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In Vitro Antimicrobial Activity of Taurolidine against Candida auris **Bloodstream Isolates from Global Sources**

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Introduction

- The Centers for Disease Control and Prevention (CDC) have recently warned that the fungus Candida auris is an emerging, resistant pathogen of concern [1], and recent epidemiological reports point to the increasing global spread of *C. auris* [2, 3].
- Recent reports highlight C. auris as an emerging cause of invasive bloodstream infections (BSI) [4-6].
- Central venous catheter use is associated with C. *auris* BSI [4].
- Taurolidine (Figure 1) has been marketed in Europe as a catheter lock solution component with the goal of reducing catheter-related bloodstream infections (CRBSI) [7].
- Taurolidine is a novel antimicrobial with broad spectrum antibacterial/antifungal activity and two mechanisms of action that does not lend itself to clinically relevant microbial resistance at concentrations contained within a central venous catheter.
- Taurolidine exerts its activity through damage to microbial cell walls by denaturing surface proteins and chemically altering membrane lipids, as well as inhibiting adherence of microorganisms to biological surfaces.
- A large, double-blind, randomized comparator-controlled study was conducted that evaluated the safety and efficacy of a catheter lock solution containing taurolidine/heparin to prevent CRBSIs in patients with advanced kidney disease on hemodialysis (*Clin J Am Soc* Nephrol, in press, https://journals.lww.com/cjasn/abstract/9900/taurolidine_heparin_lock _solution_and.236.aspx).
- The marketing application for use in the United States is currently under review by the United States Food and Drug Administration (US FDA).
- To provide additional evidence for the potential utility of taurolidine to reduce CRBSI, this study investigated the *in vitro* antimicrobial activity of taurolidine against a diverse set of C. auris strains/isolates using reference testing methods.

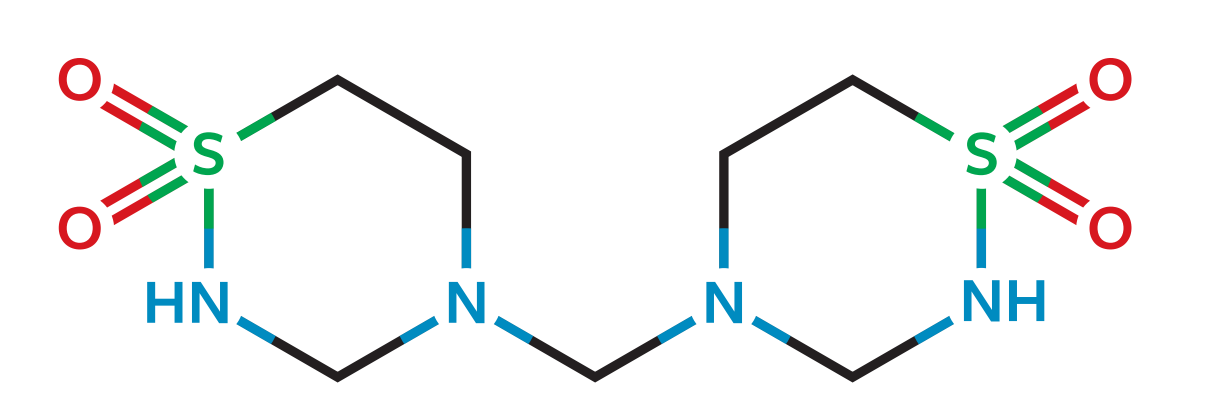
Materials and Methods

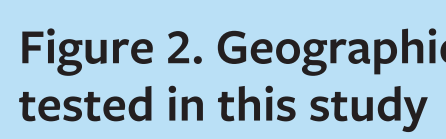
- 41 C. auris strains/isolates were collected from various sources during 2008–2019 (Table 1 and Figure 2), and the species were confirmed by matrix-assisted laser desorption ionization-time of flight mass spectrometry (Bruker Daltonics, Bremen, Germany) or other standard methods:
- The CDC and the US FDA Antimicrobial Resistance Isolate Bank (CDC AR) (n=11) [8].
- Several clades were represented (Table 1).
- The Westerdijk Fungal Biodiversity Institute (n=4) [9].
- JMI Laboratories' SENTRY Antimicrobial Surveillance Program (n=16) [10].
- A special collection of *C. auris* isolates obtained by JMI Laboratories from a hospital in Nairobi, Kenya (n=10) [11].
- Most C. *auris* isolates (90%; 37/41) were from bloodstream infections (Table 1).
- The C. auris set was tested for antifungal susceptibility using Clinical and Laboratory Standards Institute (CLSI) broth microdilution guidelines [12, 13].
- The test medium was Roswell Park Memorial Institute 1640 broth buffered with MOPS (morpholinepropanesulfonic acid) and 0.2% (w/v) glucose [12].
- JMI Laboratories produced the minimal inhibitory concentration (MIC) panels.
- CLSI-recommended quality control strains were also tested (*Candida krusei* ATCC 6258 and Candida parapsilosis ATCC 22019).
- MIC values were read after 24 hours at 35°C in ambient air.
- Taurolidine MIC values were read at both 50% and 100% growth inhibition. – No CLSI reading criteria for taurolidine have been published.
- Amphotericin B and fluconazole were also tested as control antifungal agents.
- Tentative CDC breakpoints were applied to C. *auris* [14].

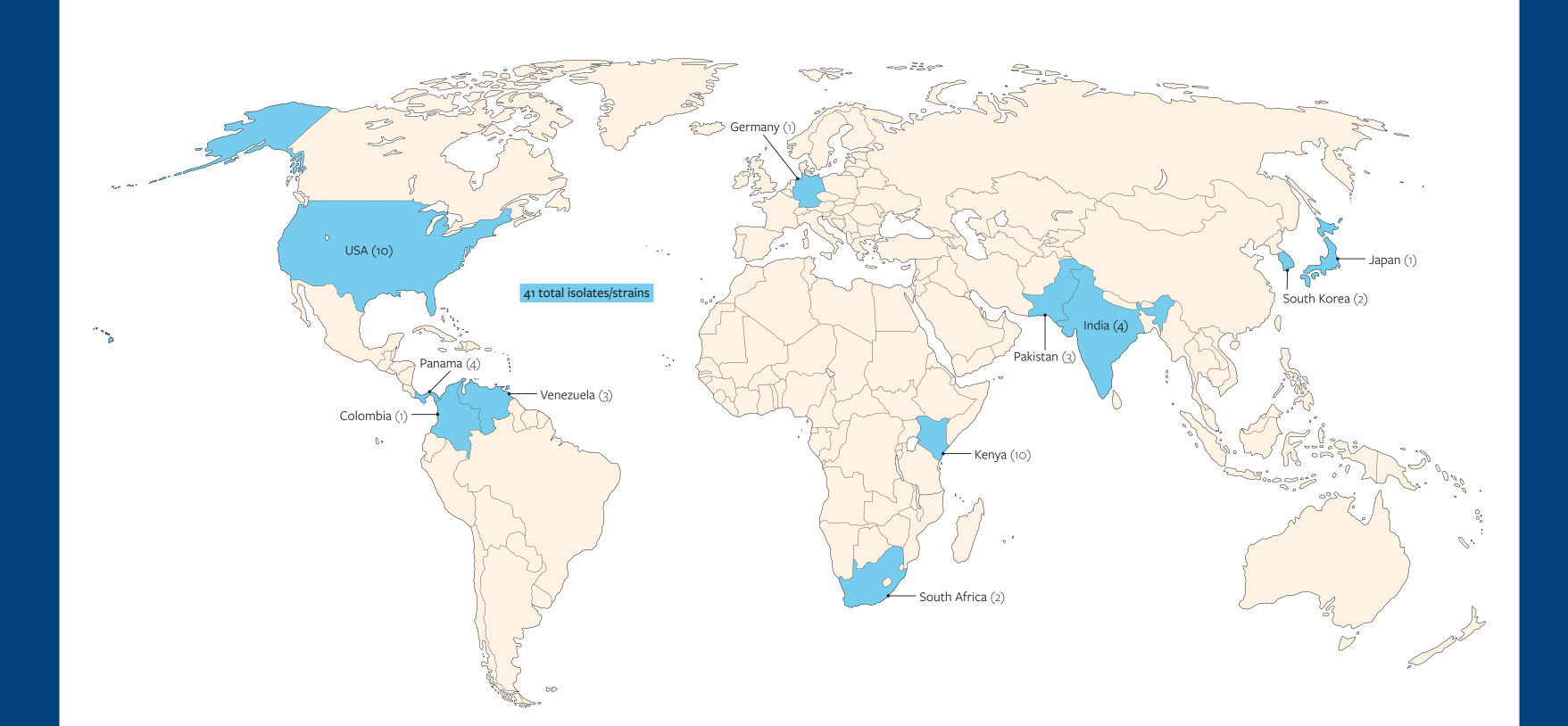
Results

- Taurolidine exhibited *in vitro* antimicrobial activity against all *C. auris* strains and isolates regardless of geographic source, year of isolation, or clade (Tables 1–3). - Overall MIC_{50/90} values were 256/512 mg/L using the 50% inhibition reading criterion and 512/512 mg/L using the 100% inhibition reading criterion (Table 3).
- The taurolidine MIC range was 128–512 mg/L and 256–1,024 mg/L using the 50% and 100% inhibition reading criteria, respectively (Table 3).
- In general, taurolidine MIC values were slightly more potent when read using the 50% inhibition criterion (Tables 1–2).
- The full set of *C. auris* isolates was 53.7% resistant to amphotericin B and 85.4% resistant to fluconazole using tentative CDC MIC breakpoints (Table 3; [14]).
- (Table 4).

Figure 1. Two-dimensional structure of taurolidine







- Where applicable, MIC data for amphotericin B and fluconazole tested against these strains/ isolates agreed well with previously published data (Table 1).
 - Because these isolates were not randomly selected, however, these resistance rates cannot be extrapolated to other sets of *C. auris* isolates.
- Taurolidine maintained activity against the amphotericin B and fluconazole-resistant subsets

Figure 2. Geographical sources of *Candida auris* strains/isolates

Table 1. Candida auris isolates/strains and their associated MIC values

									AIC (mg/L)	
Collection							Tauro	olidine	Amphotericin B	Fluconazole
No.	Source	Year	Country	Specimen type	Clade ^a	Alias	(50% inhibition)	(100% inhibition)	(published MIC value)	(published MIC value)
1	CDC AR	2019	Pakistan	Burn wound	South Asia	AR Bank# 0382	256	512	1 (0.4) ^b	4 (16) ^b
2	SENTRY	2019	Panama	Blood culture			256	512	2	64
3	SENTRY	2019	Panama	Blood culture			256	512	2	8
4	SENTRY	2019	USA	Blood culture			512	512	2	>128
5	SENTRY	2019	USA	Blood culture			256	512	2	>128
6	CDC AR	2018	India	BAL	South Asia	AR Bank# 0389	512	512	2 (4) ^b	>128 (256) ^b
7	SENTRY	2018	Panama	Blood culture			256	512	2	8
8	SENTRY	2018	Panama	Blood culture			512	512	2	4
9	SENTRY	2018	USA	Blood culture			512	512	2	>128
10	SENTRY	2018	USA	Blood culture			512	512	2	>128
11	SENTRY	2018	USA	Blood culture			512	512	2	>128
12	SENTRY	2017	USA	Blood culture			256	512	2	>128
13	SENTRY	2016	USA	Blood culture			512	512	1	>128
14	SENTRY	2016	USA	Blood culture			512	512	1	>128
15	SENTRY	2015	USA	Blood culture			512	512	1	>128
16	SENTRY	2014	Colombia	Blood culture			512	512	1	64
17	CDC AR	2014	Pakistan	Blood culture	South Asia	AR Bank# 0388	512	512	2 (1.5) ^b	>128 (>256) ^b
18	Kenya	2013	Kenya ^c	Blood culture			256	512	1	32
19	Kenya	2013	Kenya ^c	Blood culture			256	256	2	>128
20	Kenya	2013	Kenya ^c	Blood culture			256	512	1	128
21	Kenya	2013	Kenya ^c	Blood culture			512	512	1	>128
22	SENTRY	2013	USA	Blood culture			256	512	2	>128
23	CDC AR	2013	Venezuela	Blood culture	South America	AR Bank# 0931	512	1024	2 (0.75) ^b	>128 (>256) ^b
24	Kenya	2012	Kenya ^c	Blood culture			256	512	1	128
25	Kenya	2012	Kenya ^c	Blood culture			256	512	1	>128
26	Kenya	2012	Kenya ^c	Blood culture			256	512	1	>128
27	Kenya	2012	Kenya ^c	Blood culture			256	512	1	128
28	Kenya	2012	Kenya ^c	Blood culture			128	512	2	128
29	CDC AR	2012	South Africa	Blood culture	Africa	AR Bank# 0383	256	512	1 (0.4) ^b	>128 (128) ^b
30	CDC AR	2012	South Africa	Blood culture	Africa	AR Bank# 0384	256	512	1 (0.5) ^b	>128 (128) ^b
31	CDC AR	2012	Venezuela	Blood culture	South America	AR Bank# 0385	512	512	1 (0.5) ^b	>128 (>256) ^b
32	CDC AR	2012	Venezuela	Blood culture	South America	AR Bank# 0386	512	512	1 (0.5) ^b	>128 (>256) ^b
33	Kenya	2011	Kenya ^c	Blood culture			256	512	2	>128
34	SENTRY	2009	Germany	Blood culture			256	512	2	128
35	CDC AR	2009	Japan	Ear	East Asia	AR Bank# 0381	256	512	1 (0.4) ^b	2 (4) ^b
36	CDC AR	2008	Pakistan	Blood culture	South Asia	AR Bank# 0387	256	512	1 (0.75) ^b	4 (8) ^b
37	CDC AR	NR	India	Wound	South Asia	AR Bank# 0390	512	512	2 (4) ^b	>128 (>256) ^b
38	Westerdijk	NR	India	Blood culture		CBS 12768 ^d	512	1024	2 (4) ^e	>128 (32) ^e
39	Westerdijk	NR	India	Blood culture		CBS 12766 ^d	512	1024	2 (4) ^e	>128 (32) ^e
40	Westerdijk	NR	South Korea	Blood culture		CBS 12372 ^d	256	512	1 (1) ^e	128 (2) ^{e, f}
41	Westerdijk	NR	South Korea	Blood culture		CBS 12372 d	256	512	2 (1) ^e	128 (16) ^{e, f}
	,				Isolate Bank: MIC minimal inhibito	bry concentration; NR, not reported.			- (')	

Abbreviations: BAL, bronchoalveolar lavage; CDC AR, Centers for Disease Control and Prevention and US FDA Antimicrobial Resistance Isolate Bank; MIC, minimal inhibitory concentration; NR, not reported. ^a Clade categorization as reported by the CDC and FDA Antibiotic Resistance (CDC AR) isolate bank [8] ^b MIC values for amphotericin B (Etest) and fluconazole from the CDC AR bank website are shown in parentheses. he Kenyan isolates are described by Adam et al. [11].

^dCBS isolates are from the Westerdijk Fungal Biodiversity Institute [9].

e 24-hour MIC values for amphotericin B and fluconazole from Larkin et al. [15] are shown in parentheses. The 48-hour fluconazole MIC values were all >64 mg/L. The fluconazole 24-hour MIC values were measured twice for strain #40 (repeat MIC value, 64 mg/L) and strain #41 (repeat MIC value, 128 mg/L).

Table 2. Cumulative distributions of taurolidine MIC values against various Candida *auris* subsets

Candida auris set		No. an	id cun	nulativ	e % of	⁻ isolat	es inh	ibited	at MI	C (mg	/L) of:			MIC
(no. of isolates)	≤0.5	1	2	4	8	16	32	64	128	256	512	1024	MIC ₅₀	MIC ₉
All Candida auris	1		1			1								
Taurolidine 50%								0	1	22	18		256	512
inhibition (41)								0.0	2.4	56.1	100.0		230	JIZ
Taurolidine 100%									0	1	37	3	512	512
inhibition (41)									0.0	2.4	92.7	100.0	JIZ	JTZ
SENTRY Candida au	ris		I		1	1	1	1	1	1	1			
Taurolidine 50%									0	7	9		512	512
inhibition (16)									0.0	43.8	100.0		JIZ	J 12
Taurolidine 100%										0	16		512	512
inhibition (16)										0.0	100.0		JIZ	J I 2
Kenyan Candida aur	ris							1						
Taurolidine 50%								0	1	8	1		256	256
inhibition (10)								0.0	10.0	90.0	100.0		230	230
Taurolidine 100%									0	1	9		512	512
inhibition (10)									0.0	10.0	100.0		JIZ	J12
CDC AR bank Candi	da aur	is	1				1	1	1	1	1			
Taurolidine 50%									0	5	6		512	512
inhibition (11)									0.0	45.5	100.0		512	J 12
Taurolidine 100%										0	10	1	512	512
inhibition (11)										0.0	90.9	100.0	512	512
Westerdijk Candida	auris					1	1	1	1	1	1			
Taurolidine 50%									0	2	2		256	
inhibition (4)									0.0	50.0	100.0		230	
Taurolidine 100%										0	2	2	512	
inhibition (4)										0.0	50.0	100.0	512	

Table 3. Activity of taurolidine and comparators against the full Candida auris set (n=41)

Antimicrobiologont			Danca	CLSI ^a			
Antimicrobial agent	1011C ₅₀	MIC ₉₀	Range	%S	%	% R	
Taurolidine 50% inhibition	256	512	128 to 512				
Taurolidine 100% inhibition	512	512	256 to 1024				
Amphotericin B	2	2	1 to 2	46.3		53.7	
Fluconazole	>128	>128	2 to >128	14.6		85.4	

^aUsing tentative CDC antifungal susceptibility breakpoints [14]

Table 4. Activity of taurolidine against resistant Candida *auris* subsets

Resistant subset ^a	No. of	mg/L					
Taurolidine criterion	isolates	MIC ₅₀	MIC ₉₀	MIC range			
Amphotericin B MIC ≥2 mg/L							
Taurolidine 50% criterion	22	256	512	128 to 512			
Taurolidine 100% criterion	22	512	1024	256 to 1024			
Fluconazole MIC ≥32 mg/L							
Taurolidine 50% criterion	35	256	512	128 to 512			
Taurolidine 100% criterion	35	512	512	256 to 1024			
Using tentative CDC antifungal susceptibility breakpoints [14].							

Conclusions

- Taurolidine activity was similar for all *C. auris* subsets tested regardless of source or clade. – Overall MIC_{50/90} values were 256/512 mg/L using the 50% inhibition reading criterion and
- 512/512 mg/L using the 100% inhibition reading criterion.
- There was no evidence that taurolidine activity was affected by resistance to amphotericin B or fluconazole.
- Based on these data, catheter lock solutions containing the broad-spectrum antimicrobial taurolidine at 13,500 mg/L have the potential to prevent CRBSI caused by C. auris, including clinical isolates that are resistant to amphotericin B and fluconazole.
- Taurolidine MIC values are being measured against additional C. auris isolates to further explore the activity of this broad-spectrum antimicrobial.

Disclosures

JMI Laboratories received compensation for services related to the preparation of this poster.

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