






# Whole-Genome Sequence of a Stenotrophomonas maltophilia Isolate from Tap Water in an Intensive Care Unit

By	<div>Zainulabid, UA (Zainulabid, Ummu Afeera) ; Siew, SW (Siew, Shing Wei) ; Musa, SM (Musa, Siti Munirah) ; Soffian, SN (Soffian, Sharmeen Nellisa) ; Periyasamy, P (Periyasamy, Petrick) ; Ahmad, HF (Ahmad, Hajar Fauzan)</div> <div><a href="#">View Web of Science ResearcherID and ORCID</a> (provided by Clarivate)</div>
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Abstract	<div>Here, we present a 4,508,936-bp complete genome sequence of Stenotrophomonas maltophilia strain HW002Y, which was isolated from the tap water in an intensive care unit at Sultan Ahmad Shah Medical Centre at the International Islamic University of Malaysia (Kuantan, Pahang, Malaysia). Sequencing was performed using a Nanopore Flongle flow cell.</div> <div>Here, we present a 4,508,936-bp complete genome sequence of Stenotrophomonas maltophilia strain HW002Y, which was isolated from the tap water in an intensive care unit at Sultan Ahmad Shah Medical Centre at the International Islamic University of Malaysia (Kuantan, Pahang, Malaysia). Sequencing was performed using a Nanopore Flongle flow cell.</div>
Addresses	<div> <sup>1</sup> Int Islamic Univ Malaysia, Dept Internal Med, Kulliyyah Med, Kuantan, Pahang, Malaysia</div> <div> <sup>2</sup> Univ Kebangsaan Malaysia, Fac Med, Med Dept, Cheras, Kuala Lumpur, Malaysia</div> <div> <sup>3</sup> Univ Malaysia Pahang, Fac Ind Sci &amp; Technol, Gambang, Pahang, Malaysia</div>
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