In ¶679.84, revise paragraph (c)(7) to read as follows:

§679.84 Rockfish Program Recordkeeping, permits, monitoring, and catch accounting.

(7) **Pre-cruise meeting.** The Observer Program is notified by phone at 1 (907) 481–1770 (Kodiak, AK) at least 24 hours prior to departure when the vessel will be carrying an observer who has not previously been deployed on that vessel within the last 12 months. Subsequent to the vessel’s departure notification, but prior to departure, NMFS may contact the vessel to arrange for a pre-cruise meeting. The pre-cruise meeting must minimally include the vessel operator or manager and any observers assigned to the vessel.

§679.93 Amendment 80 Program recordkeeping, permits, monitoring, and catch accounting.

(7) **Pre-cruise meeting.** The Observer Program is notified by phone at 1 (907) 581–2060 (Dutch Harbor, AK) or 1 (907) 481–1770 (Kodiak, AK) at least 24 hours prior to departure when the vessel will be carrying an observer who has not previously been deployed on that vessel within the last 12 months. Subsequent to the vessel’s departure notification, but prior to departure, NMFS may contact the vessel to arrange for a pre-cruise meeting. The pre-cruise meeting must minimally include the vessel operator or manager and any observers assigned to the vessel.
Commodity Exchange Act, as added by the Dodd-Frank Wall Street Reform and Consumer Protection Act. That document inadvertently failed to remove several obsolete provisions in § 1.33(a)(2) and (b)(3). Accordingly, the Commission is making a correcting amendment to § 1.33 that removes the second paragraph (a)(2)(ii), removes paragraph (a)(2)(v), and removes the introductory clause to paragraph (b)(3).

List of Subjects in 17 CFR Part 1

Agricultural commodity, Agriculture, Brokers, Committees, Commodity futures, Conflicts of interest, Consumer protection, Definitions, Designated contract markets, Directors, Major swap participants, Minimum financial requirements for intermediaries, Reporting and recordkeeping requirements, Swap dealers, Swaps.

Accordingly, 17 CFR part 1 is corrected by making the following correcting amendments:

PART 1—GENERAL REGULATIONS UNDER THE COMMODITY EXCHANGE ACT

1. The authority citation for part 1 continues to read as follows:

Authority: 7 U.S.C. 1a, 2, 5, 6, 6a, 6b, 6c, 6d, 6e, 6f, 6g, 6h, 6i, 6k, 6l, 6m, 6n, 6o, 6p, 6r, 6s, 7, 7a-1, 7a-2, 7b, 7b-3, 8, 9, 10a, 12, 12a, 12c, 13a, 13a-1, 16, 16a, 19, 21, 23, and 24 (2012).

2. Amend § 1.33 as follows:

a. Revise paragraph (a)(2) and:

b. Revise paragraph (b)(3) introductory text.

The revisions read as follows:

§ 1.33 Monthly and confirmation statements.

(a) * * *

(2) For each commodity option position and foreign option position—

(i) All commodity options and foreign options purchased, sold, exercised, or expired during the monthly reporting period, identified by underlying futures contract or underlying commodity, strike price, transaction date, and expiration date;

(ii) The open commodity option and foreign option positions carried for such customer or foreign futures or foreign options customer as of the end of the monthly reporting period, identified by underlying futures contract or underlying commodity, strike price, transaction date, and expiration date;

(iii) All open commodity option and foreign option positions marked to the market and the amount each position is carried in such customer’s account(s) or any related foreign futures or foreign options secured amount carried in the account(s) of a foreign futures or foreign options customer.

* * * * *

(b) * * *

(3) A written confirmation of each commodity option transaction, containing at least the following information:

* * * * *


Robert N. Sidman, Deputy Secretary of the Commission.

[FR Doc. 2018–13256 Filed 6–28–18; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

21 CFR Part 514

[Docket No. FDA–2012–N–0447]

Antimicrobial Animal Drug Sales and Distribution Reporting; Small Entity Compliance Guide; Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notification of availability.

SUMMARY: The Food and Drug Administration (FDA, the Agency, or we) is announcing the availability of a final guidance for industry #252 entitled “Antimicrobial Animal Drug Sales and Distribution Reporting Small Entity Compliance Guide.” The small entity compliance guide (SECG) is intended to help small entities comply with the final rule we issued in the Federal Register of May 11, 2016, entitled “Antimicrobial Animal Drug Sales and Distribution Reporting.”

DATES: The announcement of the guidance is published in the Federal Register on June 29, 2018.

ADDRESSES: You may submit either electronic or written comments on Agency guidances at any time as follows:

Electronic Submissions

Submit electronic comments in the following way:

• Federal eRulemaking Portal: https://www.regulations.gov. Follow the instructions for submitting comments. Comments submitted electronically, including attachments, to https://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, that information will be posted on https://www.regulations.gov.

• If you want to submit a comment with confidential information that you do not wish to be made available to the public, submit the comment as a written/paper submission and in the manner detailed (see “Written/Paper Submissions” and “Instructions”).

Written/Paper Submissions

Submit written/paper submissions as follows:

• Mail/Hand Delivery/Courier (for written/paper submissions): Dockets Management Staff (HFA–305), Food and Drug Administration, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852.

• For written/paper comments submitted to the Dockets Management Staff, FDA will post your comment, as well as any attachments, except for information submitted, marked and identified, as confidential, if submitted as detailed in “Instructions.”

Instructions: All submissions received must include the Docket No. FDA–2012–N–0447 for “Antimicrobial Animal Drug Sales and Distribution Reporting; Small Entity Compliance Guide.” Received comments will be placed in the docket and, except for those submitted as “Confidential Submissions,” publicly viewable at https://www.regulations.gov or at the Dockets Management Staff between 9 a.m. and 4 p.m., Monday through Friday.

• Confidential Submissions—To submit a comment with confidential information that you do not wish to be made publicly available, submit your comments 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.” We will review this copy, including the claimed confidential information, in our consideration of comments. The second copy, which will have the claimed confidential information redacted/blacked out, will be available for public viewing and posted on https://www.regulations.gov. Submit both copies to the Dockets Management Staff. If you do not wish your name and