

GENERAL SERVICES ADMINISTRATION

Public Buildings Service

Virginia Avenue Border Crossing/San Ysidro Port of Entry, San Diego, California; Notice of Intent; Environmental Impact Statement

AGENCY: Public Buildings Service, GSA.
ACTION: Pursuant to the Council on Environmental Quality Regulations (40 CFR 1500-1508) implementing procedural provisions of the National Environmental Policy Act (NEPA), the United States General Services Administration (GSA) hereby gives notice that said agency intends to prepare an EIS on the Virginia Avenue Border Crossing/San Ysidro Port of Entry in San Diego, California. The proposed project would include construction of a small facility, four to six inspection lanes and inspection booths. The site compliments the Government of Mexico's planned new facility located at El Chaparral adjacent to Virginia Avenue to the south.

Alternatives: In addition to the proposed action, the EIS will examine two alternatives; realignment of Inter-State Highway 5 and no action or continued use of the existing San Ysidro Port Entry. Also, reasonable alternatives that may or may not be within the authority of GSA will be examined. If there are potentially a large number of alternatives, only a reasonable number of examples covering the full spectrum of alternatives shall be analyzed.

Public Involvement: There will be several public meetings including, Scoping, Critical Issue(s), Draft Review and Final EIS. There will also be public review and comment periods of the Draft EIS. Further information may be obtained from: Ms. Sheryll White, U.S. General Services Administration, Portfolio Management Division (9PT), 450 Golden Gate Avenue, 3rd Floor East, San Francisco, CA 94102-2799, Telephone: (415) 522-3488.

Dated: September 23, 1999.

Aki K. Nakao,

Deputy Regional Administrator, (9AD).

Notice of Intent To Prepare an EIS

The General Services Administration intends to prepare an Environmental Impact Statement (EIS) on the following project: Virginia Avenue Border Crossing/San Ysidro Port of Entry San Diego, California

The General Services Administration of the United States Government is proposing to expand the United States Border Crossing at Virginia Avenue in

San Diego, California in order to provide southbound vehicular inspection and to convert the existing southbound lanes at the United States San Ysidro Port of Entry at San Diego, California to northbound.

Alternatives to the proposed action include:

A. Proposed Action: Construction of a small facility, four to six inspection lanes (initially) and inspection booths. The site complements the Government of Mexico's planned new facility at El Chaparral adjacent to Virginia Avenue to the south.

B. Realignment of Inter-State Highway 5 to increase northbound inspection lanes at the San Ysidro Port of Entry. This action could affect an historical residential area in Tijuana as well as traffic access to newly aligned lanes. The site is located to the east of the Government Mexico's planned new facility El Chaparral.

C. No action-space for functions now located at the San Ysidro Port of Entry will continue.

D. Reasonable alternatives which may or may not be within the authority of GSA. If there are potentially a large number of alternatives, only a reasonable number of examples covering the full spectrum of alternatives shall be analyzed.

Public scoping will include:

Scoping Meeting

Critical Issue(s) Meeting(s)

Public Review and Comment to Draft EIS

Draft EIS Review Meeting

Final EIS Meeting

FOR FURTHER INFORMATION CONTACT:

Sheryll White, General Services Administration, Portfolio Management Division (9PT), 450 Golden Gate Avenue, 3rd Floor East, San Francisco, California 94102, (415) 522-3488, Fax: (415) 522-3215.

Email:sheryll.white@gsa.gov.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. 99N-4166]

Agency Information Collection Activities: Proposed Collection; Comment Request; Electronic Records; Electronic Signature

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing an opportunity for public comment on the proposed collection of certain information by the agency. Under the Paperwork Reduction Act of 1995 (the PRA), Federal agencies are required to publish notice in the **Federal Register** concerning each proposed collection of information, including each proposed extension of an existing collection of information and to allow 60 days for public comment in response to the notice. This notice solicits comments on information collection provisions relating to FDA's electronic records and electronic signatures.

DATES: Submit written comments on the collection of information by November 30, 1999.

ADDRESSES: Submit written comments on the collection of information to the Dockets Management Branch (HFA-305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852. All comments should be identified with the docket number found in brackets in the heading of this document.

FOR FURTHER INFORMATION CONTACT:

Karen L. Nelson, Office of Information Resources Management (HFA-250), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301-827-1482.

SUPPLEMENTARY INFORMATION:

Under the PRA (44 U.S.C. 3501-3520) Federal agencies must obtain approval from the Office of Management and Budget (OMB) for each collection of information they conduct or sponsor. "Collection of information" is defined in 44 U.S.C. 3502(3) and 5 CFR 1320.3(c) and includes agency requests or requirements that members of the public submit reports, keep records, or provide information to a third party. Section 3506(c)(2)(A) of the PRA (44 U.S.C. 3506(c)(2)(A)) requires Federal agencies to provide a 60-day notice in the **Federal Register** concerning each proposed collection of information, including each proposed extension of an existing collection of information, before submitting the collection to OMB for approval. To comply with this requirement, FDA is publishing notice of the proposed collection of information set forth in this document.

With respect to the following collection of information, FDA invites comments on: (1) Whether the proposed collection of information is necessary for the proper performance of FDA's functions, including whether the information will have practical utility; (2) the accuracy of FDA's estimate of the burden of the proposed collection of