Project Goals

MMP aims to provide a deeper understanding of the health-related experiences and needs of people living with HIV/AIDS who are receiving care in the U.S. The goals of the project are to:

1) Provide a wide array of locally and nationally representative estimates of behaviors and clinical outcomes of HIV-infected persons in care;
2) Describe health-related behaviors;
3) Determine accessibility and use of prevention and support services;
4) Increase knowledge of the care and treatment provided; and
5) Examine variations of factors by geographic area and patient characteristics

Significance

People living with HIV/AIDS, HIV prevention community planning groups, Ryan White CARE Act planning councils and consortia, providers of HIV care, and other policy makers and service planners may use MMP data for planning activities. MMP provides valuable state and national estimates of health care utilization, quality of care, severity of need, and effectiveness of prevention messages. MMP data may help estimate resource needs for treatment and services for HIV-infected persons. To be effective, programs must meet the current needs of the population. MMP data provide contextual information on prevention, care-seeking, treatment, and risk behaviors that can aid in the design and improvement of HIV programs.

What can I do to help make MMP a success?

Say, “yes” if asked to participate in MMP. Patients, providers, and the community play an important role in making MMP a success. Patients and providers who participate represent other patients and providers who were not selected to participate. The success of MMP depends on you! If you are not selected, you can still help by encouraging others to participate.

Everyone’s participation is essential.

For more information, go to:
http://www.cdc.gov/hiv/topics/treatment/MMP/index.htm

Sponsored by:
Centers for Disease Control and Prevention
Health Resources and Services Administration
National Institutes of Health
Community Advisory Board

A national Community Advisory Board (CAB), composed of a local representative from each of the 23 project areas, serves as a link between MMP staff and patients who participate in the project. The CAB includes a diverse group of individuals who are concerned about the wellbeing of persons living with HIV/AIDS and the quality of care they receive. As community advocates, CAB members convey the patient’s perspective to local and national MMP staff.

CAB members work closely with MMP staff, contributing to project activities. In addition to collaborating with MMP staff, CAB members also communicate with potential participants.

What is MMP?

The Medical Monitoring Project (MMP) is a surveillance project designed to produce nationally representative data on people living with HIV/AIDS who are receiving care in the United States. The U.S. Department of Health and Human Services’ Centers for Disease Control and Prevention (CDC), in collaboration with the National Institutes of Health (NIH), the Health Resources and Services Administration (HRSA), and state and local health departments, are conducting MMP across the nation.

Provider Advisory Board

MMP also has a national Provider Advisory Board (PAB) consisting of an HIV care provider from each project area. The PAB represents large, medium, and small HIV care facilities in both rural and urban locales. PAB members foster collaboration between local and national MMP staff and HIV care providers. Offering the provider’s perspective, PAB members also advise local and national MMP staff on project activities. They provide significant input on facility and patient recruitment strategies and increase project awareness among the medical community. In addition, PAB members inform local providers about MMP and encourage providers to participate.

Project Locations

Twenty-three (23) project areas are involved in MMP: California; Chicago, IL; Delaware; Florida; Georgia; Houston, TX; Illinois; Indiana; Los Angeles, CA; Michigan; Mississippi; New Jersey; New York; New York City, NY; North Carolina; Oregon; Pennsylvania; Philadelphia, PA; Puerto Rico; San Francisco, CA; Texas; Virginia; and Washington.

Project Participation

Maximum participation by providers and patients will increase the likelihood of obtaining information that is truly representative of patients in care for HIV. The success of MMP depends on the providers and patients selected to participate. Providers and patients who are not selected to participate in the project can still help by encouraging others to participate.

Project Procedures

To implement the project, state and local health departments identify all HIV care providers in their jurisdictions. A representative sample of these providers is then chosen. The health departments contact all sampled providers; HIV-positive patients are selected from the sampled providers that agree to participate.

MMP has two components: an interview and medical record abstraction. MMP staff invite each selected patient to participate in a face-to-face interview. The interview takes approximately 45 minutes and includes questions concerning their medical history, use of medical and social services, and risk behaviors. Trained MMP abstractors then collect additional information from the patient’s medical chart, which complements data from the interview.

MMP takes measures to ensure that the project is not burdensome to providers or participating patients. State and local health department representatives conduct all data collection activities so that they do not disrupt providers, their staff, or services their patients receive. Participants are compensated for their time. All personal and health care information collected during the project is kept secure and confidential.