Reduction of molecular oxygen by redox active thiols: comparison of glutathione, N-acetylcysteine, cysteine, and homocysteine

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Several thiol compounds are widely used as anti-oxidants in biological experiments. For example, cysteine (Cys) (Fig. 1A) is a relatively simple amino acid that has a thiol moiety. N-Acetyl-L-cysteine (NAC) (Fig. 1B) is a derivative of Cys acetylated on the amino moiety. The main endogenous antioxidant in living cells is the reduced form of glutathione (GSH) (Fig. 1C), which is a tripeptide molecule consisting of glutamic acid, Cys, and glycine. Although homocysteine (HCS) has an extra methylene bridge compared with the structure of cysteine (Fig. 1D), HCS may be an unorthodox thiol compound in a negative sense because it was reported to be an inducer of oxidative stress.1–3 Zhang et al.4 reported that the cerebrovascular effects of HCS were prevented by superoxide dismutase (SOD). HCS is biologically synthesized from methionine (Met), which is an essential amino acid for humans. Met has an extra methyl terminal on the structure of HCS (Fig. 1E).

Superoxide (O2•−) generation in an aqueous solution containing GSH at a hyperthermal temperature was reported.5,6 GSH can reduce oxygen (O2) to make hydroperoxyl radicals (HO2•), which equilibrate to O2•− in an aqueous environment. HO2• is a more efficient oxidant than O2•−, which is essentially a reductant. The GSH-induced HO2•/O2•− redox reaction at hyperthermal temperatures can cause cell death.5 The excellent biological reductant GSH can reduce oxygen to give ROS and cause cell toxicity at hyperthermal temperatures. The high reducing activity of antioxidants may cause oxidative stress.

This GSH-induced cell death at hyperthermal temperatures was amplified by the addition of catalase.5,6 HO2• molecules can react together to form hydrogen peroxide (H2O2). In addition, HO2• can be reduced by GSH to give H2O2. The simultaneously generated

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Fig. 1. Chemical structures of sulfur-containing compounds compared in this study. (A) Cys is a simple amino acid. (B) NAC is an acetylated derivative of Cys. (C) GSH is a tripeptide, consisting of glutamic acid, Cys, and glycine. (D) HCS is homologous to Cys. (E) Met is a methylated derivative of HCS.

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H₂O₂ by-product of GSH-induced HO₂-/O₂•− can absorb oxidative stress induced by HO₂-/O₂•− because it can consume O₂•−. Hydroxyl radicals (OH) may be generated by the reaction of H₂O₂ and O₂•−, whereas this extracellular single-shot OH may have no particular biological effect. As a result, the elimination of H₂O₂ by catalase may increase the oxidative stress induced in a HO₂-/O₂•− atmosphere. Anti-oxidative cooperation in an oxygenated atmosphere can cause further oxidative stress, contrary to expectation, which is termed reductive stress.

The stable nitroxy radical 4-hydroxy-2,2,6,6-tetramethylpiperidine-N-oxyl (TEMPOL) can be oxidized to an oxoammonium cation form by HO• and/or ‘OH.(7,34) The oxoammonium cations are reduced to hydroxylamine by accepting a hydrogen atom from hydrogen donors (H-donors), such as reduced GSH, reduced β-nicotinamide adenine dinucleotide (NADH), or reduced β-nicotinamide adenine dinucleotide phosphate (NADPH), in biological settings.(9,10) The oxoammonium cations can also react with GSH directly to make a binding complex.(11,12) TEMPOL can be reduced by two methods with coexisting GSH. Other thiol compounds are also expected to react with TEMPOL in two manners, similar to GSH. However, no confirmation of the reaction of other thiols with TEMPOL or the corresponding oxoammonium form has been reported.

Some biological thiol compounds, such as GSH and/or cysteine, exhibit unpredictable reactions under biological experimental conditions. There are several reports of oxidative stress by thiols, especially by GSH as an oxidant.(14–20) The purpose of this study was to find a suitable thiol-based anti-oxidant for biological experiments and to provide clear reasoning for its selection. In this study, the reaction properties of Cys, NAC, GSH, HCS, and Met were compared using TEMPOL as a redox probe. Furthermore, the availability of thiol compounds to cause reductive stress was discussed.

Materials and Methods

Chemicals. GSH, Cys, NAC, HCS, Met, and potassium ferricyanide were purchased from Wako Chemical (Tokyo, Japan). TEMPOL, SOD from human erythrocytes, hypoxanthine, xanthine oxidase, NADH, and NADPH were purchased from Sigma-Aldrich (St. Louis, MO). Other chemicals used were of analytical grade. As basic solvents for the reaction mixtures, 100 mM phosphate buffer (PB) containing 0.05 mM DTPA adjusted to pH 7.4 was prepared. Deionized water (deionization by the Milli-Q system) was used to prepare PB.

Reaction of TEMPOL with coexisting thiol compounds in the hypoxanthine-xanthine oxidase (HX-XO) reaction system. A reaction mixture containing xanthine oxidase, TEMPOL, and one of the test compounds was prepared using PB. Then, an aliquot of hypoxanthine solution was added to the reaction mixture to start the reaction. The final concentrations of hypoxanthine, xanthine oxidase, TEMPOL, and the test compound were 0.05 mM, 0.01 U/ml, 0.1 mM, and 1.0 mM, respectively. The reaction mixture was drawn into a quartz flat cell, and then fixed in the TE-mode cavity. The time course of the EPR signal intensity of TEMPOL was measured using an X-band EPR spectrometer (JEOL, Tokyo) after X-ray irradiation doses of 0, 2, 4, 8, 16, or 32 Gy.

Reaction of TEMPOL with coexisting thiol compounds in a ferricyanide reaction system. A reaction mixture containing 0.1 mM TEMPOL and 1.0 mM test compound was prepared using PB. Then, a 1/100 volume of 200 mM potassium ferricyanide, K₃[Fe(CN)₆], was added to an aliquoted volume of the reaction mixture to start the reaction. The time course of the EPR signal intensity of TEMPOL was measured using an X-band EPR spectrometer. The EPR conditions were as described above. The same experiment was performed adding 20 U/ml SOD to the reaction mixture or by bubbling the reaction mixture with N₂ gas. In addition, the same experiment was performed using ferric chloride (FeCl₃) instead of K₃[Fe(CN)₆].

Reduction of ferricyanide by a thiol compound or NAD(P)H. The solutions of K₃[Fe(CN)₆] and a test compound were prepared using PB. The K₃[Fe(CN)₆] solution and a test compound solution were mixed to make the final concentrations 2 mM and 1 mM, respectively. The time course of absorption at 420 nm of K₃[Fe(CN)₆] was observed on an Agilent 8453 photo-diode array spectrophotometer (NAD(P)H or a UNISOKU RSP-1000-02NM spectrophotometer (thiol compounds). The same experiments were repeated with 0.1 mM TEMPOL. For the oxidation of NAD(P)H by ferricyanide, time courses of the absorption at 340 nm after mixing the solutions were also observed.

Results and Discussion

TEMPOL was not reduced when it was exposed alone to the HX-XO reaction system, which induced O₂•− (Fig. 2A, open circle). The O₂•− equilibrated with HO₂• in an aqueous environment. HO₂•, which is a strong oxidant, can one-electron-oxidize the nitroxyl radical form of TEMPOL to produce the corresponding oxoammonium cation form. The oxoammonium cation can be one-electron-reduced by O₂•−, which is basically a reductant, to produce the nitroxyl radical form again. Therefore, TEMPOL was unaffected in the HX-OX reaction system. The oxoammonium cation form of TEMPOL can be two-electron-reduced to produce a relatively stable hydroxylamine form when it receives a hydrogen atom (H) from a coexisting hydrogen donor compound. The oxoammonium cation form of TEMPOL can also directly bind to a thiol moiety to make an EPR-silent complex. The reduction profile of TEMPOL in the HX-XO reaction system with a coexisting thiol compound is shown in Fig. 2A. All thiol compounds tested, including GSH (closed square), NAC (closed triangle), Cys (closed diamond), and HCS (open diamond), were able to reduce TEMPOL in the HX-OX reaction system. However, Met (Fig. 2B, closed circle), a sulfur-containing amino acid lacking a thiol, did not function as a reductant for the oxoammonium cation. NADH (Fig. 2B, open square) and NADPH (Fig. 2B, open triangle) functioned as a hydrogen donor, and reduced TEMPOL efficiently.
in the HX-OX reaction system.

The HX-OX reaction system used in this experiment continued generating HO•:O2•− during the 60-min experimental period because the TEMPOL with coexisting NADH or NADPH continued reducing. The incomplete elimination of TEMPOL with coexisting NAC, Cys, or HCS in this HX-OX reaction system may have been due to depletion of the thiol compounds by HO•, even though 10-times more thiol compound than TEMPOL was added to the reaction mixture. Thus, most thiols can be oxidized by HO• and be consumed before reducing the oxoammonium cation. However, TEMPOL was eliminated with higher concentrations of NAC, Cys, or HCS (data not shown), excluding excess Cys (8 mM), which inhibited complete TEMPOL reduction (data not shown). The oxoammonium cation form of TEMPOL and HO•− competes to oxidize thiols. Therefore, the EPR signal loss of TEMPOL was suppressed when the reaction of HO•− and a thiol compound was faster than the reaction of the oxoammonium cation. The free radical form of TEMPOL and thiols may also simultaneously compete with HO•. If this is the case, the EPR signal loss of TEMPOL may again be suppressed when the HO•−-induced oxidation of a thiol compound is faster than the oxidation of TEMPOL. As such, the relative reduction ability of thiols by HO• may be Cys > HCS > NAC > GSH.

The X-ray dose-dependent TEMPOL reduction profiles with coexisting test compounds in PB are shown in Fig. 3. X-ray induced ‘OH and/or HO•−, which can also one-electron-oxidize TEMPOL to produce the oxoammonium cation form. Almost no reduction of TEMPOL was observed when TEMPOL alone was irradiated with X-rays (Fig. 3A). All thiol compounds tested here, including GSH, NAC, Cys, and HCS, reduced TEMPOL during X-ray irradiation (Fig. 3B–E). However, coexisting Met was unable to reduce TEMPOL during X-ray irradiation (Fig. 3F). Coexisting NADH or NADPH reduced TEMPOL during X-ray irradiation, but at lower levels than the thiols (Fig. 3G and H).

X-ray irradiation of water molecules (H2O) generates ‘OH and
Fe(vivo) of K

Fig. 4A. Thiols can also function as hydrogen donors and reduce react directly with oxoammonium cations to generate a stable ions were unable to restore TEMPOL, thiol compounds mainly suggested that NAD(P)H mainly functions as a hydrogen donor to change the oxoammonium cation into the hydroxylamine form, which is the two-electron reduced form of the oxoammonium cation form or one-electron reduced form of the nitroxyl radical form. Separate from the above reaction, oxoammonium cations can directly react with thiols to make a complex. If no hydrogen donors or thiols coexist in the reaction system, oxoammonium cations rapidly one-electron reduce back to nitroxyl radicals, and no notable change is observed.

The reaction profiles of TEMPOL with the test compound in PB when potassium ferricyanide {K[Fe(CN)6]3−} was added are shown in Fig. 4. All thiol compounds were able to reduce TEMPOL immediately after the addition of ferricyanide (Fig. 4A). The behaviors of thiols in this reaction system were followed carefully when closing the reaction profiles (Fig. 4A, insertion). The reaction profile with coexisting Cys exhibited a perceptible recovery phase of TEMPOL, i.e., re-oxidation of the hydroxylamine to the nitroxyl radical form. Although the reaction profiles with coexisting GSH or HCS did not include such a clear recovery phase, an equilibrium state or slight bulging was observed on the decay slope. These may be traces of the recovery phase of TEMPOL. TEMPO decay with coexisting NAC almost fit simple first-order decay.

[FeII(CN)6]3− oxidizes TEMPOL to the oxoammonium cation, then [FeII(CN)6]3− becomes [FeII(CN)6]4−. The [FeII(CN)6]4− ion can reduce the oxoammonium cation back to TEMPOL. TEMPOL alone remained stable with ferricyanide (Fig. 4B, open circle). Met (Fig. 4B, closed circle) was non-reactive in this system. NADH (Fig. 4B, open triangle) and NADPH (Fig. 4B, open square) quickly reduced TEMPOL, whose levels recovered gradually. This suggested that NAD(P)H mainly functions as a hydrogen donor to change the oxoammonium cation into the hydroxylamine form, which can be re-oxidized to recover TEMPOL with excess [FeII(CN)6]3− ions. On the other hand, because excess [FeII(CN)6]3− ions were unable to restore TEMPOL, thiol compounds mainly react directly with oxoammonium cations to generate a stable complex. Thiols can also function as hydrogen donors and reduce a fraction of TEMPOL to the corresponding hydroxylamine form (i.e., TEMPOL-H) because the recovery phase or trace of recovery was observed in the reaction profiles of TEMPOL, as shown in Fig. 4A.

Addition of FeCl3 to the corresponding reaction mixture instead of K[Fe(CN)6]3− did not result in such reactions of TEMPOL (Fig. 5). The free FeII3+ ion may not oxidize TEMPOL to the corresponding oxoammonium cation. The free FeII3+ ion was not reduced by thiol compounds, i.e., free FeII3+ ions were unable to oxidize thiol compounds (data not shown). In addition, FeCl3 was unable to oxidize TEMPOL-H to TEMPOL (data not shown). Although FeCl3 has been widely used as an inducer of oxidized form in vivo or in cell culture experiments, the direct reaction of free FeII3+ and organic compounds is mild.

The effects of SOD or N2-bubbling on ferricyanide-induced TEMPOL decay are shown in Fig. 6. Neither SOD nor N2-bubbling suppressed the initial rapid TEMPOL decay. This suggests that the ferricyanide-induced rapid TEMPOL decay with the coexisting thiol compound was not related to oxygen-related free radicals. At the same time, the temporal recovery of TEMPOL after the initial rapid decay with coexisting Cys was increased by SOD or N2-bubbling (Fig. 6D). N2-bubbling increased temporal TEMPOL recovery with coexisting Cys more than SOD. The temporal recovery of TEMPOL observed with coexisting GSH was not affected by SOD or N2-bubbling (Fig. 6B). This suggests that oxygen-related free radicals are related, to some degree, to the relatively slow second TEMPOL decay observed with coexisting Cys.

The redox potential of TEMPOL vs the corresponding oxoammonium cation form was reported to be +0.810 V. It may be difficult for K[Fe(CN)6]3− to oxidize TEMPOL to the oxoammonium cation form based on its redox potential, which was reported to be +0.410 V. The thiol compounds examined reduced K[Fe(CN)6]3−, i.e., K[Fe(CN)6]3− oxidized thiol compounds, although the reaction was slow (Fig. 7A). The rapid TEMPOL decay induced by K[Fe(CN)6]3− also occurred with coexisting NAD(P)H. However, the reduction of K[Fe(CN)6]3− by NAD(P)H and the oxidation reaction of NAD(P)H by K[Fe(CN)6]3− were also slow (Fig. 7B and C). The addition of 0.1 mM TEMPOL to the reaction mixture quickened the reduction of K[Fe(CN)6]3− by NAD(P)H, but this cannot explain the K[Fe(CN)6]3−-induced rapid TEMPOL decay, as
shown in Fig. 4. Therefore, complex enzyme-like catalytic oxidative reactions may have occurred with $K_3[Fe(CN)_6]$, for which the detailed mechanisms are unclear. The $[Fe^{III}(CN)_6]^{3-}$ ion may function as a strong oxidant when coexisting with a hydrogen donor, and can one-electron-oxidize TEMPOL to produce the oxoammonium cation form.

As shown in Fig. 2–4, in a reaction mixture containing TEMPOL and a thiol compound in an oxidative atmosphere, TEMPOL was reduced. TEMPOL can be oxidized to the oxoammonium cation form and then the oxoammonium cation can then be reduced by the thiol. The oxoammonium cation may be reduced by either $\cdot H$ or the companion thiyl radical. The reduction of TEMPOL coexisting with a thiol compound suggests the generation of oxidants in the reaction mixture.

Based on the results of experiments shown in Fig. 2–4, we formed the hypothesis that the generation/addition of an oxidant in the reaction mixture containing TEMPOL and a thiol compound can reduce TEMPOL. If this is true, the generation of the $O_2^-$ and $HO_2^-$ redox pair, independent of its source, in a reaction mixture containing TEMPOL and a thiol compound can reduce TEMPOL. We previously reported $HO_2^-$/$O_2^-$ generation by the reduction of resolved $O_2$ in an aqueous solution containing GSH at a hyperthermal temperature.\(^{5,6}\) When the thiol compounds assessed in this paper, i.e., GSH, Cys, HCS, and NAC, reduced molecular oxygen, the reduction of TEMPOL in the reaction mixture was observed.

The reaction profiles when TEMPOL was incubated with thiols at a hyperthermal temperature (44°C) are shown in Fig. 8. As reported previously, GSH caused the EPR signal decay of TEMPOL with a characteristic profile of a time delay and sequential steep decay (Fig. 8A). GSH-dependent TEMPOL decay was slightly delayed after increasing the GSH concentra-
NAC, however, caused no EPR signal decay (Fig. 8B). Cys caused very slow decay of TEMPOL. The Cys-induced TEMPOL reduction increased with increasing Cys concentration (Fig. 8C). HCS induced no EPR signal decay of TEMPOL after hyperthermal treatment (Fig. 8D). Methionine, which is not a thiol, caused no EPR signal decay of TEMPOL after hyperthermal treatment (data not shown).

The order of reduction activity of the thiols for \( \text{O}_2 \) based on TEMPOL reduction was GSH > Cys > HCS = NAC. However, the reduction activity of thiols versus DPPH radicals was Cys > GSH = NAC, which was ordered according to the pKa of the thiol moiety.\(^{(22)}\) The order of reducing activity predicted from the results shown in Fig. 2 was similar to the reduction activity of thiols for DPPH. The reduction activity of thiols for \( \text{O}_2 \) is not necessarily the same as that for DPPH. The apparent reductive activity may change due to environment-dependent reaction mechanisms such as electron transfer, hydrogen transfer, or other combinations.

GSH-induced and Cys-induced TEMPOL decay at 44°C were inhibited by the addition of SOD and/or by bubbling \( \text{N}_2 \) gas into the reaction mixture (Fig. 9). Therefore Cys-induced TEMPOL decay was also mediated by \( \text{HO}_2^-/\text{O}_2^- \) generation, as reported...
Cys may reduce $\text{HO}_2$ coexisting Cys with increasing Cys concentration (Fig. 8C), $\text{HO}_2$ for GSH. Effects of SOD and N may result in lower concentrations of $\text{HO}_2$ reduction of TEMPOL at lower concentrations of Cys, which TEMPOL to the oxoammonium form, even if $\text{HO}_2$ were able to slightly suppress the one-electron-oxidation of competitively with TEMPOL because higher concentrations of GSH make $\text{HO}_2$ the provision of GS from O. GS was facilitated by higher temperatures. The generation of coexisting GSH was temperature dependent and increased at other thiol compounds to reduce O, but the detail mechanism for HCS cannot generate O due to the increase in TEMPOL reduction by Cys, as reported previously. Thus, GSH and Cys can reduce molecular oxygen to generate $\text{HO}_2$,$\text{O}_2$ even though Cys-induced $\text{HO}_2$,$\text{O}_2$ generation was slow. HCS cannot generate O in a direct manner, whereas HCS-induced oxidative stress is strongly related to the in vivo generation of $\text{O}_2^-$. Thus, chemically, HCS is a reductant.

As we reported previously, the reduction of TEMPOL with coexisting GSH was temperature dependent and increased at higher temperatures. Indeed the reduction of molecular oxygen by GSH was facilitated by higher temperatures. The generation of GS' may induce the reduction $\text{O}_2$, but the detail mechanism for the provision of GS' remains unclear because only the ability of other thiol compounds to reduce $\text{O}_2$ was assessed in this study.

The reduction of TEMPOL by coexisting GSH was delayed with increasing concentrations of GSH (Fig. 8A), as reported previously. This suggested that $\text{HO}_2$ can react with GSH competitively with TEMPOL because higher concentrations of GSH were able to slightly suppress the one-electron-oxidation of TEMPOL to the oxoammonium form, even if $\text{HO}_2$ generation from $\text{O}_2$ increased at higher concentrations of GSH. However, Cys may reduce $\text{HO}_2$ at a sufficiently fast rate to suppress the reduction of TEMPOL at lower concentrations of Cys, which may result in lower concentrations of $\text{HO}_2$ by reducing dissolved $\text{O}_2$ than GSH. Due to the increase in TEMPOL reduction by coexisting Cys with increasing Cys concentration (Fig. 8C), $\text{HO}_2$ generation by reducing $\text{O}_2$ may simply increase with increasing concentrations of Cys. Once an oxidative atmosphere is created by $\text{HO}_2$ generation in the reaction mixture, the EPR signal of TEMPOL may begin decreasing.

There are numerous reports on ROS generation in cells and animal models under hyperthermic conditions. Local hyperthermia may easily occur under non-physiological conditions, even during regular daily activities such as using a hair dryer, disposable body warmer, or taking a hot bath. Thus, researchers should take measures to make comfortable (less stressful) experimental conditions for animals such as controlling the body temperature under anesthesia during functional imaging experiments. Even at a physiological temperature, i.e., $37°C$, more than a 2-h incubation with GSH and O$_2$ may cause oxidative stress with increased ROS levels. Therefore, temperature-dependent and GSH-induced $\text{HO}_2$,$\text{O}_2$ generation are considered causes of cell death.

EPR signal loss by TEMPOL with coexisting thiol compounds in an aqueous reaction mixture can occur after several reactions. An oxidant may directly oxidize a thiol to produce a thyl radical (RS'), and then the RS' can react directly with the nitroxy radical (>N-O). Goldstein et al. reported that RS' and the nitroxy radical >N-O$^-$ directly react, and make a complex compound (>N-O-SR). Then, >N-O-SR becomes a stable amine form (>NH). They also proposed two alternate methods to produce >N-O-SR either by the reaction of an oxoammonium cation (>N=O) and thiolate (RS$^-$), which produces >N-O-SR directly, or by reacting >N-O' and RS', which then produce >N-O-SR. In this study, an important requirement was the irreversibleness of the reaction from >N-O' to >N-O-SR. When oxidation of the hydroxylamine form (>N-OH) to the corresponding >N-O' form, i.e., recovery of the EPR signal, was observed, it indicated the reduction of >N-O' to >N-OH. In other words, >N-O=O received hydrogen from the thiols (Fig. 4 and 6). As the reactions of thiol compounds and K$_3[\text{Fe(CN)}_6]$ were not fast (Fig. 7), and rapid TEMPOL decay was observed with coexisting NAD(P)H after adding K$_3[\text{Fe(CN)}_6]$ (Fig. 4B), the initial RS' generation by an oxidant may be not the main reaction.

Biologically reactive sulfur species, typified by thiols, mediate many pathophysiologically important redox reactions, termed Red-S-Ox. We demonstrated reductive stress caused by thiol compounds such as GSH and/or Cys.

**Conclusion**

Thiols and NAD(P)H can react with TEMPOL under oxidative conditions such as in an HX-XO reaction system, after X-ray irradiation, or in a ferricyanide reaction system. Thiols can react with the oxoammonium cation to make an EPR silent complex and function as an H-donor to provide the hydroxylamine. GSH and Cys can reduce molecular oxygen to generate $\text{HO}_2$,$\text{O}_2$ whereas Cys-induced $\text{HO}_2$,$\text{O}_2$ production was slow. Therefore, GSH and Cys can cause reductive stress. NAC, which was unable to reduce TEMPOL at hyperthermal temperatures, is a simple tractable antioxidant.

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**Conflict of Interest**

No potential conflicts of interest were disclosed.
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