Structure–activity Relationship Analysis for Antimicrobial Activities of Tryptanthrin Derivatives Using Quantum Chemical Calculations

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Tryptanthrin (T) and eight of its monosubstituted derivatives (\textbf{T2NH}, \textbf{T2Cl}, \textbf{T2Br}, \textbf{T2NO}, \textbf{T8OMe}, \textbf{T8Me}, \textbf{T8F}, and \textbf{T8Br}) were synthesized, and their antimicrobial activities were investigated against a fungus (\textit{Malassezia furfur}) and a gram-positive bacterium (methicillin-resistant staphylococcus aureus, MRSA). Antimicrobial activities of these derivatives were influenced by the substituents on tryptanthrin, with the halogen-substituted tryptanthrin derivatives (\textbf{T2Cl}, \textbf{T2Br}, \textbf{T8F}, and \textbf{T8Br}) showing the highest potency against \textit{M. furfur} and MRSA. Therefore, semiempirical molecular orbital calculations (PM3) were performed on \textbf{T} and its eight derivatives to investigate the cause of the differences in their antimicrobial activities. The results of the calculations showed that antimicrobial activities could be related to the electrophilicity of the carbonyl carbon of the five-membered ring.

**Keywords:** PM3, Antimicrobial activity, Tryptanthrin, MRSA, \textit{M. furfur}

1 Introduction

Tryptanthrin (T, Figure 1) is a weakly basic alkaloid found in a number of plant species [1]. This compound exhibits antimicrobial activity against various pathogenic bacteria and fungi [2]. In particular, the antifungal activity of tryptanthrin against \textit{Malassezia furfur}, which is the causative fungus of atopic dermatitis, is significant [3]. Tryptanthrin is also effective in the treatment of contact dermatitis (delayed-type allergy) [4]. Therefore, its use as a therapeutic drug for conditions such as atopic dermatitis or in cosmetics is anticipated. However, few reports describe the properties of chemosynthetic tryptanthrin derivatives, which are not found in nature. In particular, there are few studies on the antifungal activities of tryptanthrin derivatives against \textit{M. furfur} [3]. Therefore, we synthesized eight different tryptanthrin derivatives (Figure 2) and investigated their antimicrobial properties against a fungus (\textit{M. furfur}) and a gram-positive bacterium (methicillin-resistant Staphylococcus aureus, MRSA).

The antifungal and antibacterial activities of the derivatives were influenced by tryptanthrin substituents, with halogen-substituted tryptanthrin derivatives (\textbf{T2Cl}, \textbf{T2Br}, \textbf{T8F}, and \textbf{T8Br}) showing the highest potency against \textit{M. furfur} and MRSA [5]. Subsequently, semiempirical molecular orbital calculations were performed on tryptanthrin derivatives to investigate the structure–activity relationships of the antimicrobial activities.

![Figure 1. Structure of tryptanthrin (T).](image-url)
2 Experiment

2.1 Synthesis

Material data and preparation of tryptanthrin (T) and eight of its derivatives (T2NH2, T2Cl, T2Br, T2NO2, T8OMe, T8Me, T8F, and T8Br) have been reported [5].

2.2 Antibacterial and antifungal activity test

The antifungal and antibacterial activities of the derivatives against a fungus (M. furfur) and a gram-positive bacterium (MRSA) were investigated in vitro, using the agar plate dilution method described by the Japanese Society of Chemotherapy [6–9]. A bacterial culture (developed over 2–3 days for M. furfur or overnight for MRSA) was diluted with malt extract broth (36 g/L in distilled water; Wako Pure Chemical Industries, Ltd.) containing Tween-40 (10 g/L in distilled water; Wako Pure Chemical Industries, Ltd.), monoolein (2 g/L in distilled water; Tokyo Chemical Industry Co., Ltd.), oxgall (20 g/L in distilled water; Wako Pure Chemical Industries, Ltd.), peptone (6 g/L in distilled water; Wako Pure Chemical Industries, Ltd.), and glycerol (2 g/L in distilled water; Wako Pure Chemical Industries, Ltd.) for M. furfur and with Mueller–Hinton broth (21 g/L in distilled water; Becton, Dickinson and Co.) for MRSA to a density of 1.0 × 10^6 colony-forming units (CFU)/mL. The compounds for testing (T, T2NH2, T2Cl, T2Br, T2NO2, T8OMe, T8Me, T8F, and T8Br) were dissolved in DMSO (Wako Pure Chemical Industries, Ltd.) and diluted with malt extract broth containing Tween-40, monoolein, oxgall, peptone, and glycerol to a concentration of 4–160 µg/mL for M. furfur and with Mueller–Hinton broth to a concentration of 0.1–100 µg/mL for MRSA. Thereafter, each Petri dish was inoculated with the fungal or bacterial suspension and incubated at 37 °C for 2–4 days for M. furfur and 24 h for MRSA. The lowest concentration at which there was no visible growth was considered as the minimum inhibitory concentration (MIC).

2.3 Calculation procedure

The molecular modeling study was performed using the SPARTAN ‘10 software package (Wavefunction Inc., Irvine, CA, 2000). The minimum energy conformation of the tryptanthrin derivatives was obtained by the MMFF of the molecular mechanics calculation. We evaluated the electronic properties of the MMFF minimum energy conformation by using semiempirical molecular orbital calculations (PM3). To study the structure–activity relationships, we evaluated electronic properties of the lowest unoccupied molecular orbital (LUMO), the LUMO map, and the electrostatic potential map.

3 Results and discussion

3.1 Antifungal and antibacterial activities of Tryptanthrin and its eight derivatives

The antibacterial activities of tryptanthrin (T) and its derivatives (T2NH2, T2Cl, T2Br, T2NO2, T8OMe, T8Me, T8F, and T8Br) were investigated on a fungus (M. furfur) and a gram-positive bacterium (MRSA) in culture. MICs for M. furfur and MRSA are shown in Table 1.

<table>
<thead>
<tr>
<th></th>
<th>MIC/µg mL⁻¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>M. furfur</td>
<td>MRSA</td>
</tr>
<tr>
<td>T</td>
<td>4</td>
</tr>
<tr>
<td>T2NH2</td>
<td>20</td>
</tr>
<tr>
<td>T2Cl</td>
<td>4</td>
</tr>
<tr>
<td>T2Br</td>
<td>4</td>
</tr>
<tr>
<td>T2NO2</td>
<td>&gt;160</td>
</tr>
<tr>
<td>T8OMe</td>
<td>&gt;120</td>
</tr>
<tr>
<td>T8Me</td>
<td>&gt;120</td>
</tr>
<tr>
<td>T8F</td>
<td>1</td>
</tr>
<tr>
<td>T8Br</td>
<td>2.5</td>
</tr>
</tbody>
</table>

MIC=minimum inhibitory concentration in µg/mL.

The antifungal and antibacterial activities are enhanced for small MIC values. The MIC at which a molecule exerted an antimicrobial activity was defined as less than 800 µg/mL in accordance with the criterion specified by the Japanese Society of Chemotherapy [7–9]. Most tryptanthrin derivatives showed high antifungal (M. furfur) and antibacterial activities (MRSA). The antibacterial activity for MRSA was higher than the antifungal activity for M. furfur. The higher potency of the derivatives

![Figure 2. Structures of the eight tryptanthrin derivatives.](image-url)
against the gram-positive bacterium (MRSA) compared with
their potency against the fungus (M. furfur) can be attributed to
the structural differences between the two species. The MIC
order for M. furfur was T8F < T8Br < T2Cl = T2Br = T < T2NH2 <
T8OMe, T8Me, T2NO2, while for MRSA it was T8F = T8Br =
T2Cl < T2Br = T < T8Me < T2NH2 < T8OMe < T2NO2. For both
M. furfur and MRSA, the antifungal and antibacterial activities
of the halogen-substituted tryptanthrin derivatives were higher
when compared to the other tryptanthrin derivatives. Consider-
ing the overall efficacy, T8F was the most potent of all the
tested compounds with an MIC of 1 mg/mL for M. furfur and
0.1 mg/mL for MRSA. In contrast, the antibacterial and anti-
fungal activities of T2NO2, which has a nitro group at position
2 of tryptanthrin, were the lowest. The MIC of T2NO2 was
>160 µg/mL for M. furfur and >100 µg/mL for MRSA. Micon-
azole nitrate is currently used as a therapeutic drug for treating
atopic dermatitis. Its MIC for M. furfur, which is the causative
fungus of atopic dermatitis, is ca. 25 µg/mL. The MICs of T
and its halogen-substituted tryptanthrin derivatives T8F, T8Br,
T2Cl, T2Br, and T for M. furfur ranged from 1 to 4 µg/mL,
and their antifungal activities were more than six times that of
miconazole nitrate. T8F was found to be particularly effective.

3.2 Structure–activity relationships of
tryptanthrin and its eight derivatives

Molecular level information obtained from semiempirical
(PM3) quantum chemical calculations on the optimized elec-
tronic structure of compounds was used to explore the rela-
tionship between chemical structure and biological activity.
Structure–activity relationships are based on the fundamental
hypothesis that biological properties are functions of the mo-
lecular structure. It is reasonable to assume that certain elec-
trophilic or nucleophilic sites of a molecule can only bind to
nucleophilic or electrophilic sites of the receptor.

Previously, it was reported that the correlation between anti-
leishmanial activity of tryptanthrin derivatives and the lowest
unoccupied molecular orbital (LUMO) energy against the antifungal activity of tryptanthrin (r2 = 0.59)
indicate an excellent linear correlation (r2 = 0.71 for M. furfur;
and 0.59 for MRSA), the trend is apparent. Thus, the electron
affinity of the molecule could be an important factor associ-
ated with potent activity. The molecular properties of T, T8F
and T2NO2 retrieved by using semiempirical molecular orbital
calculations (PM3) are shown in Figure 5 as typical examples.
T8F was the most potent of all the tested compounds with an
MIC of 1 µg/mL for M. furfur and 0.1 µg/mL for MRSA. T also
showed antimicrobial activities with an MIC of 4 µg/mL for M. furfur and 0.5 µg/mL for MRSA, while T2NO2 did not show
antimicrobial activities against both M. furfur and MRSA. As
shown in Figure 5b, the shape of the LUMO of the carbonyl
carbon atom of the five-membered ring of T8F and T with high
antimicrobial activities is large. In contrast, that of T2NO2
with no antimicrobial activity is smaller, and the shape of the next
carbon atom of the carbonyl carbon atom of the five-membered
ring is larger than that of the carbonyl carbon atom of the five-
membered ring. In the LUMO map, the electron-poor sites are
indicated in blue, while the electron-rich sites are indicated

![Figure 3. The lowest unoccupied molecular orbital (LUMO) energy against the antifungal activity of tryptanthrin (r2 = 0.71).](image)

![Figure 4. The lowest unoccupied molecular orbital (LUMO) energy against the antibacterial activity of tryptanthrin (r2 = 0.59).](image)
in red. As shown in Figure 5c, the LUMO map of T8F and T shows intensely blue atoms over the carbonyl carbon atom, while that of T2NO₂ shows green atoms. The next carbon atom of the carbonyl carbon atom of the five-membered ring is intensely blue. The LUMO map shows one prominent site by the carbonyl group of the five-membered ring in tryptanthrin derivatives, the shape of the LUMO is found to be large and extended by the carbonyl carbon atom of the five-membered ring in the more potent analogs. The electrostatic potential map is an alternative approach for analyzing the electrostatic contribution to the binding of the receptor and drugs [11]. As shown in Figure 5d, the electrostatic potential map analysis revealed a lower electron density (blue color) over the carbonyl carbon atom of the five-membered ring of T8F and T than that observed in T2NO₂. Electron transfer from the receptor site to this carbonyl carbon seems a plausible path for the mechanism action of the compounds. According to the results of the theoretical calculations, the antifungal and antibacterial activities could be related to the electrophilicity of the carbonyl carbon of the five-membered ring.

4 Conclusion

Tryptanthrin (T) and its eight derivatives (T2NH₂, T2Cl, T2Br, T2NO₂, T8OMe, T8Me, T8F, and T8Br) were synthesized, and their antibacterial properties and antifungal properties against a fungus (M. furfur) and a gram-positive bacterium (MRSA) were investigated. The antifungal and antibacterial activities of the halogen-substituted tryptanthrin derivatives against M. furfur and MRSA were exceptional. Especially, T8F was the most potent of all the tested compounds with an MIC of 1 µg/mL for M. furfur and 0.1 µg/mL for MRSA. The semiempirical molecular orbital calculations (PM3) were performed on T and eight of its derivatives to investigate the cause of the differences in their antimicrobial activities. The results of the calculations showed that the antimicrobial activities could be related to the electrophilicity of the carbonyl carbon of the five-membered ring. The carbonyl groups of the five-membered rings in the tryptanthrin moiety and the electron transfer ability from a receptor could be a crucial step in the mechanism of action of these compounds.

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