Family Group acting in concert: the Dow R. Hughes Revocable Trust, Dow Hughes, trustee; the Deanne D. Hughes Revocable Trust, Deanne D. Hughes, trustee; DRH, LLC; Dave Hughes; Michelle Hughes; and the David G. Dutton Living Trust, David Dutton, trustee; all of Tulsa, Oklahoma, to retain control of Regent Capital Corporation, and thereby control Regent Bank, both in Nowata, Oklahoma:


Margaret McCloşey Shanks,
Deputy Secretary of the Board.
[FR Doc. 2013–17637 Filed 7–22–13; 8:45 am]
BILLING CODE 6210–01–P

FEDERAL RESERVE SYSTEM

Formations of, Acquisitions by, and Mergers of Bank Holding Companies

The companies listed in this notice have applied to the Board for approval, pursuant to the Bank Holding Company Act of 1956 (12 U.S.C. 1841 et seq.) (BHC Act), Regulation Y (12 CFR Part 225), and all other applicable statutes and regulations to become a bank holding company and/or to acquire the assets or the ownership of, control of, or the power to vote shares of a bank or bank holding company and all of the banks and nonbanking companies owned by the bank holding company, including the companies listed below.

The applications listed below, as well as other related filings required by the Board, are available for immediate inspection at the Federal Reserve Bank indicated. The applications will also be available for inspection at the offices of the Board of Governors. Interested persons may express their views in writing on the standards enumerated in the BHC Act (12 U.S.C. 1842(c)). If the proposal also involves the acquisition of a nonbanking company, the review also includes whether the acquisition of the nonbanking company complies with the standards in section 4 of the BHC Act (12 U.S.C. 1843). Unless otherwise noted, nonbanking activities will be conducted throughout the United States.

Unless otherwise noted, comments regarding each of these applications must be received at the Reserve Bank indicated or the offices of the Board of Governors not later than August 5, 2013.

A. Federal Reserve Bank of Kansas City (Dennis Denney, Assistant Vice President) 1 Memorial Drive, Kansas City, Missouri 64198–0001:

1. Dow R. Hughes, Tulsa, Oklahoma, individually and as fiduciary, to retain control of Regent Capital Corporation, parent of Regent Bank, both in Nowata, Oklahoma. In addition, notification by the following members of the Hughes Bank, National Association, Scottsdale, Arizona.


Margaret McCloşey Shanks,
Deputy Secretary of the Board.
[FR Doc. 2013–17637 Filed 7–22–13; 8:45 am]
BILLING CODE 6210–01–P

FEDERAL TRADE COMMISSION

Agency Information Collection Activities: Submission for OMB Review; Comment Request

AGENCY: Federal Trade Commission.

ACTION: Notice and request for comment.

SUMMARY: In compliance with the Paperwork Reduction Act (PRA) of 1995, the FTC is seeking public comments on its request to OMB for a three-year extension of the current PRA clearance for the information collection requirements contained in the Contact Lens Rule. That clearance expires on July 31, 2013 (OMB Control No. 3084–0127).

DATES: Comments must be received by August 22, 2013.

ADDRESSES: Interested parties may file a comment online or on paper, by following the instructions in the Request for Comment part of the SUPPLEMENTARY INFORMATION section below.

FOR FURTHER INFORMATION CONTACT: Requests for additional information or copies of the proposed information requirements should be addressed to Alysa S. Bernstein, Attorney, or Bonnie McGregor, Federal Trade Investigator, Division of Advertising Practices, Bureau of Consumer Protection, Federal Trade Commission, 600 Pennsylvania Avenue NW, Washington, DC 20580, (202) 326–3289 (Bernstein) and (202) 326–2356 (McGregor).

SUPPLEMENTARY INFORMATION:

Title: Contact Lens Rule (Rule), 16 CFR Part 315.

OMB Control Number: 3084–0127.

Type of Review: Extension of a currently approved collection.

Abstract: The FTC promulgated the Rule pursuant to the Fairness to Contact Lens Consumers Act (FCLCA), Public Law 108–164 (Dec. 6, 2003), which was enacted to enable consumers to purchase contact lenses from the seller of their choice. The Rule became effective on August 2, 2004. As mandated by the FCLCA, the Rule requires the release and verification of contact lens prescriptions and contains recordkeeping requirements applying to
both prescribers and sellers of contact lenses.

Specifically, the Rule requires that prescribers provide a copy of the prescription to the consumer upon completion of a contact lens fitting and verify or provide prescriptions to authorized third parties. The Rule also mandates that a contact lens seller may sell contact lenses only in accordance with a prescription that the seller either: (a) has received from the patient or prescriber; or (b) has verified through direct communication with the prescriber. In addition, the Rule imposes recordkeeping requirements on contact lens prescribers and sellers. For example, the Rule requires prescribers to document in their patients’ records the medical reasons for setting a contact lens prescription expiration date of less than one year. The Rule requires contact lens sellers to maintain for three years records of all direct communications involved in obtaining verification of a contact lens prescription, as well as prescriptions, or copies thereof, that they receive directly from customers or prescribers.

The information retained under the Rule’s recordkeeping requirements is used by the Commission to determine compliance with the Rule and may also provide a basis for the Commission to bring an enforcement action. Without the required records, it would be difficult either to ensure that entities are complying with the Rule’s requirements or to bring enforcement actions for Rule violations.

On February 8, 2013, the Commission sought comment on the Rule’s information collection requirements. One comment was received, from the American Optometric Association (“AOA”). That comment stated that the majority of the information collected by the FTC is accurate, but it provided alternate figures for some data, expressed disfavor of passive verification, and sought more effective enforcement of the Rule. Data provided by the AOA is reflected in updated burden estimates set out below and are addressed in more detail within the Agency’s “Supporting Statement for Information Collection Provisions of the Contact Lens Rule,” which is available upon request from the FTC contact officials and separately at www.reginfo.gov.

As required by OMB regulations, 5 CFR Part 1320, the FTC is providing this second opportunity for public comment.

Likely Respondents: Contact lens prescribers and contact lens sellers.

Estimated Annual Hours Burden:

1,594,981 hours (derived from 685,514 hours + 909,467 hours)

- Contact Lens Prescribers: 633,333 hours (38 million contact lens wearers × 1 minute per prescription/60 minutes) + 52,181 hours (1,043,613 wearers × 3 minutes/60 minutes) = 685,514 hours
- Contact Lens Sellers: 852,625 hours (10,231,500 wearers × 5 minutes/60 minutes) + 56,842 hours (3,410,500 wearers × 1 minute/60 minutes) = 909,467 hours.

Estimated Annual Cost Burden:

$48,991,000 (rounded to the nearest thousand), which is derived from ($52.80 × 685,514 hours) + ($14.07 × 909,467 hours) = $54,991,340.9

Request for Comment

You can file a comment online or on paper. For the Commission to consider your comment, we must receive it on or before August 22, 2013. Write “Contact Lens Rule: FTC File No. P054510” 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, such as 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 “[t]rade 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 are required 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, grans 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 comment online, or to send it to the Commission by courier or overnight service. To make sure that the Commission considers your online comment, you must file it at https://ftcpubliccommentworkshop.com/ftc/contactlensruleprocedure following the instructions on the Web-based form. If this Notice appears at https://www.regulations.gov, you also may file a comment through that Web site.

If you file your comment on paper, write “Contact Lens Rule: FTC File No. P054510” 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 J), 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. 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 August 22, 2013. 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.shtm.

Comments on the information collection requirements subject to review under the PRA should also be submitted to OMB. If sent by U.S. mail, address comments to: Office of Information and Regulatory Affairs, Office of Management and Budget, Attention: Desk Officer for the Federal Trade Commission, New Executive Office Building, Docket Library, Room 10102, 725 17th Street NW., Washington, DC 20503. Comments sent to OMB by U.S. postal mail, however,
DEPARTMENT OF HEALTH AND HUMAN SERVICES

Office of the Secretary

Notice of Interest Rate on Overdue Debts

Section 30.18 of the Department of Health and Human Services’ claims collection regulations (45 CFR Part 30) provides that the Secretary shall charge an annual rate of interest, which is determined and fixed by the Secretary of the Treasury after considering private consumer rates of interest on the date that the Department of Health and Human Services becomes entitled to recovery. The rate cannot be lower than the Department of Treasury’s current value of funds rate or the applicable rate determined from the “Schedule of Certified Interest Rates with Range of Maturities” unless the Secretary waives interest in whole or part, or a different rate is prescribed by statute, contract, or repayment agreement. The Secretary of the Treasury may revise this rate quarterly. The Department of Health and Human Services publishes this rate in the Federal Register.

The current rate of 10 3/8% as fixed by the Secretary of the Treasury, is certified for the quarter ended June 30, 2013. This rate is based on the Interest Rates for the quarter ended June 30, 2013. The rate cannot be lower than the Department of Treasury’s current value of funds rate or the applicable rate determined from the “Schedule of Certified Interest Rates with Range of Maturities” unless the Secretary waives interest in whole or part, or a different rate is prescribed by statute, contract, or repayment agreement. The Secretary of the Treasury may revise this rate quarterly. The Department of Health and Human Services publishes this rate in the Federal Register.

The current rate of 10 3/8% as fixed by the Secretary of the Treasury, is certified for the quarter ended June 30, 2013. This rate is based on the Interest Rates for the quarter ended June 30, 2013. The rate cannot be lower than the Department of Treasury’s current value of funds rate or the applicable rate determined from the “Schedule of Certified Interest Rates with Range of Maturities” unless the Secretary waives interest in whole or part, or a different rate is prescribed by statute, contract, or repayment agreement. The Secretary of the Treasury may revise this rate quarterly. The Department of Health and Human Services publishes this rate in the Federal Register.

Dated: July 12, 2013.

Margie Yanchuk,
Director, Associate Deputy Assistant Secretary.

BILLING CODE 4701–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[FR Doc. 2013–17560 Filed 7–22–13; 8:45 am]

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing an opportunity for public comment on the proposed collection of certain information by the Agency. Under the Paperwork Reduction Act of 1995 (the PRA), Federal Agencies are required to publish notice in the Federal Register concerning each proposed collection of information, including each proposed extension of an existing collection of information, and to allow 60 days for public comment in response to the notice. This notice solicits comments on the standardized format and content requirements for the labeling of over-the-counter (OTC) drug products. DATES: Submit either electronic or written comments on the collection of information by September 23, 2013.

ADDRESSES: Submit electronic comments on the collection of information to http://www.regulations.gov. Submit written comments on the collection of information to the Division of Dockets Management (HFA–305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852. All comments should be identified with the docket number found in brackets in the heading of this document.

FOR FURTHER INFORMATION CONTACT: Ila S. Mizrachi, Office of Information Management, Food and Drug Administration, 1350 Piccard Dr., P150–400B, Rockville, MD 20850, 301–796–7726, ila.mizrachi@fda.hhs.gov.

SUPPLEMENTARY INFORMATION: Under the PRA (44 U.S.C. 3501–3520), Federal Agencies must obtain approval from the Office of Management and Budget (OMB) for each collection of information they conduct or sponsor. “Collection of information” is defined in 44 U.S.C. 3502(3) and 5 CFR 1320.3(c) and includes Agency requests or requirements that members of the public submit reports, keep records, or provide information to a third party. Section 3506(c)(2)(A) of the PRA (44 U.S.C. 3506(c)(2)(A)) requires Federal Agencies to provide a 60-day notice in the Federal Register concerning each proposed collection of information, including each proposed extension of an existing collection of information, before submitting the collection to OMB for approval. To comply with this requirement, FDA is publishing notice of the proposed collection of information set forth in this document.

With respect to the following collection of information, FDA invites comments on these topics: (1) Whether the proposed collection of information is necessary for the proper performance of FDA’s functions, including whether the information will have practical utility; (2) the accuracy of FDA’s estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used; (3) ways to enhance the quality, utility, and clarity of the information to be collected; (4) ways to minimize the burden of the collection of information on respondents, including through the use of automated collection techniques, when appropriate, and other forms of information technology.

Format and Content Requirements for OTC Drug Product Labeling—(OMB Control Number 0910–0340)—Extension

In the Federal Register of March 17, 1999 (64 FR 13254) (the 1999 labeling final rule), we amended our regulations governing requirements for human drug products to establish standardized format and content requirements for the labeling of all marketed OTC drug products in part 201 (21 CFR Part 201). The regulations in part 201 require OTC drug product labeling to include uniform headings and subheadings, presented in a standardized order, with minimum standards for type size and other graphical features. Specifically, the 1999 labeling final rule added new § 201.66 to part 201. Section 201.66 sets content and format requirements for the Drug Facts portion of labels on OTC drug products.

On June 20, 2000 (65 FR 38191), we published a Federal Register final rule that required all OTC drug products marketed under the OTC monograph system to comply with the labeling requirements in § 201.66 by May 16, 2005, or sooner (65 FR 38191 at 38193). Currently marketed OTC drug products are already required to be in compliance with these labeling requirements, and thus will incur no further burden to comply with Drug Facts labeling requirements in § 201.66. Modifications of labeling already required to be in Drug Facts format are usual and customary as part of routine redesign practice, and thus do not create...