Kentucky Beef Quality Assurance Program

Program and Manual Development

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Information contained in this manual is adapted from the following publications:
Mid-Atlantic Beef Quality Assurance Program Certification Manual, 2006
Virginia Beef Quality Assurance Chuteside Record
and
Alabama Beef Quality Assurance Training Manual
Nebraska Beef Quality Assurance Training Manual
Ohio Beef Quality Assurance Manual
Milk and Dairy Beef Quality Assurance Program, Milk and Dairy Beef Residue Prevention Protocol
National Beef Quality Assurance Program Guidelines

Care and Husbandry Practices
- Follow animal care and well-being guidelines that conform to good veterinary and husbandry practices to avoid bruising, stress, or injury.
- Regularly evaluate and implement biosecurity practices.

Feedstuffs
- A quality feed control program will be maintained for all incoming feed ingredients.
- Ruminant derived protein sources will not be fed.

Feed Additives & Medications
- Only FDA-approved medicated feed additives will be used in rations.
- Proper withdrawal time for all additives and pesticide/herbicide use will be observed to avoid violative residues.

Processing/Treatment & Records
- Extra-label drug use will only be used when prescribed by a veterinarian with a valid veterinarian-client-patient relationship.
- Records will be maintained for all treatments (individual or group) following BQA suggested record keeping guidelines and will be kept for a minimum of two years.
- All processing and treatment records will be transferred with the cattle to the next production level.

Injectable Animal Health Products
- All injections will be administered in the neck region only. This includes both subcutaneous and intramuscular injections.
- All individual treatments will strictly follow only FDA/USDA/EPA guidelines. Administering products in a method that will cause tissue damage will be avoided.

BQA Code of Conduct
- I received training in BQA and use it on my cattle enterprise because I have a commitment to consumers to produce the safest, highest quality beef in the world.
- I use BQA production practices because maintaining an optimum environment for cattle to produce at their best promotes efficiency and quality at the same time. BQA training has shown me that keeping records of all my production practices is the best way for me to reduce liability, provide quality assurance to my customers, and continue to ensure a safe beef supply through strict adherence to residue avoidance practices.
- BQA has taught me to think about all of my production practices in light of their effect on the quality of the final product.
- BQA is a combination of technology, common sense, a concern for animal well-being, and a consumer oriented production system.
The Importance of Beef Quality Assurance

Beef Quality Assurance (BQA) is a program developed to ensure that beef and dairy cattle are managed in a manner that will result in safe and wholesome beef and milk products for the consumer. This statement is not only the definition but the goal of BQA. Specifically, BQA is designed to enhance carcass quality by preventing drug residues, injection-site blemishes, and bruises. The Kentucky Beef Quality Assurance Program is based on recommended national guidelines and scientific research. This program enables beef and dairy producers to enhance their product, maximize marketability, and strengthen consumer confidence.

Members of each industry sector should assume responsibility for the role they play in delivering a quality beef product to their respective markets. By working together toward continued improvement of our product and our responsiveness to consumers, we all benefit.

Sample records: See Chapter 11.
Is BQA Necessary?

From gate to plate, BQA is a positive step for producers and consumers.

Concern over food wholesomeness and safety is an important consumer issue. It is of utmost importance that the public knows beef is a safe product. A BQA program will help secure consumer confidence for expanding domestic and export markets. BQA is a good business practice, which can identify potential problem areas to avoid product defects.

All sectors of the industry—from seedstock, cow-calf, heifer growers, and dairy producers to stocker operators, backgrounders, cattle feeders, and points of sale and harvest—must take responsibility for the production of a safe food product through proper animal care, handling, and management practices.

The level of consumer confidence in beef significantly affects consumer eating habits and impacts the future of our industry. Consumer confidence is essential if we are to “steak” our claim in the meat case.

Beyond safety, the economic importance of BQA can be seen when analyzing the top quality challenges in the production of beef. The 2000 National Beef Quality Audit showed that the industry lost an average of $100 for every fed steer or heifer marketed. Quality challenges include:

- Inconsistent size of meat cuts
- Non-uniform cattle
- Injection-site blemishes
- Branding
- Excessive external fat
- Excessive seam fat
- Inadequate muscling
- Dark cutters

All meat industries face similar concerns. By following BQA guidelines and management practices, beef and dairy producers increase the value of their product in the eyes of the consumers, while enhancing their stewardship of natural and financial resources.
Chapter 1: Importance of Beef Quality Assurance

The History of Beef Quality Assurance

Consumers have always expected safe and wholesome food. In 1980, because of beef safety concerns, beef producers began investigating ways to ensure that their production practices would pass the scrutiny of the consumer.

The Beef Quality Assurance program is not a new idea. In 1982, the United States Department of Agriculture Food Safety Inspection Service (USDA-FSIS) began working with the beef industry in the United States to develop the Pre-Harvest Beef Safety Production Program. The beef industry refers to this as Beef Quality Assurance, or BQA.

Because the majority of beef is raised by small independent producers in a vast variety of environmental climates, the BQA program has been modified and adapted to meet the needs of a range of production and marketing circumstances. Presently, a BQA educational program is active in 47 states.

The Kentucky Beef Quality Assurance Program began in 2000. It is designed to bring best management practices (BMP) to the farm that, along with Hazard Analysis and Critical Control Points (HACCP) principles applied at harvest and processing facilities, will ensure safe, wholesome, uniform, and quality beef products for consumers. It is a cooperative effort among beef and dairy producers, veterinarians, Cooperative Extension agents, and other professionals representing Kentucky Veterinary Medical Association (KVMA), University of Kentucky Cooperative Extension Service, Kentucky Beef Council, Kentucky Cattlemen’s Association, Kentucky Dairy Development Council, and the Kentucky Department of Agriculture.

Implementing BQA practices provides cattle producers with an important key for avoiding additional government regulation. Producer-driven programs have proven very successful and will continue to allow the industry the flexibility needed to produce safe, wholesome food in an economical manner.
Chapter 1: Importance of Beef Quality Assurance

Meeting the Industry Quality Challenges

Four national Beef Quality Audits (NBQA) have been conducted between 1991 and 2005. In three of the audits, defects in the hide (from branding and lice) and lack of uniform size of rib eye and other meat cuts were identified. In the 2005 NBQA, inadequate tenderness, excessive external fat, insufficient marbling, and excess carcass/cut weights were identified as the major factors affecting meat quality. For the first time, the 2005 NBQA identified lack of traceability of cattle from feedlots, need for instrument grading, need for clearer market signals, and need for communication among sectors as areas that the industry must address.

Good production practices can reduce, if not eliminate, the occurrence of quality problems. This manual outlines Best Management Practices (BMP) in key areas to help producers meet the industry’s beef quality challenges. These include implementing genetic and production management systems that have been shown to reduce beef quality defects, improve beef eating quality characteristics (such as flavor, tenderness, and juiciness), and ensure food safety.

Potential Value Loss

Today’s estimated potential loss in value due to quality defects continues to exceed $100 for every fed steer and heifer marketed in the United States. The value lost due to management defects can begin to be recovered simply by evaluating and altering the management techniques used in today’s beef and dairy production systems. Current problems that producers have control over include injection-site blemishes, hide damage, bruises, and dark cutters.
Chapter 1: Importance of Beef Quality Assurance

Capturing Added Value

As the food industry develops new products and packaging processes, correct injection sites and techniques become even more critical to realizing added value. New beef products have been introduced that add value to traditionally under-utilized chuck and round primals. The popular flat iron steak, cut from the chuck, is one example. It lies 3 to 4 inches in front of the shoulder blade, therefore, producers should give intramuscular shots further forward in the shoulder blade to keep from reducing the value of the flat iron. Furthermore, the use of modified atmosphere (MA) packaging processes for case-ready beef can discolor the meat near an injection site—even if the muscle contains no blemishes from the injection.

Animal health companies continue to research and develop products with BQA-friendly routes of administration. Administering animal health products according to label directions, marketing cattle at the optimum end point, reducing stress in cattle handling, and eliminating extremes in size of breeding stock are some of the ways by which quality defects are reduced and the market value of the beef cuts is increased.

Improved awareness and implementation of BQA practices from 1991 to 2005 have reduced the incidence of injection-site blemishes.
KY-BQA Certification

It is important to understand that people become BQA certified; operations or production units do not. The people who practice the guidelines and the people who implement the requirements and recommendations impact the end product and value of the animal. All farm personnel who handle cattle should be informed about proper processing techniques and provided with training to understand cattle behavior and recommended BMPs. Therefore, certification is done on an individual basis, not by facility or production unit.

Any person who handles cattle can become eligible for certification by:
- Attending the BQA certification training program
- Passing the post-test
- Completing the training checklist and certification form

Once the appropriate forms and check are received at the Kentucky Cattlemen’s Association, they will be processed and the certification card will be provided to the producer or farm personnel.

Certifications must be renewed every three years.
Based on the 2000 National Beef Quality Audit, injection-site blemishes (lesions) cost the beef industry $188 million annually. This means producers lost an average $7.05 per head per year in the value of the steers and heifers marketed. Research sponsored by NCBA on behalf of the Beef Checkoff uncovered a negative relationship between meat tenderness and injection sites, including those injection sites that had no visible lesions. Findings concluded that all intramuscular (IM) injections, including sterile water, create permanent damage—regardless of the age of the animal at the time the product was given. At the very least, tenderness is reduced in a 3-inch area surrounding the injection site.

Lesion: an injection site blemish.

Contrary to popular belief, not all beef from market cows is sold as ground beef. For example, rib eye rolls and rounds from market cows and bulls are used as whole muscle cuts in popular consumer products such as Philly Steak and roast beef sandwiches, as well as marinated and tenderized steak products. Thus, BQA practices are just as important throughout the life of cows and bulls.

The lesions in this roast beef were not discovered until the fully cooked roast was sliced by a foodservice employee.
Chapter 2: Vaccine and Drug Practices

**Injection Sites and Techniques**

To lessen injection-site defects in economically important cuts of beef, the preferred site for all subcutaneous (SQ) or intramuscular (IM) injections is the neck region (See Figures 2-1 and 2-2.) It is particularly important to use the neck region with IM products, because even the shoulder chuck primal contains “value-added” cuts that should be protected.

Whenever possible, choose products formulated and labeled for SQ rather than IM injection. See Table 2-1 for proper needle sizes.

**Subcutaneous Injections**

SQ injections are made just under the skin but not into the muscle tissue. The side of the neck is the best area to make injections. To administer, lift the skin with your free hand and insert the needle into the raised fold of skin. This is known as the “tent technique” (Figure 2-2).

Several animal health products are now approved to be injected into the ear of cattle. This location is excellent from a BQA perspective as ears are removed at harvest and do not enter the food chain. The ear must be clean to avoid infection, and producers should take care to avoid blood vessels. Read product labels carefully. An example of an ear injection technique can be found on the internet at <http://www.excede.com/>.

**Intramuscular Injections**

IM injections are made directly into muscle tissue of the neck. Absorption of the drug is more rapid in the muscle than under the skin because of the good blood supply to muscle tissue. After the injection site is chosen, distract the animal by slapping the injection site firmly. Immediately insert the needle with a quick thrust.
Table 2-1. Determining proper needle gauge based on the route of administration, animal size, and viscosity of fluid.

<table>
<thead>
<tr>
<th>Fluid Viscosity</th>
<th>SQ Injection (¼ to 1 inch long needle)</th>
<th>IM Injection (1 to ½ inch long needle)</th>
<th>IV Injection (½ inch long needle)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Animal Size (lb)</td>
<td>Animal Size (lb)</td>
<td>Animal Size (lb)</td>
</tr>
<tr>
<td>Thin</td>
<td>&lt;300</td>
<td>300-700</td>
<td>&gt;700</td>
</tr>
<tr>
<td>Thick</td>
<td>18-16</td>
<td>16</td>
<td>16</td>
</tr>
</tbody>
</table>

Select the needle to fit the cattle size (the smallest practical size without bending).

1 An example of a thin viscosity fluid: saline; thick: oxytetracycline.

Needle Use and Handling

General Guidelines

- Select a clean injection site.
- Single-use needles are preferred.
- Keep the contents of the vaccine bottle sterile; do not store a syringe and needle in the top of a bottle.
- Do not put a needle back into the vaccine bottle once it has been used for anything else.
- Keep transfer needles in a closed container when at chute-side and after use, boil and place in a clean container (see Figure 2-4 for instructions).

Selecting the Proper Needle Gauge

Primary considerations in needle selection include route of administration, size of the animal, and site of the injection. Secondary considerations include viscosity of the fluid and volume injected. The needle size used should never be larger than necessary to adequately perform the injection (Table 2-1).

Changing Needles

- Change needles every 10 to 15 head, or with every automatic dosing syringe refill.
- Change any needle that is bent, or becomes contaminated (manure, dirt, or chemicals), or if the needle point becomes burred.
- To prevent the spread of known blood-borne infectious diseases, use a new needle for each animal.

Note: A broken needle is an emergency; it will migrate farther into the tissues. Under no circumstances should animals with broken needles be sold or sent to a packer.

Viscosity: A measure of how thick and tenacious a fluid is. High viscosity fluids (like oxytetracycline) are thicker and more sluggish; low viscosity fluids (like saline) are thinner and flow more freely.
Chapter 2: Vaccine and Drug Practices

Cleaning Needles
- Use disposable needles and syringes.
- Heat-sterilize reusable equipment by boiling.
- Do not contaminate modified live virus products with disinfectants (such as rubbing alcohol).

Proper Disposal of Sharps for Producers
- Place in a thick plastic container with a secure lid. A sharps container is best, but a liquid detergent bottle can also work.
- Place sharps container in a rigid container lined with plastic.
- Dispose of as solid waste.

Injecting into a wet or muddy site increases the risk for spreading disease as well as increasing the number of injection-site lesions.

Improper sterilization can reduce the effectiveness of future injections and result in infection at the injection site.

To sterilize:
1. Bring a container of water to a rolling boil.
2. Place equipment in the boiling water.
3. Cover the container.
4. Bring the water back up to a boil.
5. Continue boiling for three to five minutes.

Source: Center for Disease Control and Prevention

Disinfectants (like rubbing alcohol) can decrease or even eliminate the effectiveness of modified live virus products.

To reconstitute a vaccine, place one end of the transfer needle into the sterile liquid and the other into the bottle containing the freeze-dried cake of vaccine. The vacuum should pull the liquid down.

Figure 2-4. Transfer needles: SMTN (Springer Magnath Transfer Needle), disposable plastic needle, short metal needle.
Chapter 2: Vaccine and Drug Practices

Drug Management

Open and consistent communication between a dairy/livestock producer and a veterinarian is needed to assure quality control, animal welfare, and prevention of drug and chemical residues. Using animal health products exactly as they are labeled or prescribed by a veterinarian with whom the producer has a valid veterinarian-client-patient relationship (VCPR) is a requirement of a BQA program. Information about establishing a valid VCPR is contained later in this chapter.

Storage

Animal health products usually have specific storage requirements. Some require refrigeration. All should be stored in a clean place where they cannot become dirty or contaminated. Observe and obey the manufacturer’s recommended storage instructions for each product.

Where refrigeration is needed, be sure the refrigerator is kept clean and located in a safe place that is not likely to be overheated or contaminated by dirt or manure. Do not keep refrigerated drugs in door shelves because of the temperature fluctuation.

Drugs for lactating dairy cows must be stored separately from those used for nonlactating cattle. This helps prevent lactating animals from receiving drugs intended for nonlactating animals, which could cause an illegal residue in milk and meat. This restriction applies to drugs stored both at room temperature and under refrigeration.

Handling Precautions

- Always read and follow label instructions and supply them in Spanish or other languages if needed.
- Post the local poison control center number by all phones.
- Use proper restraints when injecting cattle.
- With medication known to be toxic to humans, use the one-handed SQ tent technique (Figure 2-6). Use extreme caution if using automatic syringes for these medications.

Don’t mix too much vaccine at one time. Modified live vaccines (MLV) begin to degrade after about an hour in the heat and sunlight.

Use separate syringes for each product. Even a trace amount of killed product can harm the effectiveness of the modified live product.

Figure 2-6. One-handed “tent” technique for SQ injection.
Ensuring Drug Effectiveness

**Preparation**
- Use only fresh products.
- Keep in a cooler from purchase until refrigerated or administered.
- Purchase appropriate dosage sizes for the task.
- Use transfer needles to reconstitute vaccines.

**Mixing**
- Rock bottle(s) back and forth, but do not shake.
- Do not mix too much at one time.
- After mixing, gently rock bottle(s) periodically.
- Use only approved combinations.

**Administering**
- Label syringes before processing.
- Use separate syringes for each product.

**Storage**
- Do not store partially used containers.
- Clearly label all products before storage.

**Residue Avoidance**

Drug residue in livestock products must be avoided. Consumers are concerned about the drugs used in dairy and livestock production and how they affect the food they eat. The industry can address these concerns by assuring consumers that the necessary steps are taken to prevent drug residues. Consumers expect zero tolerance.

Residue violations and condemnations can be avoided by implementing and following control systems that incorporate the following practices:
- Maintain proper individual animal identification.
- Maintain complete medical records on animals for at least two years (see sample records in Chapter 11).
- Properly store, label, and account for all medication.
- Use animal health products according to the label.
- Maintain a valid VCPR.
- Educate all employees and family members about your control systems, and emphasize the importance of keeping drug residues out of the human food chain.
Drug Classifications

The Food and Drug Administration (FDA) has the responsibility for determining the market status of animal drugs, based in part upon whether it is possible to prepare "adequate directions for use" under which a layperson can use the drugs safely and effectively. The two basic classes of drugs available to livestock producers are discussed below:

Over-the-counter Drugs

Over-the-counter (OTC) drugs can be purchased from multiple sources and must be used as directed on the label (Figure 2-7). For example, most procaine penicillin G products are labeled for use at 1 cc/cwt and are given IM. So, a 600-pound calf would get 6 cc IM. Producers are not allowed to change the dose or give the drug by any other route, such as SQ.

Prescription Drugs

A drug that has significant potential for toxicity (or other harmful effects) in humans or animals that may have a unique method of use or which requires other special considerations for its use is usually labeled as a prescription (Rx) drug. Such products can be used or dispensed only by or on the order of a licensed veterinarian, and the label must contain the legend: "Caution: Federal law restricts this drug to use by or on the order of a licensed veterinarian" (Figure 2-8).

Note: Over-the-counter drugs must be used as directed on the label.
Extra-Label Use of Drugs

Extra-label use is defined as the “actual or intended use of a drug in a manner that is not in accordance with the label.” Under the provisions of the Animal Medicinal Drug Use Clarification Act of 1994, the FDA recognizes the professional judgment of veterinarians and allows the extra-label use of drugs (either OTC or Rx) by veterinarians under certain conditions. Extra-label use is limited to situations where a failure to treat an animal would:

- Threaten the health or life of an animal
- Cause undue suffering

Veterinarians may only consider using drugs in an extra-label manner under the following conditions:

1. There is no approved drug that is labeled for such use and that contains the same active ingredient in the required dosage form and concentration.
2. A currently approved and labeled drug is clinically ineffective for its intended use (e.g., drug resistant bacterial infections).

Precautions

Prior to using or dispensing a drug in an extra-label manner, the veterinarian should take the following precautions:

- Make a careful diagnosis and evaluation of the conditions for which the drug is to be used.
- Establish a substantially extended withdrawal period prior to marketing of milk, meat, or other edible products.
- Institute procedures to assure that the identity of the treated animal(s) is carefully maintained.
- Take appropriate measures to assure that the assigned withdrawal times are met and that no illegal drug residues occur in any food-producing animal subjected to extra-label treatment.
Labeling

Drugs intended for extra-label use must have additional labeling (Figure 2-9), including at least the following information:

- The name and address of the prescribing veterinarian (not just the clinic)
- The name of the active ingredient(s)
- Directions for use, including identity of the animal being treated, dosage, frequency and duration of treatment, and route of administration
- Any cautionary statements specified by the veterinarian
- The veterinarian’s specified withdrawal time for meat and/or milk

A common mistake: Improperly administering a drug—e.g., a large dose is given all in one site instead of smaller doses in multiple sites.

Penicillin accounts for more than 20 percent of all antibiotic residue violations in beef. It is the most commonly used drug and is routinely purchased over the counter. Gentamicin and streptomycin run a close second in the number of residue violations attributed to these antibiotics.
Limitations

The extra-label use of drugs is not permitted in or on animal feeds. A veterinarian cannot use or prescribe drugs for use in feed in any manner except for the approved use and at the approved dosage. Extra-label use of drugs in treating food-producing animals for improving rate of weight gain, feed efficiency, or other production purposes is also prohibited. Some specific drugs are completely prohibited for extra-label use in food-producing animals, including:

• Chloramphenicol
• Clenbuterol
• Diethylstilbestrol
• Dimetridazole
• Ipronidazole
• Other nitroimidazoles
• Furazolidone (Furacin topical powder)
• Nitrofurazone
• Fluoroquinolones (except for approved use for beef cattle)
• Glycopeptides

Drug Withdrawal Times

A withdrawal time should be indicated on the label of medications. This is the period of time that must pass between the last treatment and the time the animal will be harvested or milk can be sold. For example, if a medication with a 14-day meat withdrawal period was last given on August 1, the withdrawal would be completed on August 15, and that would be the earliest the animal could be harvested for human consumption. Often there are separate withdrawals for milk and meat, and meat withdrawals are always longer.

It is important that you follow withdrawal time directions on the label or as prescribed by your veterinarian. From the day you acquire your animals until the day they leave your care, you should maintain feed and treatment records. This is important for the day-to-day care of your animal and for whoever may later purchase your animal.

Observe label instructions and withdrawal times carefully. When using drugs by extra label, work closely with the veterinarian on dosages and withdrawal times. Never use an approved veterinary drug in an extra-label manner without consulting the veterinarian. Treating animals in an extra-label manner without direction by a licensed veterinarian is illegal.

Unacceptable levels of drug residues detected in edible tissues collected at harvest will result in traceback, quarantine, and potential fines or jail time. Substantial economic losses may result for the individual producer as well as negative publicity for the entire beef industry. Producers are responsible for residue problems and should follow these rules:

• Do not market animals for food until the withdrawal time listed on the label or prescribed by the veterinarian has elapsed.
• Use only medications approved for cattle, and use them exactly as the label directs or as prescribed by your veterinarian.
• If ever in doubt, rely on the VCPR you have established with your veterinarian. Consult your veterinarian with all questions and concerns.
• Keep records that show drug and dosage used, animals treated, and withdrawal time.

All federally approved drugs will include the required withdrawal time for that drug on the product label or package insert. These withdrawal times can range from 0 to as many as 60 days or more. The Compendium of Veterinary Products, published by the North American Compendiums Inc., gives a comprehensive list of drugs approved for use in beef and dairy cattle as well as a description of each drug. In addition, the Compendium includes a chart of the withdrawal times for meat and also includes time of milk withholding. The drug label itself always supersedes the Compendium if there is a discrepancy. It is your responsibility to be aware of the withdrawal times of any drugs that you use on your cattle. More information is available at these Websites: <http://www.fda.gov/> and <http://www.farad.org/>.
Managing Implants

Implants may provide an economic advantage in the production of safe and wholesome beef. Beef from implanted cattle has proven to be leaner than beef from non-implanted cattle, with minute differences in hormone levels (Figure 2-10). Nevertheless, consumer concern remains high with regard to implanted beef. Administer implants properly and follow label directions, including proper sanitation and the use of antiseptic on the needle between every use. Proper sanitation results in fewer abscesses in the ear and allows for higher utilization of the implant.

Regulations governing the use of implants are set by the FDA. Always read and follow the manufacturer’s directions before implanting any cattle. The growth promotant implants approved for use in the United States are extremely safe for both production and consumption. There is no required withdrawal time for slaughter with FDA approved implants.

The only approved location for implant administration is the middle third of the backside of the ear. All implants must be located SQ within this area (Figure 2-11). Implants should never be placed in locations other than the ear.

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**Figure 2-10.** Hormone levels in beef.

<table>
<thead>
<tr>
<th>Small Amounts Found in Beef</th>
<th>The difference in levels of estrogen found in beef from cattle raised with or without growth promotants is miniscule.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Growth promotants vs. no growth promotants (in nanograms of estrogen)*</td>
<td></td>
</tr>
<tr>
<td>3-ounce serving of beef from a steer treated with growth promotants</td>
<td>......................................................... 1.9</td>
</tr>
<tr>
<td>3-ounce serving of beef from a steer raised without growth promotants, such as certified organic beef</td>
<td>......................................................... 1.3</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>FDA-Approved Safe Levels</th>
<th>One serving of beef from a steer implanted with a growth promotant has nearly 20 times less estrogen than what the FDA permits, and thousands of times less than the amount our bodies naturally produce, not to mention a fraction of the phytoestrogen levels present in foods such as soybean oil, cabbage and grains.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hormones in the Human Body</td>
<td>The human body naturally produces hormones in quantities much greater than could ever be consumed by eating any food. In fact, the average man or woman daily produces 35,000 times more hormones than could be present in beef or other food.</td>
</tr>
<tr>
<td>Male vs. Female (in nanograms of estrogen)*</td>
<td></td>
</tr>
<tr>
<td>Male child</td>
<td>......................................................... 41,500</td>
</tr>
<tr>
<td>Female child</td>
<td>......................................................... 54,000</td>
</tr>
<tr>
<td>Male adult</td>
<td>......................................................... 136,000</td>
</tr>
<tr>
<td>Female adult</td>
<td>......................................................... 480,000</td>
</tr>
<tr>
<td>Female adult (pregnant)</td>
<td>......................................................... 3,415,000</td>
</tr>
</tbody>
</table>

* A nanogram is one billionth of a gram, which is analogous to one blade of grass in an entire football field.

Sources: Food and Drug Administration; Hoffman and Evers; Scanga et al.; FSIS-USDA; Dr. Harlan Ritchie, Michigan State University.
Routine inspection of implant sites should be done every time animals are handled through a chute. Document the results of the inspection for future reference in implant management decisions.

Although there is no withdrawal period for implants, there are quality considerations in the timing. Aggressive implant strategies that maximize the response to the implant in growth and feed efficiency can compromise carcass grade. A conservative approach may not pay, however, when the Choice and Select price spread is too narrow to offset the lost value in feed efficiency and gain, which implants provide. It is as much an economic decision as it is a quality decision.

The objective is to know your options, then plan and keep records to evaluate your decisions.

<table>
<thead>
<tr>
<th>Using Implants Correctly—Implanting Mistakes and Solutions</th>
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<tbody>
<tr>
<td><strong>Problem</strong></td>
</tr>
<tr>
<td>Abscess at implant site</td>
</tr>
<tr>
<td>Bunched pellets</td>
</tr>
<tr>
<td>Retrograde abscess</td>
</tr>
<tr>
<td>In cartilage</td>
</tr>
<tr>
<td>Crushed pellet</td>
</tr>
<tr>
<td>Missing implant</td>
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<tr>
<td>Separated pellet</td>
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<tr>
<td>Partial implant</td>
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<tr>
<td>Pellet too close to the head</td>
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<tr>
<td>Walled-off implant</td>
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</tbody>
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Producer’s Guide for Judicious Use of Antimicrobials (Antibiotics) in Cattle

1. **Prevent problems.** Emphasize appropriate husbandry and hygiene, routine health examinations, and vaccinations.

2. **Select and use antibiotics carefully.** Consult with your veterinarian on the selection and use of antibiotics. Have a valid reason to use an antibiotic. Therapeutic alternatives should be considered prior to using antimicrobial therapy.

3. **Avoid using antibiotics important in human medicine as first-line therapy.** Avoid using as the first antibiotic those medications that are important for treating strategic human or animal infections.

4. **Use the laboratory to help you select antibiotics.** Cultures and susceptibility test results should be used to aid in the selection of antimicrobials, whenever possible.

5. **Avoid using broad spectrum.** Use narrow spectrum antimicrobials whenever possible. Combination antibiotic therapy is discouraged.

6. **Avoid inappropriate antibiotic use.** Confine therapeutic antimicrobial use to proven clinical indications, avoiding inappropriate uses such as for viral infections without bacterial complication.

7. **Treatment programs should reflect best use principles.** Regimens for therapeutic antimicrobial use should be optimized using current pharmacological information and principles.

8. **Treat the fewest number of animals possible.** Limit antibiotic use to sick or at-risk animals.

9. **Treat for the recommended time period.** This practice will minimize the potential for bacteria to become resistant to antimicrobials.

10. **Avoid environmental contamination with antibiotics.** Steps should be taken to minimize antimicrobials reaching the environment through spillage, contaminated ground run-off, or aerosolization.

11. **Keep records of antibiotic use.** Accurate records of treatment and outcome should be used to evaluate therapeutic regimens. Always follow proper withdrawal times.

12. **Follow label directions.** Never use antibiotics other than as labeled without a valid veterinary prescription.

13. **Extra-label antibiotic use must follow FDA regulations:** Prescriptions, including extra-label use of medications must meet the Animal Medicinal Drug Use Clarification Act (AMDUCA) amendments to the Food, Drug, and Cosmetic Act and its regulations. These regulations require a valid VCPR.

14. **Subtherapeutic antibiotic use is discouraged.** Antibiotic use should be limited to prevent or control disease and should not be used if the principle intent is to improve performance.

Source: Guidelines 1 through 13 adapted by NCBA, from AVMA, AABP, and AVC Appropriate Veterinary Antibiotic Use Guidelines.