A. INTRODUCTION

This regulatory guide directs the reader to the type of information acceptable to the U.S. Nuclear Regulatory Commission (NRC) staff for preparing and reviewing an application for a medical use license. Title 10, Part 35, “Medical Use of Byproduct Material,” of the Code of Federal Regulations (10 CFR Part 35) (Ref. 1) regulates the medical use of byproduct material. In addition to the requirements of 10 CFR Part 35, medical use licensees may be subject to those portions of 10 CFR Part 20, “Standards for Protection Against Radiation” (Ref. 2), that relate to radiation safety and the sections of 10 CFR Part 30, “Rules of General Applicability to Domestic Licensing of Byproduct Material” (Ref. 3), that relate to licensing and the noncommercial transfer of specific radioactive drugs to medical use licensees within a consortium.

This regulatory guide endorses the methods and procedures for medical licensing applications contained in the current revision of NUREG-1556, Volume 9, “Consolidated Guidance about Materials Licenses: Program-Specific Guidance about Medical Use Licenses” (Ref. 4), as a process that the NRC staff finds acceptable for meeting the regulatory requirements.

Volume 9 of NUREG-1556 addresses the issues that an applicant must respond to when preparing a license application on NRC Form 313, “Application for Materials License,” and the NRC Form 313A series of forms. The NUREG also includes (for clarification purposes) descriptions of certain key elements of a medical use program that do not require a response on NRC Form 313.

This regulatory guide contains information collection requirements covered by 10 CFR Parts 20, 30, and 35 and NRC Form 313 that the Office of Management and Budget (OMB) approved under OMB
control numbers 3150-0014, 3150-0017, 3150-0010, and 3150-0120, respectively. The NRC may neither conduct nor sponsor, and a person is not required to respond to, an information collection request or requirement unless the requesting document displays a currently valid OMB control number.

**B. DISCUSSION**

As part of its redesign of the materials licensing program, the NRC consolidated and updated numerous guidance documents for materials licenses into the multivolume NUREG-1556. Various volumes in the NUREG-1556 series provide current, program-specific guidance on testing, licensing, decommissioning, and terminating materials licenses.

Volume 9 of NUREG-1556 provides program-specific guidance about medical use licenses. It identifies the information needed to complete NRC Form 313 and the NRC Form 313A series of forms. NUREG-1556 provides an overview of the types of licenses issued by the NRC, the commitments and responsibilities that a licensee must undertake, applicable regulations, the process for filing a license application, and the contents of applications for different medical uses of byproduct material. In particular, Volume 9 of NUREG-1556 gives an item-by-item description of the information the applicant should provide. Because of the wide variety in the types of medical use programs, NUREG-1556 contains indicators to alert applicants for particular types of medical use licenses to information that pertains to those types of uses. The NRC is placing added emphasis on conducting its regulatory activities in a risk-informed and performance-based manner. The agency intends for this approach to be less prescriptive and to allow licensees the flexibility to implement the agency’s regulations in a manner that is more specific to their needs yet still meets the regulatory requirements. By supplying examples, the NRC seeks to provide information to meet the needs of applicants for licensure without being prescriptive. Guidance in NUREG-1556 represents one means of complying with NRC regulations. It is not the only means of satisfying the regulatory requirements.

**C. REGULATORY POSITION**

This regulatory guide endorses the method(s) of preparing a medical use program license application or revision request described in the current revision of NUREG-1556, Volume 9, as a process that the NRC has found to be acceptable for meeting the regulatory requirements.

**D. IMPLEMENTATION**

The purpose of this section is to provide information to applicants and licensees regarding the NRC’s plans for using this regulatory guide. The NRC does not intend or approve any imposition or backfit in connection with its issuance.

In some cases, applicants or licensees may propose or use a previously established acceptable alternative method for complying with specified portions of the NRC’s regulations. Otherwise, the methods described in this guide will be used in evaluating compliance with the applicable regulations for license applications, license amendment applications, and amendment requests.
REFERENCES


¹ All NRC regulations listed herein are available electronically through the Electronic Reading Room on the NRC’s public Web site, at http://www.nrc.gov/reading-rm/doc-collections/cfr/. Copies are also available for inspection or copying for a fee from the NRC’s Public Document Room (PDR) at 11555 Rockville Pike, Rockville, MD; the mailing address is USNRC PDR, Washington, DC 20555; telephone (301) 415-4737 or (800) 397-4209; fax (301) 415-3548; and email PDR@nrc.gov.

² All NUREG-series reports listed herein are published by the U.S. Nuclear Regulatory Commission. These volumes are available electronically through the Electronic Reading Room on the NRC’s public Web site, at http://www.nrc.gov/reading-rm/doc-collections/nuregs/. Copies are also available for inspection or copying for a fee from the NRC’s Public Document Room (PDR) at 11555 Rockville Pike, Rockville, MD; the mailing address is USNRC PDR, Washington, DC 20555; telephone (301) 415-4737 or (800) 397-4209; fax (301) 415-3548; and email PDR@nrc.gov. In addition, copies are available at current rates from the U.S. Government Printing Office, P.O. Box 37082, Washington, DC 20402-9328, telephone (202) 512-1800; or from the National Technical Information Service (NTIS), at 5285 Port Royal Road, Springfield, VA 22161, online at http://www.ntis.gov, by telephone at (800) 553-NTIS (6847) or (703) 605-6000, or by fax to (703) 605-6900.