—Evaluate whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information will have practical utility;
—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;
—Enhance the quality, utility, and clarity of the information to be collected; and
—Minimize the burden of the collection of information on those who are to respond, including through the use of appropriate automated, electronic, mechanical, or other technological collection techniques or other forms of information technology, e.g., permitting electronic submission of responses.

**Overview of this information**

1. **Type of Information Collection:** Renewal of a currently approved collection (1121–0329 and 1121–0188).
2. **Title of the Form/Collection:** OJP Solicitation Template
3. **Agency Form Number, if any, and the Applicable Component of the Department Sponsoring the Collection:** No form number available. Office of Justice Programs, Department of Justice.
4. **Affected Public Who Will be Asked to Respond:** The primary respondents are state agencies, tribal governments, local governments, colleges and universities, non-profit organizations, for-profit organizations, and faith-based organizations. The purpose of the solicitation template is to provide a framework to develop program-specific announcements soliciting applications for funding. A program solicitation outlines the specifics of the funding program; describes requirements for eligibility; instructs an applicant on the necessary components of an application under a specific program (e.g., project activities, project abstract, project timeline, proposed budget, etc.); outlines program evaluation and performance measures; explains selection criteria and the review process; and provides registration dates, deadlines, and instructions on how to apply within the designated application system. This collection is also incorporating the previously approved collection for the OJP Budget Detail Worksheet (1121–0188). The Budget Detail Worksheet is only required during the application process, and therefore should be included in this collection with the solicitation template, reducing the number of OMB PRA reviews and approvals needed. The primary respondents are the same, as listed above, and the worksheet provides auto calculated fields and instructions for the necessary budget information required for each application submission (e.g. personnel/benefits, travel, indirect cost rates, etc.). The form is not mandatory and is recommended as guidance to assist the applicant in preparing their budget as authorized in 28 CFR part 66 and 28 CFR part 70.
5. **An Estimate of the Total Number of Respondents and the Amount of Time Estimated for an Average Respondent to Respond:** It is estimated that information will be collected annually from approximately 10,000 applicants. Annual cost to the respondents is based on the number of hours involved in preparing and submitting a complete application package. Mandatory requirements for an application include a program narrative and budget details and narrative (formerly 1121–0188). Optional requirements can be imposed depending on the type of program to include, but not limited to: Project abstract, indirect cost rate agreement, tribal authorizing resolution, timelines, logic models, memos of understanding, letters of support, resumes, disclosure of pending applications, and research and evaluation independence and integrity. Public reporting burden for this collection of information is estimated at up to 32 hours per application. The 32-hour estimate is based on the amount of time to prepare a research and evaluation proposal, one of the most time intensive types of application solicited by OJP. The estimate of burden hours is based on OJP’s prior experience with the research application submission process.
6. **An Estimate of the Total Public Burden (in hours) Associated with the Collection:** The estimated public burden associated with this application is 320,000 hours.

If additional information is required contact: Jerri Murray, Department Clearance Officer, United States Department of Justice, Justice Management Division, Policy and Planning Staff, Two Constitution Square, 145 N Street NE., Room 3W–1407–B, Washington, DC 20530.


**Jerri Murray.**

Department Clearance Officer for PRA, U.S. Department of Justice.

**DEPARTMENT OF JUSTICE**

**Drug Enforcement Administration**

[Docket No. 12–57]

**Sanjay Trivedi, M.D.: Decision and Order**

On September 25, 2012, Administrative Law Judge (ALJ) Gail A. Randall issued the attached recommended decision. Neither party filed exceptions to the decision. Having reviewed the entire record, I have decided to adopt the ALJ’s rulings, findings of fact, conclusions of law, and recommended Order.

**Order**

Pursuant to the authority vested in me by 21 U.S.C. 823(f) and 824(a), as well as 28 CFR 0.100(b), I order that DEA Certificate of Registration FT0896754, issued to Sanjay Trivedi, M.D., be, and it hereby is, revoked. I further order that any pending application of Sanjay Trivedi, M.D., to renew or modify his registration, be and it hereby is, denied. This Order is effective immediately.¹


Michele M. Leonhart, Administrator.


**Recommended Rulings, Findings of Fact, Conclusions of Law, and Decision of the Administrative Law Judge**

**I. Facts**

Gail A. Randall, Administrative Law Judge. The Administrator, Drug Enforcement Administration (“DEA” or “Government”), issued an Order to Show Cause and Immediate Suspension of Registration (“Order”) dated June 25, 2012, proposing to revoke the DEA Certificate of Registration, No. FT0896754, of Sanjay Trivedi, M.D. (“Respondent”), as a practitioner, pursuant to 21 U.S.C. 824(a)(4) (2006), and deny any pending applications for renewal or modification of such registration pursuant to 21 U.S.C. 823(f) (2006), because the continued registration of the Respondent would be inconsistent with the public interest, as that term is used in 21 U.S.C. 823(f). The Respondent’s registration will expire by its own terms on November 30, 2013.

Specifically, the Order alleged that the Respondent dispensed at least

¹ For the same reason I ordered that Respondent’s registration be immediately suspended, I conclude that the public interest necessitates that this Order be effective immediately. See 21 CFR 1319.07.
On September 19, 2012, the Government filed Government’s Reply to Respondent’s Response to Motion for Summary Disposition and Motion to Stay Proceedings and Request for Extension of Time for Further Response (“Government’s Reply”). Therein, the Government argues that the only due process that need be afforded to the Respondent is an “opportunity to oppose a motion for summary disposition by showing that his state authority has not been suspended or revoked.” [Government’s Reply at 1]. The Government further argues that because there has not been a showing that Respondent’s state license is valid, the Respondent currently lacks state authority to handle controlled substances and thus, the Respondent cannot remain registered by the DEA. [Id. at 2].

For the reasons set forth below, I will grant the Government’s Motion and recommend that the Administrator revoke the Respondent’s DEA Certificate of Registration: But, I note that, pursuant to 21 CFR 1301.13(a) (2012), the Respondent may apply for a new DEA Certificate of Registration at any time.

II. Discussion

A. Respondent Currently Lacks Authority To Handle Controlled Substances In Florida

The DEA will not maintain a controlled substances registration if the registrant is without state authority to handle controlled substances in the state in which the registrant practices. The Controlled Substances Act (“CSA”) provides that obtaining a DEA registration is conditional on holding a state license to handle controlled substances. See 21 U.S.C. 802(21) (2006) (defining “practitioner” as "a physician * * * licensed, registered, or otherwise permitted, by * * * the jurisdiction in which he practices * * * to distribute, dispense, or administer * * * a controlled substance in the course of professional practice"); 21 U.S.C. 823(f) (2006) (“the Attorney General shall register practitioners * * * if the applicant is authorized to dispense * * * controlled substances under the laws of the State in which he practices”). The DEA, therefore, has consistently held that the CSA requires the DEA to revoke the registration of a practitioner who no longer possesses a state license to handle controlled substances. See 21 U.S.C. 824(a)(3) (2006) (stating “a registration may be suspended or revoked by the Attorney General upon a finding that the registrant has had his State license or
registration suspended, revoked or denied by competent State authority’’); Beverley P. Edwards, M.D., 75 FR 49,991 (DEA 2010); Joseph Baumstarck, M.D., 74 FR 17,525 (DEA 2009).

In this case, the Respondent does not dispute that he currently lacks state authority to handle controlled substances. However, the Respondent argues that his current state medical license suspension is temporary, as he and the Florida Department of Health are currently involved in settlement negotiations in which he anticipates that he will regain his Florida medical license. [Respondent’s Response at 1–3].

Respondent argues that his DEA registration should not be revoked because he will soon likely regain his state medical license in the state of Florida. [Id. at 2–3]. However, the Emergency Suspension from the Florida Department of Health effectively suspends the Respondent’s license to practice medicine in the state of Florida. Regardless of whether the Respondent and the Florida Department of Health eventually decide upon a settlement agreement in which the Respondent’s state license is reinstated, the Respondent currently lacks the necessary state authority to practice medicine and handle controlled substances in Florida. Consequently, his DEA registration must be revoked. See Joseph Baumstarck, M.D., 74 FR 17,525, 17,527 (DEA 2009) (stating that “a practitioner may not maintain his DEA registration if he lacks state authority to handle controlled substances under the laws of the state in which he practices”); Treasure Coast Specialty Pharmacy, 76 FR 66,965 (DEA 2011); Roy Chi Lung, M.D., 74 FR 20,346 (DEA 2009); Gabriel Sagun Orzame, M.D., 69 FR 58,959 (DEA 2004).

While the Respondent argues that his state license may be reinstated in the future, this possibility is immaterial in light of the Respondent’s current lack of state registration. Indeed, the CSA and Agency precedent make clear that as a prerequisite to registration the Respondent must have state authority to handle controlled substances, and that without such authority all other issues before this forum are moot. See 21 U.S.C. 802(21); 21 U.S.C. 823(f); Joseph Baumstarck, M.D., 74 FR at 17,527 (DEA 2009). Thus, because there is no dispute that the Respondent lacks state authority to handle controlled substances, the Respondent’s registration must be revoked.

B. Respondent Is Entitled To Reapply For Registration With the DEA

Any person who is required to register with the DEA may apply for registration at any time. 21 CFR 1301.13(a) (2012) (“Any person who is required and who is not registered may apply for registration at any time. No person required to be registered shall engage in any activity for which registration is required until the application for registration is granted and a Certificate of Registration is issued by the Administrator to such person”). The Respondent is permitted to reapply for a Certificate of Registration with the DEA at any time in the future. 21 CFR 1301.13(a). However, the Respondent will not be permitted to engage in activity for which a registration is required until his application is granted by the DEA. Id.

III. Conclusion, Order, and Recommendation

Consequently, there is no genuine dispute of material fact regarding the Respondent’s lack of state authority to handle controlled substances. Thus, summary disposition for the Government is appropriate. It is well settled that when there is no question of material fact involved, there is no need for a plenary, administrative hearing. See Michael G. Dolin, M.D., 65 FR 5,061 (DEA 2000). Here, there is no genuine dispute that the Respondent currently lacks state authority to practice medicine and to handle controlled substances in Florida.

Accordingly, I hereby grant the Government’s Motion for Summary Disposition. I also forward this case to the Administrator for final disposition. I recommend that the Respondent’s DEA Certificate of Registration, Number FT0896754, be revoked.2


Gail A. Randall,
Administrative Law Judge.

[FR Doc. 2013–02232 Filed 2–1–13; 8:45 am]

BILLING CODE 4410–09–P

DEPARTMENT OF LABOR
Office of the Secretary
Agency Information Collection Activities; Submission for OMB Review; Comment Request; Gamma Radiation Surveys

ACTION: Notice.

SUMMARY: The Department of Labor (DOL) is submitting the Mine Safety and

Health Administration (MSHA) sponsored information collection request (ICR) titled, “Gamma Radiation Surveys,” to the Office of Management and Budget (OMB) for review and approval for continued use in accordance with the Paperwork Reduction Act (PRA) of 1995 (44 U.S.C. 3501 et seq.).

DATES: Submit comments on or before March 6, 2013.

ADDRESSES: A copy of this ICR with applicable supporting documentation; including a description of the likely respondents, proposed frequency of response, and estimated total burden may be obtained from the RegInfo.gov Web site, http://www.reginfo.gov/public/do/PRAMain on the day following publication of this notice or by contacting Michel Smyth by telephone at 202–693–4129 (this is not a toll-free number) or sending an email to DOL_PRA_PUBLIC@dol.gov.

FOR FURTHER INFORMATION CONTACT: Michel Smyth by telephone at 202–693–4129 (this is not a toll-free number) or by email at DOL_PRA_PUBLIC@dol.gov.


SUPPLEMENTARY INFORMATION:

Regulations 30 CFR 57.5047 requires a covered mine operator to maintain a record of cumulative individual gamma radiation exposure to ensure that annual exposure does not exceed five (5) Rems. This requirement protects the health of workers in mines with radioactive ores.

This information collection is subject to the PRA. A Federal agency generally cannot conduct or sponsor a collection of information, and the public is generally not required to respond to an information collection, unless it is approved by the OMB under the PRA and displays a currently valid OMB Control Number. In addition, notwithstanding any other provisions of law, no person shall generally be subject to penalty for failing to comply with a collection of information that does not display a valid Control Number. See 5 CFR 1320.5(a) and 1320.6. The DOL obtains OMB approval for this information collection under Control Number 1219–0039. The current approval is scheduled to expire on February 28, 2013; however, it should be noted that existing information