RESEARCH ARTICLE

Effects of salt stress on antioxidant and ascorbate glutathione cycle enzyme activities in Pokkali rice varieties – Vytilla 1-9

Lins Simon & Yusuf A*
Inter University Centre for Plant Biotechnology, Department of Botany, University of Calicut, Kerala 673 635, India
*Email: akkara69@yahoo.co.in

ARTICLE HISTORY
Received: 12 December 2019
Accepted: 08 April 2020
Published: 01 July 2020

KEYWORDS
antioxidant enzymes
ascorbate - glutathione cycle
pokkali rice
rice
salt tolerance

ABSTRACT
The enzymatic and non-enzymatic antioxidant levels in the released salt tolerant Pokkali, (vytilla, VTL 1-9) varieties were studied under different NaCl concentrations (0-150 mM NaCl). The specific activity of superoxide dismutase (SOD), catalase (CAT) and ascorbate-glutathione cycle enzymes and non-enzymatic antioxidants like superoxide (O$_2^-$), hydrogen peroxide (H$_2$O$_2$), malondialdehyde (MDA), glutathione (GSH) and ascorbic acid (AsA) was determined in plants exposed to salt stress. IR-28 was used as positive control and the VTL varieties were used as negative control. The H$_2$O$_2$ and superoxide (O$_2^-$) contents were higher in IR-28 at all the applied concentrations of NaCl. The VTL varieties without salt treatment did not evoke any response substantiating the role of salt priming in antioxidant signalling. The MDA contents were higher in the positive and negative control. MDA content was reduced in the NaCl treated VTL varieties. In the positive and negative control varieties, the quantity of ascorbate and glutathione contents were lesser and upregulated in salt treated VTL varieties. Highest H$_2$O$_2$ content was observed in 150 mM NaCl treatment. The H$_2$O$_2$ contents decreased with the increase in all concentrations of NaCl and lowest H$_2$O$_2$ contents was observed in VTL-1 and highest in VTL-2 and VTL-8 treated with 150 mM NaCl. Superoxide contents varied in all the nine varieties depending on the salt concentration. The SOD levels in all the varieties showed a positive correlation with the superoxide and H$_2$O$_2$ content. Lesser quantities of SOD, CAT and the ascorbate - glutathione cycle enzymes were expressed in the positive and negative control. The increased NaCl concentration (25-150 mM) upregulated antioxidant and ascorbate-glutathione cycle enzymes in the VTL varieties. The APX activity was lower in the control and salt treated plants. The GR activity increased linearly in all the varieties with respect to salt concentrations. The MDHAR and DHAR activities showed marginally linear increase, with all concentrations of NaCl. The APX activity was similar or lower to MDHAR activity while DHAR activity was similar to MDHAR activity. The results of the present study reveals the higher levels of enzymatic and non-enzymatic antioxidants under salt stress reflect the salt tolerance potential of pokkali varieties mediated by the up regulation of ROS scavenging enzymes.

Introduction
Rice (Oryza sativa L.) is one of the important cereal crops throughout the world. The global climate change is reducing the productivity by increasing salinity in agricultural land by about 50% (1). In this context, food security issues can be addressed by developing salt tolerant varieties of rice suitable for arid, semi-arid and waterlogged regions. Kerala has a long coastal area of about 580 km, which has several lagoons and backwaters covering a very large area linked to the sea. The deltaic regions are either at the sea level or below affecting the intrusion of sea water to agriculture lands (2). The salt tolerance mechanism is obscure in most of the salt tolerant rice varieties developed through conventional breeding techniques. The survival of plants in high salt habitat is depending on the production of reactive oxygen species (ROS) and highly efficient scavenging of this anion by the antioxidant and ascorbate-glutathione cycle enzymes. The production of ROS, the by-product of...
oxidative metabolism in plants causes damage to DNA, proteins and chlorophyll etc. (3-5). It is enhanced in response to abiotic stresses like drought (6, 7), elevated temperature (8) and xenobiotics (9).

Studies suggest that antioxidant enzymes are upregulated in plants exposed to abiotic stress (10). Charged superoxide radical cannot move freely across the membranes, hence the subcellular compartmentalisation of the antioxidant system is necessary for efficient quenching of this anion and its immediate product \( \text{H}_2\text{O}_2 \), at the site of production. Synchronous action of superoxide dismutase (SOD), catalase (CAT) and ascorbate-glutathione cycle enzymes, ascorbate peroxidase (APX), monodehydroascorbate reductase (MDHAR), dehydroascorbate reductase (DHAR), glutathione reductase (GR) is responsible for quenching of \( \text{O}_2^- \) and the reduction of \( \text{H}_2\text{O}_2 \). The APX, MDHAR and DHAR are the most important peroxidase in detoxifying \( \text{H}_2\text{O}_2 \) (13). Correlation between the ascorbate-glutathione cycle enzymes and salt tolerant is compared in the sensitive and resistant varieties of rice (14). However, the salt tolerance capacity of the pokkali rice variety, vytilla (VTL) with respect to the antioxidant and ascorbate-glutathione cycle enzymes is not investigated so far. The present study was undertaken to evaluate the salt tolerance capacity and the levels of antioxidant and ascorbate-glutathione cycle enzymes in the released pokkali rice varieties, vytilla (VTL) 1-9 (negative control) and IR-28 variety (positive control) under different salt concentrations. This study will help in understanding the ROS scavenging mechanism and the adaptation to salt stress in these released varieties.

**Materials and Methods**

**Plant materials**

Seeds of the released Pokkali rice varieties (VTL 1-9), were collected from Rice Research station, Kerala Agriculture University, Vytilla, Kerala while, IR-28 seeds were obtained from Rice Research Station, Pattambi, Kerala.

**Experimental design**

The experiment was set up in a completely randomized design in 7 tanks filled equal volume of water containing NaCl concentrations of 0, 25, 50, 75, 100, 125 and 150 mM. The seeds were germinated in seed beds. Twenty-one day old seedlings were planted in pots containing a soil mixture comprising of loam soil and cow dung mixture in the ratio 1:1. Three replications were performed, each containing 10 seedlings. NaCl treatment started on 25\(^{th}\) day of germination by incrementing 25 mM per day to attain a final concentration of 150 mM. One set of plantlets were kept as control without NaCl application and watered with tap water. Leaf samples for the enzyme assay were harvested after 21 days (46 days from seed germination) of salt application and frozen in Liquid Nitrogen (LN\(_2\)) and stored at -80°C.

**Quantification of ROS and non-enzymatic antioxidants**

The superoxide radical (\( \text{O}_2^- \)) was determined (15). A standard curve was prepared with nitrogen dioxide (NO\(_2\)) to calculate the \( \text{O}_2^- \) generation rate. Hydrogen peroxide (H\(_2\)O\(_2\)) content was determined (16). The lipid peroxidation was determined using malondialdehyde (MDA) content as per standard method (17). The concentration of the MDA/TBA (malondialdehyde/thiobarbituric acid) complex was calculated and the level of lipid peroxidation was determined.

Ascorbic acid (AsA) content was assayed using the protocol (18). Optical density was recorded at 525 nm. A standard curve was prepared using a gradient concentration of AsA.

Total and reduced glutathione were determined using previously described protocol (19). The GSH concentration was calculated using a standard curve prepared using gradient concentrations of GSH. The spectrophotometric reading was taken at 412 nm. The quantity of GSSG is determined using the formula, GSSG = Total GSH – GSH.

**Quantification of enzymatic antioxidants**

Superoxide dismutase (SOD) and catalase (CAT) was extracted from the frozen sample (0.5 g), homogenized in LN\(_2\) by adding 100 mM phosphate buffer (4 ml, pH 7.0), 1 mM EDTA (16 µl) and 1% (w/v) polyvinyl pyrrolidone (PVP). The homogenate was centrifuged at 10000 rpm for 15 min at 4 °C and the supernatant was collected for further studies. Ascorbate peroxidase (APX) was extracted from the leaf sample (0.5 gm) homogenized in extraction medium containing 50 mM potassium phosphate buffer (pH 7.0), 1 mM EDTA, 2% (w/v) PVP and 1 mM ascorbic acid. The homogenate was centrifuged at 15000 rpm for 20 min at 4 °C. For the glutathione reductase (GR) assay, the sample (0.5 gm) was powdered in LN\(_2\), homogenized in 100 mM potassium phosphate buffer (pH 7.8) containing 2 mM EDTA and 1% (w/v) PVP-40 at 4 °C. The homogenate was centrifuged at 15000 rpm for 20 min at 4 °C and the supernatant was collected. Monodehydroascorbate reductase (MDHAR) and dehydroascorbate reductase (DHAR) was assayed using powdered plant sample (0.5 g) homogenized in 50 mM potassium phosphate buffer (pH 7.8) and 1 mM EDTA. Homogenate was centrifuged at 14000 rpm for 20 min at 4 °C and the supernatant was used for enzyme assay.

Superoxide dismutase (SOD, EC 1.15.1.1) activity was measured according to the modified protocol (20). One unit SOD was calculated as the enzyme activity required for 50% photoreduction of nitrobluetetrazolium to blue formazan.

Catalase (CAT; EC 1.11.1.6) activity was measured (21). Change in absorbance at 240 nm due to the degradation of H\(_2\)O\(_2\) was recorded at an interval of 15 s for 2 min.
Ascorbate peroxidase (APX; EC 1.11.1.11) activity was measured (22). The decrease in absorbance due to ascorbate oxidation was recorded at 290 nm at 15 s intervals for 2 min. The specific activity was calculated and expressed as µmol mg protein⁻¹ min⁻¹.

Glutathione reductase (GR; EC 1.11.1.9) activity was determined (23) with some modifications. Change in absorbance at 412 nm due to the oxidation of NADPH was recorded in 15-s intervals for 2 min and the specific activity of the enzyme was calculated and expressed as µmol mg protein⁻¹ min⁻¹.

Monodehydroascorbate reductase (MDHAR; EC 1.6.5.4) and Dehydroascorbate reductase (DHAR; EC,1.6.5.4) activities were assayed (24). The decrease in absorbance in MDHAR was monitored at 340 nm owing to the oxidation of NADH while, DHAR activity was determined by monitoring the rate of AsA formation at 265 nm.

Total protein from the experimental samples was determined (25). The absorbance was measured at 660 nm using a UV-VIS spectrophotometer and the protein were quantified using gradient concentrations of BSA (1.0 mg/l).

Statistical analysis
All the enzymatic and non-enzymatic assays were repeated thrice and the data was subjected to analysis of variance (p ≤ 0.05) and the mean values were compared using ANOVA.

Results

Quantification of ROS contents
The superoxide (O₂⁻) contents in the VTL varieties increased proportionate to the NaCl concentration (Fig. 1A). In the positive control IR-28, the O₂⁻ content was 8-10 times higher compared to the tolerant varieties. Among the Pokkali varieties, VTL-4 showed lesser O₂⁻ content and VTL-9 showed the highest levels of O₂⁻ in all the concentrations of NaCl used. The varieties VTL-4, VTL-5, VTL-6 and VTL-7 showed a decrease in O₂⁻ content with increase in NaCl concentration. All the other varieties showed a steady increase in O₂⁻ content proportionate with NaCl concentration. The positive control IR-28 produced 3-4 times higher H₂O₂ (0.789±0.08 mMol g⁻¹) compared to salt tolerant Pokkali varieties (Fig. 1B). The H₂O₂ content in the negative control VTL varieties were lesser compared to salt treated VTL varieties. During NaCl treatment the highest H₂O₂ content was observed in VTL-2 (0.267 mMol g⁻¹) and the lowest quantity was observed in VTL-4 (0.188 mMol g⁻¹) in 150 mM NaCl. All the other varieties showed an increase in the H₂O₂ content upon exposure to NaCl, however, the response was dependent on NaCl concentration and time of exposure.

MDA content
The MDA quantity was higher (0.712 mMol g⁻¹) in the positive control IR-28 (Fig. 1C). The salt tolerant VTL varieties used as negative control produced lesser quantity of MDA in 150 mM NaCl. Higher MDA content was observed in the salt treated VTL varieties comparing to IR-28 and a maximum quantity was observed in VTL-9 (0.502 mMol g⁻¹). The lowest quantity in VTL-3 (0.323 mMol g⁻¹) treated with 150 mM NaCl on the 21st day.

Ascorbic acid (AsA) content
The AsA content in the positive control IR-28, sharply declined with increasing in NaCl concentration and duration of NaCl exposure (Fig. 1D). In the negative control, the VTL varieties produced higher AsA, while VTL varieties under salt stress maintained steady level of AsA. Among the VTL varieties, the highest AsA content was observed in VTL-2 (0.928 mMol g⁻¹) and the lowest in VTL-8 (0.667 mMol g⁻¹). The AsA content decreased proportionate with the increase in NaCl concentration and the duration of exposure.

GSH contents and GSH/GSSG ratio
The GSH contents in the positive control IR-28 was lesser compared to the negative control VTL varieties (Fig. 1E). In the salt treated varieties, the higher quantity of glutathione was measured in VTL-1 treated with 150 mM NaCl on the 21st day. The GSH/GSSG ratio (Fig. 1F) decreased with increase in NaCl concentration suggesting the presence of a higher quantity of oxidised glutathione.

SOD activity
The SOD activity was upregulated on exposure to different concentrations of NaCl. The highest SOD activity was expressed in 150 mM NaCl treated VTL-2 with 43.99±4.36 U SOD mg protein⁻¹. In rest of the varieties the SOD activity was upregulated and showed consistent increase with increase in salt concentration. In the control experiment, the variety VTL-9 showed the highest SOD activity of 21.34±1.38 U SOD mg protein⁻¹ followed by VTL8 > VTL7 > VTL6 > VTL1 > VTL5 > VTL2 > VTL4 > VTL3 (Fig. 2A). In the positive control IR-28, the SOD expression decreased with increasing NaCl concentration and duration of exposure.

Catalase activity
Catalase activity in the salt treated VTL varieties followed the same pattern of SOD, though the expression level varies, the highest enzyme activity was observed in VTL-1 treated with 150 mM NaCl with a specific activity of 0.506±0.0087 µmole mg protein⁻¹ min⁻¹ and in the negative control plants, the CAT activity was lesser than the salt treated varieties. The salt treated varieties showed an increased CAT activity in all the concentrations of NaCl used, though the enzyme activity varied slightly in salt treatment (Fig. 2B). The CAT activity in the positive control IR-28, was approximately 4-6 times lesser than the salt treated varieties.

APX activity
In the control experiment, highest APX specific activity was measured in VTL-6 (0.249±0.018 µmole mg protein⁻¹ min⁻¹). The other varieties showed almost similar APX activity upregulated in response to salt concentration and the specific activity increased proportionate with the salt concentration. The
highest specific activity (0.527±0.018 μmole mg protein⁻¹min⁻¹) was observed in 150 mM NaCl treated VTL-3 and the lowest in VTL-7. The specific activity of APX in all the varieties treated with different concentrations of NaCl is shown in (Fig. 2C). The APX activity in the positive control (IR28) showed lesser activity of APX in all the concentrations of NaCl used.

**GR activity**

The highest GR activity (0.274±0.006 μmole mg protein⁻¹ min⁻¹) was observed in VTL-1 compared to the other varieties in the negative control. Among the salt stressed varieties the highest GR activity of 0.553±0.008 μmole mg protein⁻¹ min⁻¹ was observed in VTL-1, grown in 150 mM NaCl. All the other varieties expressed an increased GR activity expressing concentrations and varietal differences (Fig. 2D). In the positive control IR-28 the GR activity was 5-7 times lesser than the VTL varieties.

**MDHAR activity**

In the negative control, the variety VTL-3 showed the highest MDHAR activity of 1.19±0.02 μmole mg protein⁻¹ min⁻¹. In the salt treatments the highest MDHAR activity was observed in VTL-2 with a specific activity of 1.67±0.49 μmole mg protein⁻¹ min⁻¹. The MDHAR activity in the salt stressed and control did not show much difference, except in VTL-2 and VTL-6 suggesting that the Pokkali varieties used for the study have high inherent MDHAR activity (Fig. 2E). The positive control, IR-28 showed 2-3 times lesser MDHAR activity compared to the VTL varieties treated with different concentrations of NaCl.

**DHAR activity**

In the negative control, the variety VTL-2 expressed the highest DHAR activity of 1.19±0.08 μmole mg protein⁻¹ min⁻¹. The DHAR activity was upregulated in salt treated varieties and the highest DHAR activity was observed in 150 mM NaCl treated VTL-6 variety with an enzyme activity of 1.30±0.015 μmole mg protein⁻¹ min⁻¹ (Fig. 2F). The positive control IR-28 showed 2-3 times lesser DHAR activity compared to the salt treated VTL varieties.

**Discussion**

The salt tolerant Pokkali varieties, (Vyttila-1-9) developed through conventional breeding methods, were capable of constitutive activation of peroxide scavenging pathways in order to adapt to salt stress. This suggests that the salt tolerant pathways were initiated by scavenging the stress related compounds, thus maintaining a stable Na⁺/K⁺ ratio in plants (26). Under salt stress conditions, VTL varieties produced lesser quantity of H₂O₂ compared to other pokkali varieties like CSR-1 (27) and positive control IR-28 suggesting the adaptability of the VTL varieties to salt stress. The reduced H₂O₂ in the salt tolerant varieties is related to the upregulation of SOD and CAT activity that assist in scavenging the generated H₂O₂. The increased O₂⁻ contents in the negative control plants suggest the inherent capacity of the salt tolerant Pokkali varieties in alleviating salt stress. However, the superoxide radical emissions were dependent on NaCl concentration. Over production of ROS during salt stress leads to the toxicity in cells causing cellular damages. The incidence of salinity stress in plants produces ROS and several adverse effects that have a destructive influence on different plant processes (28). In accordance with the prevention from oxidative damage caused due to lipid peroxidation, plants have developed specific strategies comprising upregulation of specific enzyme pathways, metabolite production and gene expression (29). Plants strongly control the production and elimination of ROSs by many enzymatic and non-enzymatic processes to the alleviation of their damages (30). These non-enzymatic and enzymatic antioxidants play a vital role by interacting with each other in order to scavenge the reactive oxygen species mediated by the over expression of different genes and thus helping the plants to acclimatise to the adverse environmental conditions (31, 32). MDA content is considered as a sign of the extent of oxidative damage under stress conditions (33). Higher quantity of MDA in the negative and positive control during salt stress suggested membrane peroxidation and the MDA content did not show any upregulation, thus preventing the lipid peroxidation; congruent results are reported in Beta maritima, a salt tolerant variety during salt stress (34). Ascorbate content in the positive and negative control were upregulated, whereas in the salt tolerant varieties, the AsA content decreased in the VTL varieties attributing the up regulated activity of the APX enzyme which utilises ascorbate. It may also possible that the reduced AsA content can be due to the increased catabolism of the reduced or oxidised ascorbate. The two fold increase in GSH content in the salt stressed VTL varieties is related to the higher levels of GSH synthesis and the considerable increase in GR activity that assists in maintaining the GSH/GSSG ratio, the decrease in the GSH/GSSG ratio specifies the oxidative stress related cellular redox buffer under salt stress.

The upregulated SOD and CAT activities in the VTL varieties suggests that the superoxide radical generation due to salt stress is also alleviated by the participation of NADPH and the signalling cascade involving the ascorbate glutathione cycle enzymes that help to scavenge the ROS. Combined action of SOD and CAT in alleviating the stress responses were reported in other plant species (35). The H₂O₂ dependant oxidation of AsA is active in the tolerant varieties and the upregulation of APX specific activity suggest that, the H₂O₂ produced in the tolerant plants during salt stress is effectively scavenged through the ascorbate-glutathione cycle enzymes. The upregulated activity of CAT and APX enzymes also suggests that the competition for the substrate H₂O₂ is not affecting the activity of these enzymes as reported earlier (14).

Catalase is one of the H₂O₂ detoxifying enzymes which is mostly associated with peroxisomes and helps in removing H₂O₂ formed during photorespiration. The differential behaviour of these cultivars on time scale indicates the early adaptive potential of Pokkali in terms of CAT activity. The upregulation of H₂O₂ and CAT activity is correlated with the increased protection from the damage associated with oxidative stress (4, 36). This
corroborates the significant differences in the CAT activity of the salt tolerant and salt-sensitive cultivars of pea reported during salt stress (37). The upregulated APX activity suggests that the VTL varieties develop inherent characteristics for protection against salt stress by maintaining the H$_2$O$_2$ concentration inside the cell to a non-toxic level as the combined activities of APX and CAT are responsible for H$_2$O$_2$ scavenging (38). APX is one of the most important antioxidant enzymes in plants that detoxify H$_2$O$_2$ using ascorbate as reductant. It exists in several isoforms in various compartments of plant cells and shows differential expression and regulation under environmental stress factors. In the VTL varieties, the APX activity decreased without salt treatment, however, the SOD to APX activity differed significantly in the different VTL varieties. The higher ratio of SOD to APX correlate with H$_2$O$_2$ scavenging, suggesting that in the VTL varieties the H$_2$O$_2$ produced is actively detoxified.

Glutathione reductase (GR) is essential for the elimination of ROS and for the maintenance of reduced glutathione (GSH) by the ascorbate/glutathione cycle. The enzyme GR mediates the reduction of oxidized glutathione (GSSG) produced by the action of DHAR, which in turn produced by the APX during the peroxidation of

![Fig. 1. A) Superoxide content, B) Hydrogen peroxide content, C) MDA content, D) Ascorbate content, E) Glutathione content, F) GSH/GSSG ratio in the IR-28 and VTL varieties treated with different concentrations of NaCl on the 21st day. The values are the mean of three independent experiments and are subjected to one-way ANOVA. The values are significant at p > 0.05.](image-url)
ascorbate, to the reduced GSH pool using NADPH as an electron donor. Thus, a highly reduced state of GSH/GSSG ratio is maintained at the intracellular level by this reaction during oxidative stress. It was observed that the activity of GR increase linearly to the salt concentration in all the VTL varieties compared to negative control VTL varieties. These results are in corroboration with the earlier results that GR activity increases during salt treatment in order to generate glutathione, an intracellular non-enzymatic antioxidant that protects the plant membranes and prevents the oxidative denaturation of proteins under stress by protecting their thiol groups (39).

MDHAR is the major enzyme in AsA-GSH cycle contributing to ASC generation and the APX and MDHAR activities are higher or the MDHAR activity is one order of magnitude lower than the APX activity and the DHAR activity one order lesser than the MDHAR activity under salt stress in most of the plant species (40). The results obtained in VTL varieties under salt stress suggest that the capacity to regenerate ASC was similar or higher than the capacity of APX to oxidise MDA. The lower activities of GR and DHAR support this view. The increased MDHAR activity in the salt stressed VTL varieties compared to negative control VTL varieties suggest the higher capacity of ASC generation under salt stress conditions. As observed in some other plants, the decreased MDHAR activity in the negative control suggests the level of ascorbate regeneration can be correlated with salt concentration among VTL varieties (41, 42). Furthermore, the higher activity of various

![Fig. 2. A) Superoxide dismutase activity, B) Catalase activity, C) Ascorbate peroxidase activity, D) Glutathione reductase activity, E) Monodehydroascorbate reductase activity, F) Dehydroascorbate reductase activity in the IR-28 and vytilla varieties treated with different concentrations of NaCl on the 21st day. The values are the mean of three independent experiments and are subjected to one-way ANOVA. The values are significant at p > 0.05.](image-url)
antioxidant enzymes (SOD, CAT, APX, MDHAR, DHAR and GR) in a coordinated manner in the salt tolerant cultivars suggests that they are the major determinants in elucidating salt tolerance.

The Pokkali varieties (VTL-1-9) could not withstand NaCl concentration above 150 mM as the NaCl treated plants turn yellow from the 25th day of salt treatment, indicating the tolerance threshold of 150 mM for these varieties. Most of the studies conducted on salt tolerance in rice varieties were based on salt treatment for 24-48 hrs (14, 43), however, the present study, the salt tolerance upto 30 days in 75 day old plants with 25-150 mM NaCl provide conclusive results on the salt tolerance potential of these varieties. The negative control VTL varieties, the quantity of antioxidant enzymes match with the salt treated varieties, demonstrating that the salt tolerant varieties have inherently higher expression of antioxidant enzymes.

Authors’ contributions
LS carried out the enzyme assays and prepared the first draft of the manuscript. AY conceived of the research idea designed the experiments and edited the manuscript. All authors read and approved the final manuscript.

Acknowledgements
The authors are grateful to the Director, Interuniversity Centre for Plant Biotechnology for providing facilities and support. Financial support from Kerala State Council for Science Technology and Environment for this study under the SRS scheme No. 001/SRSLS/2013/CSTE is acknowledged.

Conflict of interest
The authors declare that they have no conflict of interests.

References
23. Smith IK, Vierheller TL, Thorne CA. Assay of glutathione reductase in crude tissue homogenates using 5, 5’ dithiobis (2-
