”³²⁴ Section 1320d(6), a portion of the Social Security Act created by HIPAA, defines “individually identifiable health information” as “any information . . . that is created or received by a health care provider, health plan, employer, or health care clearinghouse.”³²⁵ Section 1320d(3), another section of the Social Security Act created by HIPAA, defines “health care provider” as, first, “a provider of services” as defined in section 1395x(u);³²⁶ second, “a provider of medical or other health services” as defined in section 1395x(s);³²⁷ and, third, “any other person furnishing health care services or supplies.”³²⁸

The term “health care services or supplies,” undefined in the statute, is defined in the Final Rule as follows:

Health care services or supplies means any online service such as a website, mobile application, or internet-connected device that provides mechanisms to track diseases, health conditions, diagnoses or diagnostic testing, treatment, medications, vital signs, symptoms, bodily functions, fitness, fertility, sexual health, sleep, mental health, genetic information, diet, or that provides other health-related services or tools.³²⁹

The dissent argues that this definition violates certain canons of statutory construction.³³⁰ But its effort to cabin the third category of HIPAA's “health care provider” reads it out of existence, violating the canon that holds interpretations giving effect to every clause of a statute are superior to those that render distinct clauses superfluous.³³¹ Specifically, the second

³²³ Health Information Technology for Economic and Clinical Health Act, Public Law 111–5, Div. A, Title XIII, Subtitle D, section 13407 (codified at 42 U.S.C. 17937(a)).

³²⁴ 42 U.S.C. 17937(f)(2).

³²⁵ 42 U.S.C. 1320d(6).

³²⁶ See 42 U.S.C. 1395x(u) (“The term “provider of services” means a hospital, critical access hospital, rural emergency hospital, skilled nursing facility, comprehensive outpatient rehabilitation facility, home health agency, hospice program, or, for purposes of section 1395(g) and section 1395n(e) of this title, a fund.”).

³²⁷ 42 U.S.C. 1395x(s) (listing a vast array of services, tests, supplies, and measurements, comprising over 2000 words and 15 categories, one of which has over 30 subcategories).

³²⁸ 42 U.S.C. 1320d(3) (emphasis added).

³²⁹ HBNR Final Rule § 318.2(e).

³³⁰ Dissent at 2 (“When a statute contains a list, “each word in that list presumptively has a ‘similar’ meaning” under the canon of *noscitur a sociis*. And when a general term follows a list of specific terms, the *ejusdem generis* canon teaches that the general term “should usually be read in light of those specific words to mean something ‘similar.’” Together, these canons instruct that the final category of health care provider that includes the general term “other person” must be similar to the more specific terms that precede it.” (citations omitted)).

³³¹ *Marx v. Gen. Revenue Corp.*, 568 U.S. 371, 386 (2013) (Thomas, J.) (“Finally, the canon against

category of “health care provider” already comprises a vast array of “provider[s] of medical and other services.”³³² If the Commission were to interpret the third category as comprising, as the dissent recommends, only “traditional forms of health care providers,” this distinct provision would be entirely redundant.

The dissent’s approach also fails to give meaning to other textual differences between the second and third category. The second category in the definition of “health care provider” discusses a “provider” and “medical” services.³³³ The third category, by contrast, drops the terms “provider” in favor of “person furnishing” and drops “medical” in favor of “health care.”³³⁴ Honoring the materially different words of the statute requires us to read these two categories as covering distinct, not entirely overlapping, entities.³³⁵ The Final Rule faithfully follows these textual markers and identifies specific services and tools that comprise “health care services or supplies.”³³⁶ Contrary to this plain reading of the text, the dissent claims that Congress must have meant for this provision to apply only to “traditional forms of health care providers.”³³⁷ But we cannot subordinate the text of the statute to speculative accounts of what Congress intended.

The dissent also notes that the Department of Health and Human Services (“HHS”) “has never interpreted the term ‘health care provider’ to reach the expansive, creative conclusion that the Commission does today.”³³⁸ HHS has, however, interpreted “health care provider,” and its interpretation of this term is consistent with the

surplusage is strongest when an interpretation would render superfluous another part of the same statutory scheme.”).

³³² 42 U.S.C. 1320(d)(3) (citing 42 U.S.C. 1395x(u)).

³³³ 42 U.S.C. 1320(d)(3).

³³⁴ *Id.*

³³⁵ See *Southwest Airlines Co. v. Saxon*, 596 U.S. 450, 458 (2022) (Thomas, J.) (“Where a document has used one term in one place, and a materially different term in another, the presumption is that the different term denotes a different idea” (cleaned up)).

³³⁶ In addition to defining this term by identifying specific services, the Final Rule actually also narrowed the definition originally proposed in the NPRM, by eliminating “includes” from the definition. SBP at 27 (“[T]he Commission has substituted the word ‘means’ for ‘includes’ to avoid implying greater breadth than the Commission intends.”).

³³⁷ Dissent at 3. This rejection of the text of the statute, in favor of vague speculation about what Congress intended, mirrors the argument advanced by the Chamber of Commerce (“the Chamber”). The Chamber purports to rely on a “plain text reading” of the statute but immediately switches—in the very same sentence—to vague notions of Congressional intent: “It is clear from a plain text reading of both the HITECH Act and HIPAA [*sic*] that Congress intended for the HBNR to cover health records more aligned with the provision of health services provided by traditional health providers at a time when it was attempting to digitize traditional health records.” Comment submitted by U.S. Chamber of Com., Health Breach Notification Rule, *Regulations.gov* (Aug. 8, 2023) at 3, <https://www.regulations.gov/comment/FTC-2023-0037-010>.

³³⁸ Dissent at 3.

³³⁹ Dissent at 3.

Commission’s definition.³³⁹ In the HIPAA Privacy Rule, HHS defines first two categories of “health care provider” using the same language as the statute, but the third category is changed from “any other person furnishing health care services or supplies” to “any other person or organization who furnishes, bills, or is paid for health care in the normal course of business.”³⁴⁰ HHS also defines “health care” broadly, as any “care, services, or supplies related to the health of an individual.”³⁴¹

Notably, in its 1999 Notice of Proposed Rulemaking for the HIPAA Privacy Rule, HHS originally had proposed to define the term “health care” as constituting “the provision of care, services, or supplies. . . .”³⁴² But, in its final rule, HHS eliminated the concept of “provision” in order to distinguish the broader term of “health care” from the narrower term “treatment.”³⁴³ HHS explained: “We delete the term ‘providing’ from the definition [of health care] to delineate more clearly the relationship between ‘treatment,’ as the term is defined in § 164.501, and ‘health care.’”³⁴⁴ HHS defined “treatment,” in contrast to “health care,” as “the provision, coordination, or management of health care and related services.”³⁴⁵ In short, HHS defines “health care” broadly, covering all aspects related to the health of an individual, and defines “treatment” more narrowly, referring to the provision of medical care to an individual. The dissent’s proposal to narrow the third category of “health care provider” to “traditional forms of health care providers” closely mirrors the approach that HHS rejected when it defined this term.³⁴⁶

The dissent also claims that changing the phrase “can be drawn” to “has the technical capacity to draw” violates the surplusage canon because it renders the limitation meaningless as to health apps, because “virtually every app has the technical capacity to draw some information from more than one source.”³⁴⁷ This argument

³³⁹ That the HIPAA Privacy rule has a narrower overall scope does not change this fact.

³⁴⁰ 45 CFR 160.103.

³⁴¹ *Id.* (emphasis added). The dissent asserts that we “mischaracterize[] the HIPAA Privacy Rule, which only applies to HIPAA ‘covered entities’ and their ‘business associates,’—*i.e.*, to traditional health care providers, that do not include the broad swath of app developers the Final Rule will encompass.” Dissent at 4 n.24 (internal citations omitted). It is not clear how this qualifies as a mischaracterization. Indeed, this is precisely the stated purpose of the Health Breach Notification Rule: To cover entities that HIPAA does not. The dissent also notes that we fail to recognize that HHS provides two examples of “health care.” But, HHS expressly states that the definition “includes, but is not limited to” these categories. 45 CFR 160.103. In any case, the breadth of these categories further underscores the expansive scope of HHS’s definition of health care. *Id.*

³⁴² Dissent at 2.

³⁴³ Proposed Rule, Standards for Privacy of Individually Identifiable Health Information, 64 FR 59918, 60049 (Nov. 3, 1999) (emphasis added).

³⁴⁴ 65 FR 82462, 82477.

³⁴⁵ *Id.*

³⁴⁶ 45 CFR 164.501.

³⁴⁷ Dissent at 2.

³⁴⁸ Dissent at 4.

fails for two reasons. First, as the Statement of Basis and Purpose (“SBP”) explains, there are products and services that do not satisfy this requirement.³⁴⁸ Second, even if the definition did reach every health app, that would not itself suggest that the Final Rule’s definition was wrongly crafted. Rather, it would reflect the rapid growth in digital applications and services related to consumers’ health.³⁴⁹

The practical ramifications of the dissent’s legal shortcomings are significant.

Just last year, the Commission brought an action against Easy Healthcare Corporation, alleging privacy violations by its fertility tracking application Premom.³⁵⁰ As laid out in the complaint, Premom—which encourages users to provide information about their menstrual cycles, fertility, and pregnancy, as well as to import their data from other services, such as Apple Health—shared information with advertisers and China-based companies through software development kits (“SDKs”) embedded in the application. The Commission’s eight-count complaint against Easy Healthcare reflected the seriousness of this misconduct, charging the business with deceptive and unfair practices, as well as a violation of the Health Breach Notification Rule, which triggered civil penalties.

Under the dissent’s analysis of health care services or supplies, the developer of the Premom application—Easy Healthcare—would not be covered by the Health Breach Notification Rule. This reading would mean that when companies like Easy Healthcare suffer a breach that may divulge health information to companies located in China, the Health Breach Notification Rule would not require them to disclose the breach to its users. It would also mean that when Easy Healthcare broadcasts women’s sensitive health data across the vast commercial surveillance network propped up by SDKs and ad networks, the Health Breach Notification Rule would not require Easy Healthcare to alert women. Today’s Final Rule rejects this atextual and cramped reading of the law, ensuring that businesses that hold themselves out as health care services companies—like Easy Healthcare—

³⁴⁸ SBP at 29–30.

³⁴⁹ The dissent’s argument anachronistically assumes that Congress intended for the Rule to cover some health apps, but not other health apps. But, in fact, the Apple and Google app stores were in their infancy when Congress drafted this legislation in 2009, and so there is no indication that Congress was thinking about specific health apps at all. To the extent the dissent’s argument is that Congress simply did not anticipate the vast number of products that would end up covered by the broad category of “supplies and services,” it is not within the Commission’s authority to re-write the statute based on the Commission’s belief of what Congress would have wanted. *MCI Telecomms. Corp. v. Am. Telephone & Telegraph Co.*, 512 U.S. 218, 229 (1994) (holding that FCC’s authority to “modify” does not extend to eliminating altogether a statutory requirement).

³⁵⁰ Press Release, Fed. Trade Comm’n, Ovulation Tracking App Premom Will Be Barred from Sharing Health Data for Advertising Under Proposed FTC Order (May 17, 2023), <https://www.ftc.gov/news-events/news/press-releases/2023/05/ovulation-tracking-app-premom-will-be-barred-sharing-health-data-advertising-under-proposed-ftc>.

are considered “health care services” companies under the law.

Lastly, the dissent claims that the Final Rule introduces ambiguity where previous there was none. But *GoodRx* suggests otherwise. In a unanimous action, the Commission charged *GoodRx* with making unauthorized disclosures of people’s health data to Facebook and Google, among others.³⁵¹ *GoodRx*, meanwhile, disputed the applicability of the HBNR to its practices, calling it a “novel” application.³⁵² By codifying how HBNR applies to online platforms and applications, today’s Final Rule provides market participants with more clarity about what entities are covered—thereby providing greater certainty and notice.³⁵³

GoodRx marked the first time the Commission had ever enforced the Health Breach Notification Rule. A top priority for us at the Commission is ensuring we are faithfully discharging our statutory duties, rather than letting the authorities that Congress has granted us sit dormant, and we are proud of the work the Commission and the staff are doing to take care that the full set of laws assigned to the FTC are being faithfully executed.³⁵⁴ We agree with the

³⁵¹ Press Release, Fed. Trade Comm’n, FTC Enforcement Action to Bar *GoodRx* from Sharing Consumers’ Sensitive Health Info for Advertising (Feb. 1, 2023), <https://www.ftc.gov/news-events/news/press-releases/2023/02/ftc-enforcement-action-bar-goodrx-sharing-consumers-sensitive-health-info-advertising>; See also, Concurring Statement of Comm’r Christine S. Wilson, *GoodRx* Holdings, Inc. (Feb. 1, 2023), https://www.ftc.gov/system/files/ftc_gov/pdf/2023090_goodrx_final_concurring_statement_wilson.pdf (“Today’s settlement marks the first enforcement matter in which the FTC has invoked the HBNR. I congratulate staff on this important step—the agency rightly is focused on protecting the privacy of sensitive health data and empowering consumers to make informed choices about the goods and services they use.”); see also *id.* at 5 (describing the *GoodRx* case as “an important milestone in the Commission’s privacy work.”). The dissent suggests that Commissioners Holyoak and Ferguson would have supported the application of HBNR to *GoodRx*.

³⁵² See *GoodRx*, *GoodRx* Response to FTC Settlement (Feb. 1, 2023) (“We believe this is a novel application of the Health Breach Notification Rule by the FTC. . . . We do not agree with the assertion that this was a violation of the HBNR.”).

³⁵³ The dissent concedes that it does support an update to the rule that provides more clarity—and specifically an update that provides clarity to show that the rule covers *GoodRx*. Dissent at 7 (“I would support changes to the Rule that clarify the Rule’s application to companies like *GoodRx*.”). That is precisely what today’s Final Rule does. Previously, the rule did not define “health care services or supplies,” and today’s Final Rule does. Previously, health apps like *GoodRx* stated that it was unclear whether the rule applies to them, and today’s Final Rule makes clear that it does. This concession from the dissent suggests a more modest disagreement with the contours of how the Rule defines “health care services or supplies,” though—notably—the dissent does not provide an alternative definition.

³⁵⁴ See, e.g., Press Release, Fed. Trade Comm’n, FTC Hits *R360* and its Owner With \$3.8 Million Civil Penalty Judgment for Preying on People Seeking Treatment for Addiction (May 17, 2022), <https://www.ftc.gov/news-events/news/press-releases/2022/05/ftc-hits-r360-its-owner-38-million-civil-penalty-judgment-preying-people-seeking->

dissent that we must look out for the institutional integrity of the Commission. Failing to use the full scope of our statutory tools to protect Americans—and failing to update our application of these tools even as technologies change—would undermine the agency’s integrity and credibility alike.

We are deeply grateful to the Division of Privacy and Identity Protection for leading the Commission’s work to activate the Health Breach Notification Rule and for finalizing this Rule update. In an environment rife with new and evolving threats to Americans’ health data, ensuring we are faithfully harnessing all of our statutory tools to protect people from data breaches is paramount.

Dissenting Statement of Commissioner Melissa Holyoak, Joined by Commissioner Andrew Ferguson

The Health Breach Notification Rule (“Final Rule”) that the Commission adopts today exceeds the Commission’s statutory authority, puts companies at risk of perpetual non-compliance, and opens the Commission to legal challenge that could undermine its institutional integrity. I share the majority’s goal of protecting the privacy and security of consumers’ identifiable health information,¹

treatment-addiction (the Commission’s first action brought under the Opioid Addiction Recovery Fraud Prevention Act); Harris Jewelry, Press Release, Fed. Trade Comm’n, FTC and 18 States Sue to Stop Harris Jewelry from Cheating Military Families with Illegal Financing and Sales Tactics (Jul. 20, 2022), <https://www.ftc.gov/news-events/news/press-releases/2022/07/ftc-18-states-sue-stop-harris-jewelry-cheating-military-families-illegal-financing-sales-tactics> (the Commission’s first action brought under the Military Lending Act); Press Release, Fed. Trade Comm’n, Smart Home Monitoring Company Vivint Will Pay \$20 Million to Settle FTC Charges That It Misused Consumer Credit Reports (Apr. 29, 2021), <https://www.ftc.gov/news-events/news/press-releases/2021/04/smart-home-monitoring-company-vivint-will-pay-20-million-settle-ftc-charges-it-misused-consumer> (the Commission’s first action brought under the Red Flags Rule, brought under Acting Chair Slaughter); Press Release, Fed. Trade Comm’n, FTC Sues Burger Franchise Company That Targets Veterans and Others With False Promises and Misleading Documents (Feb. 8, 2022), <https://www.ftc.gov/news-events/news/press-releases/2022/02/ftc-sues-burger-franchise-company-targets-veterans-others-false-promises-misleading-documents> (the Commission’s first action under the Franchise Rule since 2007); Press Release, Fed. Trade Comm’n, FTC Issues Rule to Deter Rampant Made in USA Fraud (Jul. 1, 2021), <https://www.ftc.gov/news-events/news/press-releases/2021/07/ftc-issues-rule-deter-rampant-made-usa-fraud> (issuance of the Made in the USA Rule, more than 25 years after Congress authorized the Commission to promulgate a rule).

¹ Like the majority, and other Commissioners before me, I support federal privacy legislation, particularly where such legislation could address gaps in sector-specific laws and level the playing field for companies navigating a patchwork of laws. And like the majority, and other Commissioners before me, I care deeply about protecting the privacy and security of consumers’ health information, particularly where it falls outside the bounds of the Health Insurance Portability and Accountability Act (“HIPAA”). For more than two decades, the FTC has been in a leader in protecting consumers’ health information. See, e.g., *Eli Lilly*, FTC File No. 0123214 (May 10, 2002), <https://www.ftc.gov/legal-library/browse/cases-proceedings/012-3214-eli-lilly-company-matter>. I

and I support vigorous enforcement of laws protecting sensitive personal information with which Congress has entrusted the FTC.² I would support finalizing a rule that extends and clarifies the scope of the Commission’s enforcement in this important area of consumer protection if that rule were consistent with our grant of authority from Congress. But, no matter how the majority attempts to shoehorn its desired policy goal into a “plain reading” of the statute,³ I cannot support a rule that exceeds the bounds Congress clearly established. Indeed, a core principle guiding my tenure at the Commission will be that our rules must effectuate the law as it is—not as the Commission may wish it to be. For these reasons, I respectfully dissent.

The American Recovery and Reinvestment Act of 2009 (“Recovery Act”)⁴ authorized the Commission to issue a rule requiring vendors of “personal health records” (“PHRs”) and related entities that are not covered by HIPAA to notify individuals and the FTC of a “breach of security” of “unsecured PHR identifiable health information.”⁵ The Commission issued the Health Breach Notification Rule in 2009,⁶ initiated a routine review of the Rule in 2020,⁷ issued a policy statement re-interpreting the then-current Rule in 2021 (“2021 Policy Statement”),⁸ issued a Notice of Proposed Rulemaking on June 9, 2023 (“NPRM”),⁹ and today issues the Final Rule.¹⁰

I am encouraged that today the Commission is acting by rulemaking, as authorized by statute and following a period of notice and comment that elicited a range of views, rather than acting by fiat in a policy statement, as the Commission did in 2021.¹¹ I cannot endorse any policy statement that either displaces Congress’s authority to make law or subverts the rulemaking process. The 2021 Policy Statement did both. The majority clearly recognizes this overreach. After all, if the 2021 Policy Statement had any force, today’s rulemaking would be unnecessary.

Setting aside this troubling history, I turn to the Final Rule itself, which, unfortunately, I find equally troubling in its extension beyond the parameters established by Congress.

look forward to continuing the Commission’s important work in this area.

² See, e.g., Children’s Online Privacy Protection Rule, 16 CFR part 312, as authorized by the Children’s Online Privacy Protection Act of 1998, 15 U.S.C. 6501 *et seq.*

³ Joint Statement of Chair Lina M. Khan, Comm’r Rebecca Kelly Slaughter, and Comm’r Alvaro M. Bedoya at 2 (Apr. 24, 2024) (“Majority Statement”).

⁴ Am. Recovery and Reinvestment Act of 2009, Public Law 111–5, 123 Stat. 115 (2009).

⁵ 42 U.S.C. 17937(a), (g).

⁶ 74 FR 42962 (Aug. 25, 2009).

⁷ 85 FR 31085 (May 22, 2020).

⁸ See Statement of the Comm’n on Breaches by Health Apps and Other Connected Devices (Sept. 15, 2021), https://www.ftc.gov/system/files/documents/public_statements/1596364/statement_of_the_commission_on_breaches_by_health_apps_and_other_connected_devices.pdf (“2021 Policy Statement”).

⁹ 88 FR 37819 (June 9, 2023).

¹⁰ See Statement of Basis and Purpose (“SBP”) accompanying the Final Rule, Section I (summarizing procedural history).

¹¹ See 2021 Policy Statement, *supra* note 8.

Some background first. Under the Recovery Act, PHR identifiable health information means “individually identifiable health information,” as defined by the Social Security Act, 42 U.S.C. 1320d(6).¹² The Social Security Act defines “individually identifiable health information” as information that is “created or received by a health care provider, health plan, employer, or health care clearinghouse.”¹³ The Social Security Act then defines “health care provider” to include three categories: “[1] a provider of services (as defined in section 1395x(u) of this title), [2] a provider of medical or other health services (as defined in section 1395x(s) of this title), and [3] any other person furnishing health care services or supplies.”¹⁴

The Commission takes liberties with the final category in that definition (“any other person furnishing health care services or supplies”) to adopt a new, capacious definition of “covered health care provider” and a new, similarly capacious definition of “health care services and supplies,” whose joint effect is to sweep a large swath of apps and app developers under the purview of the Final Rule. These expansive definitions are not consistent with the statute. Under longstanding principles of statutory interpretation, the final category of provider (“any other person . . .”) must be understood in relation to the first two categories (“provider of services” and “provider of medical or other health services”).¹⁵ When a statute contains a list, “each word in that list presumptively has a ‘similar’ meaning” under the canon of *noscitur a sociis*.¹⁶ And when a general term follows a list of specific terms, the *eiusdem generis* canon teaches that the general term “should usually be read in light of those specific words to mean something ‘similar.’”¹⁷ Together, these canons instruct that the final category of health care provider that includes the general term “other person” must be similar to the more specific terms that precede it.

The first two categories of health care provider incorporate the definitions of sections 1395x(u) and 1395x(s) of the Social Security Act, respectively.¹⁸ The first category of provider includes “a hospital, critical access hospital, rural emergency hospital, skilled nursing facility, comprehensive outpatient rehabilitation

facility, home health agency, hospice program, or . . . a fund.”¹⁹ The second category of provider includes an extensive list (section 1395x(s) includes 17 paragraphs and over 35 subparagraphs) of medical professionals including physicians, physician assistants, nurse practitioners, clinical psychologists, clinical social workers, and others, and the specific services administered by medical professionals.²⁰ These two categories comprise traditional forms of health care providers.

The final category, addressing “any other person furnishing health care services or supplies,” must therefore only include persons that are “similar in nature” to these first two categories.²¹ The majority argues that my “effort to cabin the third category . . . reads it out of existence, violating the canon that holds interpretations giving effect to every clause of a statute are superior to those that render distinct clauses superfluous.”²² This application of the canon is incorrect. Requiring similarity among categories does not result in superfluity; it merely prevents interpretations that extend beyond what the text permits. A catch-all’s limited application due to its context is not a reason to expand that phrase to encompass dissimilar applications.

The Final Rule’s definition of “covered health care provider” is not remotely similar, because it incorporates a new, astonishingly broad definition of “health care services or supplies,” which means “any online service such as a website, mobile application, or internet-connected device that provides mechanisms to track diseases, health conditions, diagnoses or diagnostic testing, treatment, medications, vital signs, symptoms, bodily functions, fitness, fertility, sexual health, sleep, mental health, genetic information, diet, or that provides other health-related services or tools.”²³ Thus, the Commission transforms “health care provider,” which both under common usage and in context of the statutory provision means entities such as physicians and hospitals, to now include any company “furnishing” a health-related app.²⁴ As a result, the Final Rule creates a tautology: Health app developers may be “vendors of personal health records” by offering an app containing health information that has been created or received by a health care provider,

where the health app developer is itself the health care provider that creates or receives that health information by virtue of offering the app.

Notably, even though the Department of Health and Human Services (“HHS”) interprets this same provision of the Social Security Act, HHS has—notwithstanding the majority’s assertion to the contrary²⁵—never interpreted the term “health care provider” to reach the expansive, creative conclusion that the Commission does today.²⁶ The majority’s argument misstates the scope and language of the HIPAA Privacy Rule, which only applies to HIPAA “covered entities” and their “business associates,”²⁷—i.e., to traditional health care providers that do not include the broad swath of app developers the Final Rule will encompass. Significantly, the majority omits from its characterization of the term “health care” HHS’s own illustrations of that term, which highlight the proximity to traditional forms of health care by different kinds of medical professionals:

(1) Preventive, diagnostic, therapeutic, rehabilitative, maintenance, or palliative care, and counseling, service, assessment, or procedure with respect to the physical or mental condition, or functional status, of an individual or that affects the structure or function of the body; and

(2) Sale or dispensing of a drug, device, equipment, or other item in accordance with a prescription.²⁸

The Majority Statement repeatedly says that HHS defines “health care” broadly,²⁹ but the language it cites provides no such support.

Aware of this incongruity, the Commission seeks to differentiate its use of “health care provider” from that of “other government agencies.”³⁰ Yet the Commission provides no explanation why its definition should differ, particularly where it is unclear whether the Commission has interpretative authority over the Social Security Act’s definition of health care provider and where other agencies *are* delegated such interpretative authority.³¹

²⁵ Majority Statement at 3.

²⁶ See NPRM at 37823.

²⁷ 45 CFR 160.102 through 103.

²⁸ *Id.* § 160.103.

²⁹ Majority Statement at 3–4.

³⁰ SBP at 26.

³¹ *Id.* at 13 (noting that HHS interprets these provisions of the Social Security Act). Cf. *City of Arlington, Tex. v. F.C.C.*, 569 U.S. 290, 323 (2013) (Roberts, C.J., dissenting) (“When presented with an agency’s interpretation of such a statute, a court cannot simply ask whether the statute is one that the agency administers; the question is whether authority over the particular ambiguity at issue has been delegated to the particular agency.”).

¹² 42 U.S.C. 17937(f)(2).

¹³ 42 U.S.C. 1320d(6).

¹⁴ *Id.* 1320d(3).

¹⁵ See *Yates v. United States*, 574 U.S. 528, 549–51 (2015) (Alito, J., concurring); Antonin Scalia & Bryan A. Garner, *Reading Law: The Interpretation of Legal Texts* 195–196, 199–200 (2012).

¹⁶ *Yates*, 574 U.S. at 549.

¹⁷ *Id.* at 550.

¹⁸ 42 U.S.C. 1320d(3).

¹⁹ 42 U.S.C. 1395x(u).

²⁰ *Id.* 1395x(s).

²¹ *Yates*, 574 U.S. at 545 (internal quotation marks omitted).

²² Majority Statement at 2.

²³ Final Rule at 98.

²⁴ The SBP explains that an app developer (or any company “furnishing” a health app) would be covered as a health care provider because its health app is a health care service or supply. SBP at 7, 22–28.

The Commission also takes troubling liberties with the statute’s definition of “personal health record,” which are evident

from a side-by-side comparison of the statute and the Final Rule:

Recovery act	Final rule
“an electronic record of PHR identifiable health information . . . on an individual that can be drawn from multiple sources and is managed, shared, and controlled by or primarily for the individual.” ³² .	“an electronic record of PHR identifiable health information on an individual that has the technical capacity to draw information from multiple sources and that is managed, shared, and controlled by or primarily for the individual.” ³³

Under the Final Rule, a PHR need not actually draw health information from multiple sources, as the statute contemplates (because the statutory phrase “that can be drawn” modifies its immediate antecedent, “health information”). Rather, under the Final Rule, a single source of health information will render an app a PHR as long as the “PHR” has the “technical capacity” to draw some other information elsewhere.³⁴ The implications of this change, in conjunction with the expansion of “health care provider,” are significant. Any retailer that offers an app that tracks health-related purchases (e.g., bandages, vitamins, dandruff shampoo) may be a vendor of a PHR covered by the Rule if the app draws health information (e.g., purchasing information) from the consumer and the app has the “technical capacity” to draw any information from any other source. As the Statement of Basis and Purpose notes, commenters warned that virtually every app has the technical capacity to draw some information from more than one source.³⁵ That expansive scope could be appropriate if Congress’s language permitted it. But the Commission’s interpretation, which effectively renders the Recovery Act’s “multiple sources” requirement meaningless, ignores longstanding principles of statutory interpretation that require each provision of a statute to be given effect.³⁶

The Commission’s expansive definitions of “covered health care provider,” “health care services and supplies,” and “personal health record” have a profound effect on the scope of the Rule: Most companies that offer or disseminate health-related apps or similar products would be treated as “covered health care providers” that therefore hold “PHR identifiable health information” in their apps (i.e., PHRs), such that they are vendors of PHRs—even if their app is merely health-adjacent.

Remarkably, the Commission imposes no limit on this extraordinary breadth in the Rule itself. Rather, in a post-NPRM attempt to check the scope, the Commission fashions a limiting principle: Apps are covered only if they are “more than tangentially relating to health.”³⁷ This extra-statutory, extra-

regulatory limit has several significant problems.

First, if the majority were correct, from where would it draw the authority to impose this “more than tangentially relating to health” limitation? If Congress in fact commanded us to cover all the apps the majority claims, this extra-textual limitation would be beyond our power to impose.³⁸ Why, then, does the majority blink in the face of what it understands Congress to have required? There may be good policy reasons not to follow Congress’s language—as the majority understands it—wherever it leads, but we do not have power to shortchange Congress’s commands. That even the majority feels compelled to adopt this extra-textual limitation—again, as the majority understands the text—on the statute’s reach suggests that the language probably does not mean what the majority says.

The second problem is substantive: What does this language mean? When does an app cross the line between tangentially related to health and more than tangentially related? If a gas station with a loyalty app sells Advil, is the app only tangentially related to health and outside the Final Rule’s purview? If the gas station adds Robitussin and pregnancy tests to its inventory, does it cross the line to more than tangentially related to health? If a clothing store with an e-commerce app sells a handful of maternity shirts, is the app only tangentially related to health? If the store adds more maternity clothes, nursing bras, and some anti-nausea ginger tea to its in-app offerings, is the app more than tangentially related to health? If vitamins, over-the-counter medicines, acne creams, bandages, and similar items comprise 0.1% or 1% or 10% of a superstore’s inventory, when is the retailer’s e-commerce app more than tangentially related to health? I see no clear answers to any of these hypotheticals in today’s Final Rule, which suggests that the marketplace will see no clear answers either.³⁹

The third problem is procedural. The Commission did not propose this ambiguous

but impactful limitation in a Notice of Proposed Rulemaking—likely because there is no statutory basis for this newly-created language. Rather, it introduces this crucial concept for the first time in a Statement of Basis and Purpose (a purely interpretive document) as a *post hoc* fix to the problem the Commission itself created with its expansive definitions. As a result, the Commission did not provide notice or receive public comment on the efficacy or propriety of this limitation, depriving the public of its opportunity to meaningfully participate in the rulemaking process and depriving itself of potentially valuable input from commenters.

The final problem is that this *post hoc*, extra-regulatory limitation renders the Commission’s burden analysis inadequate. The Paperwork Reduction Act (“PRA”) requires the Commission to estimate the reportable breaches by entities covered by the Rule and compliance costs.⁴⁰ The Regulatory Flexibility Act (“RFA”) requires the Commission to assess the economic impact on small businesses.⁴¹ Apparently relying on the SBP’s “more than tangentially related to health” limitation, the PRA and RFA analyses only address breaches by apps categorized as “Health and Fitness.”⁴² Because the Rule itself contains no such limitation, general retailers with e-commerce apps, gas stations with loyalty apps, and other similar generalists that sell any health-related items do not factor into these analyses. As a result, they likely dramatically underestimate the numbers of regulated entities, number of breaches, and costs to businesses.

Perhaps the breath of the Final Rule would be more of a theoretical than practical concern to businesses, if they could adopt practices sufficient to avoid any breach that would trigger notice obligations under the Final Rule, or, in the event of a breach, err on the side of notification. But § 318.3(b) of the Final Rule imposes affirmative obligations on companies to notify their service providers if they are covered by the Final Rule, regardless of whether they experience a breach.⁴³ To comply with this requirement, companies must know whether they are covered by the Rule—that is, which side of “more than tangentially relating to health” they fall on. Without clarity on that line, companies run the risk of being in

³² 42 U.S.C. 17921(11).

³³ Final Rule at 99.

³⁴ See SBP at 32 (“Next, adding the phrase ‘technical capacity to draw information’ clarifies that a product is a personal health record if it can draw any information from multiple sources, even if it only draws health information from one source.”).

³⁵ See *id.* at 34.

³⁶ Scalia & Garner, *supra* note 15 at 174 (discussing surplusage canon).

³⁷ SBP at 28.

³⁸ See *Nat’l Fed’n of Indep. Business v. Dep’t of Labor*, 595 U.S. 109, 117 (2022) (per curiam) (“Administrative agencies are creatures of statute. They accordingly possess only the authority that Congress has provided.”).

³⁹ The expansive coverage increases the likelihood of creating unintended consequences. Will the gas station decline to add over-the-counter medicines to its inventory to avoid crossing the line of “more than tangentially related to health”? Will the clothing retailer shy away from maternity apparel? Will the e-commerce giant avoid selling bandages and dandruff shampoo? These potentially detrimental outcomes undermine a Rule intended to benefit consumers.

⁴⁰ See generally 44 U.S.C. 3501 *et seq.*; SBP at 86.
⁴¹ 5 U.S.C. 601 through 612.

⁴² SBP at 86, 93.

⁴³ This may have been a sensible requirement in 2009, when the scope of the Rule was much narrower, but it has dramatic consequences in this much-expanded Rule.

perpetual violation of the Final Rule and, therefore, perpetually at the mercy of the Commission's enforcement discretion. The Commission, at this moment, may not intend to pursue such technical violations. But any expression of intended restraint will be cold comfort to companies that have seen the Commission's self-imposed restraint wax and wane in other areas.⁴⁴

I find the majority's liberties with the statute particularly troubling because they are unnecessary to reach health apps. Indeed, the Commission's own recent enforcement action against digital healthcare platform GoodRx makes that clear. Only last year, a bipartisan Commission applied the 2009 Rule to GoodRx's online platform and app because the company received identifiable health information on prescription medications (among other things) from pharmacy benefit

⁴⁴ Significantly, the Majority Statement is silent as to the propriety and consequences of its "tangentially related" limiting principle, likely because this approach is indefensible.

managers and pharmacies, among other sources, so that consumers could manage their information.⁴⁵ The majority argues that today's changes are necessary to provide clarity to the market about the Rule's scope,⁴⁶ but *GoodRx* has already done that—and I would support changes to the Rule that are consistent with the statute. In short, I agree with the majority's goals—safeguarding consumers' sensitive health information and implementing a Congressional mandate to put consumers on notice of the breach of that data—but I believe that we must effectuate those goals within the scope of the law as it

⁴⁵ See *Concurring Statement of Commissioner Christine S. Wilson, GoodRx*, Matter No. 2023090 1 n.2 (Feb. 1, 2023) ("GoodRx has violated the HBNR based on a plain reading of the text, setting aside any gloss the Commission sought to add in its September 2021 Statement on Breaches by Health Apps and Other Connected Devices."), https://www.ftc.gov/system/files/ftc_gov/pdf/2023090_goodrx_final_concurring_statement_wilson.pdf.

⁴⁶ Majority Statement at 5.

is, rather than legislating in the guise of applying the law.

The FTC is a venerable institution that does vital work to protect consumers and promote competition, thanks to its hardworking and devoted career staff. I commend the staff attorneys, economists, and technologists who worked on the rule for their careful and thoughtful consideration of difficult issues. Ultimately, while I am sympathetic to the majority's goal, I fear that adopting a Final Rule that is irreconcilable with the statute and that puts companies in an untenable position puts the Commission at risk. Legal challenges may undermine the Commission's institutional integrity, and Congress may be reluctant to trust the Commission with other authority—even the much-needed authority to protect the privacy of consumers' sensitive personal information. I therefore respectfully dissent.

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