Direct detection of diclofenac radical produced by ultraviolet irradiation using electron spin resonance method

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Diclofenac, a nonsteroidal anti-inflammatory drug, is commonly used as an antipyretic analgesic owing to its strong anti-inflammatory action in clinical treatment. However, diclofenac can cause injury, with gastrointestinal mucosal lesions and skin photosensitivity as the main side effects. In general, photosensitive drugs contain photosensitive chemical sites, and form free radicals under ultraviolet irradiation, leading to phototoxic reactions. Therefore, this study focuses on free radical production in photosensitive reactions of diclofenac. The free radical production mechanism of diclofenac under ultraviolet irradiation, which might result in photo-toxicity, was clarified using a direct electron spin resonance method. When diclofenac was irradiated with ultraviolet light (254 nm), diclofenac radicals were generated depending on the ultraviolet irradiation time and stably present for 30 min at room temperature. Diclofenac radicals were produced by the ultraviolet irradiation system depending on the dose of diclofenac until 2 mM. Therefore, diclofenac radicals might directly or indirectly react with various biomolecules to cause phototoxicity, other side effects, and new diclofenac pharmacology owing to its stability of diclofenac radicals.

Key Words: ESR, diclofenac (DCF), nonsteroidal anti-inflammatory drugs (NSAIDs), UV irradiation, phototoxicity, DCF radical

Free radicals such as superoxide (\(\text{O}_2^–\)), hydroxyl radical (‘OH), nitric oxide are considered to play important roles in various diseases, including acute lung injury, renal disorder with dialysis, and periodontal disease.\(^{(1–7)}\) As controlling inflammation in these pathological states is important, nonsteroidal anti-inflammatory drugs (NSAIDs) are frequently used as general therapeutic agents.\(^{(8,9)}\) Although much research on NSAIDs and radicals has been conducted, the detailed reaction mechanism of NSAIDs about radical has yet to be clarified.\(^{(10–17)}\)

Recently, NSAIDs, especially propionic acid derivatives, have shown phototoxicity.\(^{(18–20)}\) Some drugs used to treat human diseases are known to be activated under light irradiation to cause skin rashes, including redness, swelling, and pigmentation. Phototoxic reactions are known to be among the causes of photosensitivity in these drugs, in which agents activated by photochemical reactions cause damage to biochemical components directly or through reactive oxygen species (ROS).\(^{(18,19)}\) When a drug molecule absorbs photon energy, electrons are excited from the ground state to the excited state, depending on the bond type and associated energy level. As the electrons enter different orbitals through photoexcitation and electron pairs are eliminated, the excited electrons have radical properties. Energy transfer from excited drug molecules to oxygen (type-II photochemical reaction) generates singlet oxygen (\(\text{O}_2^+\)), which might participate in the oxidation of membrane lipids and proteins, or induce DNA damage.\(^{(20,21)}\) Furthermore, these excited drug molecules might react directly with \(\text{in vivo}\) molecules (such as DNA, proteins, and cell membranes) through electron or hydrogen transfer (type-I photochemical reaction).\(^{(20)}\) Excessive ROS production in the body, can result in the oxidation of nucleic acids, proteins, sugars, and lipids to cause various biological disorders.\(^{(22–24)}\)

Diclofenac (DCF) is an acetic acid derivative and NSAID that is often used clinically. The action of DCF is particularly strong among NSAIDs, and is characterized by the rapid suppression of pain and heat generation.\(^{(25–28)}\) In addition to a strong action, DCF has shown clinical side effects, such as causing gastrointestinal and kidney disorders, and phototoxicity.\(^{(26–28)}\) However, the detailed mechanism of DCF phototoxicity has yet to be clarified.

Therefore, this study aimed to clarify the reaction mechanism of DCF radical (‘DCF) production under ultraviolet (UV) irradiation, which can cause DCF phototoxicity, using a direct electron spin resonance (ESR) method.

Materials and Methods

Chemicals. DCF was purchased from Wako Pure Chemical Ind. (Tokyo, Japan). 1-Oxyl-2,2,6,6-tetramethyl-4-hydroxypiperidine (TEMPOL) was purchased from TOCRIS Bioscience (Ellisville, MO). Superoxide dismutase from bovine erythrocytes (SOD) was purchased from Wako Pure Chemical Ind. (Osaka, Japan). Water used in these experiments was treated to remove the trace metals by passing through Chelex 100 Resin (Bio-Rad Laboratories, Inc. Hercules, CA) after distillation.

Measurement of diclofenac radical (‘DCF) by direct ESR method. DCF was diluted with distilled water in a quartz flat cell (160 \(\mu\)l) and, irradiated with UV (254 nm) using a Handy UV Lamp (SUV-6) (AS ONE, Osaka, Japan). After UV irradiation, ESR spectra were immediately recorded at room temperature in a quartz flat cell using a JEOL JES-FR 30 EX Free Radical Monitor (JEOL, Tokyo, Japan). The operating conditions of the ESR spectrometer were as follows: frequency, 9.42 GHz; field, 335.618 ± 5 mT; microwave power, 16.0 mW; modulation frequency, 100 kHz; modulation width, 0.32 mT; amplitude, 7.9 × 100; time constant, 0.3 s; and sweep time, 1 min. To identify and determine the amount of radical species ‘DCF, the g value was corrected using the internal Mn marker of the ESR instrument used in this study. The ‘DCF concentration was calculated from a calibration curve prepared from the integrated value of the ESR signal of TEMPOL aqueous solution, for which the spin concentration is known.

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Statistical analysis. The statistical significance of the difference was determined by an unpaired Student’s t-test. Data are expressed as means ± SE. Differences between groups were considered statistically significant at the level of p < 0.05.

Result

ESR spectra and production mechanisms of •DCF. A prominent ESR spectrum of •DCF was observed after UV irradiation (254 nm) for 3 min in the absence of H₂O₂ (Fig. 1A–D). This represents the first report of •DCF detection using a direct ESR method at room temperature. •DCF increased until 2 mM DCF in the reactive mixture and then gradually decreased to 10 mM DCF in the reactive mixture (Fig. 1E). When DCF (0.1 mM) was irradiated with UV (254 nm), the intensity of •DCF radical generation increased with increasing UV irradiation time (Fig. 2). Increased the amount of •DCF production depending on UV irradiation time could be also observed with other concentrations (0.1, 0.2, 0.5, 1.0, 2.0, 5.0, and 10 mM) of DCF (data not shown). To evaluate the •DCF production mechanism, we attempted to reduce the amount of dissolved oxygen in the •DCF production system by bubbling with N₂. With reduced dissolved oxygen in the DCF solution (10 mM), the ESR peak of •DCF significantly decreased in intensity, but did not disappear (Fig. 3). These results indicated that •DCF was generated even when the oxygen concentration was low.

Time course of •DCF. To study the lifetime of •DCF, the amount of •DCF was measured after UV irradiation (254 nm) for 3 min. The •DCF peak height obtained by the direct ESR method gradually decreased, but did not disappear, for 30 min (Fig. 4). •DCF was found to be a longer life radical in this instance than O₂⁻ and •OH.

Fig. 1. ESR spectra was measured in various concentration of DCF irradiated with UV (254 nm) for 3 min. (A) Spectrum obtained in the reaction mixture of 0 mM DCF. (B) Spectrum obtained in the reaction mixture of 1 mM DCF. (C) Spectrum obtained in the reaction mixture of 2 mM DCF. (D) Spectrum obtained in the reaction mixture of 10 mM DCF. (E) •DCF production with dose-dependent manner of DCF. ESR measurement conditions were described in materials and methods. All spectra were recorded after DCF irradiation with UV (254 nm) for 3 min.

Fig. 2. •DCF production with time dependent manner of UV irradiation. ESR signal intensity of •DCF were obtained in the reaction system of DCF (0.1 mM), irradiated with UV (254 nm). ESR measurement condition were as described in materials and methods. Amounts of •DCF are expressed as a relative intensity (R.I.) by normalization of the •DCF signal height to the standard signal intensity of manganese oxide (MnO) and are the means ± SD of three independent experiments (n = 3). The waveform and R.I. of the ESR spectra from the •DCF spin adducts at each UV irradiation time was shown on the figure. Intensity: mean ± SD.
Reactivity of 'DCF with O$_2^-$'. To study the effect of O$_2^-$ on 'DCF' production, the amount of 'DCF' was measured after UV irradiation (254 nm) for 5 min. The relative intensity of 'DCF' measured using the direct ESR method increased with the addition of superoxide dismutase (SOD; 100 U/ml) (Fig. 5). This result confirmed that 'DCF' was more likely to be generated without O$_2^-$.

Characteristics of 'DCF'. The spin adducts of radical species 'DCF' were identified, and their amounts determined. A sharp single P$_1$ signal was obtained at a g value of 2.0038 when DCF (1 mM) was irradiated with UV light (254 nm) for 10 min (Fig. 6). These results showed that 'DCF' detected under UV irradiation was an oxygen radical. The amount of radicals at this time was determined to be $5.39 \times 10^7$ spins. The 'DCF' concentration in this system was calculated, to be 1 µM comparison with the spins of TEMPO radicals. The 'DCF' concentration was significantly higher, indicating that 'DCF' might cause tissue injury directly.

Discussion

In this study, we obtained direct evidence that DCF forms extremely stable 'DCF' under UV irradiation. This represents the first evidence of stable 'DCF' detection in a UV irradiation system using direct ESR method.

Kawaguchi et al. previously estimated that the photolysate of DCF was cyclized and radicalized at the center of the molecule by LC-NMR analysis, which was in agreement with our results. LC-NMR analysis provides detailed information on substance structure, but information on unpaired electrons is difficult to
obtain using this method. Therefore, in this study, we elucidated the mechanism of DCF formation using a direct ESR method.

Our results showed that DCF generation was dependent on the DCF concentration until 2 mM, but on the UV irradiation time, and that the radical species might be an alkoxyl radical or a nitrogen radical based on the g value (2.0038). In order to identify radical species, it need to be further examination of g value using more useful internal marker as well as of the hyperfine structure of the ESR spectrum with smaller modulation width.

Assuming that DCF is an alkoxyl radical, we showed that electron transfer from DCF to O2- hardly occurred because the amount of DCF production decreased with N2 bubbling [see equation (1)].

\[
\text{DCF} + \text{O}_2^- \rightarrow \text{DCF} + \text{O}_2^- \quad (1)
\]

Furthermore, we showed that electron transfer from DCF to O2- occurred because the amount of DCF produced increased with the addition of SOD [see equation (2)]. Besides, when ascorbic acid (ASA) as a scavenger of \('\text{OH}\) and \(\text{O}_2^-\) was added to the complete reaction mixture, ASA strongly suppressed DCF production. Therefore, it might be suggested that \('\text{OH}\) play important role in the production of \('\text{DCF}\) (data not shown).

\[
\text{DCF} + \text{O}_2^- + 2\text{H}^+ \rightarrow \text{DCF} + \text{H}_2\text{O}_2 \quad (2)
\]

Accordingly, DCF can be assumed to exist for a unusually long time compared with other radicals. Free radicals with unpaired electrons are known to be very unstable, for example, \(\text{O}_2^-\) and \('\text{OH}\) have half-lives of \(10^6\) and \(10^9\) s, respectively. In this study, the stability of DCF did not change, even at low \(\text{O}_2^-\) concentrations, suggesting that DCF would be present not only in blood vessels with relatively high \(\text{O}_2^-\) levels, but also in tissues with low \(\text{O}_2^-\) levels. Therefore, it is possible that DCF acts as a toxic radical, both in the periphery and in the blood vessels. In contrast, DCF, reducing radicals, scavenge oxidative radicals and act as proactive.

In any other possible knowledge, DCF is a secondary amine, which generally act as an electron donor. It is possible that the excited state of DCF may reduce molecular oxygen to produce the DCF radical cation and \(\text{O}_2^-\). The DCF radical cation may undergo deprotonation to produce \('\text{DCF}\) which is a nitrogen radical. In such a case, molecular oxygen act as an oxidant to generate \('\text{DCF}\). Nitrogen bubbling may decrease the oxidant, i.e., molecular oxygen, leading to the decrease of \('\text{DCF}\) production. SOD converts \(\text{O}_2^-\) to molecular oxygen and water. The oxidant (molecular oxygen) to produce \('\text{DCF}\) may be recycled in the presence of SOD, leading to the increase of \('\text{DCF}\) production. In future examination, the hyperfine structure of the ESR spectrum of \('\text{DCF}\) will provide detailed information about the electronic structure as well as production mechanism of \('\text{DCF}\).

DCF generation in vivo is still unknown, which indicate the need for further studies. We propose that DCF is generated in vivo because in a previous study by Miura37) and Muraoka38) showed that NSAIDs react with peroxidase to possibly generate NSAID radicals (NSAID). Previously, for both the intended and side effects of NSAID inhibition of cyclooxygenase (COX), the main mechanism has been considered to involve changes in the balance of eicosanoids.39-41) However, this report suggests that radical formation, not only by DCF, but also by other NSAIDs is needed for physiological effects other than COX inhibition.

In conclusion, DCF might generate via one-electron reduction of DCF in vivo or in human. Furthermore, the generated DCF might affect the various diseases where DCF is frequently used for treatment, even such as inflammation, cancer, or orthopedic disorders. Therefore, it was suggested that our detected DCF might have important roles in various diseases.

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Conflict of Interest

No potential conflicts of interest were disclosed.

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