Is South Africa at risk for Zika virus disease?

Zika virus (ZIKV) was originally isolated from a sentinel rhesus monkey in the Zika forest in Uganda in 1947. Twenty-one years later the virus was isolated from naturally infected humans in Nigeria. ZIKV belongs to the genus *Flavivirus*, family Flaviviridae, comprising enveloped viruses with RNA genomes, mostly arthropod borne. For decades ZIKV remained relatively unknown, affecting mainly monkeys and occasionally causing a mild disease in humans residing across a narrow equatorial belt in Africa and Asia. The virus is transmitted to humans by daytime active *Aedes aegypti* mosquitoes, usually through mosquito bites, no specific antivirals are available, and treatment is supportive. Infection rarely leads to hospitalisation, and it is believed that up to 80% of infected individuals remain asymptomatic. While investigations to determine the underlying pathology of ZIKV infection are ongoing, pregnant women are discouraged from travelling to affected areas at any stage of pregnancy. Two cases of possible sexual transmission of ZIKV have been reported. The most recent is associated with the first case of ZIKV disease diagnosed in a traveller (returning from Venezuela) and diagnosis of ZIKV disease in his sexual partner who did not travel outside the USA (unofficial report at time of publication). It would be reasonable to consider precautions (i.e. abstinence, condom use) for pregnant women with sexual partners who have travelled to affected areas and could be affected or are confirmed with ZIKV infection. Two cases of probable perinatal transmission are also on record from French Polynesia (2013). The potential for ZIKV transmission through blood transfusion has been demonstrated from the outbreak in French Polynesia (2013).

The risk of ZIKV transmission in any given area partly depends on the availability of competent mosquito vectors. *A. aegypti* mosquitoes are the major transmitters of ZIKV, as for dengue viruses. While causing an increased number of outbreaks in Africa and Asia in recent years, none of the known dengue virus types is endemic in South Africa (SA). Considering that the risk of transmission of the two related viruses is likely to be the same, the prediction models based on the distribution of *A. aegypti* therefore do not place SA at high risk for transmission of ZIKV. It should be emphasised here that different populations of *A. aegypti* mosquitoes may have dissimilar competence in vectoring these viruses to humans. No human cases of infection with ZIKV related to local transmission have been diagnosed in SA to date. The first imported case of ZIKV disease in SA was confirmed in the week of 15 February 2016, in a traveller from Colombia with mild illness. Further imported cases in travellers visiting affected areas can be expected. Dengue cases have only been diagnosed in travellers returning home from countries with active circulation of the virus. *A. aegypti* mosquitoes are found in SA, particularly in the eastern coastal plain but also in the cities of the inland plateau. In urban areas these mosquitoes breed in small collections of water such as in discarded tyres and buckets, or the leaf axils of banana trees. There is a possibility that a traveller infected with ZIKV may return to SA. The transient presence of virus in the blood (viraemia), however, decreases the likelihood of human-to-mosquito transmission and the establishment of an autochthonous transmission cycle.

Patients presenting with the symptoms described above and with a history of recent (<2 weeks) travel to an affected area should consider ZIKV disease, but also related infections such as chikungunya or dengue, as a possible diagnosis. Infection with these viruses can be confirmed by specialised referral laboratory testing at the National Institute for Communicable Diseases (NICD). Malaria should also be considered. Laboratory testing for confirmation of ZIKV infection at the NICD requires the submission of serum or clotted blood along with a comprehensive clinical and travel history. Blood/serum collected up to day 5 after the onset of disease is most suitable for confirmation of acute infection by virus culture and detection of ZIKV virus nucleic acid by reverse transcription-polymerase chain reaction (RT-PCR) assay. Although RT-PCR is highly sensitive and specific, its application in laboratory confirmation of ZIKV infection is limited by the short duration of viraemia (2 - 4 days). Serological diagnosis of ZIKV infection is complicated by the high level of cross-reaction with other flaviviruses. For this reason laboratory confirmation of infection based on serological results is only feasible through the testing of paired serum samples taken at least 14 days apart, to demonstrate a four-fold rise in antibody titre. Parallel serological testing for dengue virus antibodies is also important and diagnostic interpretation hinges on the quantitative comparison of these results. Rapid diagnostic tests (point-of-care assays) are not currently available.

In response to the increasing spread of ZIKV and the potential high health threats it poses, the following measures need to be considered: (i) detection and monitoring of virus dissemination...
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