providers accurately report data on all Medicare enrolled patients; and all patient confidentiality, privacy, and other Federal laws must be followed. Consistent with section 1142 of the Social Security Act (the Act), the Agency for Healthcare Research and Quality (AHRQ) supports clinical research studies that CMS determines meets specified standards and addresses the specified research questions. To qualify for payment, providers must prescribe certain NaF–18 PET scans for beneficiaries with a set of clinical criteria specific to each solid tumor. The statutory authority for this policy is section 1862(a)(1)(E) of the Act. The need to prospectively collect information at the time of the scan is to assist the provider in decision making for patient management. Form Number: CMS–10152 (OCN: 0938–0968); Frequency: Annual; Affected Public: Private Sector (Business or other for-profits); Number of Respondents: 25,000; Total Annual Responses: 25,000; Total Annual Hours: 2,084 hours. (For policy questions regarding this collection contact Stuart Caplan at 410–736–8564.)

Dated: March 1, 2016.

William N. Parham, III,
Director, Paperwork Reduction Staff, Office of Strategic Operations and Regulatory Affairs.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA–2016–N–0321]

Risk Assessment of Foodborne Illness Associated With Pathogens From Produce Grown in Fields Amended With Untreated Biological Soil Amendments of Animal Origin; Request for Scientific Data, Information, and Comments

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice; request for comments and for scientific data and information.

SUMMARY: The Food and Drug Administration (FDA or we) is requesting scientific data, information, and comments that would assist the Agency in its plan to develop a risk assessment for produce grown in fields or other growing areas amended with untreated biological soil amendments of animal origin (including raw manure). The risk assessment will evaluate and, if feasible, quantify the risk of human illness associated with consumption of produce grown in fields or other growing areas amended with untreated biological soil amendments of animal origin that are potentially contaminated with enteric pathogens, such as Escherichia coli O157:H7 or Salmonella. The risk assessment will also evaluate the impact of certain interventions, such as use of a time interval between application of the soil amendment and crop harvest, on the predicted risk. The risk assessment is intended to inform policy decisions with regard to produce safety.

DATES: Submit either electronic or written comments and scientific data and information by May 3, 2016.

ADDRESSES: You may submit comments and scientific data and information as follows:

Electronic Submissions

Submit electronic comments and scientific data and information in the following way:

• Federal eRulemaking Portal: http://www.regulations.gov. Follow the instructions for submitting comments. Comments and scientific data and information submitted electronically, including attachments, to http://www.regulations.gov will be posted to the docket unchanged. Because your comment will be made public, you are solely responsible for ensuring that your comment does not include any confidential information that you or a third party may not wish to be posted, such as medical information, your or anyone else’s Social Security number, or confidential business information, such as a manufacturing process. Please note that if you include your name, contact information, or other information that identifies you in the body of your comments and scientific data and information, that information will be posted on http://www.regulations.gov.

• If you want to submit a comment with confidential information that you do not wish to be made publicly available, submit your comments and scientific data and information only as a written/paper submission. You should submit two copies total. One copy will include the information you claim to be confidential with a heading or cover note that states “THIS DOCUMENT CONTAINS CONFIDENTIAL INFORMATION”. The Agency will review this copy, including the claimed confidential information, in its consideration of comments and scientific data and information. The second copy, which will have the claimed confidential information redacted/blacked out, will be available for public viewing and posted on http://www.regulations.gov. Submit both copies to the Division of Dockets Management. If you do not wish your name and contact information to be made publicly available, you can provide this information on the cover sheet and not in the body of your comments and scientific data and information and you must identify this information as “confidential.” Any information marked as “confidential” will not be disclosed except in accordance with 21 CFR 10.20 and other applicable disclosure law. For more information about FDA’s posting of comments and scientific data and information to public dockets, see 80 FR 56469, September 18, 2015, or access the information at: http://www.fda.gov/regulatoryinformation/dockets/default.htm.

Docket: For access to the docket to read background documents or the electronic and written comments and scientific data and information received, go to http://
A. What are the food safety concerns related to untreated biological soil amendments of animal origin?

Biological soil amendments of animal origin (BSAAO) can be a source of contamination of produce with pathogens that can cause human illness. Human pathogens in BSAAO, once introduced to the growing environment, may be inactivated at a rate that is dependent upon a number of environmental, regional, and other agricultural and ecological factors. The rate of pathogen population decline over time is also influenced by the types of BSAAO and application methods. Furthermore, the types of produce and whether or not BSAAO may come into contact with a harvestable portion of the crop influences the likelihood of pathogen transfer from the amended soil to produce (Ref. 1).

Some produce farms use untreated BSAAO for various reasons, including that they are inexpensive, readily available, and rich nutrient sources for growing crops. Whether it is feasible for a farm to use untreated BSAAO as a principal nutrient source depends on numerous factors, including whether there is a required time interval between application and harvest and the length of such an interval (which may affect the nutrients retained or available from BSAAO), and crop nutrient demand (i.e., the nutrients needed to support crop growth). Typical examples of untreated BSAAO are raw cattle manure, poultry litter, swine slurry, and horse manure. FDA acknowledges that required application intervals for certain uses of untreated BSAAO could influence the number of crop cycles a farm is able to undertake each year and/or the choices farms make regarding which type of amendment to apply (e.g., raw manure, composted manure, or other nutrient sources).

B. How did FDA’s rule on produce safety address BSAAO?

In January 2013, based in part upon authority provided by the FDA Food Safety Modernization Act, we published a proposed Produce Safety Rule entitled “Standards for the Growing, Harvesting, Packing, and Holding of Produce for Human Consumption” (78 FR 3504, January 16, 2013). Among other provisions related to BSAAO, the proposed rule included at § 112.56(a)(1)(i) (21 CFR 112.56(a)(1)(i)) a 9-month minimum application interval for untreated BSAAO applied in a manner that does not contact covered produce during application and minimizes the potential for contact with covered produce after application (78 FR 3504 at 3637). In response to public comments, we withdrew this proposed 9-month minimum application interval in a supplemental proposed rulemaking that we published on September 29, 2014 (79 FR 58434 at 58457 through 58461). In the supplemental proposed rule, we acknowledged the limited body of currently available scientific evidence related to the proposed 9-month interval and the need for additional research in this area, and described our planned risk assessment and research agenda (79 FR 58434 at 58460 through 58461). Accordingly, we deferred our decision on an appropriate minimum application interval.

On November 27, 2015, we published a final Produce Safety Rule entitled “Standards for the Growing, Harvesting, Packing, and Holding of Produce for Human Consumption,” (80 FR 74354). The final rule is now codified at 21 CFR part 112. In the preamble to the final rule, we restated our decision with respect to the appropriate minimum BSAAO application interval (80 FR 74354 at 74463). We reserved one of the provisions in the final rule’s Subpart F (Biological Soil Amendments of Animal Origin and Human Waste) because we continue to believe that a quantitative application interval standard is necessary and anticipate locating such a future standard in that provision. As finalized, the Produce Safety Rule establishes that there is no minimum application interval required when untreated BSAAO are applied in a manner that does not contact covered produce during or after application (§ 112.56(a)(1)(ii)), and the minimum application interval is [reserved] when applied in a manner that does not contact produce during application and minimizes the potential for contact with produce after application (§ 112.56(a)(1)(i)).

II. FDA’s Risk Assessment

FDA, in consultation with the U.S. Department of Agriculture, is conducting a risk assessment to evaluate the risk of human illness associated with the consumption of produce grown in growing areas amended with untreated BSAAO that are potentially contaminated with enteric pathogens such as E. coli O157:H7 or Salmonella. The risk assessment will evaluate the impact of different agricultural and ecological conditions and certain interventions, such as use of a time interval or intervals between application of untreated BSAAO and crop harvest, on the predicted risk. The risk assessment will take into account available data and information on relevant steps in the produce food safety continuum including: The initial prevalence and levels of pathogens in untreated BSAAO; the methods used to apply untreated BSAAO to soils; pathogen survival (and growth) in untreated BSAAO and soils amended with untreated BSAAO; pathogen transfer to produce grown in amended soils; pathogen survival and growth on produce; and pathogen survival, growth, and cross-contamination during storage and other steps in the supply chain (e.g., washing). The risk assessment will include characterization of the variability and uncertainty of pathogen survival and growth under different agricultural and ecological conditions (e.g., soil types, application methods, or geographic locations/climatic factors) and time intervals between application of untreated BSAAO and crop harvest. The risk assessment is intended to inform policy decisions with regard to produce safety.

III. Issues for Consideration

FDA is requesting comments and scientific data and other information relevant to this risk assessment. We are particularly interested in scientific data and information concerning, but not limited to, the following factors that may affect the risk of human illness associated with the consumption of produce grown in fields or other growing areas amended with untreated BSAAO (including raw manure):

1. Data on the prevalence and levels of pathogens.
   a. The frequency of detecting the presence of pathogens in untreated BSAAO and soil amended with BSAAO, such as Salmonella in poultry litter, and E. coli O157:H7 and other pathogenic Shiga-toxin producing E. coli in cattle manure. Samples may be obtained at different stages of untreated BSAAO storage prior to application, or after
application. If available, for each data point, we also invite information regarding the following:

1. The type of untreated BSAAO (e.g., animal origin and content);
2. how the untreated BSAAO, including raw manure, was sampled and handled prior to analysis;
3. the size of the analytical unit (i.e., detection limit) and test method;
4. the number of positives, the total number of samples, and the time period in which the testing was conducted; and
5. sampling protocol (e.g., simple random, stratified random, targeted).

b. The pathogen concentration, i.e., the number of pathogen cells per amount (unit volume or weight), in contaminated untreated BSAAO or soil amended with untreated BSAAO, especially cattle manure and poultry litter. If available, for each data point, we ask that the data be provided in unaggregated form and that Most Probable Number (MPN) patterns as well as raw data (e.g., number of positive and negative tubes per serial dilution) be provided.

c. Kinetic data that describe the survival (or inactivation) or growth of pathogens in soil amended with untreated BSAAO, especially cattle manure and poultry litter.

d. The pathogen concentration, i.e., the number of pathogen cells per amount (unit volume or weight), in contaminated untreated BSAAO or soil amended with untreated BSAAO, especially cattle manure and poultry litter. If available, for each data point, we ask that the data be provided in unaggregated form and that Most Probable Number (MPN) patterns as well as raw data (e.g., number of positive and negative tubes per serial dilution) be provided.

e. The survival of pathogens on produce in the field or other growing area before harvest; and

f. The variability in the survival of different Salmonella serotypes, different subtypes of E. coli O157:H7, or other pathogens of public health significance in amended soils under field, greenhouse, or laboratory conditions.

3. On-farm practices with regard to the use of untreated BSAAO, including, but not limited to, the following aspects.

a. The extent to which untreated BSAAO are used in different regions in the United States, as well outside the United States in regions that export produce to the United States;

b. The types of untreated BSAAO and the soil type, and associated physical and chemical parameters (including but not exclusive to nutrient content, moisture and pH); and the crops typically grown in each BSAAO-amended soil type;

c. Characterization of the proportion of produce farms that have one or more soil types per geographical location;

d. The amount of untreated BSAAO applied per unit surface (e.g., per acre) or the ratio of untreated BSAAO/soil, including typical ratio and variability by commodity type, including, for example, row crops such as leafy greens;

e. The time of year, number of applications, and amount of untreated BSAAO that are applied;

f. The method of application (e.g., surface, incorporated), and whether or not the amended soil is covered (e.g., with plastic mulch);

g. Produce commodity type and cropping cycles;

h. Climate conditions and irrigation practices after soil is amended, before and after planting; and

i. The crop density (e.g., the number of rows per bed and the distance between adjacent rows in a bed), distance between two crop beds (furrow width), and the influence of such factors on pathogen transfer.

4. Harvesting, handling, and storage conditions that may affect pathogen detection and levels, survival, growth, or inactivation between harvest and retail sale along the farm-to-fork continuum.

a. The harvesting practices and the average conditions as well as the range of climactic conditions prior to harvesting (e.g., time and temperature, rain events) under which produce is handled in the field and in packing operations;

b. The survival, growth, or inactivation of pathogens on produce (including, for example, specific commodities or categories such as leafy greens, or produce generally) during transportation and storage:

c. Typical storage conditions (e.g., time, temperature) for produce (including, for example, specific commodities or categories, such as leafy greens, or produce generally), from harvest until consumer purchase and whether and how those storage conditions affect pathogen levels; and

d. The types and concentration of antimicrobial chemicals or other treatments, if any, applied to the water used for wash or transport of produce during farm or other distribution operations prior to retail, and the efficacy of these treatments in reducing pathogen levels, as well as the likelihood of cross-contamination during wash or transport.

5. Storage conditions such as times and temperatures that may affect pathogen growth and/or survival during transportation and storage of produce in the consumer’s home, and consumer handling practices with respect to produce after purchase, including data and information on consumer washing practices.

We are also interested in other comments concerning, but not limited to, the types of untreated BSAAO, produce commodities, relevant agricultural and ecological conditions, and appropriate mitigation strategies that the Agency should consider in the risk assessment.

IV. Reference

The following reference is on display in the Division of Dockets Management (see ADDRESSES) and is available for viewing by interested persons between 9 a.m. and 4 p.m., Monday through Friday; they are also available electronically at http://www.regulations.gov. FDA has verified the Web site address, as of the date this document publishes in the Federal Register, but Web sites are subject to change over time.


Leslie Kux,
Associate Commissioner for Policy.

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