Measles virus causes an acute infection characterised by rash and fever. Measles infection is preventable by vaccination, but remains a significant cause of childhood mortality in the developing world with an estimated number of approximately 242,000 deaths by measles in 2006 [1].

The World Health Organization (WHO) is coordinating a global control programme for measles, and all WHO regions have identified targets for control or elimination of transmission of this disease. In the WHO European Region (WHO/Europe) the goal is to eliminate measles by 2010 [2]. Surveillance to document the elimination of transmission is in place, using case-based laboratory confirmation by detection of measles-specific IgM and by sequence characterisation of measles strains from outbreaks and chains of transmission. Measles virus is isolated and/or sequenced by the participating laboratories in the ‘Global Measles and Rubella Laboratory Network’ [3].

Following the success of HepSEQ [4], a public health database generated by the Centre for Infections of the United Kingdom Health Protection Agency, and a European Union-funded measles network called ‘Enhanced Laboratory Surveillance of Measles’ (ELSM) [5], we developed a web-based, quality-controlled database with epidemiological and nucleotide data for measles infection in the WHO/Europe region (MeaNS). The major objectives of the MeaNS initiative are to function as an epidemiological surveillance tool and to monitor progress of the measles control programme.

Sequence data from the 450 nucleotide region encoding the C-terminal region of the measles virus nucleoprotein (N) and, optionally, the complete nucleotide sequence of the haemagglutinin (H) gene are deposited into MeaNS, together with epidemiological data. The data is quality checked and curated, first automatically by the database application and then manually by a curator. During the curation, specific identifiers called ‘WHO names’ [8] are created for each sample unless the names were provided by the depositors. In addition, the deposited sequences are assigned a genotype and a cluster identifying number by matching, respectively, against WHO reference sequences deposited into MeaNS, together with epidemiological and nucleotide data. The existing tools and the tools that are now being developed will be useful in the surveillance, sequence analysis, evolution, and genome annotation of measles.

Currently, MeaNS is the only known publicly available global database on measles nucleotide sequences. After registering the purpose of their interest, researchers can access MeaNS at www.hpa-bioinformatics.org.uk/Measles/Public/Web_Front/main.php.

References