sporotrichosis (10). Cats with sporotrichosis should be kept indoors and apart from other cats in the home to reduce the potential for further transmission. In conclusion, increased awareness of sporotrichosis in cats and the potential for zoonotic transmission could help veterinary professionals more quickly recognize and treat feline cases and take precautions to prevent human acquisition in the veterinary setting.

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This activity was reviewed by CDC and was conducted consistent with applicable federal law and CDC policy (e.g., 45 C.F.R. part 46, 21 C.F.R. part 56; 42 U.S.C. §241(d); 5 U.S.C. §552a; 44 U.S.C. §3501 et seq).

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References

- Rossow JA, Queiroz-Telles F, Caceres DH, Beer KD, Jackson BR, Pereira JG, et al. A One Health approach to combatting *Sporothrix brasiliensis*: narrative review of an emerging zoonotic fungal pathogen in South America. J Fungi (Basel). 2020;6:247. https://doi.org/10.3390/ jof6040247
- 2. Rees RK, Swartzberg JE. Feline-transmitted sporotrichosis: a case study from California. Dermatol Online J. 2011;17:2. https://doi.org/10.5070/D30459K1JB
- Reed KD, Moore FM, Geiger GE, Stemper ME. Zoonotic transmission of sporotrichosis: case report and review. Clin Infect Dis. 1993;16:384–7. https://doi.org/10.1093/ clind/16.3.384
- de Miranda LHM, Silva JN, Gremião IDF, Menezes RC, Almeida-Paes R, Dos Reis ÉG, et al. Monitoring fungal burden and viability of *Sporothrix* spp. in skin lesions of cats for predicting antifungal treatment response. J Fungi (Basel). 2018;4:92. https://doi.org/10.3390/jof4030092
- Reis ÉG, Schubach TMP, Pereira SA, Silva JN, Carvalho BW, Quintana MSB, et al. Association of itraconazole and potassium iodide in the treatment of feline sporotrichosis: a prospective study. Med Mycol. 2016;54:684–90. https://doi.org/10.1093/mmy/myw027
- 6. Gremião IDF. Martins da Silva da Rocha E, Montenegro H, Carneiro AJB, Xavier MO, de Farias MR, et al. Guideline for the management of feline sporotrichosis caused by

Sporothrix brasiliensis and literature revision. Braz J Microbiol. 2021;52:107-24. https://doi.org/10.1007/s42770-020-00365-3

- Barnacle JR, Chow YJ, Borman AM, Wyllie S, Dominguez V, Russell K, et al. The first three reported cases of *Sporothrix brasiliensis* cat-transmitted sporotrichosis outside South America. Med Mycol Case Rep. 2023;39:14–7. https://doi.org/10.1016/j.mmcr.2022.12.004
- Yingchanakiat K, Limsivilai O, Sunpongsri S, Niyomtham W, Lugsomya K, Yurayart C. Phenotypic and genotypic characterization and antifungal susceptibility of *Sporothrix schenckii sensu stricto* isolated from a feline sporotrichosis outbreak in Bangkok, Thailand. J Fungi (Basel). 2023;9:590. https://doi.org/10.3390/jof9050590
- Welsh RD. Sporotrichosis. J Am Vet Med Assoc. 2003 ;223:1123-6. https://doi.org/10.2460/javma.2003.223.1123
- Crothers SL, White SD, Ihrke PJ, Affolter VK. Sporotrichosis: a retrospective evaluation of 23 cases seen in northern California (1987–2007). Vet Dermatol. 2009;20:249–59. https://doi.org/10.1111/j.1365-3164.2009.00763.x

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Burkholderia thailandensis Isolated from Infected Wound, Southwest China, 2022

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We report a clinical isolate of *Burkholderia thailandensis* 2022DZh obtained from a patient with an infected wound in southwest China. Genomic analysis indicates that this isolate clusters with *B. thailandensis* BPM, a human isolate from Chongqing, China. We recommend enhancing monitoring and surveillance for *B. thailandensis* infection in both humans and livestock.

Burkholderia thailandensis is a member of the Bur-kholderia pseudomallei complex and is generally considered nonpathogenic (1). Initially identified in Thailand, B. thailandensis was distinguished from B. pseudomallei by its ability to assimilate arabinose (2). Similar to B. pseudomallei, B. thailandensis is frequently found in soil and water, especially in rice paddies (3,4). Although invasive infections caused by B. thailandensis are rare worldwide, recent reports have documented cases of suppurative infections, such as wound infections, cellulitis, and tissue abscesses (5,6). Previously, we identified a strain of B. thailandensis BPM that caused a human infection in Chongqing, southwest China (7). In this study, we report another clinical isolate of B. thailandensis that we obtained in Dazhu, Sichuan, southwest China, from an infected wound resulting from a cut inflicted by a farm tool in 2022.

A 61-year-old male farmer who had untreated type 2 diabetes mellitus reported a 1-month history of pain and swelling in his left knee. He had injured the middle toe of his left foot with a plow a month earlier, and redness, swelling, and pain developed below the left knee joint. Despite a week of antimicrobial treatment at a local community health center, his symptoms did not improve. In December 2022, the patient began to experience weakness in his right lower limb, and he later fell, sustaining an injury to his left lower limb. During this period, he experienced lower-limb weakness and exhibited symptoms related to the central nervous system. He sought care and was admitted to the orthopedics department of Dazhu County People's Hospital (Dazhou, China) for treatment of a left lower-limb injury and a left-foot diabetic foot infection. However, because his central nervous system symptoms worsened, he was transferred to the neurology department and receive a diagnosis of osteomyelitis and demyelinating disease. B. thailandensis strain 2022DZh was obtained from a deep-tissue specimen during surgical debridement of the infected wound on the left foot (Appendix Figure 1, https:// wwwnc.cdc.gov/EID/article/30/5/23-0743-App1. pdf). The initial empirical antimicrobial therapy consisted of cefradine. However, the hospital laboratory tested the isolate 2022DZh using the VITEK 2 COMPACT system (bioMérieux, https://www. biomerieux.com) and identified it as B. pseudomallei, leading to a switch to intravenous meropenem treatment (Appendix Table 1). Despite treatment with meropenem, the patient's condition continued to deteriorate; he died 3 days after applying for discharge.

Subsequently, the isolate 2022DZh was submitted to the laboratory for confirmatory identification.

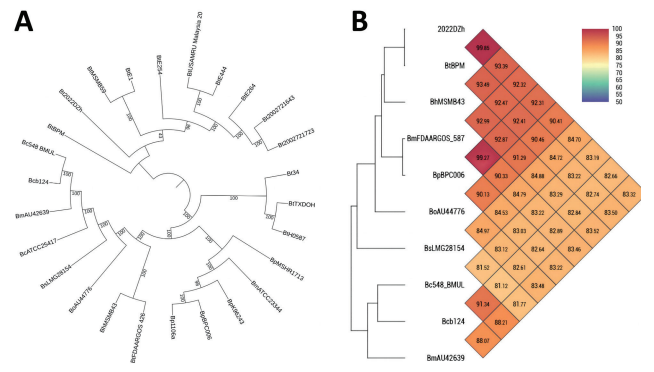


Figure. Analysis of the single-copy gene phylogenetic tree and average nucleotide identity for genomes of *Burkholderia thailandensis* 2022DZh from a patient in Dazhu, Sichuan, China, and other isolates from *Burkholderia* species. A) Single-copy gene phylogenetic tree created using the genomes of *B. thailandensis* 2022DZh and 25 other isolates from various *Burkholderia* species. B) Average nucleotide identity heatmap developed using genomes of *B. thailandensis* 2022DZh and 9 other isolates from various *Burkholderia* species.

Results of arabinose assimilation testing of isolate 2022DZh by API 20NE system (bioMérieux) were positive, consistent with the biochemical characteristics of B. thailandensis (Appendix Figure 2). We extracted DNA from the isolate 2022DZh for confirmation and further characterization. We performed 16S rRNA gene sequencing of 2022DZh using nucleotide (https://blast.ncbi.nlm.nih.gov/Blast.cgi), BLAST revealing 100% similarity with B. thailandensis BPM (Appendix Figure 3). The nucleotide sequences of the 2 chromosomes of isolate 2022DZh are >99.5% consistent with those of B. thailandensis E264 and B. thailandensis E254 (Appendix Table 2). We conclusively identified isolate 2022DZh as B. thailandensis based on our phenotypic and molecular data. We deposited the genome sequences of B. thailandensis strain 2022DZh into GenBank (accession nos. CP141809.1 and CP141811.1).

For phylogenetic analysis, we compared the genome of B. thailandensis 2022DZh with a reference panel of publicly available Burkholderia species genomes (Appendix Table 3). The single-copy gene phylogenetic tree analysis indicated that B. thailandensis 2022DZh clusters with B. thailandensis BPM (Figure, panel A). The results of average nucleotide identity revealed that B. thailandensis 2022DZh also clusters with B. thailandensis BPM; genome identity was 99.85%. However, when compared with B. pseudomallei BPC006, genome identity was 92.31% (Figure, panel B), consistent with the commonly used 95% threshold for distinguishing species. Through multilocus sequence type analysis (https://pubmlst. org/organisms/burkholderia-pseudomallei) (8), we determined that B. thailandensis 2022DZh and B. thailandensis BPM both belong to sequence type 76. In addition, there appears to be no known epidemiologic link between B. thailandensis 2022DZh and B. thailandensis BPM; they are geographically separated by a significant distance of ≈200 km (Appendix Table 3).

One limitation of this study is that we did not attempts to identify related isolates of *B. thailandensis* from environmental samples in this region of southwest China. Further studies are needed to identify the primary molecular mechanisms underlying the pathogenicity of *B. thailandensis* 2022DZh and to determine its molecular and evolutionary relationships with other strains of *B. thailandensis* (9).

In conclusion, our findings underscore that *B. thailandensis* can cause serious infections, and clinical practitioners should be aware of this type of infection (10). Therefore, we strongly recommend enhancing monitoring and surveillance for *B. thailandensis* infection in both humans and livestock.

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References

- Ronning CM, Losada L, Brinkac L, Inman J, Ulrich RL, Schell M, et al. Genetic and phenotypic diversity in *Burkholderia*: contributions by prophage and phage-like elements. BMC Microbiol. 2010;10:202. https://doi.org/ 10.1186/1471-2180-10-202
- Lertpatanasuwan N, Sermsri K, Petkaseam A, Trakulsomboon S, Thamlikitkul V, Suputtamongkol Y. Arabinose-positive *Burkholderia pseudomallei* infection in humans: case report. Clin Infect Dis. 1999;28:927–8. https://doi.org/10.1086/517253
- Birnie E, van 't Hof S, Bijnsdorp A, Mansaray Y, Huizenga E, van der Ende A, et al. Identification of *Burkholderia thailandensis* with novel genotypes in the soil of central Sierra Leone. PLoS Negl Trop Dis. 2019;13:e0007402. https://doi.org/10.1371/journal.pntd.0007402
- Hall CM, Stone NE, Martz M, Hutton SM, Santana-Propper E, Versluis L, et al. *Burkholderia thailandensis* isolated from the environment, United States. Emerg Infect Dis. 2023;29:618– 21. https://doi.org/10.3201/eid2903.221245
- Gee JE, Elrod MG, Gulvik CA, Haselow DT, Waters C, Liu L, et al. *Burkholderia thailandensis* isolated from infected wound, Arkansas, USA. Emerg Infect Dis. 2018;24:2091–4. https://doi.org/10.3201/eid2411.180821
- Glass MB, Gee JE, Steigerwalt AG, Cavuoti D, Barton T, Hardy RD, et al. Pneumonia and septicemia caused by *Burkholderia thailandensis* in the United States. J Clin Microbiol. 2006;44:4601–4. https://doi.org/10.1128/JCM.01585-06
- Chang K, Luo J, Xu H, Li M, Zhang F, Li J, et al. Human infection with *Burkholderia thailandensis*, China, 2013. Emerg Infect Dis. 2017;23:1416–8. https://doi.org/10.3201/ eid2308.170048
- Tuanyok A, Mayo M, Scholz H, Hall CM, Allender CJ, Kaestli M, et al. Burkholderia humptydooensis sp. nov., a new species related to Burkholderia thailandensis and the fifth member of the Burkholderia pseudomallei complex. Appl Environ Microbiol. 2017;83:e02802–16. https://doi.org/ 10.1128/AEM.02802-16
- Li J, Zhong Q, Li J, Chong HM, Wang LX, Xing Y, et al. Genomic features and virulence characteristics of a rare *Burkholderia thailandensis* strain causing human infection. J Med Microbiol. 2023;72:72. https://doi.org/10.1099/jmm.0.001688
- Cossaboom CM, Marinova-Petkova A, Strysko J, Rodriguez G, Maness T, Ocampo J, et al. Melioidosis in a resident of Texas with no recent travel history, United States. Emerg Infect Dis. 2020;26:1295–9. https://doi.org/10.3201/ eid2606.190975

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Burkholderia thailandensis Isolated from Infected Wound, Southwest China, 2022

Appendix

Appendix Table 1. Antibiotic susceptibility testing results of *B. thailandensis* 2022DZh

Antimicrobial	MIC	Interpretation
Amoxicillin/clavulanate	<u><</u> 4	S
Ceftazidime	2	S
Imipenem	<u><</u> 1	S
Tetracycline	4	S
Doxycycline	<u><</u> 0.5	S
Trimethoprim/sulfamethoxazole	40	S

	Query_	Query_	Query_							
Query_name	length	start	end	Hit_name	Hit_description	Hit_length	Hit_start	Hit_end	Aln_length	Identity
chr1	3804194	627187	941171	CP008785.1	Burkholderia	3986340	1022844	1336839	314053	99.90320105
					thailandensis E264					20517
					chromosome 1, complete					
					sequence					
chr2	2860001	831328	941171	CP004382.1	Burkholderia	2870750	2145935	2478653	332805	99.87410044
	200001	031320	941171	CF004382.1	thailandensis E254	2010130	2143933	2470033	332003	99.87410044
										5212
					chromosome 2, complete					
					sequence					
plasmid1	215545	135181	157561	CP013426.1	<i>Burkholderia</i> sp.	249331	8541	30930	22395	98.04420629
					MSMB0856 plasmid					60482
					pMSMB0856, complete					
					sequence					

Appendix Table 2	. The Nucleotide Sequence Database	e (NT) identified strain 2022DZh as <i>B. thailandensis</i> *
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*The NT database is the Nucleotide Sequence Database, and the link is http://www.ncbi.nlm.nih.gov

solate	Abbreviation	Origin	Year	Source	MLST	GenBank accession no
B. thailandensis H0587	BtH0587	USA	1997	Clinical	101	NZ_CP004089.1;
				(Human)		NZ_CP004090.1
3. thailandensis BPM	BtBPM	China	2013	Clinical	76	NZ_CP050020.1;
				(Human)		NZ_CP050021.1
3. thailandensis E264	BtE264	Thailand	1994	Environmental	80	NZ_CP008785.1;
				(soil)		NZ_CP008786.1
3. thailandensis E254	BtE254	Thailand	1992	Environmental	345	NZ_CP004381.1;
				(soil)		NZ_CP004382.1
B. thailandensis E444	BtE444	Thailand	2002	Environmental	79	NZ_CP004117.1;
				(soil)		NZ_CP004118.1
3. thailandensis MSMB59	BtMSMB59	Australia	2006	Environmental	669	NZ_CP004385.1;
				(soil)		NZ_CP004386.1
3. thailandensis USAMRU	BtUSAMRU Malaysia	Malaysia	2015	Unknown	80	NZ_CP004383.1;
Malaysia 20	20					NZ_CP004384.1
3. thailandensis 34	Bt34	USA	2002	Unknown	73	NZ_CP010017.1;
						NZ_CP010018.1
3. thailandensis 2002721723	Bt2002721723	USA	2013	Clinical	80	NZ_CP004098.1;
				(Human)		NZ_CP004097.1
3. thailandensis 2002721643	Bt2002721643	USA	2002	Unknown	80	NZ_CP009601.1;
						NZ_CP009602.1
3. thailandensis E1	BtE1	Papua New	1995	Environmental	669	NZ_LOXF00000000.1
		Guinea		(soil)		
3. thailandensis TXDOH	BtTXDOH	USA	2003	Clinical	101	NZ_ABBD00000000.1
				(Human)		
3. thailandensis	BtFDAARGOS_426	Malaysia	2017	Environmental	1023	NZ_CP023499.1;
DAARGOS_426				(soil)		NZ_CP023498.1
3. pseudomallei K96243	BpK96243	Thailand	1993	Clinical	10	NZ_CP009538.1;
				(Human)		NZ_CP009537.1
3. pseudomallei BPC006	BpBPC006	China	2008	Clinical	70	NZ_CP003781.1;
				(Human)		NZ_CP003782.1
3. pseudomallei 1106a	Bp1106a	Thailand	2008	Clinical	70	NZ_CP008758.1;
				(Human)		NZ_CP008759.1
3. pseudomallei MSHR1713	BpMSHR1713	Australia	2003	Clinical	131	NZ_CP111138.1;
				(Human)		NZ_CP111139.1
		Australia	2000	Clinical	131	NZ_CP111140.1;
B. pseudomallei MSHR1046	BpMSHR1046	Australia	2000	Olimodi	101	h<u><u></u></u>u

Appendix Table 3. Genomes of additional *Burkholderia* species isolates used for single-copy gene phylogenetic tree analysis and average nucleotide identity (ANI) analysis in the study of *B. thailandensis* 2022DZh

Isolate	Abbreviation	Origin	Year	Source	MLST	GenBank accession no.
B. mallei ATCC23344	BmATCC23344	USA	1942	Clinical	40	NZ_CP124608.1;
				(Human)		NZ_CP124607.1
B. singularis LMG 28154	BsLMG 28154	Canada	2003	Clinical	166	NZ_FXAN01000000
				(Human)		
B. oklahomensis AU44776	BoAU44776	USA	2022	Clinical	1772	NZ_JAUJRD00000000.1
				(Human)		
<i>B. cepacia</i> b124	Bcb124	Bangladesh	2017	Clinical	140	NZ_SNSI01000000
				(Human)		
B. cenocepacia 548_BMUL	Bc548_BMUL	USA	2016	Clinical	162	NZ_JVCQ01000000
				(Human)		
B. multivorans AU42639	BmAU42639	USA	2020	Clinical	79	NZ_JAHPLO01000000
				(Human)		
B. cocovenenans ATCC25417	BcATCC25417	USA	2012	Environmental	101	NZ_JPGG0000000.1
				(leaves)		
B. mallei FDAARGOS_587	BmFDAARGOS_587	USA	1961	Environmental	40	NZ_RKJX00000000.1
				(Horse)		
B. humptydooensis MSMB43	BhMSMB43	Australia	1995	Environmental	318	NZ_AJXB00000000.1
				(Water)		



Appendix Figure 1. Morphologic features of left-foot diabetic foot infection caused by *B. thailandensis* 2022DZh in a patient in Dazhu, Sichuan, China. A) Result of surgical debridement

of a wound in the 61-year-old male patient. B) Colony morphology of *B. thailandensis* 2022DZh

from the patient on a Columbia blood plate.



Appendix Figure 2. The biochemical profiles of the API 20NE system, including arabinose

assimilation, identified isolate 2022DZh as *B. thailandensis*.

Sec	uences producing significant alignments	Downloa	ad ~	S	elect	column	s ~ S	how 1	00 ~ 00
•	select all 100 sequences selected	GenE	GenBank			Distance tree of results			MSA Viewer
	Description	Scientific Name	Max Score		Query Cover	E value	Per. Ident	Acc. Len	Accession
	Burkholderia thailandensis strain BPM chromosome 2, complete sequence	Burkholderia thailandensis	2542	2542	100%	0.0	100.00%	2875717	CP050021.1
	Burkholderia thailandensis strain BPM chromosome 1, complete sequence	Burkholderia thailandensis	2542	7828	100%	0.0	100.00%	3725540	CP050020.1
	Burkholderia thailandensis strain FDAARGOS 242 chromosome 1, complete sequence	Burkholderia thailandensis	2542	7606	100%	0.0	100.00%	3801254	CP022217.1
	Burkholderia thailandensis strain FDAARGOS 242 chromosome 2, complete sequence	Burkholderia thailandensis	2542	2542	100%	0.0	100.00%	2852057	CP022216.1
	Burkholderia thailandensis strain FDAARGOS 237 chromosome 1, complete sequence	Burkholderia thailandensis	2542	7618	100%	0.0	100.00%	3819771	CP020390.1
	Burkholderia thailandensis strain MSMB59 chromosome 2, complete sequence	Burkholderia thailandensis	2542	2542	100%	0.0	100.00%	2822193	CP013408.1
	Burkholderia thailandensis strain MSMB59 chromosome 1, complete sequence	Burkholderia thailandensis	2542	7626	100%	0.0	100.00%	3917337	CP013407.1
	Burkholderia thailandensis 2002721843 chromosome II, complete seguence	Burkholderia thailandensis 2002721643	2542	2542	100%	0.0	100.00%	2914772	CP009602.1
	Burkholderia thailandensis 2002721843 chromosome I, complete sequence	Burkholderia thailandensis 2002721643	2542	7626	100%	0.0	100.00%	3808029	CP009601.1
	Burkholderia thailandensis E254 chromosome 2, complete seguence	Burkholderia thailandensis E254	2542	2542	100%	0.0	100.00%	2870750	CP004382.1
	Burkholderia thailandensis E254 chromosome 1, complete seguence	Burkholderia thailandensis E254	2542	7620	100%	0.0	100.00%	3805980	CP004381.1
	Burkholderia thailandensis MSMB59 chromosome 2, complete sequence	Burkholderia thailandensis MSMB59	2542	2542	100%	0.0	100.00%	2823176	CP004388.1
	Burkholderia thailandensis MSMB59 chromosome 1, complete sequence	Burkholderia thailandensis MSMB59	2542	7620	100%	0.0	100.00%	3916324	CP004385.1
	Burkholderia thailandensis E264 chromosome 2, complete sequence	Burkholderia thailandensis E264	2542	5076	100%	0.0	100.00%	2735759	CP008788.1
	Burkholderia thailandensis E284 chromosome 1, complete sequence	Burkholderia thailandensis E264	2542	5076	100%	0.0	100.00%	3986340	CP008785.1
	Burkholderia thailandensis USAMRU Malaysia #20 chromosome 2, complete seguence	Burkholderia thailandensis USAMRU	2542	2542	100%	0.0	100.00%	2821713	CP004384.1
	Burkholderia thailandensis USAMRU Malaysia #20 chromosome 1, complete sequence	Burkholderia thailandensis USAMRU	2542	7615	100%	0.0	100.00%	3882848	CP004383.1
	Burkholderia thailandensis E444 chromosome 2, complete sequence	Burkholderia thailandensis E444	2542	2542	100%	0.0	100.00%	2852101	CP004118.1
	Burkholderia thailandensis E444 chromosome 1, complete sequence	Burkholderia thailandensis E444	2542	7618	100%	0.0	100.00%	3799595	CP004117.1
	Burkholderia thailandensis 2002721723 chromosome 2, complete seguence	Burkholderia thailandensis 2002721723	2542	2542	100%	0.0	100.00%	2914718	CP004098.1
	Burkholderia thailandensis 2002721723 chromosome 1, complete sequence	Burkholderia thailandensis 2002721723	2542	7626	100%	0.0	100.00%	3662415	CP004097.1
	Burkholderia thailandensis strain AW34-19p chromosome 2, complete sequence	Burkholderia thailandensis	2542	2542	100%	0.0	100.00%	2826723	CP068710.1
	Burkholderia thailandensis strain AW34-19p chromosome 1, complete sequence	Burkholderia thailandensis	2542	7484	100%	0.0	100.00%	3830943	CP068709.1
	Burkholderia thailandensis E264 chromosome I, complete sequence	Burkholderia thailandensis E284	2542	7626	100%	0.0	100.00%	3809201	CP000088.1
	Burkholderia thailandensis E264 16S ribosomal RNA, partial sequence	Burkholderia thailandensis E284	2538	2538	100%	0.0	99.93%	1529	NR 074312.2
	Burkholderia thailandensis E284 chromosome II, complete seguence	Burkholderia thailandensis E284	2538	2536	100%	0.0	99.93%	2914771	CP000085.1
	Burkholderia thailandensis strain FDAARGOS 241 chromosome 2, complete sequence	Burkholderia thailandensis	2534	2534	100%	0.0	99.93%	2821560	CP022215.1
	Burkholderia thailandensis strain FDAARGOS 237 chromosome 2, complete sequence	Burkholderia thailandensis	2534	2534	100%	0.0	99.93%	2853041	CP020389.1
	Burkholderia thailandensis strain BD10-00323 16S ribosomal RNA, partial sequence	Burkholderia thailandensis	2534	2534	99%	0.0	100.00%	1459	NR 118629.1
	Burkholderia thailandensis strain BD11-00023 16S ribosomal RNA gene, partial seque	Burkholderia thailandensis	2534	2534	99%	0.0	100.00%	1485	KF444905.1
	Burkholderia thailandensis strain BD11-00022 16S ribosomal RNA gene, partial seque	Burkholderia thailandensis	2534	2534	99%	0.0	100.00%	1457	KF444904.1
	Burkholderia thailandensis E264 16S ribosomal RNA, partial sequence	Burkholderia thailandensis E284	2531	2531	99%	0.0	100.00%	1541	NR 116051.1
	Burkholderia thailandensis strain 2002721627 16S ribosomal RNA gene, partial seque	Burkholderia thailandensis	2531	2531	99%	0.0	100.00%	1488	AY268182.1
	Burkholderia thailandensis strain FDAARGOS 241 chromosome 1, complete sequence	Burkholderia thailandensis	2529	7554	100%	0.0	99.85%	3882584	CP022214.1
-					10000	1000	10000000000000000000000000000000000000		

Appendix Figure 3. The 16S rRNA gene sequencing of strain 2022DZh using nucleotide BLAST.