



Synthesis of novel [1,2]-diamines with antituberculosis activity

Qingyi Meng^a, Huibing Luo^{b,*}, Yilang Chen^b, Tiancai Wang^a, Qizheng Yao^{a,*}

^a Department of Medicinal Chemistry, China Pharmaceutical University, Nanjing 210009, China

^b Shanghai Sun-sail Pharmaceutical Science and Technology Company, Shanghai 201203, China

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ABSTRACT

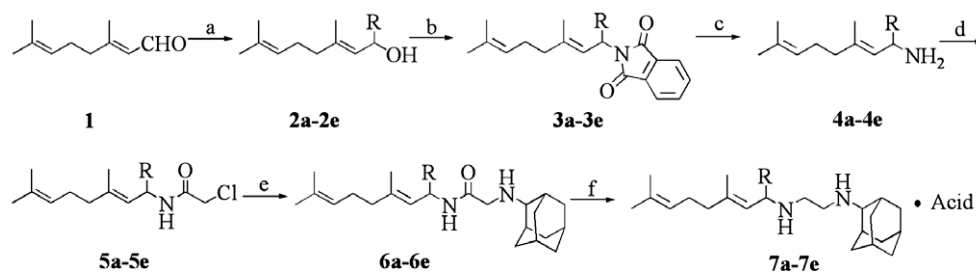
Guided by the metabolism information of **SQ109**, derivatives with substituted geranylamine moiety or substituted adamantane ring of **SQ109** were synthesized and evaluated as antituberculosis agents. Among all tested compounds, compound **11c** showed the most potent antituberculosis activity with MIC value of 0.3 μ M against *Mycobacterium tuberculosis* H37Rv.

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Despite a 5000 year history, tuberculosis (TB) remains the leading single-agent infectious disease killer in the world. Almost a third of the world's population is infected with TB bacilli, and each year approximately 8 million people develop active TB and 2 million die as a result.¹ The major challenges for tuberculosis control are the development of multidrug-resistant tuberculosis (MDR-TB) strains and the increasing numbers of immunocompromised individuals with HIV infectious who are highly susceptible to the disease. Consequently, there is a pressing need for new antituber-

cular agents acting with greater potency and efficacy than the current existing drugs. No novel antituberculosis drugs have been introduced into clinical practice over the past 4 decades. Only within the last few years have several promising drug candidates emerged.^{2,3}

N-Geranyl-*N'*-(2-adamantyl)ethane-1,2-diamine (**SQ109**), a second-generation agent from the first-line drug ethambutol, entered clinical trial stage in 2005 and completed its phase Ia trial in 2007.^{4–6} **SQ109** exhibited potent activity against *Mycobacterium*

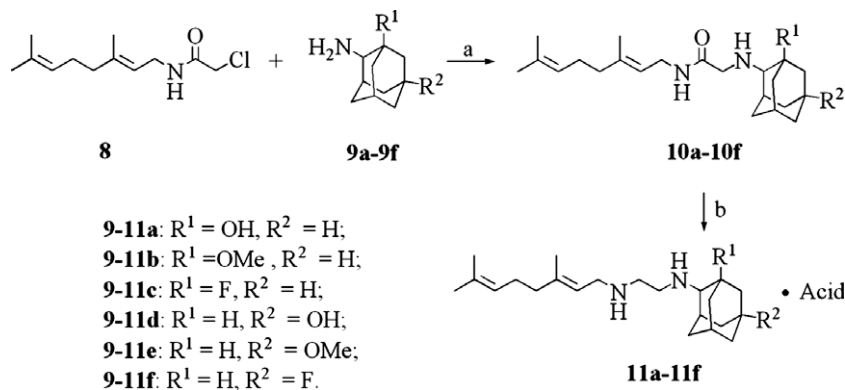


1-7a: R = methyl; **1-7b**: R = Et; **1-7c**: R = benzyl;
1-7d: R = 4-fluoro-benzyl; **1-7e**: R = 4-trifluoromethyl-benzyl

Scheme 1. Reagents and conditions: (a) RMgBr, THF, rt, 1 h; (b) Ph₃P, DIAD, phthalimide, THF, 8 h; (c) MeNH₂, MeOH, reflux, 12 h; (d) chloroacetyl chloride, pyridine, THF, 0 °C, 0.5 h; (e) 2-adamantylamine, K₂CO₃, THF, reflux, 12 h; (f) Red-Al, THF, reflux, 16 h, then HCl or maleic acid, MeOH.

* Corresponding authors. Tel.: +86 21 80278492; fax: +86 21 50278493 (H.L.), tel.: +86 25 83271445; fax: +86 25 86634730 (Q.Y.).

E-mail addresses: hb_luo@yahoo.com.cn (H. Luo), qz_yao@yahoo.com.cn (Q. Yao).



Scheme 2. Reagents and conditions: (a) K₂CO₃, THF, reflux, 12 h; (f) Red-Al, THF, reflux, 16 h, then HCl or maleic acid, MeOH.

tuberculosis including multi-drug resistant strains both in vitro and in vivo.⁷ Unfortunately, **SQ109** showed poor oral bioavailability, only 12% in rats and 3.8–5% in dogs, resulting from serious first-pass effect in liver. The metabolism study of **SQ109** by Jia et al. in human liver microsomes suggested that the predominant metabolisms of **SQ109** were oxidation and epoxidation of geranyl-

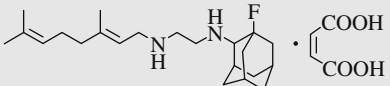
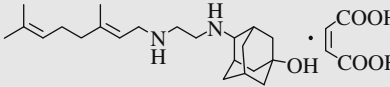
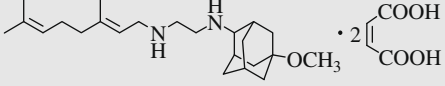
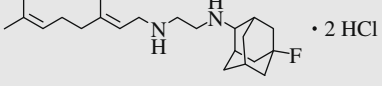
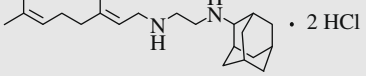
amine moiety and adamantane ring.⁸ Guided by the metabolism information of **SQ109**, derivatives of **SQ109** with substituted geranylamine moiety and adamantane ring were investigated to improve anti-tuberculosis activity and pharmacokinetic property. Herein, we described the synthesis and biological activity of these novel agents.

Table 1
The structures of synthesized compounds and their MICs against *M. tuberculosis* H37Rv

Compound	Structure	MIC (μM)
7a		1.2
7b		1.2
7c		1.0
7d		3.0
7e		5.2
11a		0.6
11b		0.5

(continued on next page)

Table 1 (continued)

Compound	Structure	MIC (μM)
11c		0.3
11d		8.6
11e		1.7
11f		0.6
SQ109		0.6

The first objective of this investigation was to immediately synthesize a variety of derivatives with alkyl attached geranylamine moiety or substituted adamantylamine moiety. Synthesis of compounds **7a–e** were outlined in Scheme 1. The starting material (*E*)-3,7-dimethylocta-2,6-dienal **1** was converted to **2a–e** by treatment with corresponding Grignard reagents in yields 93–97%. And these alcohols were treated with phthalimide in the presence of Ph_3P and DIAD to give intermediates **3a–e**, which were converted to corresponding amines **4a–e** by treatment with methylamine. The yields of these two steps from **2a–e** to **4a–e** were 37–49%. Chloroacylation of **4a–e** using chloroacetyl chloride and pyridine in THF gave **5a–e** in 96–98% yield. Reaction of 2-admantylamines with **5a–e** afforded intermediates **6a–e** with yields of 64–73%. Finally, **6a–e** were treated with Red-Al in THF to provide the 1,2-diamines, and these diamines were treated with maleic acid or hydrochloride to afford target compounds **7a–e** in 52–63% yield. Derivatives with substituted adamantane ring were synthesized as Scheme 2. The starting material **8** was synthesized according to the method described by Lee et al.⁴ and it was converted to **10a–f** by treatment with corresponding substituted adamantylamines **9a–f** and K_2CO_3 in THF at reflux temperature with yields of 67–75%. Then **10a–f** were treated with Red-Al in THF to provide the 1,2-diamines, and these diamines were treated with maleic acid or hydrochloride to give target compounds **11a–f** in 55–67% yield. The chemical structures of compounds **7a–e** and **11a–f** were determined by ^1H NMR and HR-MS.⁹

All of the synthesized compounds **7a–e** and **11a–e** were evaluated for tuberculosis inhibition against *M. tuberculosis* H37Rv strain (ATCC 27294, susceptible both to rifampin and isoniazid) using microbroth dilution assay described by Lee et al.⁴ and results are summarized in Table 1.

Our initial work focused on exploring variants to the geranylamine moiety of SQ109. As shown in Table 1, by introducing an alkyl group, such as methyl, ethyl, to the geranylamine moiety of SQ109, compounds **7a** and **7b** displayed similar potency against *M. tuberculosis* H37Rv with the 1.2 μM compared with 0.6 μM of SQ109. However when substituted benzyl was attached, for exam-

ple compound **7d** and **7e**, the antituberculosis potency reduced about 5–9 fold. It was interesting that removal of the substituted group of benzyl (**7c** vs **7e**) led to a fivefold increase in antituberculosis potency.

To further improve the antituberculosis potency, we also investigated the requirement for the 2-admantylamine moiety of SQ109. Guided by the metabolism information of SQ109, we attached the hydroxyl, halide, methoxyl group to the 1 and 5 position of the 2-admantylamine nucleus. As shown in Table 1, for the 5-substituted 2-admantylamine, 5-fluoro-2-admantylamine derivative **11f** showed MIC value of 0.6 μM , the same potency to the SQ109, whereas 14 fold increase in antituberculosis potency over 5-hydroxyl-2-admantylamine derivative **11d**. For the 1-substituted 2-admantylamine, as exemplified by compound **11c**, no significant change was observed compared to SQ109, MIC values of those compounds ranged from 0.3 to 0.6 μM . As far as the substitution of 2-admantylamine nucleus was concerned the fluorine appears to be the most favorable group. Among all tested compounds, compound **11c** showed the most potent antituberculosis activity with MIC value of 0.3 μM against *M. tuberculosis* H37Rv strain.

In summary, guided by the metabolism information of SQ109, derivatives of SQ109 with substituted geranylamine moiety or substituted adamantane ring were synthesized and evaluated as an anti-tuberculosis agent. Among all tested compounds, compound **11c** showed the most potent antituberculosis activity with MIC value of 0.3 μM against *M. tuberculosis* H37Rv.

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9. *Selected data: Compound 7a*: mp 194–195 °C; HRESIMS 345.3264, C₂₃H₄₁N₂ ([M+H]⁺ of free base) requires 345.3270; ¹H NMR (DMSO-*d*₆ 400 MHz) δ: 1.30 (d, *J* = 6.5 Hz, 3 H), 1.51 (s, 1H), 1.55 (s, 5H), 1.63 (s, 4H), 1.67 (s, 6H), 1.73 (s, 2H), 1.78–1.88 (m, 5H), 1.96–2.08 (m, 4H), 2.17 (s, 4H), 2.21 (s, 1H), 3.96–4.04 (m, 1H), 5.02–5.08 (m, 1H), 5.16 (d, *J* = 9.8 Hz, 1H), 9.38 (br s, 2H), 9.74 (br s, 2H). *Compound 7c*: mp 62–64 °C; HRESIMS 421.3591, C₂₉H₄₅N₂ ([M+H]⁺ of free base) requires 421.3583; ¹H NMR (DMSO-*d*₆ 400 MHz) δ: 1.03 (d, *J* = 6.23 Hz, 1H), 1.22 (s, 2H), 1.53–1.62 (m, 7H), 1.68–1.78 (m, 4H), 1.82–1.97 (m, 8H), 2.11–2.20 (m, 4H), 2.76 (t, *J* = 11.6 Hz, 1H), 3.30–3.42 (m, 6H), 4.18 (m, 1H), 4.95 (m, 1H), 5.12 (d, *J* = 10.2 Hz, 1H), 7.20–7.30 (m, 5H), 9.30 (br s, 1H), 9.85 (br s, 1H). *Compound 11b*: mp 109–111 °C; HRESIMS 361.3212, C₂₃H₄₁N₂O ([M+H]⁺ of free base) requires 361.3219; ¹H NMR (DMSO-*d*₆ 400 MHz) δ: 1.36–1.38 (m, 1 H), 1.48–1.58 (m, 5H), 1.58–1.64 (m, 5H), 1.65–1.73 (m, 5H), 1.96–2.12 (m, 7H), 2.90 (br s, 3H), 3.05 (s, 2H), 3.10 (s, 3H), 3.59 (d, *J* = 7.3 Hz, 2H), 5.04–5.07 (m, 1H), 5.17–5.22 (m, 1H), 6.02 (s, 2H). *Compound 11c*: mp 109–110 °C; HRESIMS 349.3024, C₂₂H₃₈N₂F ([M+H]⁺ of free base) requires 349.3019; ¹H NMR (DMSO-*d*₆ 400 MHz) δ: 1.28 (d, *J* = 10.6 Hz, 2H), 1.56–1.71 (m, 17H), 1.80–1.88 (m, 3H), 2.34 (s, 1H), 2.81–2.83 (m, 3H), 2.97 (t, *J* = 6.7 Hz, 2H), 3.60 (d, *J* = 7.3 Hz, 2H), 5.07–5.09 (m, 1H), 5.20–5.24 (m, 1H), 6.04 (s, 2H).