EARLY TERMINATIONS GRANTED—Continued
DECEMBER 1, 2013 THRU DECEMBER 31, 2013

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<td>G Henry Schein, Inc.; HealthPoint Capital Partners II, LP; Henry Schein, Inc.</td>
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<td>LKQ Corporation; Platinum Equity Capital Partners II, L.P.; LKQ Corporation.</td>
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FOR FURTHER INFORMATION CONTACT:

By direction of the Commission.

Donald S. Clark,
Secretary.

[FTR Doc. 2014–00240 Filed 1–10–14; 8:45 am]

BILLING CODE 6750–01–M

FEDERAL TRADE COMMISSION

[File No. 122 3077]

Accretive Health, 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 of Proposed Consent Order 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 January 30, 2014.

ADDRESSES: Interested parties may file a comment at https://ftcpublic.commentworks.com/ftc/accretiveconsent online or on paper, by following the instructions in the Request for Comment part of the SUPPLEMENTARY INFORMATION section below. Write “Accretive Health, Inc.-Consent Agreement; File No. 122 3077” on your comment and file your comment online at https://ftcpublic.commentworks.com/ftc/accretiveconsent by following the instructions on the web-based form. If you prefer to file your comment on paper, mail or deliver your comment to the following address: Federal Trade Commission, Office of the Secretary, Room H–113 (Annex D), 600 Pennsylvania Avenue NW, Washington, DC 20580.


SUPPLEMENTARY INFORMATION: Pursuant to Section 6(f) of the Federal Trade Commission Act, 15 U.S.C. 46(f), and FTC Rule 2.34, 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 December 31, 2013), on the World Wide Web, at http://www.ftc.gov/os/actions.shtm. 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.

You can file a comment online or on paper. For the Commission to consider your comment, we must receive it on or before January 30, 2014. Write “Accretive Health, Inc.-Consent...
Agreement; File No. 122 3077” on your comment. Your comment—including your name and your state—will be placed on the public record of this proceeding, including, to the extent practicable, on the public Commission Web site, at http://www.ftc.gov/os/publiccomments.shtm. As a matter of discretion, the Commission tries to remove individuals’ home contact information from comments before placing them on the Commission Web site.

Because your comment will be made public, you are solely responsible for making sure that your comment does not include any sensitive personal information, like anyone’s Social Security number, date of birth, driver’s license number or other state identification number or foreign country equivalent, passport number, financial account number, or credit or debit card number. You are also solely responsible for making sure that your comment does not include any sensitive health information, like medical records or other individually identifiable health information. In addition, do not include any “trade secret or any commercial or financial information which . . . is privileged or confidential,” as discussed in Section 6(f) of the FTC Act, 15 U.S.C. 46(f), and FTC Rule 4.10(a)(2), 16 CFR 4.10(a)(2). In particular, do not include competitively sensitive information such as costs, sales statistics, inventories, formulas, patterns, devices, manufacturing processes, or customer names.

If you want the Commission to give your comment confidential treatment, you must file it in paper form, with a request for confidential treatment, and you have to follow the procedure explained in FTC Rule 4.9(c), 16 CFR 4.9(c). Your comment will be kept confidential only if the FTC General Counsel, in his or her sole discretion, grants your request in accordance with the law and the public interest.

Postal mail addressed to the Commission is subject to delay due to heightened security screening. As a result, we encourage you to submit your comments online. To make sure that the Commission considers your online comment, you must file it at https://ftcpublic.commentworks.com/ftc/accreteveconsent by following the instructions on the web-based form. If this Notice appears at http://www.regulations.gov/#/home, you also may file a comment through that Web site.

If you file your comment on paper, write “Accretive Health, Inc.- Consent Agreement; File No. 122 3077” on your comment and on the envelope, and mail or deliver it to the following address: Federal Trade Commission, Office of the Secretary, Room H–113 (Annex D), 600 Pennsylvania Avenue NW., Washington, DC 20580. If possible, submit your paper comment to the Commission by courier or overnight service.

Visit the Commission Web site at http://www.ftc.gov to read this Notice and the news release describing it. The FTC Act and other laws that the Commission administers permit the collection of public comments to consider and use in this proceeding as appropriate. The Commission will consider all timely and responsive public comments that it receives on or before January 30, 2014. You can find more information, including routine uses permitted by the Privacy Act, in the Commission’s privacy policy, at http://www.ftc.gov/ftc/privacy.htm.

Analysis of Proposed Consent Order To Aid Public Comment

The Federal Trade Commission has accepted, subject to final approval, a consent order applicable to Accretive Health Systems, Inc.

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 and take appropriate action or make final the agreement’s proposed order.

Accretive Health enters into service agreements with hospital systems around the country to provide services related to the hospital systems’ “revenue cycle” operations. Revenue cycle operations include registration, transcription, coding and medical documentation, billing, pricing, and collection of past due accounts. In exchange for these services, hospital systems pay Accretive Health both fixed fees and incentive payments based on a percentage of the monetary benefit from increased revenues. Accretive Health employees work at hospital facilities to assist with these services. As part of its service to client hospitals, Accretive Health collects, maintains, and has access to information about hospitals’ patients, including sensitive health and personal information. This information may include patient names, dates of birth, billing information, diagnostic information, and Social Security numbers.

The Commission’s complaint alleges that Accretive Health unfairly failed to provide reasonable and appropriate security for consumers’ personal information it collected and maintained by engaging in a number of practices that, taken together, unreasonably and unnecessarily exposed consumers’ personal data to unauthorized access. Among other things, Accretive Health created unnecessary risks of unauthorized access or theft of personal information by:

a. Transporting laptops containing personal information in a manner that made them vulnerable to theft or other misappropriation;

b. Failing to adequately restrict access to, or copying of, personal information based on an employee’s need for information;

c. Failing to ensure that employees removed information from their computers for which they no longer had a business need; and

d. Using consumers’ personal information in training sessions with employees and failing to ensure that the information was removed from employees’ computers following the training.

The complaint further alleges that these failures contributed to a July 2011 incident in Minneapolis, Minnesota in which an Accretive Health laptop containing over 600 files with over 20 million pieces of information related to 23,000 patients was left in the locked passenger compartment of the employee’s car and stolen. The laptop included sensitive health and personal information, including patient names, dates of birth, billing information, diagnostic information, and Social Security numbers. The user of this laptop had data that was not necessary to perform his job.

The proposed order contains provisions designed to prevent Accretive Health from engaging in the future in practices similar to those alleged in the complaint.

Part II of the proposed order requires Accretive Health to establish and maintain, or continue to maintain, a comprehensive information security program that is reasonably designed to protect the security, confidentiality, and integrity of personal information collected from or about consumers. The security program must contain administrative, technical, and physical safeguards appropriate to Accretive Health’s size and complexity, nature and scope of its activities, and the sensitivity of the information collected.

1 In particular, the written request for confidential treatment that accompanies the comment must include the factual and legal basis for the request, and must identify the specific portions of the comment to be withheld from the public record. See FTC Rule 4.9(c), 16 CFR 4.9(c).
from or about consumers. Specifically, the proposed order requires Accretive Health to:

- Designate an employee or employees to coordinate and be accountable for the information security program;
- Identify material internal and external risks to the security, confidentiality, and integrity of personal information that could result in the unauthorized disclosure, misuse, loss, alteration, destruction, or other compromise of such information, and assess the sufficiency of any safeguards in place to control these risks;
- Design and implement reasonable safeguards to control the risks identified through risk assessment, and regularly test or monitor the effectiveness of the safeguards’ key controls, systems, and procedures;
- Develop and use reasonable steps to select and retain service providers capable of appropriately safeguarding personal information they receive from Accretive Health, and require service providers by contract to implement and maintain appropriate safeguards; and
- Evaluate and adjust its information security program in light of the results of testing and monitoring, any material changes to operations or business arrangements, or any other circumstances that it knows or has reason to know may have a material impact on its information security program.

Part III of the proposed order requires Accretive Health to obtain within the first one hundred eighty (180) days after service of the order, and on a biennial basis thereafter for a period of twenty (20) years, an assessment and report from a qualified, objective, independent third-party professional, certifying, among other things, that: (1) It has in place a security program that provides protections that meet or exceed the protections required by Part II of the proposed order; and (2) its security program is operating with sufficient effectiveness to provide reasonable assurance that the security, confidentiality, and integrity of sensitive consumer, information has been protected.

Parts IV through VIII of the proposed order are reporting and compliance provisions. Part IV requires Accretive Health to retain documents relating to its compliance with the order. For most records, the order requires that the documents be retained for a five-year period. For the third-party assessments and supporting documents, Accretive Health must retain the documents for a period of five years after the date that each assessment is prepared. Part V requires dissemination of the order now and in the future to all current and future principals, officers, directors, and managers, and to persons with responsibilities relating to the subject matter of the order. Part VI ensures notification to the FTC of changes in corporate status. Part VII mandates that Accretive Health submit a compliance report to the FTC within 60 days, and periodically thereafter as requested. Part VIII is a provision “sunsetting” the order after twenty (20) years, with certain exceptions.

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 complaint or order or to modify the order’s terms in any way.

By direction of the Commission.

Janice Podoll Frankle,
Acting Secretary.

[FR Doc. 2014–00373 Filed 1–10–14; 8:45 am]
BILLING CODE 6750–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA–2013–N–0001]

Request for Notification From Industry Organizations Interested in Participating in the Selection Process for Nonvoting Industry Representatives and Request for Nominations for Nonvoting Industry Representatives on the Pharmacy Compounding Advisory Committee

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is requesting that industry organizations interested in participating in the selection of nonvoting industry representatives to represent the interests of the pharmacy compounding industry and the pharmaceutical manufacturing industry on the Pharmacy Compounding Advisory Committee for the Center for Drug Evaluation and Research notify FDA in writing. A nominee may either be self-nominated or nominated by an organization to serve as a nonvoting industry representative. Nominations will be accepted for two vacancies effective with this notice.

FDA seeks to include the views of women and men, members of all racial and ethnic groups, and individuals with and without disabilities on its advisory committees, and therefore, encourages nominations of appropriately qualified candidates from these groups. Specifically, in this document, nominations for nonvoting representatives of industry interests are encouraged from the pharmacy compounding industry and the pharmaceutical manufacturing industry.

DATES: Any industry organization interested in participating in the selection of appropriate nonvoting members to represent the interests of the pharmacy compounding industry and the pharmaceutical manufacturing industry should send a letter stating the interest to FDA by February 12, 2014, for the vacancies listed in this notice. Concurrently, nomination materials for prospective candidates should be sent to FDA by February 12, 2014.

ADDRESSES: All letters of interest and nominations should be submitted electronically to PCAC@fda.hhs.gov, or in writing by mail to Janey E. Peterson, Center for Drug Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 31, Rm. 2417, Silver Spring, MD 20993.

FOR FURTHER INFORMATION CONTACT: Janey E. Peterson, Center for Drug Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 31, Rm. 2417, Silver Spring, MD 20993, 301–796–9001, FAX: 301–847–8533, email: PCAC@fda.hhs.gov.

SUPPLEMENTARY INFORMATION: The Agency requests nominations for nonvoting industry representatives on the Pharmacy Compounding Advisory Committee (the Committee) to represent the interests of the pharmacy compounding industry and the pharmaceutical manufacturing industry.

I. Pharmacy Compounding Advisory Committee

The Committee advises the Commissioner of Food and Drugs (the Commissioner) or designee in discharging responsibilities as they relate to the regulation of compounded drug products. The Committee provides advice on scientific, technical, and medical issues concerning drug compounding under sections 503A and 503B of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 353a and 353b), and as required, any other product for which FDA has regulatory responsibility. The Committee also makes appropriate recommendations to the Commissioner.

The Committee will include one or more nonvoting members who represent industry interests. The Committee will include one representative of the pharmacy compounding industry and