Mumps in Children

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Abstract

The Mumps Virus (MuV) is an enveloped, non-segmented, negative-sense RNA virus that belongs to the Paramyxoviridae family. Mumps is a contagious disease that causes severe inflammatory symptoms like parotitis and orchitis. The virus is highly neurotropic, with laboratory evidence of infection in the Central Nervous System (CNS) in around half of the cases. Although symptomatic CNS infection is rare, MuV was a prominent cause of aseptic meningitis and viral encephalitis in many developed nations previous to the introduction of widespread vaccination. Despite being one of the oldest recognised diseases with a global distribution, it has received relatively little research attention. Aseptic meningitis instances linked to specific vaccine strains, as well as a global resurgence of cases, especially in well vaccinated populations, have reignited interest in the virus, particularly its pathophysiology and the need for clinically relevant disease modelss.

Introduction

Prior to the introduction of widespread mumps vaccination programmes, 95% of people had serological signs of exposure, with peak acquisition occurring during childhood. Mumps vaccination became widely used in the United States in the late 1960s, and by the 1980s, only a few instances had been reported. By 2001, the disease had almost completely disappeared, with only 0.1 occurrences per 100,000 people documented, showing a 99.9% reduction in disease incidence relative to the pre-vaccination era. Other countries have had similar results with immunisation to control mumps. Large, occasional mumps outbreaks began to develop internationally within a few years after these historic lows, involving a high percentage of people who had previously been vaccinated. The cause why mumps vaccinations’ lower-than-expected efficiency is a source of significant discussion, with theories ranging from warning immunity to the introduction of virus strains capable of eluding the vaccine’s immunity. In addition to vaccination efficacy concerns, instances of meningitis connected to particular vaccine strains used outside the United States have raised safety concerns. As a result, some vaccine strains have been withdrawn, and mumps immunisation has been discontinued in some situations. Mumps vaccination, for example, has been phased out of Japan's national immunisation programme. With nearly a million cases recorded annually, Japan today has one of the highest rates of mumps among affluent countries. A study of MuV pathogenesis is essential given the myriad concerns facing mumps, including current outbreaks. MuV has only one natural host: humans. The disease causes severe swelling of the parotid glands, but it can also affect other tissues and organs, causing a variety of inflammatory reactions such as encephalitis, meningitis, orchitis, myocardiitis, pancreatitis, and nephritis. Mumps is usually self-limiting, with full recovery occurring within a few weeks after commencement; however, long-term complications such as paralysis, convulsions, cranial nerve palsies, hydrocephalus, and deafness can develop. The condition is infrequently fatal, and the paucity of autopsy tissue limits the pathophysiology and pathology of the disease. As a result, our present understanding of MuV pathogenesis is mostly based on animal research, which frequently employs non-natural infection methods. As a result, the virus’s pathophysiology in humans is yet unknown.

Mumps was first described by Hippocrates in his first Book of Epidemics in the fifth century BC, but a viral aetiology was not demonstrated until the 1930s, when Johnson and Goodpasture proved Koch's hypotheses by transferring the disease from experimentally infected rhesus macaques (Macaca mulatta) to children in his neighbourhood using a bacteria-free, virus-free method, filter-sterilized preparation of macerated monkey parotid tissue. The virus is an enclosed particle with a non-segmented negative strand RNA molecule of 15,384 nucleotides that belongs to the Paramyxoviridae family. Measles virus, canine distemper virus, parainfluenza virus, Newcastle disease virus, respiratory syncytial virus, and metapneumovirus are all important paramyxoviruses that infect humans and livestock.
Nucleo-(N), V/P/I (V/phospho-/I proteins), Matrix (M), Fusion (F), Small Hydrophobic (SH), Haemagglutinin-Neuraminidase (HN), and Large (L) proteins are all tandemly linked transcription units in the encapsidated genome. The Ribonucleoprotein (RNP) complex, which is made up of negative-strand viral RNA encapsulated by N protein, serves as a template for viral replication and transcription. By entering at a single promoter at the 3’ end of the genome, the RNA-dependent RNA polymerase, a complex of the L and P proteins, operates as a replicase to copy the negative sense (−) RNA to a positive sense (+) RNA and as a transcriptase to create mRNAs from the (−) RNP.