ARTICLE IN PRESS

Bioorganic & Medicinal Chemistry Letters xxx (2018) xxx-xxx



Contents lists available at ScienceDirect

Bioorganic & Medicinal Chemistry Letters

journal homepage: www.elsevier.com/locate/bmcl



Thioether-bridged arylalkyl-linked *N*-phenylpyrazole derivatives: Design, synthesis, insecticidal activities, structure-activity relationship and molecular-modeling studies

Chengcheng Fei^a, Yanfei Chen^a, Zhiyan Jiang, Dingxin Jiang*

Key Laboratory of Natural Pesticide and Chemical Biology, Ministry of Education, Laboratory of Insect Toxicology, South China Agricultural University, Guangzhou 510642, PR China

ARTICLE INFO

Article history: Received 9 December 2017 Revised 4 April 2018 Accepted 11 April 2018 Available online xxxx

Keywords: N-Phenylpyrazole derivatives Thioether bridge Insecticidal activities Structure-activity relationship Molecular-modeling studies

ABSTRACT

Owing to thioether diverse physicochemical properties by non-covalent interactions with bio-macromolecules, thioether derivatives containing heterocyclic moiety are known for their interesting insecticidal bioactivities and attracting considerable attention as neuroactive insecticides. Here we synthesis a series of novel thioether bridged N-phenylpyrazole derivatives incorporating various (hetero)aromatic substituents into 4-position of the pyrazole ring. Structure-activity relationship (SAR) studies resulted in compounds 6d and 7d with the most potent insecticidal activity among the series containing various substituted benzene substituents ($LC_{50} = 13.70-25.47 \mu g/g$). Further optimization to increase the lipophilicity and charge density of aromatic substituents of compounds 6d and 7d resulted in compounds 12d, 14d and 16d with sulfur-containing heterocycle substituents possessing good insecticidal activity against Musca domestica L. among the series ($LC_{50} = 0.67 - 1.30 \mu g/g$). The thioether bridge N-phenylpyrazole derivatives, which exhibit different length of the spacer arm introduced between N-phenylpyrazole moiety and the (hetero)aromatic substituents, were also prepared and evaluated. By contrast, the insecticidal activities of compounds containing the short thioether bridge, 1,2-bis((hetero)aromatic thio) ethane, are higher than that containing the long thioether bridge, 1,3-bis((hetero)aromatic thio) propane. The results of molecular docking and pharmacophore analyses indicated A299, T303, and L306 of a subunit were essential to form non-covalent interactions contacts with the ligands. Specially, the sulfur-containing heterocycle substituent derivatives 12d and 14d as the sterically favored areas could form the important hydrophobic interactions with the deeper residue P295.

© 2018 Elsevier Ltd. All rights reserved.

N-Phenylpyrazole moiety has been shown to bestow biological activity, including insecticidal, miticidal, and herbicidal activity. More specifically, *N*-phenylpyrazoles with a cyano or acetyl groups on 3-position, a sulfenyl, sulfinyl, thioalkyl, alkyl, acyl, alkynyl or cyano groups on 4-position, and an amino (or substituted amino) on 5-position of the pyrazole ring exhibit potent insecticidal activity.²⁻⁶ Fipronil, an *N*-phenylpyrazole insecticide with a trifluoromethylsulfinyl substituent at 4-position, has attracted particular attention of researchers due to their more favorable selective toxicity in invertebrates relative to mammals.⁷ The selective toxicity of fipronil is due in part to differ in binding between insect and mammalian GABA receptors⁸, but is also dependent on the relative rates of conversion to the less selective sulfone fipronil metabolite at 4-position of the pyrazole ring.⁹ The change

malian toxicity of *N*-phenylpyrazole insecticides was observed such as vaniliprole (Fig. 1A), pyrafluprole (Fig. 1B) and pyriprole (Fig. 1C) with 4-SCF₃, 4-SCH₂F and 4-SCF₂H instead of 4-SOCF₃ at 4-position of the pyrazole ring. However, fipronil with a trifluoromethylsulfinyl moiety probably exists in the cyclic hydrogenbonded form allowing interaction of the sulfinyl and amino moieties, and may be the first step in extrusion of SO from fipronil to give higher toxic and persistent desulfinylfipronil, then lead to more toxic and persistent detrifluoromethylsulfinyl product than parent fipronil by photolysis reactions. But the sulfide analogs of fipronil at 4-position of the pyrazole ring are much more stable and do not undergo an analogous photoextrusion reaction. In addition, the sulfur-aromatic interaction is often described as

improving the stability for modulation of protein interactions^{12,13}

in single substituent, i.e., thioalkyl vs sulfinyl (Fig. 1), alters the

lipophilicity and electronic properties and potentially also the photochemical and metabolic fate, effectiveness, and toxicology. ¹⁰ For

example, high insecticidal activity and selective toxicity, low mam-

https://doi.org/10.1016/j.bmcl.2018.04.022 0960-894X/© 2018 Elsevier Ltd. All rights reserved.

^{*} Corresponding author.

E-mail address: dxj2005@scau.edu.cn (D. Jiang).

^a The authors contributed equally to the work.

C. Fei et al./Bioorganic & Medicinal Chemistry Letters xxx (2018) xxx-xxx

Fig. 1. Structure of thioalkyl fipronil derivatives.

and one-third of all known protein structures contain an energetically stabilizing methionine-aromatic motif. ¹⁴ Considering fipronil (trifluoromethylsulfinyl phenylpyrazole) have higher insecticidal activity than sulfone fipronil (trifluoromethylsulfonyl phenylpyrazole) and intramolecular thioether-bridge formation is an effective way to protect compound against mixed function oxidases degradation *in vivo*, we speculated that thioalkyl fipronil derivatives may be increasing frequency near aromatic side-chains of γ -aminobutyric acid receptor proteins and altering bioavailability of drugs due to ligand(drug)-proteins interaction, and then improved insecticidal activity.

Meanwhile, after decades of intensive use, many target pests have developed higher resistance to fipronil, and owing in part to its high toxicity to beneficial organisms^{16,17}, fipronil was greatly limited to be used as a pesticide in China since 2009 and the European Union followed suit in 2013.¹⁸ To reduce resistance and toxicological risk, we developed several fipronil derivatives by modifying the amino group at 3-position of the pyrazole ring with salicylide, substituted phenoxyacetyl, amino acid and inner salt groups.^{19–24}

Inspired by these reports, herein we report a series of fipronil derivatives containing an arylalkyl thioether moiety at 4-position of the pyrazole ring (Scheme 1). In addition, the binding site of meta-diamides was demonstrated to be distinct from that of conventional noncompetitive antagonists such as fipronil. Thus, it is expected to become a prominent insecticide against pests with resistance to cyclodienes and fipronil. So, different modes of actions of novel fipronil derivatives containing an arylalkyl thioether moiety at 4-position of the pyrazole ring may have appeared and it is expected to be effective against resistant pest insects.

The bioactivities of the target compounds were evaluated, the structure-activity relationships and molecular docking studies of these compounds are shown in this paper.

The target thioalkyl fipronil derivatives were prepared from 5amino-3-cyano-1-(2,6-dichloro-4-trifluoromethylphenyl)pyrazole (b) as shown in Scheme 1. The thiophenol or aromatic thiol was reacted with dibromoalkane in the presence of potassium carbonate and potassium iodide to afford the intermediates 1a-29a. In addition, 1H-benzo[d]imidazole-2-thiol, 2-methoxybenzenethiol, 1H-1,2,4-triazole-3-thiol, thiazole-2-thiol and 1,3-dibromopropane; 4-methoxybenzenethiol and 1,2-dibromoethane were also tried but failed to give target products under similar conditions. We have tried to react with a more laborious reaction pathway, unfortunately the reaction was unsuccessful due to the complex formation between substrate and the catalyst. Compound **b** was converted to the key intermediate, bis(5-amino-3-cyano-1-(2,6-dichloro-4-trifluoromethylphenyl)pyrazol-4-yl) disulfide (\mathbf{c}), according to the reported method.³ The cleavage of disulfide \mathbf{c} by intermediates 1a-29a assisted with an alkali and a reducing reagent afforded the target thioalkyl fipronil derivatives **1d–29d**. 15

According to experimental result, the aromatic halides such as phenyl bromide could not react with pyrazolyl disulfide. We

add a spacer linker between the *N*-phenylpyrazole moiety and thiophenol/aromatic thiol. The structure notably included a phenylpyrazole ring connected to the aromatic moiety via two thioether bond. Using the cleavage of S–S bond chemical reaction by intermediates **1a–29a**, the target fipronil derivatives were synthesized easily and quickly in good yields.

The structures of the synthetic compounds were confirmed by melting points, ¹H NMR, ¹³C NMR and the structures of the title compounds **1d–29d** were confirmed by HRMS spectroscopic data additionally.

The preliminary insecticidal activities of compounds **1d–29d** were assessed against *Musca domestica* L. by the artificial diet dipping methods as the final mortality rates at 20 μg/g and fipronil as positive controls. The mortalities of *Musca domestica* L. were shown in **Tables 1** and **2**. Among all the tested compounds, compounds **12d**, **13d**, **14d**, **16d** and fipronil showed potent insecticidal activity (96.67%–100%); compounds **c**, **2d**, **3d**, **6d**, **7d** and **29d** showed moderate insecticidal activity (40.00%–53.33%). Insecticidal activities of other compounds were very low (<36.67%).

To gain further insight into potent toxicity of these compounds, fipronil and the ten compounds **2d**, **3d**, **6d**, **7d**, **12d**, **13d**, **14d**, **16d**, **29d** and **c** were investigated further at serial concentration gradient to determine their LC₅₀s (Table 3). The bioactivities of compounds **12d**, **14d** and **16d** being equipotent to fipronil against *Musca domestica* L. The LC₅₀ values of fipronil and compounds **12d**, **14d** and **16d** were 0.68, 0.67, 0.90 and 1.30 μ g/g, respectively. Comparable to fipronil, compound **13d** dramatically reduced the insecticidal activity (LC₅₀ = 6.00 μ g/g). Compounds **2d**, **3d**, **6d**, **7d**, **29d** and **c** displayed inferior activity as well, their LC₅₀s were 22.79, 22.47, 13.70, 25.47, 22.00 and 19.07 μ g/g, respectively.

Initially, in order to find the optimal aromatic substitution, a series of thioether N-phenylpyrazole derivatives substituted with naphthalene, 2-methoxybenzene, 4-methoxybenzene, 2-fluorobenzene, 4-fluorobenzene, 2-chlorobenzene, 4-chlorobenzene, 2-bromobenzene, 4-bromobenzene were prepared and evaluated. Insecticidal activity data showed that derivatives containing the short thioether bridge, 1,2-bis((hetero)aromatic thio) ethane, far better than that of the long thioether bridge, 1,3-bis((hetero)aromatic thio) propane (4d–11d vs 19d–26d). In particular, fluoro substituted benzene derivatives 6d and 7d possessed the most potent bioactivities, with the LC_{50} s of 13.70 μ g/g and 25.47 μ g/g, respectively. Inspired by the potent activity possessed by the fluoro substituted benzene derivatives, further research on heteroaromatic motif with strong lipophilicity and charge density were carried out.

Based on a rationally conceived pharmacophore model to increase the lipophilicity and charge density of aromatic substituents, nitrogen, oxygen, sulfur-containing heterocycle substituents were introduced to explore the SAR (Tables 2 and 3). The target compounds of nitrogen, oxygen-containing heterocycle substituents (except compound 13d) showed much less potency compared to fipronil against *Musca domestica* L. The target compounds of sulfur-containing heterocycle substituents displayed

Download English Version:

https://daneshyari.com/en/article/7778735

Download Persian Version:

https://daneshyari.com/article/7778735

Daneshyari.com