Register list for 5 new names including Methylophilus medardensis sp. nov.

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Table 1: Complete list of names proposed in the current register list.

Proposed Taxon	Etymology	Description	Parent Taxon	Туре	Registry URL
Genus Methylosemipumilus dw oxi pe	L. neut. n. methyl, ertaining to the methyl roup; L. pref. semi-, elf; L. masc. n. pumilus, warfish; N.L. masc. n. dethylosemipumilus, half- warfish methyl (group kidizing) organism, ertaining to an termediated genome	Consists of one species, Methylosemipumilus turicensis (GCF_000953015.1), that was initially proposed as 'Candidatus Methylopumilus turicensis' in Salcher et. al. 2015 and later changed to Methylosemipumilus turicensis in Salcher et al. 2019. Also known as PRD01a001B from 16S rRNA gene based studies. Methylosemipumilus has a relatively small genome size (<1.8 Mb) but not as small as Methylopumilus (1.3-1.4 Mb), which - together with improved genomic analyses - led to the reclassification to Methylosemipumilus. The closest cultivated relatives are Methylotenera sp. L2L1 (GCF_000744605.1) with an average amino acid identity of 67.45% and average nucleotide identity of 70.55% and Methylovorus sp. MP688 (GCF_000183115.1) with an AAI of 69.37% and an ANI of 69.42%, while the genus Methylopumilus is more distantly related (AAI of 61.5-62.2% and ANI of 66.4-66.9%). Methylosemipumilus are aerobic methylotrophs containing pathways for methanol oxidation (Xox), the RuMP cycle and the tetrahydromethanopterin (H4MPT) pathway for formaldehyde oxidation. Methylosemipumilus are found in the plankton of lakes in relatively low abundances. The so far only strain	Methylophilaceae	Methylosemipumilus turicensis ^{Ts}	seqco.de/i:49942

Proposed Taxon	Etymology	was isolated via dilution-to extinction cultivation by using autoclaved lake	Parent Taxon	Туре	Registry URL
		water, no growth is observed in rich medium or on agar plates (Salcher et al. 2015). Therefore, the strain was not submitted to a culture collection because these bacteria are hard to maintain, i.e., they are very slowly growing, reach low densities in liquid culture, and do not grow on agar plates.			
Species <i>Methylophilus</i> <i>medardensis</i>	[me.dar.den'sis] N.L. masc. adj. <i>medardensis</i> , pertaining to Lake Medard (Czechia), the isolation source of the type strain.	Type genome is <i>Methylophilus medardensis</i> MMS-M-51 (GCF_006363915.1), that was isolated by dilution-to-extinction cultivation from the water column of Lake Medard, Czechia. MMS-M-51 has a genome size of 2.6 Mb with a genomic GC content of 49.8%, contains 6 rRNA genes and 42 tRNAs. The genome is complete, consisting of 1 circular chromosome. The closest cultivated relatives are <i>Methylophilus</i> sp. Q8 (GCF_000800115.1) with an average amino acid identity of 85.53% and average nucleotide identity of 78.73% and <i>Methylophilus</i> sp. OH31 (GCF_000576615.1) with an AAI of 85.28% and an ANI of 78.71%.	Methylophilus	NCBI Assembly: GCF_006363915.1 Ts	seqco.de/i:39350
		Type genome is <i>Methylopumilus rimovensis</i> MMS-RI-1 (GCF_006364615.1), an axenic bacterial strain isolated by dilution-to-extinction cultivation from the water column of the Římov Reservoir (Czechia). MMS-RI-1 has a genome size of 1.28 Mb with a genomic GC content of 36.7%, contains 3 rRNA genes and 37 tRNAs. The genome is complete, consisting of 1 circular chromosome. <i>Methylopumilus rimovensis</i> are very small, aerobic, non-motile and methylotrophic. The genome contains genes encoding two rhodopsins (proteorhodopsin and			

Proposed Taxon	[ri.mo.vetysnolngy	xantho-like rhodopsin) and the biosynthetic pathway for retinal	Parent Taxon	Туре	Registry URL
Species Methylopumilus rimovensis	masc. adj. rimovensis, pertaining to the Římov Reservoir, Czech Republic, the isolation source of the species.	biosynthesis. No genes for flagellar or pilus assembly and chemotaxis were annotated. Pathways for methanol oxidation (Xox) and the RuMP cycle for methylotrophy and the biosynthesis of all amino acids were predicted. Methylopumilus rimovensis are highly abundant only in their isolation source (the Římov Reservoir in Czechia) and recorded in very low abundances in other freshwater lakes, which contrasts with members of other species of the genus that have a global distribution (Methylopumilus planktonicus and Methylopumilus universalis). None of the isolated strains were yet submitted to a culture collection because these bacteria are hard to maintain, i.e., they are very slowly growing, reach low densities in liquid culture, and do not grow on agar plates.	Methylopumilus	NCBI Assembly: GCF_006364615.1 Ts	seqco.de/i:42658
Species Methylopumilus universalis	[u.ni.ver.sa'lis] L. adj. masc. <i>universalis</i> , universal, pertaining to the widespread distribution of the species	Type genome is <i>Methylopumilus universalis</i> MMS-RVI-51 (GCF_006363895.1), an axenic bacterial strain isolated by dilution-to-extinction cultivation from the water column of the Římov Reservoir (Czechia). MMS-RVI-51 has a genome size of 1.27 Mb with a genomic GC content of 37.2%, contains 3 rRNA genes and 36 tRNAs. The genome is complete, consisting of 1 circular chromosome. <i>Methylopumilus universalis</i> are very small, aerobic, non-motile and methylotrophic. The genome contains genes encoding two rhodopsins (proteorhodopsin and xantho-like rhodopsin) and the biosynthetic pathway for retinal biosynthesis. No genes for flagellar or pilus assembly and chemotaxis were annotated. Pathways for methanol oxidation (Xox) and the RuMP cycle for	Methylopumilus	NCBI Assembly: GCF_006363895.1 Ts	seqco.de/i:41611

Proposed Taxon	Etymology	methylotrophy and the biosynthesis of all amino acids were predicted.	Parent Taxon	Туре	Registry URL
		Methylopumilus universalis are			
		abundant in the plankton of			
		freshwater lakes with a world-wide			
		distribution and a preference for lakes			
		with a higher nutrient level than members of another species,			
		Methylopumilus planktonicus. None of			
		the isolated strains were yet			
		submitted to a culture collection			
		because these bacteria are hard to			
		maintain, i.e., they are very slowly			
		growing, reach low densities in liquid			
		culture, and do not grow on agar			
		plates.			
		Type genome is <i>Methylosemipumilus</i> turicensis MMS-10A-171			
		(GCF 000953015.1), that was initially			
		proposed as 'Candidatus			
		Methylopumilus turicensis' in Salcher			
		et. al. 2015 and later changed to			
		<i>Methylosemipumilus turicensis</i> in			
		Salcher et al. 2019. The strain was			
		isolated by dilution-to-extinction cultivation from the water column of			
		Lake Zurich, Switzerland. MMS-10A-			
		171 has a genome size of 1.76 Mb			
		with a genomic GC content of 44.52%,			
		contains 6 rRNA genes and 40 tRNAs.			
		The genome is complete, consisting of			
		1 circular chromosome.			
		Methylosemipumilus turicensis MMS- 10A-171 has a relatively small			
	The of contain N. I	genome size (1.76 Mb) but not as			
	[tu.ri.cen'sis] N.L. masc. adj. <i>turicensis</i> , referring	small as <i>Methylopumilus</i> (1.3-1.4 Mb),			
Species	to Turicum, the latin	which - together with improved			
<i>Methylosemipumilus</i>		genomic analyses - led to the	Methylosemipumilus	NCBI Assembly:	segco.de/i:4414
turicensis ^{™s}	pertaining to Lake Zurich,	reclassification to		GCF_000953015.1 Ts	
	the isolation source of the	Methylosemipumilus. The genome			
	type strain	contains genes encoding proteorhodopsins and the biosynthetic			
		pathway for retinal biosynthesis.			
		Pathways for methanol oxidation			
		(Xox), the ribulose monophosphate			
		(RuMP) cycle and the			

Proposed Taxon	Etymology	tetrahydromethanopterin (H4MPT) pathway for formaldehyde oxidation	Parent Taxon	Туре	Registry URL
		were predicted. The closest cultivated relatives are <i>Methylotenera</i> sp. L2L1 (GCF_000744605.1) with an average amino acid identity of 67.45% and average nucleotide identity of 70.55% and <i>Methylovorus</i> sp. MP688 (GCF_000183115.1) with an AAI of 69.37% and an ANI of 69.42%, while the genus <i>Methylopumilus</i> is more distantly related (AAI of 61.5-62.2% and ANI of 66.4-66.9%).			