Spectroscopic properties of rubber oxygenase RoxA from *Xanthomonas* sp., a new type of dihaem dioxygenase

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Natural rubber (poly-*cis*-1,4-isoprene) is cleaved to 12-oxo-4,8-dimethyltrideca-4,8-diene-1-al (ODTD) by rubber oxygenase A (RoxA) isolated from *Xanthomonas* sp. RoxA has two *c*-type haem centres that show two distinct *α*-bands at 549 and 553 nm in the dithionite-reduced state. A well-resolved midpoint potential (*E_<0>*) of −65 mV was determined for one haem by spectrophotometric titrations in the absence of dioxygen with dithionite and ferricyanide as reductant and oxidant, respectively. The midpoint potential of the second haem was not resolvable (*E_<0>*) about −130 to −160 mV). One of the two haems was reduced by NADH (549 nm *α*-band), similar to bacterial dihaem peroxidases. Evidence for an electron transfer between the two haems was provided by slow reduction of the second haem (553 nm *α*-band) upon incubation of the partially reduced enzyme at room temperature. Addition of imidazole or related compounds to RoxA led to UV/vis spectral features similar to those observed for partially reduced RoxA. Notably, reduction of RoxA with dithionite or NADH, or binding of compounds such as imidazole, resulted in a reversible inactivation of the enzyme, unlike dihaem peroxidases. In line with this result, RoxA did not show any peroxidase activity. EPR spectra of RoxA as isolated showed two low-spin Fe(III) haem centres, with apparent *g*-values of 3.39, 3.09, 2.23, 1.92 and 1.50. A weak signal in the *g* = 6 region resulting from a high-spin Fe(III) haem was also observed with a preparation-dependent intensity that disappeared in the presence of imidazole. Attempts to provide spectroscopic evidence for binding of the natural substrate (polyisoprene latex) to RoxA failed. However, experimental data are presented that RoxA is able to subtract redox equivalents from its substrate or from model compounds. In conclusion, RoxA is a novel type of dihaem dioxygenase with features clearly different from classical cytochrome *c* peroxidases.

INTRODUCTION

Natural rubber (NR) or caoutchouc is a biopolymer that is synthesized by many plant species and by some fungi. NR has been commercially exploited for more than 100 years by cultivating and tapping the rubber tree (*Hevea brasiliensis*) at a level of several million tons per year.

The natural polymer has an average molecular mass of about one million and is composed of two *trans* isoprene units at one end of the molecule followed by several hundreds to thousands of *cis*-1,4-isoprene units (Tanaka & Sakdapipanich, 2001). The linear, non-branched polymer chains can be cross-linked by treatment with sulfur and heat (vulcanization), generating a material with superior physical and mechanical properties. Despite the development of chemosynthetic rubbers, the natural polymer is still required as a basic material for tyres, latex gloves, condoms, seals and many other items.

Naturally occurring NR is subject to biological mineralization cycles, and many reports on the isolation and characterization of rubber-degrading micro-organisms have been published (for overview, see Rose & Steinbüchel, 2005, and references cited therein). Rubber-degrading bacteria either produce clearing zones on opaque latex agar, such as *Xanthomonas* sp. 35Y (Tsuchii...
& Takeda, 1990) and many actinomycetes (Jendrossek et al., 1997), or grow adhesively on rubber without clearing-zone formation, such as several *Gordonia* strains (Linos et al., 1999, 2002). Based on identified intermediates that had been isolated from rubber-grown bacteria (Bode et al., 2000; Tsuchii et al., 1985; Tsuchii & Takeda, 1990), a biochemical degradation route has been proposed (Bode et al., 2001; Rose & Steinbüchel, 2005). Meanwhile, two candidate proteins have been described that are involved in the primary attack of the polyisoprene carbon backbone. One is the latex-clearing protein (Lcp) of *Streptomyces* sp. K30 (Rose et al., 2005) and the other is rubber oxygenase RoxA of *Xanthomonas* sp. Lcp and RoxA are apparently completely different polypeptides with no significant amino acid similarities. RoxA is a dihaem oxygenase releasing low-molecular-mass oligoisoprene units as products. 12-Oxo-4,8-dimethyltrideca-4,8-diene-1-al (ODTD) has been identified as the major product under *in vitro* conditions together with a homologous series of minor metabolites that differ from the major degradation product only in the number of repetitive isoprene units between terminal functions, CHO- and -CH$_2$-COCH$_3$ (Braaz et al., 2004). Isotope labelling experiments have revealed that RoxA is a dioxygenase (Braaz et al., 2005). A large set of experiments was carried out to investigate the reactivity of the RoxA haem centres towards substrates, reductants, oxidants, inhibitors and well-known haem ligands, including imidazole and related compounds.

**METHODS**

**Cultivation of bacteria and purification of rubber oxygenase (RoxA).** *Xanthomonas* sp. 35Y (Tsuchii & Takeda, 1990) was grown in nutrient broth (NB) or in a mineral salts medium (MSM) with 0.01–0.2% purified rubber latex at 30 °C. For isolation of RoxA from *Xanthomonas* sp. wild-type, 24 250 ml cultures were grown for about 10–14 days, and RoxA was purified from the combined cell-free culture fluid as described previously (Braaz et al., 2005). For purification of recombinantly expressed RoxA, *Xanthomonas* sp. harbouring p4782.1::roxA (p4782.1) is a mobilizable broad-host-range vector suitable for rhannose-dependent expression of cloned genes and confers resistance against kanamycin) was grown in 15 0.5 l cultures of nutrient broth (NB) or modified Luria–Bertani (LB) broth (yeast extract of LB replaced by NB) supplemented with 0.1% l-rhamnose for 60 h at 30 °C by continuous shaking, as described previously (Hambsch et al., 2010). Purification of RoxA was performed from the combined cell-free culture fluid.

**Purification of RoxA** RoxA was purified at room temperature using an AKTA FPLC system (GE Healthcare). Cell-free supernatant of *Xanthomonas* sp. p4782.1::roxA culture was concentrated by ultrafiltration (30 kDa cut-off) and passed through a Q-Sepharose Fast Flow column (Q-FF 50/11; GE Healthcare) pre-equilibrated with 20 mM Tris/HCl (pH 7.7) at a flow rate of 3 ml min$^{-1}$. RoxA was collected in the flow-through and separated from proteins that bound to the column at pH 7.7. In a second step, the RoxA-containing flow-through was passed through the same column but equilibrated with 20 mM Tris/HCl (pH 8.5). RoxA was bound at alkaline pH and eluted at ~50 mM NaCl. RoxA-containing fractions were combined and stored at ~70 °C. The RoxA was pure (~95%), as judged by SDS-PAGE analysis and subsequent staining with silver. If highly purified RoxA was necessary, an additional step involving a hydroxyapatite column [several runs with a Bio-Scale CHT5-1 column (Bio-Rad)] equilibrated with potassium phosphate (KP) buffer (5 mM, pH 6.2) was used after changing the buffer of the RoxA pool with a HiPrep 26/10 desalting column (GE Healthcare) after the second step of Q-Sepharose gel-filtration chromatography. RoxA was eluted with a linear gradient of 5–200 mM KP at ~40 mM. RoxA fractions were pooled and concentrated by ultrafiltration (30 kDa cut-off) to ~5 mg ml$^{-1}$. The addition of 300 mM NaCl was necessary to prevent precipitation of RoxA. Purity was tested by SDS-PAGE and by determination of the ratio of absorption at 406 and 280 nm, which was 1.35 for highly pure RoxA after hydroxyapatite chromatography and final concentration. Purified RoxA was frozen in liquid nitrogen and stored at ~70 °C.

**Assay of RoxA.** The following conditions were used for product analysis of RoxA-catalysed rubber degradation by HPLC analysis: the reaction mixture contained 100 μl purified RoxA (2 μg ml$^{-1}$), rubber latex (0.2%, w/v, emulsion) and KP buffer (100 mM, pH 7.0) in a total volume of 1 ml. The reaction was carried out at 40 °C for 3 h in a test tube sealed with Parafilm. The mixture was extracted with ethyl acetate or diethyl ether, dried, dissolved in 100–200 μl methanol, and then subjected to HPLC analysis and/or to carbonyl content determination (Katz & Keeney, 1966). Mixtures without RoxA and with heat-inactivated RoxA (10 min, 95 °C) served as controls. One unit (U) of RoxA activity corresponds to one micromole carbonyl function generated per minute.

**Peroxidase assay.** Peroxidase activity was assayed as described by Mason et al. (2001). RoxA was incubated in 100 mM sodium phosphate buffer (pH 7.0) containing 5 mM 2,4-dichlorophenol, 3.2 mM 4-aminoantipyrine and 1 mM hydrogen peroxide, and the increase in $\text{A}_{510}$ was followed. Alternatively, pyrogallol (420 nm), guaiacol (436 nm) and Fe(II) (horse heart) cytochrome c (decrease at 550 nm) were used as electron donors. Horseradish peroxidase (HRP) was used as a reference.

**Transformation of Xanthomonas sp.** A 100 ml volume of NB medium in a 500 ml Erlenmeyer flask was inoculated with 0.01 volumes of an NB seed culture and grown overnight at 30 °C until the OD$_{600}$ reached 0.4–0.6. Cells were centrifuged in 50 ml Falcon tubes aseptically at 4 °C and 5100 g for 15 min. The supernatant was discarded and the pellet was resuspended in about 0.2 ml of the supernatant. It was important to ensure that the culture was kept on ice during all manipulation steps. Cells in Falcon tubes were diluted with 40 ml ice-cold 0.1 M HEPES buffer that had been titrated to pH 7.0 by the addition of 1 M Tris base, and were centrifuged (15 min, 5100 g, 4 °C) and resuspended as described above. The washing procedure was repeated with 40 ml 10% (v/v) glycerol that had been adjusted to neutral pH by the addition of a few drops of 0.1 M HEPES-Tris, pH 7. Cells were resuspended in about 0.2 ml of the remaining supernatant and were then ready for use in electro-transformation experiments. Cells were kept on ice until use. A 75 μl volume of electrocompetent cells was mixed with 50–500 ng DNA in an ice-cold electroporation cell. Electroporation was performed at 25 μF, 200 Ω, 1.5 kV. Cells were transferred to 0.6 ml fresh NB medium and incubated for up to 3 h at 30 °C before aliquots were plated on selective solid media (NB supplemented with 20–30 μg kanamycin ml$^{-1}$). Determination of c.f.u. ml$^{-1}$ after electroporation by plating appropriate dilutions on NB agar was useful because a high loss of viability after electroporation was frequently observed and resulted in insufficient electroporation efficiency. Alternatively, mobilizable plasmids were transferred to *Xanthomonas* sp. via conjugation with *Escherichia coli* S17-1. The efficiency of conjugative transfer of DNA was about two to three orders of magnitude higher than that of electro-transformation.
**Haem staining.** Haem staining was performed after separation of RoxA samples by SDS-PAGE and subsequent assay for pseudoperoxidase activity of the RoxA haem groups. The gels were washed with water and incubated at room temperature in a solution consisting of 15 ml 3,3',5,5'-tetramethylethylenebenzidine (1.5 mM in methanol), 35 ml 0.25 mM sodium acetate, pH 5, and 26 mM H$_2$O$_2$. A blue colour indicated the presence of haem-dependent pseudoperoxidase activity. Protein concentration was determined by the Bradford method (Bradford, 1976) and by commercial bicinchoninic acid (BCA) assay (Pierce).

**Iron analysis.** Iron was determined by Spuren-analytisches Laboratorium Dr Baumann (Maxhütte-Haidhof, Germany) using inductively coupled plasma MS (ICP-MS).

**UV/vis-monitored oxidation-reduction.** Potentiometric titrations were performed under the exclusion of dioxygen (Argon) in a Thunberg cell, at pH 7.0 and 25 °C, by stepwise addition of sodium dichitron (reduction) or potassium ferricyanide (oxidation) to 8 μM RoxA in 5 mM potassium phosphate buffer, containing 1 μM of the following redox mediators: N,N,N',N'-tetramethylphenylenediamine (E$_{1/2}$ = +260 mV), 2,6-dichlorophenolindophenol (+217 mV), phenazine methosulfate (PMS) (+80 mV), indigo trisulfonate (−70 mV), indigo disinoluate (−125 mV), 2-hydroxy-1,4-naphthoquinone (−152 mV), antracquinone-2-sulfonate (−225 mV),phenosafranine (−252 mV) and methyl viologen (−440 mV). All solutions were made anoxic through several cycles of argon and vacuum; the values of the potentials were recorded when they changed by less than ±2 mV within 5 min. The UV/vis spectra were corrected for turbidity and for contributions from the mediators; potentials are reported versus the standard hydrogen electrode (SHE).

Alternatively, the haem midpoint potentials were determined by the xanthine/xantheine oxidase reox system (Hopkins & Williams, 1995) in an atmosphere of dinitrogen, with resorufin (2 versus the standard hydrogen electrode (SHE)). and for contributions from the mediators; potentials are reported mV within 5 min. The UV/vis spectra were corrected for turbidity and by commercial bicinchoninic acid (BCA) assay (Pierce).

**RESULTS**

**Biochemical characterization of RoxA purified from Xanthomonas sp.**

RoxA was isolated from a polysisoprene latex-grown Xanthomonas sp. culture and purified to homogeneity by the criterion of SDS-PAGE, as described previously (Braaz et al., 2005). The purified enzyme contained two (1.9) haems and two (2.3) iron atoms per molecule of RoxA, as determined by the haem–pyridine chromophore assay (e$_{551}$ =29 mM$^{-1}$ cm$^{-1}$) and by ICP-MS, respectively. These analytical data are in line with the presence of the two classical c-type haem-binding motifs CXXCH found in the amino acid sequence of RoxA (accession no. Y230855). Specific activity (300 μU mg$^{-1}$) and UV/vis absorption properties were almost identical to those reported previously (Braaz et al., 2004, 2005). The yield of pure RoxA varied from preparation to preparation (1–5 mg l$^{-1}$) and depended on the number of remaining latex particles left at the end of the growth phase, typically after 11–14 days of incubation in latex medium. Most likely, a portion of the RoxA remained bound to latex particles, thus reducing the yield of RoxA. Numerous attempts to express recombinant RoxA in E. coli or other Gram-negative bacteria were not successful. Even co-expression of roxA with the cytochrome c maturation (ccm) genes (Arslan et al., 1998; Thöny-Meyer, 1997) did not result in significant expression of RoxA. Thus, a homologous expression system in Xanthomonas sp. was established by construction of a recombinant Xanthomonas sp. strain with roxA cloned in a broad-host-range plasmid, p4782.1, under the control of a rhamnose-dependent promoter. About 6 mg purified RoxA was obtained from 1 l supernatant of recombinant Xanthomonas sp. (p4782.1::roxA) cultures on NB medium supplemented with 0.1% rhamnose. RoxA purified from recombinant Xanthomonas sp. was active and did not differ significantly in its biochemical properties from RoxA that had been purified from latex-grown Xanthomonas sp. wild-type.

**Spectroscopic and redox properties of RoxA**

**RoxA haem centres and relationship to bacterial cytochrome c peroxidases (CCPs)**

UV/vis spectra of pure RoxA as isolated (oxidized) and after addition of dithionite in the absence of dioxygen showed absorption maxima typical for c-type haem centres in the oxidized Fe(III) and reduced Fe(II) state, respectively.
Dithionite-reduced RoxA had a Soret band at 418 nm, the $\beta$-band was at 522 nm, and the $\alpha$-band was split into two separate peaks at 549 and 553 nm. These optical properties, as well as some similarity between the amino acid sequences of RoxA and bacterial CCPs, including the presence of two $\epsilon$-type haem-binding motifs and a conserved region around His517 (Jendrossek & Reinhardt, 2003), suggest that RoxA is distantly related to CCPs. However, the RoxA polypeptide is much larger (71.5 kDa) in comparison with most bacterial CCPs (30–40 kDa). UV/vis-monitored potentiometric titrations of RoxA with dithionite (reduction) and ferricyanide (oxidation) showed fully reversible behaviour of the haem centres in RoxA. The redox mediators used in the titrations did not interfere with the absorption spectra of RoxA. Two Nernst equations with separately variable midpoint potentials had to be used to fit the experimental data, with a first midpoint potential $E^{0}$ at −65 mV. The redox step at the more negative potential was not resolved ($E^{0}$ about −130 to −160 mV; Supplementary Fig. S1). Several repetitions of the experiment did not result in a better resolution of the more negative potential (results not shown).

Experiments using the xanthine/xanthine oxidase reduction system with different redox dyes were performed to obtain a better resolution. Best results were obtained when indigo-disulfate ($E^{0}$ ≈ −125 mV) or hydroxynaphthoquinone ($E^{0}$ ≈ −152 mV) was used, and the midpoint potential of RoxA was determined to be between −110 and −160 mV, which is in agreement with the results from spectrophotometric determinations. However, a more precise determination and separation of the two potentials was not possible. In conclusion, RoxA differs from bacterial CCPs in its redox properties; CCPs generally have clearly separated midpoint potentials ($E^{0}$ ≈ −650 mV; see Table 1 for a comparison of RoxA and CCP properties) (Fülöp et al., 1995).

Determination of the number of electrons that can be taken up by RoxA was performed by UV/vis-monitored titration of RoxA using the 5-deazaflavin photoreduction system. RoxA again showed fully reversible redox behaviour (not shown). After complete photoreduction by the addition of stoichiometric amounts of ferricyanide (with respect to the number of haem groups present) the original (oxidized) spectrum was restored by almost 90%, indicating that approximately two electrons can be abstracted from reduced RoxA (one electron per haem). However, complete reoxidation of RoxA required at least a threefold molar excess of ferricyanide and long incubation times.

**EPR spectroscopy**

The EPR technique is a valuable tool to characterize the spin states of haem centres. The low-temperature EPR spectrum of RoxA (as isolated) recorded at the X-band

![Fig. 1. UV/vis spectra of RoxA. (a) RoxA as isolated from latex culture. (b) RoxA as isolated from latex culture (ox, oxidized; black line), after complete reduction with sodium dithionite (red, reduced; red line) and after incubation under a N₂ atmosphere (N₂; green line) for 48 h. The double maximum of the $\alpha$-band at 549/553 nm indicates the presence of two different haem groups. A partial, slowly continuing reduction of both haem groups (one more readily than the other) was observed after anaerobic incubation for longer periods. The inset figure shows an enlargement of the $\alpha$-region. (c) UV/vis spectrum of RoxA in the region of the $\alpha$-bands upon incubation with NADH/PMS. Both haem centres could be reduced after prolonged incubation times. A time-dependent (from black to blue) increase of the 549 nm peak was observed.](image-url)
(Fig. 2) showed major resonances at $g = 3.09, 2.23$ and $\sim 1.50$, which are typical for hexacoordinate low-spin Fe(III) haem centres (Palmer, 1983), with histidine residues as the fifth and sixth ligands to the iron, having nearly parallel imidazole planes similar to other $c$-type cytochromes. A second set of resonances was found at $g = 3.39$ and $g = 1.92$, originating from a second low-spin Fe(III) centre, with the imidazole planes of the axial histidine ligands being nearly perpendicular (Walker, 1999). The $g$-value of this centre at high magnetic field was extremely broad and thus hard to detect under the experimental conditions. The complete set of resonances was assigned to two magnetically isolated haem centres, since their $g$-values were frequency-independent. Furthermore, there was no signal in the parallel-mode X-band spectrum originating from an integer spin system (data not shown). Spin quantification (CuSO$_4$ standard) of the former signal yielded 0.8 low-spin Fe(III) haem centre.

Table 1. Comparison of RoxA with bacterial CCPs

<table>
<thead>
<tr>
<th>Parameter</th>
<th>RoxA (Xanthomonas sp. 35Y)</th>
<th>CCP</th>
</tr>
</thead>
<tbody>
<tr>
<td>$M_r$ of mature protein (amino acids)</td>
<td>71.5 (648)</td>
<td>35.0 (323)</td>
</tr>
<tr>
<td>Location</td>
<td>Extracellular</td>
<td>Periplasm</td>
</tr>
<tr>
<td>Function</td>
<td>Poly(cis-1,4)-isoprene dioxygenase</td>
<td>Cytochrome $c_{551}$ peroxidase</td>
</tr>
<tr>
<td>Specific activity</td>
<td>0.3 U mg$^{-1}$</td>
<td>$-120$ U mg$^{-1}$</td>
</tr>
<tr>
<td>Haem-binding motifs/haem centres</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Haem iron coordination</td>
<td>His-Fe-(His)</td>
<td>His-Fe-Met</td>
</tr>
<tr>
<td>Haem midpoint potentials</td>
<td>$-65$ mV, $(-130$ to $-160$ mV)</td>
<td>+ (His$^{517}$)</td>
</tr>
<tr>
<td>Conserved ‘MauG’ region</td>
<td>+ (His$^{261}$)</td>
<td></td>
</tr>
<tr>
<td>Peroxidase activity</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>$H_2O_2$</td>
<td>Inhibitor</td>
<td>Substrate</td>
</tr>
<tr>
<td>$Ca^{2+}$</td>
<td>No effect</td>
<td>Activator</td>
</tr>
<tr>
<td>EDTA</td>
<td>No effect</td>
<td>Inhibitor</td>
</tr>
<tr>
<td>External cofactors (electron donor)</td>
<td>None</td>
<td>Cytochrome $c_{551}$</td>
</tr>
<tr>
<td>Activity</td>
<td>+</td>
<td></td>
</tr>
<tr>
<td>Oxidized state</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Mixed valence state</td>
<td>-</td>
<td>+</td>
</tr>
</tbody>
</table>

Properties of rubber oxygenase RoxA

Fig. 2. (a) Experimental (exp) and simulated (sim) perpendicular-mode X-band EPR spectra of RoxA (as isolated) from Xanthomonas sp. at 9.38 GHz, 10 K, 2 mW. Protein at 0.03 mM in 5 mM 1,3-diaminopropane, pH 8.0. Simulated parameters are given in Supplementary Table S1.
of Fe(III) by dithionite was reversible by the addition of air. The high-spin signal at g = 6.1 was often more intense in dithionite-reduced RoxA after reoxidation under air than in RoxA as isolated (results not shown).

**Effects of dioxgen removal and of reducing compounds on UV/vis spectra and activity of RoxA**

Dioxgen was removed from RoxA preparations by eight to 10 cycles of vacuum and flushing with dinitrogen gas. No significant changes were observed in the EPR spectrum (results not shown). However, UV/vis spectroscopy of RoxA under N₂ atmosphere showed a time-dependent increase in the absorbance at 549–553 nm and a concomitant shift of the Soret band from 406 nm to higher wavelengths (418 nm). The observed effects corresponded to the effects of a partial reduction of RoxA (Fig. 1b). Repetition of this experiment and recording of the UV/vis spectra after different incubation periods with dinitrogen gas (1 min to 72 h) indicated that the 549 nm z-band was reproducibly the first to increase, within hours; the 553 nm z-band only increased after prolonged incubation times (up to days). Chemical removal of dioxgen by addition of pyrogallol resulted in similar but more rapid spectral changes. Controls under air showed no significant changes in the UV/vis spectrum (results not shown).

Most known bacterial CCPs (e.g. those of *Pseudomonas aeruginosa*, with the exception of *Nitrosomonas europaea* (Arciero & Hooper, 1994) and *Methylococcus capsulatus* (Zahn et al., 1991) CCPs, require a reduction of one of the two haem centres for activity (mixed-valence state). RoxA, however, was active in the fully oxidized state as isolated in the presence of dioxygen. To investigate the effect of the redox state of the haem groups in RoxA, mild chemical reduction was performed. About a 100-fold excess of NADH was added to RoxA as isolated (2–5 μM) in the presence of the electron mediator PMS. Dioxgen was removed and replaced by dinitrogen. A time-dependent increase of absorbance in the area of the z-bands was detected (Fig. 1c) that was not present in controls without PMS. This increase was significantly stronger and appeared much faster than that obtained without NADH/PMS (Fig. 1b). Apparently, the haem centres of RoxA can be reduced by NADH. Similar to the results obtained for dithionite-treated RoxA, one haem started to be reduced first and at lower amounts of NADH/PMS, which was visible by the earlier and more pronounced increase of the 549 nm band relative to the increase at 553 nm (Fig. 1c). It was not possible temporally to separate the increase at both wavelengths. As observed with dithionite-reduced RoxA, exposure of NADH/PMS-treated RoxA to air led to reoxidation within a few minutes (results not shown). An almost complete reduction of RoxA was found even under oxic conditions when a high molar excess of NADH (10 mM) was used (results not shown). Treatment of RoxA with NADH/PMS resulted in strong inactivation (10 % residual activity at 1 mM NADH/PMS) and indicated that RoxA, unlike most bacterial CCPs, is inactive in the partially reduced state (Table 2 and Supplementary Table S2).

Addition of external nitrogen-containing compounds that could serve as potential haem ligands, such as imidazole, pyridine and related compounds, led to remarkable effects on the activity of RoxA and its UV/vis spectra: incubation of RoxA with 1 mM imidazole, 1-methylimidazole, pyrazole, pyrazine, 2-methylpyrazine, pyridine, 4-methylpyrimidine or pyridazine resulted in significant inactivation of RoxA and caused significant spectral changes (i.e. a shift of the Soret band to longer wavelengths and an increase at 549 nm; Fig. 3a, Supplementary Table S3). The individual velocity of each effect was concentration-dependent for each compound and the reaction proceeded to a concentration-independent end point (molar excess of the compound). Depending on the chemical structures of the test compounds, the spectral changes in difference spectra (UV/vis spectrum of RoxA as isolated minus the UV/vis spectrum recorded after incubation with the test compound) differed slightly (i) in the wavelength of the Soret

<table>
<thead>
<tr>
<th>Compound</th>
<th>Residual activity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydrogen peroxide</td>
<td>≤5</td>
</tr>
<tr>
<td>Ferricyanide</td>
<td>25</td>
</tr>
<tr>
<td>Potassium permanganate</td>
<td>21</td>
</tr>
<tr>
<td>Sodium dithionite</td>
<td>26</td>
</tr>
<tr>
<td>NADH/PMS</td>
<td>10</td>
</tr>
<tr>
<td>Ascorbate</td>
<td>16</td>
</tr>
<tr>
<td>Squalene</td>
<td>5</td>
</tr>
<tr>
<td>x-Tocopherol</td>
<td>≤5</td>
</tr>
<tr>
<td>δ-Tocotrienol</td>
<td>62</td>
</tr>
<tr>
<td>β-Carotene</td>
<td>≤5</td>
</tr>
<tr>
<td>Quinone</td>
<td>11</td>
</tr>
<tr>
<td>Hydroxylamine</td>
<td>≤5</td>
</tr>
<tr>
<td>Hydrazine</td>
<td>≤5</td>
</tr>
<tr>
<td>Phenylhydrazine</td>
<td>≤5</td>
</tr>
<tr>
<td>Sodium azide</td>
<td>86</td>
</tr>
<tr>
<td>Ammonium hydroxide</td>
<td>&gt;95</td>
</tr>
<tr>
<td>CaCl₂</td>
<td>&gt;95</td>
</tr>
<tr>
<td>NaCl</td>
<td>&gt;95</td>
</tr>
<tr>
<td>NaF</td>
<td>92</td>
</tr>
<tr>
<td>FeSO₄</td>
<td>91</td>
</tr>
<tr>
<td>FeCl₂</td>
<td>89</td>
</tr>
<tr>
<td>Imidazole</td>
<td>50</td>
</tr>
<tr>
<td>Histidine</td>
<td>80</td>
</tr>
<tr>
<td>Pyrogallol</td>
<td>43</td>
</tr>
<tr>
<td>DMSO</td>
<td>80</td>
</tr>
<tr>
<td>Sodium hypochlorite</td>
<td>85</td>
</tr>
<tr>
<td>Formic acid</td>
<td>&gt;95</td>
</tr>
<tr>
<td>Oxalic acid</td>
<td>86</td>
</tr>
</tbody>
</table>
Properties of rubber oxygenase RoxA

When the low-molecular-mass test compounds (imidazole, 1-methylimidazole, pyrazole, pyridine, pyrazine and pyridazine) were removed from the buffer by gel filtration, RoxA activity was recovered, although the observed changes in the UV/vis spectra remained (except for pyrazole), which indicated that the compounds were still bound to the enzyme. Removal of imidazole and 1-methylimidazole resulted in only partial recovery (~50 and 70%, respectively) of the activity.

Linear nitrogen-containing compounds such as hydrazine, phenylhydrazine and hydroxylamine, which are known as catalase/peroxidase inhibitors (Wariishi et al., 2000), resulted in inactivation of RoxA (Table 2 and Supplementary Table S2). However, only minimal spectral changes were observed in the wavelength of the signals and a general decrease of haem absorption was detected, suggesting that these compounds led to partial destruction of the enzyme. This conclusion was supported by the detection of a significant increase of the signal for free iron \((g=4.3)\) in EPR measurements (results not shown).

**Effect of oxidizing compounds**

The results described above showed that compounds which can reduce or bind to one of the haem groups are potent inhibitors of the RoxA-mediated cleavage reaction. Next, oxidizing compounds were tested for their effect on RoxA (Table 2 and Supplementary Table S2). Hydrogen peroxide turned out to be a potent inhibitor of RoxA. Concentrations of 0.1 mM or greater completely inhibited the reaction and 10 \(\mu\)M hydrogen peroxide still inhibited the reaction by 80%. Hydrogen peroxide did not change the UV/vis spectrum of RoxA (results not shown). Treatment of RoxA with hydrogen peroxide at 1 mM or higher resulted in slow destruction of the protein, similar to the effect of hydrazine. An interesting effect was found when catalase or HRP was added to RoxA. Peroxidase at 1 \(\mu\)M inhibited the cleavage reaction by 90% (catalase) or 65% (HRP). When hydrogen peroxide at a completely inhibitory concentration and catalase at a partially inhibitory concentration were added simultaneously, significant residual activities were determined that were of the same order as those in the presence of catalase alone. This result indicates that a peroxy compound could be an intermediate of the cleavage reaction. Other oxidizing compounds, such as potassium permanganate and potassium ferricyanide (1 mM), partially inhibited RoxA by 70 and 80%, respectively (Supplementary Table S2 and Table 2).

**Peroxidase activity of RoxA**

No significant peroxidase activity could be detected for purified RoxA in peroxidase assays using cytochrome \(c\) or aromatic phenols such as pyrogallol, guaiacol and 4-aminoantipyrine/2,4-dichlorophenol (Mason et al., 2001) as electron donors. All assays were negative in comparison to wild-type RoxA. When 1 mM hydrogen peroxide and pyrogallol were added simultaneously to RoxA, a significant residual activity was determined, which was of the same order as that of RoxA alone. This result indicates that a peroxy compound could be an intermediate of the cleavage reaction. Other oxidizing compounds, such as potassium permanganate and potassium ferricyanide (1 mM), partially inhibited RoxA by 70 and 80%, respectively (Supplementary Table S2 and Table 2).
with HRP, which served as positive control and which resulted in strong signals. A trace peroxidase activity was detected when 10^3-fold or higher molar amounts of RoxA with respect to HRP were used. Assays of other enzymes containing metal ions, such as catalase and lipoxygenase, also resulted in minor activity at high enzyme concentrations that was even stronger than the one detected for RoxA. We conclude that RoxA has no significant peroxidase activity and that the trace activity can be attributed to a non-specific background that results simply from the presence of metal centres.

**DISCUSSION**

The recently discovered extracellular rubber-oxidizing enzyme RoxA of *Xanthomonas* sp. 35Y cleaves poly-(cis-1,4-isoprene) to give ODTD as the major degradation product of the haem signals was only observed after reduction with dithionite or by NADH in the presence of substrate and potential ligands.

EPR of RoxA in the presence of substrate and potential ligands

The addition of imidazole to RoxA as isolated resulted in the disappearance of the high-spin signal at g = 6.1 (Fig. 4a) and confirmed that imidazole is able to react with one haem centre. Incubation of RoxA with latex did not result in detectable qualitative changes of the EPR spectrum. In particular, the high-spin signal was still present regardless of whether high latex concentrations and/or long incubation times were applied (range of minutes to hours tested). However, the intensity of the haem signals decreased slightly in the presence of latex. This was more pronounced when dioxygen was removed and replaced by argon (Fig. 4b). Interestingly, a weak signal at g = 2.16 appeared in the absence of dioxygen after several hours of incubation and indicated the presence of an organic radical. When EPR measurements were performed in the presence of α-tocopherol or β-carotene (instead of latex), the same signal at g = 2.16 was detected after short incubation times (Fig. 4c). No combination of RoxA assay conditions was found (e.g. presence or absence of latex, dioxygen, dinitrogen or time of incubation) that resulted in a decrease of any of the high- or low-spin signals in EPR measurements by 50% or more. A complete disappearance of the haem signals was only observed after reduction with dithionite in the absence of dioxygen.

**DISCUSSION**

The recently discovered extracellular rubber-oxidizing enzyme RoxA of *Xanthomonas* sp. 35Y cleaves poly-(cis-1,4-isoprene) to give ODTD as the major degradation product by a yet unknown reaction mechanism. The amino acid sequence of RoxA (71.5 kDa for the mature protein) harbours two motifs for covalently attached haems (CXXCH) and has an additional approximately 20 amino acid-long sequence that is conserved in CCPs. This amino acid sequence includes a conserved histidine residue in the consensus sequence ‘P***H517NGSVP’, where an asterisk signifies an amino acid with a hydrophobic side chain (Jendrossek & Reinhardt, 2003). His_517 corresponds to His_261, His_275 and His_244 in *P. aeruginosa*, *Paracoccus pantotrophus* (previously *Paracoccus denitrificans*) and *N. europaea* CCPs, respectively (McGinnity et al., 1996; Fülöp et al., 1995; Ellfolk et al., 1983; Shimizu et al., 2001). These similarities suggest that RoxA is somehow related to bacterial CCPs and that His_517 could have a role in electron transfer between the two centres. On the other hand, all attempts to demonstrate peroxidase activity of RoxA failed, despite the use of a set of different electron donors. Another difference from CCPs is the finding that RoxA is active only in the fully oxidized state (Table 1). Most CCPs must be converted to the mixed-valence state to become active (Echalier et al., 2006) or, as in the case of *N. europaea* CCP, are active in both the oxidized and the half-reduced form (Arciero & Hooper, 1994). RoxA also differs from bacterial CCPs in the small difference in the redox potentials of the two haem centres (Table 1). While the high- and low-potential haem redox potentials of conventional bacterial CCPs differ by more than 0.6 V (Ellfolk et al., 1983; Arciero & Hooper, 1994), a difference of less than 100 mV was found for RoxA. The high-potential haem is His–Met-coordinated in bacterial CCPs (Shimizu et al., 2001). In RoxA, there is probably a bis-His coordination that causes a much more negative potential. This result resembles findings reported for the redox titration of the dihaem protein DHC2 from *Geobacter sulfurreducens* (Heitmann & Einsle, 2005), which has two haem centres with axial bis-histidinyl coordination. The potential of the second haem was also not resolvable. The nona-haem cytochrome c from *Desulfovibrio desulfuricans* (Fritz et al., 2001) contains nine haem groups with midpoint potentials of −100 to −200 mV, whose single potentials are barely distinguishable. The small difference in midpoint potentials of the RoxA haem centres is apparently the reason why it was not possible to obtain the half-reduced state of RoxA in which one haem is completely reduced while the other haem is still oxidized. Similar results have been described for the 42 kDa dihaem c-type protein MauG, which is required for the biosynthesis of tryptophan tryptophylquinone, the prosthetic group of methylamine dehydrogenase (MADH) (Li et al., 2006). In comparison with RoxA, MauG has comparable amino acid similarities to CCPs and also has no peroxidase activity. The authors attribute the MauG midpoint potential values of −159 and −244 mV to a first and second one-electron reduction in an electron-connected system with two equivalent haems (Wang et al., 2003). However, unlike RoxA, MauG has only one x-band signal at 550 nm in the fully reduced state instead of a split signal originating from two separate haems, as in RoxA. Therefore, and because of the preferential reduction of one haem (549 nm x-band) by dithionite or by NADH/PMS (Fig. 1b, c), we conclude that the two haem centres of RoxA are not equivalent.

It should be emphasized that RoxA, unlike many bacterial peroxidases, is an extracellular enzyme that is secreted to the environment and is active without hydrogen peroxide or other cofactors (Table 1) (Braaz et al., 2004; Gilmour et al., 1994; Kobayashi et al., 1989; Pauleta et al., 2001; Thomas & Stocker, 1999). Hydrogen peroxide even inhibits
the reaction. RoxA does not need reductants for activity such as cytochrome c in CCPs. Ions such as Ca$^{2+}$ that are important for P. aeruginosa (Echalier et al., 2008) and Paracoccus pantotrophus CCPs (Pauleta et al., 2008) are not essential for RoxA and do not change the EPR signals, in contrast to peroxidases. RoxA is insensitive to high concentrations (10 mM) of chelating agents such as EDTA (Braaz et al., 2004). All the data mentioned above clearly show that RoxA is not a peroxidase and that it apparently uses a different reaction mechanism for polyisoprene cleavage. This is reasonable, as the function of RoxA is the oxidative cleavage of a carbon–carbon double bond of polyisoprene and not the reduction of hydrogen peroxide or another compound.

Noteworthy results were obtained when low-molecular-mass compounds such as imidazole, pyrazine and related compounds were tested for their effects on the UV/vis spectrum and activity of RoxA. Imidazole and related compounds strongly inhibited RoxA activity (Supplementary Tables S2 and S3) and led to rapid spectral changes. Interestingly, pyrrole, in contrast to the structurally related pyrazole, had no significant effect on RoxA. In pyrrole, the free electron pair of the nitrogen atom is part of the aromatic ring and cannot serve as an electron donor for the haem iron. In pyrazole, the second nitrogen atom has a free electron pair that can react with the haem. Compounds with large side groups such as methyl groups in the direct vicinity of the nitrogen atom possessing the free electron pair (e.g. 2-methylimidazole, Fig. 3a) did not inhibit or only poorly inhibited RoxA and had only a small effect on the UV/vis spectrum. Apparently, a voluminous side group in the direct vicinity of the atom providing the free electron pair prevents binding to haem due to steric hindrance. When the side group is further away from the nitrogen atom, as in 1-methylimidazole, the compound again has a strong effect on activity and the UV/vis spectrum. The optical effects summarized in Supplementary Table S3 mimic a partial reduction of one haem and are probably a consequence of orbital overlapping. This interpretation is supported by the finding that imidazole and the related compounds listed in Supplementary Table S3 affected only one of the two $\alpha$-bands (549 nm) (Fig. 1b). In contrast, the addition of dithionite or NADH/PMS increased both $\alpha$-bands (549 and 553 nm) and indicated a true reduction of the 549 nm haem and a subsequent intramolecular electron transport to the 553 nm haem (Fig. 1c). Since EPR analysis showed that one haem is (partially) present in the high-spin state, it is likely that imidazole and related compounds become

![Overlay EPR spectra of RoxA as isolated (~5 mg ml$^{-1}$) and RoxA with imidazole. The high-spin signal at $g$=6.1 (arrow) disappeared completely. Line 'a' (green), RoxA as isolated; line 'b' (red), RoxA + imidazole (10 mM) incubated for 1 h at room temperature under aerobic conditions. (b) Overlay EPR spectra of RoxA as isolated (a, black), RoxA with rubber latex incubated for 10 min (b, red) or for 90 min (c, green) under argon atmosphere. (c) Part of the EPR spectrum of RoxA as isolated (a, black), with $\beta$-carotene (b, red) and with $\alpha$-tocopherol (c, green) incubated for 10 min at room temperature. An organic radical signal appears in the spectrum of RoxA with substrate analogues (red arrow) at around $g$=2.1 that is barely present in the as-isolated spectrum. The spectrum of RoxA with ascorbate remained unchanged. Ascorbate is an antioxidant, similar to $\beta$-carotene and $\alpha$-tocopherol, but has no structural similarity to isoprene compounds.](http://mic.sgmjournals.org)
bound to this high-spin haem. The disappearance of the high-spin signal at g=6 in the presence of imidazole is in agreement with this conclusion. However, we did not observe the formation of a new Fe(III) low-spin signal.

The presence of two ferric Fe(III) ions in RoxA contradicts previous assumptions that the polyisoprene cleavage reaction begins with binding of dioxygen to an Fe(II) ion. Some evidence for partial and reversible binding of dioxygen to one haem iron was obtained from the weak spectral changes at 549 nm after replacement of an oxicy anoxic (dinitrogen) atmosphere or by removal of dioxygen (vacuum). We assume that the reaction is initiated by binding of the polymeric substrate to RoxA. The detection of a haem centre that was partially high spin, and the finding that polyisoprene cleavage cannot proceed if the sixth coordination site of this haem is occupied (e.g. by imidazole), suggest that the free coordination sphere is the location of the substrate cleavage reaction. Consequently, the polymeric substrate has to be placed in close proximity to the catalytic iron. This would require, however, either that the high-spin haem is located at the surface of the protein or that RoxA has a channel or cleft large enough to take up the polymeric substrate. This haem is accessible principally to small ligands (see above, Fig. 4a). The observation that RoxA activity in the presence of imidazole or related compounds was strongly decreased in a concentration-dependent manner, but was not completely inhibited even by high imidazole concentrations (10 mM, Supplementary Table S2), could be indicative of a competition between polyisoprene and/or dioxygen and the low-molecular-mass compounds for the substrate-binding site. Alternatively, polyisoprene could be bound at the surface of RoxA at a so far unknown rubber-binding site. Polymer-specific substrate-binding sites are well-known structures of other biopolymer-degrading enzymes, such as cellulases, chitinases, other glycohydrolases and polyhydroxalkanoate depolymerases (Behrends et al., 1996; Gilkes et al., 1991; Hiraishi et al., 2010; Rodriguez-Sanoja et al., 2005; Schrempf, 1999; Shinomiya et al., 1998).

EPR data recorded for RoxA after aerobic incubation with latex did not lead to significant disappearance of the high-spin signal, and accordingly a direct binding of latex to the high-spin haem is unlikely or could represent a short transient state that is difficult to observe experimentally. Experiments using reductants such as NADH/PMS or small compounds with free electron pairs showed that RoxA has an intrinsic capacity to abstract electrons from exogenous compounds. We do not know in which form these electrons are stored. One possibility is that the ferric iron is reduced to Fe(II); another is that an electron is stored by selected amino acids such as tyrosine and tryptophan. Reduction of ferric iron was clearly observed by reduction of RoxA with dithionite. In this case, both haems became diamagnetic and all the EPR signals disappeared.

When RoxA was incubated with latex in the absence of dioxygen, the EPR signals tentatively became less intense. This could indicate a reduction of Fe(III) to Fe(II) ions that give no EPR signals. However, quantification of EPR signals is problematic, as repeated recording of EPR data from the same RoxA preparation resulted in variable EPR signal intensities. Because of the low difference in midpoint potentials of the two haems (ΔE°’ ~80 mV) it is reasonable to suppose that a reduction at one haem would cause a subsequent reduction of the other haem instead of a preferred reduction of only one haem, which would explain why EPR signals became less intense. Data obtained from UV/vis spectroscopy during stepwise reduction were in agreement with this assumption (results not shown).

When EPR data were recorded with latex in the absence of dioxygen or with inhibiting substrate analogues such as β-carotene and α-tocopherol, a minor signal at g=2.16 was detected (Fig. 4c). When the same experiment was performed with latex in the presence of air the signal indicative of an organic radical could barely be detected. These results indicate that RoxA is able to partially abstract an electron from a substrate and stabilize the resulting radical. The intermediate formation of organic radicals during catalysis is common for many oxygenases. The radical can be enzyme-bound or can be an enzyme (tyrosyl) radical (Ryle et al., 2003; Siegbahn & Haeffner, 2004; Su et al., 1998). The remaining polyisoprene radical could combine with a dioxygen molecule and resume an electron, resulting in an intermediate peroxide that subsequently dissociates into the keto and aldehyde product ODTD. If we assume that the formation of the radical is a slow process relative to the reaction with dioxygen and to dissociation, the consequence is that no or only very little effect would be seen by EPR measurements in the presence of latex and that the signal indicative of the polyisoprene radical would be seen only in the absence of dioxygen. Note that the RoxA-catalysed cleavage of polyisoprene is a very slow process, with the specific activity of purified RoxA as low as 300 nmol min⁻¹ mg⁻¹ (Braaz et al., 2005). Accordingly, cultivation of Xanthomonas sp. on NR latex takes almost 2 weeks until a significant portion of the substrate has been consumed. We propose that a polyisoprenyl radical is formed in the rate-limiting step triggered by interaction of the substrate possibly with the high-spin haem centre or with an unknown part of RoxA followed by a reaction with dioxygen. The intermediate formation of a radical and of a peroxide is in agreement with the inhibition of the reaction by radical scavengers (ascorbate) and by peroxidases (catalase) (Table 2 and Supplementary Table S2). Future experiments should address the role of each of the haem centres and the importance of potentially electron-transferring amino acid residues such as His517.

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