National Vaccine Plan and updates from other Working Groups. If there is a change in meeting dates this information will be posted on the NVAC Web site (http://www.hhs.gov/nvpo/nvac/) as soon as the pertinent information becomes available.

For these special meetings, members of the public are invited to attend by telephone via a toll-free call-in phone number. The call-in number will be operator assisted to provide members of the public the opportunity to provide comments to the Committee. Public participation and ability to comment will be limited to space and time available. Public comment will be limited to no more than three minutes per speaker. Pre-registration is required for public comment only. Individuals who plan to attend and need special assistance, such as accommodation for hearing impairment or other reasonable accommodations, should notify the designated contact person at least one week prior to the meeting.

Any members of the public who wish to have written material distributed to NVAC should submit materials to the Executive Secretary, NVAC, through the contact person listed above prior to close of business one week before each meeting (conference call). A draft agenda and any additional materials will be posted on the NVAC Web site (http://www.hhs.gov/nvpo/nvac/) prior to the meeting.

Dated: June 1, 2010.

Bruce Gellin,
Deputy Assistant Secretary for Health, Director, National Vaccine Program Office, Executive Secretary, NVAC.

For further information contact:
Elizabeth Berbakos, Office of Information Management, Food and Drug Administration, 1350 Piccard Dr., PI50–400B, Rockville, MD 20850, 301–1061, Rockville, MD 20852. All comments should be identified with the docket number found in brackets in the heading of this document.

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA–2010–N–0266]

Agency Information Collection Activities; Proposed Collection; Comment Request; Study of Clinical Efficacy Information in Professional Labeling and Direct-to-Consumer Print Advertisements for Prescription Drugs

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, and to allow 60 days for public comment in response to the notice. This notice solicits comments on the Study of Clinical Efficacy Information in Professional Labeling and Direct-to-Consumer (DTC) Print Advertisements for Prescription Drugs. This study is designed to investigate efficacy and effectiveness of prescription drugs as conveyed to healthcare providers through approved labeling and to consumers through print advertisements.

DATES: Submit either electronic or written comments on the collection of information by August 16, 2010.

ADDRESSES: Submit electronic comments on the collection of information to http://www.regulations.gov. Submit written comments on the collection of information to 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:
Elizabeth Berbakos, Office of Information Management, Food and Drug Administration, 1350 Piccard Dr., PI50–400B, Rockville, MD 20850, 301–796–3792.
Elizabeth.Berbakos@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 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; and (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.

Study of Clinical Efficacy Information in Professional Labeling and Direct-to-Consumer (DTC) Print Advertisements for Prescription Drugs—New

FDA regulations require that an advertisement that makes claims about a prescription drug include a “fair balance” of information about the benefits and risks of the advertised product, in terms of both content and presentation (§ 202.1(e)(5)(ii) (21 CFR 202.1(e)(5)(ii))). In past research, FDA has focused primarily on the risk component of the risk-benefit ratio. In the interest of thoroughly exploring the issue of fair balance, however, the presentation of effectiveness, or benefit, information is equally important.

The act requires that manufacturers, packers, and distributors (sponsors) who advertise prescription human and animal drugs, including biological products for humans, disclose in advertisements certain information about the advertised product’s uses and risks.1 By its nature, the presentation of this risk information is likely to evoke active tradeoffs by consumers, i.e., comparisons with the perceived risks of not taking treatment, and comparisons with the perceived benefits of taking a treatment.2 Because FDA has an interest in fostering safe and proper use of prescription drugs, an activity that engages both risks and benefits, an indepth understanding of consumers’ processing of this information is central to this regulatory task.

Research and guidance to sponsors on how to present benefit and efficacy information in prescription drug advertisements is limited. For example, “benefit claims,” broadly defined, appearing in advertisements are often presented in general language that does not inform patients of the likelihood of efficacy and are often simply variants of

1 For prescription drugs and biologics, the act requires advertisements to contain “information in brief summary relating to side effects, contraindications, and effectiveness” (21 CFR 202.1(e)(1)).

an “intended use” statement. In a content analysis of DTC advertising, the researchers classified the “promotional techniques” used in the advertisements. Emotional appeals were observed in 67 percent of the ads while vague and qualitative benefit terminology was found in 87 percent of the ads. Only 9 percent contained data. For risk information, however, half the advertisements used data to describe side-effects, typically with lists of side-effects that generally occurred infrequently.

FDA regulations require that prescription drug advertisements that make (promotional) claims about a product also include risk information in a “balanced” manner (§ 202.1(e)(5)(ii)), both in terms of the content and presentation of the information. This balance applies to both the front (a.k.a. “display”) page of an advertisement, as well as the brief summary page. However, beyond the “balance” requirement limited guidance and research exists to direct or encourage sponsors to present benefit claims that are informative, specific, and reflect clinical effectiveness data.

The purpose of this project is to: (1) Understand how physicians process clinical efficacy information and how they interpret approved product label information; (2) determine physician preferences for alternative presentations of clinical efficacy information in DTC advertising, and (3) examine how different presentations of clinical efficacy information in DTC advertising affect consumers’ perceptions of efficacy and safety. Specifically, we are interested in how physicians and consumers make risk/benefit assessments and particularly, how consumers make such judgments in response to variations in the efficacy presentations in the “display” (first) page of a DTC print ad. A particular concern is whether certain presentations cause consumers to form skewed perceptions or unfounded risk/benefit tradeoffs. Therefore, we will investigate to what extent consumers, when provided with efficacy information, form perceptions that correspond with clinically-based physicians’ assessments of the benefits, risks, and benefit/risk tradeoffs of the same drugs. These studies will inform FDA’s thinking regarding how manufacturers may provide useful and non-misleading efficacy information in DTC print advertisements.

**Design Overview**

This study will be conducted in two concurrent, independent parts. The first part will involve 2,500 consumers in an experimental examination of variations of the display page of print DTC ads for two fictitious drugs, closely approximating existing drugs for overactive bladder (OAB) and benign prostatic hyperplasia (BPH). In the second part, 600 general practitioners will review and evaluate a fictitious “approved” label for the same conditions. This design will allow us to compare consumers’ perceptions of efficacy with a more objective measure of the true efficacy of the drug as measured by physician perceptions of clinical efficacy from labeling.

**Consumer Experiment.** In this part of the study, we have been interested in consumers who have been diagnosed with or are at risk for OAB (self-designated based on relevant symptoms) will be recruited and will view one version of a DTC ad for a drug to treat OAB. Men who have been diagnosed with or are at risk for BPH (self-designated based on relevant symptoms) will be recruited and will view one version of a DTC ad for a drug to treat BPH. Although the two conditions are somewhat specific to gender (men can suffer from OAB but it is much more prevalent in women), they share many of the same symptoms and characteristics. These medical conditions afford us the ability to maintain various realistic manipulations of placebo level and type of claim, as explained in the following paragraphs. The graphical elements and construction of the two ads will be comparable yet still realistic.

Consumers will be randomly assigned to see 1 of 12 DTC print ads within their respective medical condition and will answer questions about the effectiveness and safety of the fictitious drug advertised in them. These 12 experimental conditions will be created by examining three independent variables in the following manner: Type of claim (2 levels: treatment, prevention), placebo rate (3 levels: high, low, none), and framing (2 levels: single, mixed). Please note that the numbers describing efficacy seen in the table are for illustration only. Actual numbers used will be determined by pretesting.

![Table](image_url)

---


4 As part of this effort, a qualitative mental models procedure was completed that helped us determine how physicians think about the efficacy of potential pharmaceutical options (OMB control no. 0910–0649).
We will investigate variations of numerical presentation in two different types of claims: Treatment and prevention. Treatment claims usually involve symptoms that may be alleviated by taking a given prescription drug. This type of claim is directly observable and somewhat testable by patients. If bothersome symptoms do not go away, a patient can return to the healthcare provider with this information and pursue additional options for treatment. In general, drugs that treat symptoms typically show clinical meaningful but objectively noticeable (e.g., risk of bladder cancer) improvements. While we will examine the current issues in both treatment and prevention claims, we do not intend to make comparisons between the two.

The second variable of interest is communication of a placebo rate. Three levels will be examined. In addition to testing a control condition with no placebo information, we will utilize a high and low placebo rate to better understand if and how consumers use placebo information. We see three possibilities: (1) People use placebo numbers correctly, such that the low placebo group demonstrates higher perceived efficacy than the high placebo group, (2) people use the placebo numbers as a peripheral cue to mean “science” so there are no differences between high and low placebo groups on perceived efficacy but both are higher than the no placebo group and (3) people do not find the numbers meaningful or cannot process them, so the high and low groups do not differ from one another and they do not differ from the no placebo group. In an attempt to make our claims as realistic as possible, we will maintain fairly low rates of prevention in the prevention conditions. For this reason, in addition to the 12 cells in the table previously illustrated in this document, we will also have an additional control cell in which the effectiveness rates are quite high—higher than could reasonably be expected but high enough to be objectively noticeable (e.g., risk of bladder cancer on Drug X, 4/100; risk of bladder cancer on placebo, 15/100).

This additional condition will provide confidence that our research manipulations are operating as we expect.

Finally, we will examine the addition of mixed framing to the traditional use of a single positive frame in a DTC ad. Mixed framing provides the number of people who benefited and the number of people who did not benefit, whereas positive framing provides only the number of people who benefited. Only a few studies have actually measured

<table>
<thead>
<tr>
<th>Extra High Efficacy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Treatment Claim Study</td>
</tr>
<tr>
<td>Frame</td>
</tr>
<tr>
<td>Low</td>
</tr>
<tr>
<td>None</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
</tbody>
</table>
this mixed approach although risk communication guides recommend the use of mixed framing to create more accurate perceptions. Although a completely balanced design would also include a negative framing condition (which would provide only the number of people who did not benefit), we feel it is unrealistic to create an ad that would suggest, for example, that “Drug X did not work for 70% of people in clinical trials,” so we have chosen not to include negative framing in our investigation.

In this part of the project, we are most interested in consumers’ perceived efficacy and safety, which we can then compare with ratings physicians will provide based on the prescribing information, described in the next section. We will also ask consumers questions to measure their accuracy with regard to claims, their recall of the information in the ad, and demographic questions that may influence their responses, such as knowledge about their medical condition and their level of numeracy.

Physician Study. Six hundred general practitioners will participate in an Internet survey lasting no longer than 20 minutes. They will complete two tasks during this time. In the first task, they will evaluate a prescription drug label (also known as the prescribing information, written for healthcare practitioners) for one of the two fictitious drugs described in the consumer study located in the following paragraphs. To provide a match for the variations of information in the DTC ads the consumers will observe, physicians will be randomly assigned to see prescribing information that varies in terms of claim type, placebo rates in clinical trials, and the medical condition the drug treats (OAB or BPH).

As part of this task, we will obtain timing and sequence information on which sections of the label physicians examine. This will enable us to have a deeper understanding of physicians’ processing of the prescribing information. We are not aware of existing literature on this topic. Additionally, physicians will answer questions about the efficacy and safety of the drug and quantitative questions about the benefit shown in the clinical studies (as described in the label). These questions have been designed such that they can be reasonably compared with the responses of consumers who will answer the same questions after viewing a corresponding DTC ad.

In the second task, physicians will see four versions of a print DTC ad for a fictitious product for high cholesterol and will rank the ads in order of how representative of the clinical data as the physicians know it the ads are and how useful they believe the ads would be for their patients. The four versions will be selected to mirror the versions of the OAB/BPH drug that consumers will see in the consumer experiment (i.e., low placebo, frame).

Thus, this research will provide us with a rich data set in order to address several questions: (1) How physicians process clinical efficacy information and how they use approved product label information, (2) how physicians’ interpretations of clinical efficacy information relate to their preferences for alternative DTC ad presentations, and (3) which variations of information in DTC ads bring consumers closer to or farther away from the conclusions of the physicians regarding the same drugs.

The total respondent sample for this data collection is 3,400. We estimate the response burden to be 20 minutes in the first part and 15 minutes in the second part, for a burden of 906 hours.

The response burden chart is listed below.

<table>
<thead>
<tr>
<th>TABLE 1.—ESTIMATED ANNUAL REPORTING BURDEN1</th>
</tr>
</thead>
<tbody>
<tr>
<td>21 CFR Section</td>
</tr>
<tr>
<td>----------------</td>
</tr>
<tr>
<td>Physician survey-pretest</td>
</tr>
<tr>
<td>Physician survey-main study</td>
</tr>
<tr>
<td>Consumer experiment-pretest</td>
</tr>
<tr>
<td>Consumer experiment-main study</td>
</tr>
<tr>
<td>Total</td>
</tr>
</tbody>
</table>

1 There are no capital costs or operating and maintenance costs associated with this collection of information.


7 Including internists, general practitioners, and family practitioners.

8 To reduce burden, the physician sample will be split in this task, so that half of the physicians see the four ad versions with treatment claims and the other half see the four ad versions with prevention claims. Type of claim is described in greater detail in the consumer experiment section.
DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Proposed Collection; Comment Request Resource for the Collection and Evaluation of Human Tissues and Cells From Donors With an Epidemiology Profile (NCI)

SUMMARY: In compliance with the requirement of Section 3506(c) (2)(A) of the Paperwork Reduction Act of 1995, for opportunity for public comment on proposed data collection projects, the National Cancer Institute [NCI], the National Institutes of Health [NIH] will publish periodic summaries of proposed projects to be submitted to the Office of Management and Budget [OMB] for review and approval. Proposed Collection: Title: Resource for the Collection and Evaluation of Human Tissues and Cells From Donors With an Epidemiology Profile [NCI]. Type of Information Collection Request: Now. Need and Use of Information Collection: Under the auspices of three NCI IRB-approved protocols and instruments, the Laboratory of Human Carcinogenesis conducts case-control studies to investigate the relations between biomarkers, the environment, and human cancer. Human subjects recruited from the general population are needed as controls (Population Controls) for bio-specimens and personal histories (social, occupational and health) that serve as references for the significance of the frequency and prevalence of bio-markers found in cancer patients and thought to be important in the development, progression, and/or response to treatment of the malignant growths in cancer patients. The questionnaires will be used to obtain the personal histories to compare to the life styles and exposures and the biospecimens will serve as controls for the assay results obtained from cancer patients. The collection of information and specimens from the cancer cases received NIH Clinical Exemption [Request #2009–09–002] on October 28, 2009. Frequency of Response: Once. Affected Public: Adult and senior members of the licensed driver population in Baltimore, Maryland and eleven nearby counties, including the Eastern Shore. Type of Respondents: Responders will be English speaking, male and female, Caucasian, African-American and Asian. The total annual reporting burden is estimated to be 692 (see table below). There are no Capital Costs, Operating Costs, and/or Maintenance Costs to report.

<table>
<thead>
<tr>
<th>Type of respondents</th>
<th>Survey instrument</th>
<th>Number of respondents</th>
<th>Frequency of response</th>
<th>Average time per response (minutes/hour)</th>
<th>Annual burden hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adults (40–79 years old)</td>
<td>Telephone Screener (Attachment 2)</td>
<td>1700</td>
<td>1</td>
<td>10/60 (0.17)</td>
<td>283</td>
</tr>
<tr>
<td></td>
<td>Main Questionnaire (Attachment 6)</td>
<td>225</td>
<td>1</td>
<td>60/60 (1)</td>
<td>225</td>
</tr>
<tr>
<td></td>
<td>Prostate Supplemental Questionnaire (Attachment 7)</td>
<td>125</td>
<td>1</td>
<td>30/60 (0.5)</td>
<td>63</td>
</tr>
<tr>
<td></td>
<td>Liver Supplement (Attachment 8)</td>
<td>225</td>
<td>1</td>
<td>30/60 (0.5)</td>
<td>113</td>
</tr>
<tr>
<td></td>
<td>Refusal Questionnaire Form (Attachment 21)</td>
<td>225</td>
<td>1</td>
<td>2/60 (0.03)</td>
<td>8</td>
</tr>
<tr>
<td>Totals</td>
<td></td>
<td>2500</td>
<td></td>
<td></td>
<td>692</td>
</tr>
</tbody>
</table>

Request for Comments: Written comments and/or suggestions from the public and affected agencies should address one or more of the following points: (1) Evaluate whether the proposed collection of information is necessary for the proper performance of the function of the agency, including whether the information will have practical utility; (2) Evaluate the accuracy of the agency’s estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used; (3) Enhance the quality, utility, and clarity of the information to be collected; and (4) Minimize the burden of the collection of information on those who are to respond, including the use of appropriate automated, electronic, mechanical, or other technological collection techniques or other forms of information technology.

FOR FURTHER INFORMATION CONTACT: To request more information on the proposed project or to obtain a copy of the data collection plans and instruments, contact Glenwood E. Trivers or Elise Bowman, Center for Cancer Research, NCI, NIH, 37 Convent Drive, Room 3060–C or 3060–A, Building 37, Bethesda, Maryland 30893–4258 or call non-toll-free number 301–496–2094 or 301–496–2090 or e-mail your request, including your address to triversg@mail.nih.gov or bowmane@mail.nih.gov.

Comments Due Date: Comments regarding this information collection are best assured of having their full effect if received within 60 days of the date of this publication.

Dated: June 9, 2010.

Vivian Horovitch-Kelley,
NCI Project Clearance Liaison, National Institutes of Health.

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Draft Guideline for the Prevention and Control of Norovirus Gastroenteritis Outbreaks in Healthcare Settings

AGENCY: Centers for Disease Control and Prevention [CDC], Department of Health and Human Services [DHHS].

ACTION: Notice of availability and request for public comment.

SUMMARY: This notice is a request for review of and comment on the Draft Guideline for the Prevention and Control of Norovirus Gastroenteritis Outbreaks in Healthcare Settings, available on the following Web site: http://www.cdc.gov/publiccomments/. This document is for use by infection prevention staff, healthcare epidemiologists, healthcare administrators, nurses, other healthcare...