- 1 <u>International Journal of Antimicrobial Agents</u>, <u>Volume 41, Issue 5</u>, 2013, Pages
- 2 488–489
- 3 <u>http://dx.doi.org/10.1016/j.ijantimicag.2013.01.001</u>
- 4 Title
- 5 Activity of the efflux pump inhibitor SILA 421 against drug-resistant tuberculosis

7 Authors

6

11

- 8 Sami O. Simons#¹, Jette E. Kristiansen², Gyorgy Hajos³, Tridia van der Laan⁴, József
- 9 Molnár⁵, Martin J. Boeree¹, Jakko van Ingen⁶, Jørn B. Christensen⁷, Miguel Viveiros⁸,
- 25 Zsuzsanna Riedl³, Leonard Amaral⁸, and Dick van Soolingen⁶

12 Affiliations

- ¹ Department of Pulmonary Medicine, Radboud University Nijmegen Medical Centre,
- 14 Nijmegen, the Netherlands
- ²Centre for Biomembrane Physics (Memphys), University of Southern Denmark, Odense,
- 16 Denmark
- ³Institute of Organic Chemistry, Research Centre for Natural Sciences, Hungarian Academy
- of Sciences, Budapest, Hungary
- ⁴National Tuberculosis Reference Laboratory, National Institute for Public Health and the
- 20 Environment (RIVM), Bilthoven, the Netherlands
- ⁵Department of Medical Microbiology and Immunobiology, University of Szeged, Szeged,
- 22 Hungary
- ⁶Department of Clinical Microbiology, Radboud University Nijmegen Medical Centre,
- Nijmegen, the Netherlands
- ⁷Department of Chemistry, University of Copenhagen, Copenhagen, Denmark

26	⁸ Unit of Mycobacteriology, Institute of Hygiene and Tropical Medicine, New University of
27	Lisbon, Lisbon, Portugal
28	
29	(#)Corresponding author:
30	Sami O. Simons, MD
31	Department of Pulmonary Medicine
32	Radboud University Nijmegen Medical Centre
33	PO Box 9101, 6500 HB Nijmegen
34	the Netherlands
35	Tel: +31 (0)243610325
36	Fax: +31 (0)243610324
37	E-mail: s.simons@long.umcn.nl
38	
39	
40	
41	
42	
43	
44	
45	
46	
47	
48	
49	
50	

51 Dear editor,

52

53

54

55

56

57

58

59

60

61

62

63

64

65

66

67

68

69

70

71

72

73

74

75

Organosilicon compounds are efflux pump inhibitors with potency as an antituberculosis drug. Of the organisilicon compounds tested, SILA 421 has been shown to have a highest potency as an antituberculosis drug (1). It shares the common pathways for antimycobacterial killing with other efflux pump inhibitors: it revealed direct in vitro activity against M. tuberculosis (1), it has been shown to modify resistance by inhibiting mdr-1 efflux pumps and has shown to enhance killing of M. tuberculosis by macrophages (1). However, the activity of SILA 421 has only been tested against the M. tuberculosis H37Rv type strain and one clinical XDR M. tuberculosis isolate so far (1). We therefore assessed the antimycobacterial activity of SILA 421 against an extended set of M. tuberculosis strains with varying drug susceptibility profiles. Secondly, we compared antimycobacterial activity of SILA 421 with that of thioridazine and other phenothiazines. We selected 21 clinical *M. tuberculosis* isolates with varying drug susceptibility profiles: five isolates were pansusceptible, 11 were monoresistant (6 isoniazid monoresistant, 4 rifampin monoresistant and 1 streptomycin monoresistant) and 5 were MDR M. tuberculosis isolates. All isolates were extracted from a -70 °C freezer collection and sub-cultured on Ogawa medium. Inoculum preparation was performed as described previously aiming at 2 x 10³ to 10 x 10³ CFU (2). To determine the Minimal Inhibitory Concentration (MIC) we performed drug-susceptibility testing using a well-standardized Middlebrook 7H10 agar dilution method as described previously (2) using the following drug concentrations: 1, 2, 4, 8, and 16 mg/L. All drugs were received as chemically pure powder. SILA 421 was provided by Dr. G. Hajos (Institute for Biomolecular Chemistry, Budapest, Hungary). Chlorpromazine, desipramine, promazine and thioridazine were obtained from Sigma-Aldrich (Zwijndrecht, the Netherlands). Because the thioridazine S-enantiomer might induce less side-effects (4) we

- studied both thioridazine enantiomers separately. The two thioridazine enantiomers were
- prepared by Dr. J.B. Christensen (Dept. Chemistry, Copenhagen, Denmark). All drugs were
- 78 dissolved in distilled water to a stock solution of 10 g/L
- 79 The MIC of SILA 421 could be determined after a median of 14 days (range 14-18 days). The
- 80 MIC₅₀ was 4 mg/L and the MIC₉₀ was 8 mg/L. Table 1 shows the comparison of the MIC₅₀
- and MIC₉₀ of SILA 421 with the various phenothiazines. The MICs found for the
- phenothiazines were comparable with those found in the literature (3). The MIC_{50} of SILA
- 421 was similar to the MIC₅₀ of thioridazine and chlorpromazine. On the contrary, MIC₅₀ of
- both desipramine and promazine were both much higher than the MIC₅₀ of SILA 421. When
- comparing the MIC₉₀, lowest MIC₉₀ was measured for SILA 421.
- These data show that the SILA 421 is active in vitro against *M. tuberculosis* isolates with a
- wide variety of drug susceptibility patterns, thereby providing incremental evidence of its
- antimycobacterial activity. Moreover, we showed that SILA 421 is equally active in vitro as
- 89 thioridazine, another well-known efflux pump inhibitor.
- In a previous study, SILA 421 has been proposed to be more potent that thioridazine (1).
- However, only two M. tuberculosis isolates had been tested (1). The present study confirms
- 92 that SILA 421 may be a promising antituberculosis drug with activity against a wide variety
- 93 of M. tuberculosis strains comparable to that of thioridazine. Further studies are warranted to
- investigate the potency of SILA 421 as an antituberculosis drug in animals and in humans.
- 95 Besides its antimycobacterial activity, SILA 421 might also have another appealing feature.
- Like all efflux pump inhibitors, SILA 421 will inhibit the activity of efflux pumps of MDR
- 97 mycobacteria, presumptively rendering mycobacteria more susceptible to the antituberculosis
- 98 drugs to which it was initially resistant as a consequence of their extrusion from the cell (5).
- 99 Thus, these drug resistance modifiers might be of interest not only for their immediate

antimycobacterial activity but also as an adjunctive agent to be added to new or existing antituberculosis regimens. Of additional interest is our finding that the S-enantiomer of thioridazine is as effective as thioridazine itself against M. tuberculosis. It is thought this S-enantiomer might induce less neurological side effects owing to a lower affinity to the D2 dopamine receptor (4), putatively favoring the S-enantiomer over the racemate as an antituberculosis drug. However, it is unknown if the D2 dopamine receptor is the only target for neurological side-effects of thioridazine and it is unclear whether the pharmacokinetic and pharmacodynamic properties of the thioridazine enantiomers are equal. Notwithstanding these limitations, our findings raise hope that structural optimization of thioridazine-derivatives is possible without losing antimycobacterial activity (3.4). In conclusion, the present study provides confirmatory and incremental evidence of the antimycobacterial activity of SILA 421 and the phenothiazines, such as thioridazine. Therapeutic opportunities lay in the structural optimization of these drugs as well as in their drug resistance modifying effect. Future studies should bring SILA 421 and the phenothiazines further up the drug development pipeline.

100

101

102

103

104

105

106

107

108

109

110

111

112

113

114

115

117		
118	1.	Martins M, Viveiros M, Ramos J, Couto I, Molnar J, Boeree M, et al. SILA 421, an
119		inhibitor of efflux pumps of cancer cells, enhances the killing of intracellular extensively
120		drug-resistant tuberculosis (XDR-TB). Int. J. Antimicrob. Agents. 2009;33:479-482.
121		
122	2.	Van Klingeren B, Dessens-Kroon M, Van der Laan T, Kremer K, Van Soolingen D.
123		Drug susceptibility testing of Mycobacterium tuberculosis complex by use of a high-
124		throughput, reproducible, absolute concentration method. J. Clin. Microbiol.
125		2007;45:2662-2668.
126		
127	3.	Amaral L, Viveiros M. Why thioridazine in combination with antibiotics cures
128		extensively drug-resistant Mycobacterium tuberculosis infections. Int. J. Antimicrob.
129		Agents 2012;39:376-380.
130		
131	4.	Thanacoody HK. Thioridazine: resurrection as an antimicrobial agent? Br. J. Clin.
132		Pharmacol. 2007;64:566-574.
133		
134	5.	Amaral L, Viveiros M. Why thioridazine in combination with antibiotics cures
135		extensively durg-resistant Mycobacterium tuberculosis infections. Int J Antimicrob
136		Agents 2012;39:376-80
137		
138		

References

139140 **Tables**

141

147

142 Table 1

The MIC₅₀ and MIC₉₀ of SILA 421 and the various phenothiazines^(a).

Efflux pump inhibitor	MIC ₅₀ (mg/L)	MIC ₉₀ (mg/L)	Range (mg/L)
SILA 421	4	8	2 - 16
Thioridazine	4	16	2 - 16
Thioridazine, S-enantiomer	8	16	4 - 16
Thioridazine, <i>R</i> -enantiomer	8	16	4 - 16
Chlorpromazine	4	16	<1 - 16
Desipramine ^(b)	>16	>16	16 - >16
Promazine ^(b)	>16	>16	16 - >16

 ⁽a) MIC₅₀, MIC at which ≥ 50% of isolates are inhibited. MIC₉₀, MIC at which ≥ 90% of isolates are inhibited.
145 isolates are inhibited.

146 (b) MIC distribution was based on 20 instead of 21 isolates.