

hereby given that the above-captioned consent agreement containing a consent order to cease and desist, having been filed with and accepted, subject to final approval, by the Commission, has been placed on the public record for a period of thirty (30) days. The following Analysis to Aid Public Comment describes the terms of the consent agreement, and the allegations in the complaint. An electronic copy of the full text of the consent agreement package can be obtained from the FTC Home Page (for September 17, 2007), on the World Wide Web, at <http://www.ftc.gov/os/2007/09/index.htm>. A paper copy can be obtained from the FTC Public Reference Room, Room 130-H, 600 Pennsylvania Avenue, NW., Washington, DC 20580, either in person or by calling (202) 326-2222.

Public comments are invited, and may be filed with the Commission in either paper or electronic form. All comments should be filed as prescribed in the **ADDRESSES** section above, and must be received on or before the date specified in the **DATES** section.

Analysis of Agreement Containing Consent Order to Aid Public Comment

The Federal Trade Commission has accepted, subject to final approval, an agreement containing a consent order from Ingenix, Inc. ("respondent" or "Ingenix").

The proposed consent order has been placed on the public record for thirty (30) days for receipt of comments by interested persons. Comments received during this period will become part of the public record. After thirty (30) days, the Commission will again review the agreement and the comments received, and will decide whether it should withdraw from the agreement or make final the agreement's proposed order.

Ingenix markets MedPoint, a data aggregation service that provides individual medical profiles to health and life insurance companies. Insurance companies use MedPoint for underwriting or claims review purposes. The medical profile generated by MedPoint analyzes the individual's prescription drug history, and provides, based on that analysis, potential medical conditions that may be present and predictive scores for the individual.

The Commission's complaint alleges that the medical profile generated for the MedPoint service is a consumer report and that respondent is a consumer reporting agency, as those terms are defined in Sections 603(d) and (f) of the Fair Credit Reporting Act, 15 U.S.C. §§ 1681a(d) and (f). The complaint alleges that the respondent's failure to provide the "Notice To Users

of Consumer Reports: Obligations of Users Under the FCRA" ("Notice to Users"), the required content of which is found in 16 CFR 698, Appendix H, is a violation of Section 607(d) of the Fair Credit Reporting Act, 15 U.S.C. § 1681e(d).

The proposed consent order contains provisions designed to prevent respondent from engaging in similar acts and practices in the future.

Part I of the proposed order requires respondent to provide the Notice To Users to any user or prospective user of any medical profile generated by MedPoint that constitutes a consumer report, or of any other consumer report.

Part II.A. of the proposed order requires respondent to maintain or continue to maintain reasonable procedures to limit the furnishing of consumer reports to those with a permissible purpose, as required by Section 607(a) of the Fair Credit Reporting Act, 15 U.S.C. § 1681e(a).

Part II.B. of the proposed order requires respondent to follow or continue to follow reasonable procedures to assure maximum possible accuracy of the information concerning the individuals about whom the reports relates, as required by Section 607(b) of the Fair Credit Reporting Act, 15 U.S.C. § 1681e(b).

Part II.C. of the proposed order requires respondent to maintain or continue to maintain reasonable procedures to ensure compliance with Section 611 of the Fair Credit Reporting Act, 15 U.S.C. § 1681i, "Procedure in case of disputed accuracy."

Part II.D. of the proposed order requires respondent to conduct or continue to conduct a reasonable reinvestigation in cases of disputed accuracy, as required by Section 611 of the Fair Credit Reporting Act, 15 U.S.C. § 1681i.

Part II.E. of the proposed order requires respondent to comply or continue to comply with the Disposal of Consumer Report Information and Records Rule, 16 C.F.R. Part 682.

Part III of the proposed order contains a document retention requirement. It requires respondent to maintain and upon request make available to the Commission for inspection and copying documents demonstrating compliance with the requirements of Parts I and II of the proposed order.

Part IV of the proposed order requires respondent to distribute copies of the order to various principals, officers, directors, and managers, employees, agents, and representatives having decision-making responsibilities with respect to MedPoint or any other consumer report.

Part V of the proposed order requires respondent to notify the Commission of any changes in corporate structure that might affect compliance with the order.

Part VI of the proposed order requires respondent to file with the Commission one or more reports detailing its compliance with the order.

Part VII of the proposed order is a "sunset" provision, dictating the conditions under which the order will terminate twenty years from the date it is issued or twenty years after a complaint is filed in federal court, by either the United States or the FTC, alleging any violation of the order.

The purpose of this analysis is to facilitate public comment on the proposed order. It is not intended to constitute an official interpretation of the proposed order or to modify in any way its terms.

By direction of the Commission.

Donald S. Clark,

Secretary.

[FR Doc. E7-19152 Filed 9-27-07; 8:45 am]

[Billing Code: 6750-01-S]

FEDERAL TRADE COMMISSION

[File No. 062 3189]

Milliman, Inc.; Analysis of Proposed Consent Order to Aid Public Comment

AGENCY: Federal Trade Commission.

ACTION: Proposed Consent Agreement.

SUMMARY: The consent agreement in this matter settles alleged violations of federal law prohibiting unfair or deceptive acts or practices or unfair methods of competition. The attached Analysis to Aid Public Comment describes both the allegations in the draft complaint and the terms of the consent order—embodied in the consent agreement—that would settle these allegations.

DATES: Comments must be received on or before October 17, 2007.

ADDRESSES: Interested parties are invited to submit written comments. Comments should refer to "Milliman, File No. 062 3189," to facilitate the organization of comments. A comment filed in paper form should include this reference both in the text and on the envelope, and should be mailed or delivered to the following address: Federal Trade Commission/Office of the Secretary, Room 135-H, 600 Pennsylvania Avenue, NW., Washington, DC 20580. Comments containing confidential material must be filed in paper form, must be clearly labeled "Confidential," and must comply with Commission Rule 4.9(c).

16 CFR 4.9(c) (2005).¹ The FTC is requesting that any comment filed in paper form be sent by courier or overnight service, if possible, because U.S. postal mail in the Washington area and at the Commission is subject to delay due to heightened security precautions. Comments that do not contain any nonpublic information may instead be filed in electronic form as part of or as an attachment to email messages directed to the following e-mail box: consentagreement@ftc.gov.

The FTC Act and other laws the Commission administers permit the collection of public comments to consider and use in this proceeding as appropriate. All timely and responsive public comments, whether filed in paper or electronic form, will be considered by the Commission, and will be available to the public on the FTC Web site, to the extent practicable, at <http://www.ftc.gov>. As a matter of discretion, the FTC makes every effort to remove home contact information for individuals from the public comments it receives before placing those comments on the FTC website. More information, including routine uses permitted by the Privacy Act, may be found in the FTC's privacy policy, at <http://www.ftc.gov/ftc/privacy.htm>.

FOR FURTHER INFORMATION CONTACT:

Rebecca E. Kuehn, Bureau of Consumer Protection, 600 Pennsylvania Avenue, NW., Washington, DC 20580, (201) 326-2252.

SUPPLEMENTARY INFORMATION: Pursuant to section 6(f) of the Federal Trade Commission Act, 38 Stat. 721, 15 U.S.C. 46(f), and § 2.34 of the Commission Rules of Practice, 16 CFR 2.34, notice is hereby given that the above-captioned consent agreement containing a consent order to cease and desist, having been filed with and accepted, subject to final approval, by the Commission, has been placed on the public record for a period of thirty (30) days. The following Analysis to Aid Public Comment describes the terms of the consent agreement, and the allegations in the complaint. An electronic copy of the full text of the consent agreement package can be obtained from the FTC Home Page (for September 17, 2007), on the World Wide Web, at <http://www.ftc.gov/os/2007/09/index.htm>. A paper copy can be obtained from the

FTC Public Reference Room, Room 130-H, 600 Pennsylvania Avenue, NW., Washington, DC 20580, either in person or by calling (202) 326-2222.

Public comments are invited, and may be filed with the Commission in either paper or electronic form. All comments should be filed as prescribed in the **ADDRESSES** section above, and must be received on or before the date specified in the **DATES** section.

Analysis of Agreement Containing Consent Order to Aid Public Comment

The Federal Trade Commission has accepted, subject to final approval, an agreement containing a consent order from Milliman, Inc. ("respondent" or "Milliman").

The proposed consent order has been placed on the public record for thirty (30) days for receipt of comments by interested persons. Comments received during this period will become part of the public record. After thirty (30) days, the Commission will again review the agreement and the comments received, and will decide whether it should withdraw from the agreement or make final the agreement's proposed order.

Milliman markets IntelliScript, a data aggregation service that provides individual medical profiles, including but not limited to prescription drug purchase histories of insurance applicants, to health and life insurance companies. Insurance companies use IntelliScript for underwriting or claims review purposes. The medical profile generated by IntelliScript analyzes the individual's prescription drug history, and provides a 'map' of the risk levels associated with each drug, based on information provided by the insurer.

The Commission's complaint alleges that the medical profile generated for the IntelliScript service is a consumer report and that respondent is a consumer reporting agency, as those terms are defined in Sections 603(d) and (f) of the Fair Credit Reporting Act, 15 U.S.C. §§ 1681a(d) and (f). The complaint alleges that the respondent's failure to provide the "Notice To Users of Consumer Reports: Obligations of Users Under the FCRA" ("Notice To Users"), the required content of which is found in 16 CFR 698, Appendix H, is a violation of Section 607(d) of the Fair Credit Reporting Act, 15 U.S.C. § 1681e(d).

The proposed consent order contains provisions designed to prevent respondent from engaging in similar acts and practices in the future.

Part I of the proposed order requires respondent to provide the Notice To Users to any user or prospective user of any medical profile generated by

IntelliScript that constitutes a consumer report or of any other consumer report.

Part II.A. of the proposed order requires respondent to maintain or continue to maintain reasonable procedures to limit the furnishing of consumer reports to those with a permissible purpose, as required by Section 607(a) of the Fair Credit Reporting Act, 15 U.S.C. § 1681e(a).

Part II.B. of the proposed order requires respondent to follow or continue to follow reasonable procedures to assure maximum possible accuracy of the information concerning the individuals about whom the reports relates, as required by Section 607(b) of the Fair Credit Reporting Act, 15 U.S.C. § 1681e(b).

Part II.C. of the proposed order requires respondent to maintain or continue to maintain reasonable procedures to ensure compliance with Section 611 of the Fair Credit Reporting Act, 15 U.S.C. § 1681i, "Procedure in case of disputed accuracy."

Part II.D. of the proposed order requires respondent to conduct or continue to conduct a reasonable reinvestigation in cases of disputed accuracy, as required by Section 611 of the Fair Credit Reporting Act, 15 U.S.C. § 1681i.

Part II.E. of the proposed order requires respondent to comply or continue to comply with the Disposal of Consumer Report Information and Records Rule, 16 C.F.R. Part 682.

Part III of the proposed order contains a document retention requirement. It requires respondent to maintain and upon request make available to the Commission for inspection and copying documents demonstrating compliance with the requirements of Parts I and II of the proposed order.

Part IV of the proposed order requires respondent to distribute copies of the order to various officers, directors, and managers, employees, agents, and representatives having decision-making responsibilities with respect to IntelliScript or any other consumer report.

Part V of the proposed order requires respondent to notify the Commission of any changes in corporate structure that might affect compliance with the order.

Part VI of the proposed order requires respondent to file with the Commission one or more reports detailing its compliance with the order.

Part VII of the proposed order is a "sunset" provision, dictating the conditions under which the order will terminate twenty years from the date it is issued or twenty years after a complaint is filed in federal court, by

¹ The comment must be accompanied by an explicit request for confidential treatment, including the factual and legal basis for the request, and must identify the specific portions of the comment to be withheld from the public record. The request will be granted or denied by the Commission's General Counsel, consistent with applicable law and the public interest. See Commission Rule 4.9(c), 16 CFR 4.9(c).

either the United States or the FTC, alleging any violation of the order.

The purpose of this analysis is to facilitate public comment on the proposed order. It is not intended to constitute an official interpretation of the proposed order or to modify in any way its terms.

By direction of the Commission.

Donald S. Clark,

Secretary.

[FR Doc. E7-19159 Filed 9-27-07; 8:45 am]

Billing Code: 6750-01-S

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institute for Occupational Safety and Health; Designation of a Class of Employees for Addition to the Special Exposure Cohort

AGENCY: National Institute for Occupational Safety and Health (NIOSH), Department of Health and Human Services (HHS).

ACTION: Notice.

SUMMARY: The Department of Health and Human Services (HHS) gives notice of a decision to designate a class of employees at the Ames Laboratory, Ames, Iowa, as an addition to the Special Exposure Cohort (SEC) under the Energy Employees Occupational Illness Compensation Program Act of 2000. On September 12, 2007, the Secretary of HHS designated the following class of employees as an addition to the SEC:

Sheet metal workers, physical plant maintenance and associated support staff (including all maintenance shop personnel), and supervisory staff who were monitored or should have been monitored for potential internal radiation exposures associated with the maintenance and renovation activities of the thorium production areas in Wilhelm Hall (a.k.a. the Metallurgy Building or "Old" Metallurgy Building) at the Ames Laboratory from January 1, 1955, through December 31, 1970, for a number of work days aggregating at least 250 work days or in combination with work days within the parameters established for one or more other classes of employees in the Special Exposure Cohort.

This designation will become effective on October 12, 2007, unless Congress provides otherwise prior to the effective date. After this effective date, HHS will publish a notice in the **Federal Register** reporting the addition of this class to the SEC or the result of any provision by Congress regarding the decision by HHS to add the class to the SEC.

FOR FURTHER INFORMATION CONTACT: Larry Elliott, Director, Office of

Compensation Analysis and Support, National Institute for Occupational Safety and Health (NIOSH), 4676 Columbia Parkway, MS C-46, Cincinnati, OH 45226, Telephone 513-533-6800 (this is not a toll-free number). Information requests can also be submitted by e-mail to OCAS@CDC.GOV.

Dated: September 24, 2007.

John Howard,

Director, National Institute for Occupational Safety and Health.

[FR Doc. E7-19297 Filed 9-27-07; 8:45 am]

BILLING CODE 4160-17-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institute for Occupational Safety and Health; Designation of a Class of Employees for Addition to the Special Exposure Cohort

AGENCY: National Institute for Occupational Safety and Health (NIOSH), Department of Health and Human Services (HHS).

ACTION: Notice.

SUMMARY: The Department of Health and Human Services (HHS) gives notice of a decision to designate a class of employees at the Hanford Engineer Works, Richland, Washington, as an addition to the Special Exposure Cohort (SEC) under the Energy Employees Occupational Illness Compensation Program Act of 2000. On September 12, 2007, the Secretary of HHS designated the following class of employees as an addition to the SEC:

Employees of the Department of Energy (DOE), its predecessor agencies, or DOE contractors or subcontractors who were monitored or should have been monitored for internal radiological exposures while working at the Hanford Engineer Works in: the 300 Area fuel fabrication and research facilities from October 1, 1943 through August 31, 1946; the 200 Area plutonium separation facilities from November 1, 1944 through August 31, 1946; or the 100 B, D, and F reactor areas from September 1, 1944 through August 31, 1946; for a number of work days aggregating at least 250 work days or in combination with work days within the parameters established for one or more other classes of employees in the Special Exposure Cohort.

This designation will become effective on October 12, 2007, unless Congress provides otherwise prior to the effective date. After this effective date, HHS will publish a notice in the **Federal Register** reporting the addition of this class to the SEC or the result of any provision by Congress regarding the decision by HHS to add the class to the SEC.

FOR FURTHER INFORMATION CONTACT:

Larry Elliott, Director, Office of Compensation Analysis and Support, National Institute for Occupational Safety and Health (NIOSH), 4676 Columbia Parkway, MS C-46, Cincinnati, OH 45226, Telephone 513-533-6800 (this is not a toll-free number). Information requests can also be submitted by e-mail to OCAS@CDC.GOV.

Dated: September 24, 2007.

John Howard,

Director, National Institute for Occupational Safety and Health.

[FR Doc. E7-19243 Filed 9-27-07; 8:45 am]

BILLING CODE 4160-17-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Agency for Healthcare Research and Quality

AHRQ Health Care Innovations Exchange

AGENCY: Agency for Healthcare Research and Quality (AHRQ), HHS.

ACTION: Notice of Submission of Innovations.

SUMMARY: To support its objective of accelerating the diffusion and adoption of innovative health care delivery changes, the Agency for Healthcare Research and Quality (AHRQ) recently launched version 1.0 of the AHRQ Health Care Innovations Exchange (HCIE) Web site, <http://www.innovations.ahrq.gov>. The HCIE is a new initiative designed to support health care professionals in sharing and adopting innovations that improve health care quality. Version 1.0 of the Web site is focused on stimulating creativity and innovation and will serve as a virtual place to which innovators will be encouraged to submit their innovations and experiences from which potential adopters can begin learning about the nuances of implementation.

In Spring 2008, AHRQ will deploy version 2.0 of its Health Care Innovations Exchange site making hundreds of profiles of health care service innovations of varying degrees of novelty and scientific rigor accessible to the public. Version 2.0 will also offer expert commentary; stories; tools; lessons learned; "change packages"—sets of innovations implemented simultaneously; expanded content on implementation; and opportunities to learn and network.

To build the database of innovations profiles, AHRQ invites submissions of