Sustained Formation of Nitroglycerin-Derived Nitric Oxide by Aldehyde Dehydrogenase-2 in Vascular Smooth Muscle without Added Reductants: Implications for the Development of Nitrate Tolerance


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ABSTRACT

According to current views, oxidation of aldehyde dehydrogenase-2 (ALDH2) during glyceryl trinitrate (GTN) biotransformation is essential for vascular nitrate tolerance and explains the dependence of this reaction on added thiols. Using a novel fluorescent intracellular nitric oxide (NO) probe expressed in vascular smooth muscle cells (VSMCs), we observed ALDH2-catalyzed formation of NO from GTN in the presence of exogenously added dithiothreitol (DTT), whereas only a short burst of NO, corresponding to a single turnover of ALDH2, occurred in the absence of DTT. This short burst of NO associated with oxidation of the reactive C302 residue in the active site was followed by formation of low-nanomolar NO, even without added DTT, indicating slow recovery of ALDH2 activity by an endogenous reductant. In addition to the thiol-reversible oxidation of ALDH2, thiol-refractive inactivation was observed, particularly under high-turnover conditions. Organ bath experiments with rat aortas showed that relaxation by GTN lasted longer than that caused by the NO donor diethylamine/NONOate, in line with the long-lasting nanomolar NO generation from GTN observed in VSMCs. Our results suggest that an endogenous reductant with low efficiency allows sustained generation of GTN-derived NO in the low-nanomolar range that is sufficient for vascular relaxation. On a longer time scale, mechanism-based, thiol-refractive irreversible inactivation of ALDH2, and possibly depletion of the endogenous reductant, will render blood vessels tolerant to GTN. Accordingly, full reactivation of oxidized ALDH2 may not occur in vivo and may not be necessary to explain GTN-induced vasodilation.

Introduction

Aldehyde dehydrogenase-2 (ALDH2) catalyzes formation of nitric oxide (NO) from the antianginal drug nitroglycerin (GTN), resulting in vasodilation mediated by cGMP (Mayer and Beretta, 2008). Efforts to detect GTN-derived NO in blood vessels have consistently failed (Kleschyov et al., 2003; Núñez et al., 2005; Miller et al., 2008). Using a novel fluorescent probe expressed in vascular smooth muscle cells (VSMCs), we have recently demonstrated that NO formation is necessary and sufficient to explain GTN bioactivity (Opelt et al., 2016). Since the reaction of GTN with ALDH2 results in oxidation of the reactive cysteine residue C302 in the catalytic site, reactivation of the enzyme by a reductant is required for sustained turnover (Chen et al., 2002; Beretta et al., 2008). Dithiothreitol (DTT) has proven most efficient in vitro, but the identity of the putative endogenous reductant remains elusive. Dihydrolipoic acid (LPA-H₂) has been suggested as a candidate (Wenzel et al., 2007), although its efficiency appears rather low. In addition to the inactivation of ALDH2 by C302 oxidation that is reversed by reducing agents, particularly DTT, part of the ALDH2/GTN reaction results in thiol-refractive, irreversible inactivation (Beretta et al., 2008). The irreversible component of ALDH2 inactivation is apparently turnover-dependent since it requires the combined presence of GTN and DTT. Based on these observations, we have previously suggested a role for this reaction in the development of nitrate tolerance (Beretta et al., 2008).

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ABBREVIATIONS: ALDH2, aldehyde dehydrogenase-2; C-geNOp, cyan genetically encoded NO probe; DEA/NO, 2,2-diethyl-1-nitrosooxyhydrazine, diethylamine/NONOate; DTT, dithiothreitol; GTN, glyceryl trinitrate, nitroglycerin; LPA-H₂, dihydrolipoic acid; NO, nitric oxide; PETN, pentaerythrityl tetranitrate; PX-12, 2-[(1-methylpropyl)dithio]-1H-imidazole; sGC, soluble guanylate cyclase; VSMC, vascular smooth muscle cell.
To clarify the mechanism of ALDH2-catalyzed GTN biotransformation, we previously characterized several mutants of the enzyme expressed in and purified from *Escherichia coli* (Wenzl et al., 2011). Results obtained with a double mutant in which the cysteine residues adjacent to the active C302 had been replaced by serine residues (C301S/C303S) were of special interest. This mutant lacks the clearance-based pathway that leads to the formation of inorganic nitrite, which is predominant in the wild-type enzyme, whereas the rates of initial NO formation and decay in the absence of DTT are significantly higher than for the wild-type. Thus, mutation of C301 and C303 caused a shift from clearance-based GTN biotransformation to the NO pathway that mediates GTN bioactivity but did not interfere with oxidative inactivation of the enzyme.

Recently, highly selective fluorescent sensors allowing quantification of intracellular NO have become available (Eroglu et al., 2016). Using one of these sensors (cya genetically encoded NO probe (C-geNOp)), we demonstrated that ALDH2 catalyzes a burst of GTN-derived NO in VSMCs, followed by a rapid decrease in the signal in the absence of exogenously added thiols (Opelt et al., 2016). These observations confirmed the essential role of C302 oxidation in the enzymatic reaction but were in apparent conflict with the relatively long-lasting relaxation of isolated blood vessels exposed to a single dose of GTN in organ bath experiments. Similarly, in vivo nitrate tolerance develops within several hours but not within minutes. Conceivably, a redundant present in intact blood vessels could have been lost during isolation and culture of VSMCs. Alternatively, tissues adjacent to the smooth muscle layer could be involved in ALDH2 reactivation.

In the present study, we applied the novel NO detection method with C301S/C303S-ALDH2 expressing VSMCs to compare the kinetics of ALDH2-catalyzed NO formation from GTN with the time course of GTN-induced aortic relaxation. Surprisingly, we found that ALDH2 produces low-nanomolar concentrations of GTN-derived NO in VSMCs, even in the absence of added thiols. Low-level NO formation appears to be sufficient to trigger sustained relaxation of GTN-exposed isolated blood vessels. Thus, GTN bioactivity may be explained without invoking a mysterious endogenous reductant with DTT-like efficiency that is capable of fully restoring the activity of oxidized ALDH2.

**Materials and Methods**

**Materials.** Nitro POHL ampoules (G. Pohl-Boskamp GmbH & Co., Hohenlockstedt, Germany) containing 4.4 mM GTN in 250 mM glucose were obtained from a local pharmacy and diluted in distilled water. 2,2-Diethyl-1-nitroso-oxahydrine (diethyline/NOOate, DEA/NO) was from Enzo Life Sciences (Lausen, Switzerland) and purchased through Eubio (Vienna, Austria). DEA/NO was dissolved and diluted in 10 mM NaOH. Antibiotics and fetal calf serum were obtained from PAA Laboratories (Linz, Austria). Adeno-X293 cells were obtained from Takara Bio Europe (Saint-Germain-en-Laye, France). AdEasy Viral Titer Kit was obtained from Agilent Technologies (Vienna, Austria). Adenovirus encoding C-geNOp and iron(II)ferrocene solution were obtained from NGFI (Graz, Austria). Chloral hydrate was from Fluka Chemie (Vienna, Austria). 2-(I-Methylpropyl)dithio)-1H-imidazole (PX-12) was obtained from Tocris (Bristol, United Kingdom). PX-12 was dissolved in dimethylsulfoxide and diluted in distilled water. Culture media and all other chemicals were from Sigma-Aldrich (Vienna, Austria).

**Generation of Recombinant Adenoviral Vectors.** Mutation of C301 and C303 to serine in human ALDH2 was performed as described by Wenzl et al. (2011). The adenovector encoding C301S/C303S-ALDH2 under the control of the cytomegalovirus promotor was generated as described (Beretta et al., 2012). The adenovector encoding the genetically encoded cyan fluorescent NO probe was generated as described (Eroglu et al., 2016). Adenoviral vectors were propagated in Adeno-X293 cells, purified by CsCl centrifugation, and the titer was determined by using the AdEasy Viral Titer kit (Agilent Technologies).

**Cell Culture and Adenoviral Transfection.** Vascular smooth muscle cells (VSMCs) were isolated from ALDH2 KO mice and immortalized as described (Beretta et al., 2012). Cells were cultured in Dulbecco’s modified Eagle’s medium, supplemented with 10% (v/v) heat-inactivated fetal calf serum, 100 U/ml penicillin, 0.1 mg/ml streptomycin, and 1.25 µg/ml amphotericin in a humidified atmosphere (95% O2/5% CO2) at 37°C. For adenoviral transfection, subconfluent VSMCs were incubated with AdV-CgeNOp (MOI 3) and AdV-C301S/C303S-ALDH2 (MOI 7) in Dulbecco’s modified Eagle’s medium containing 10% fetal calf serum at 37°C for 48 hours.

**Single-Cell NO Measurements with C-geNOp.** GTN-derived NO formation catalyzed by VSMCs expressing C-geNOp and C301S/C303S-ALDH2 was determined by live-cell imaging as described recently (Eroglu et al., 2016). Before fluorescence microscopy, cells were incubated for 10 minutes with nonoxic iron(II) booster solution containing 1 mM iron(II)ferrocene and 1 mM ascorbic acid. During the experiments, cells were perfused in the same physiological buffer without iron(II)ferrocene and ascorbic acid, pH 7.4, containing 140 mM NaCl, 5 mM KCl, 2 mM CaCl2, 1 mM MgCl2, 10 mM Na-glucose, and 10 mM HEPES. Intracellular NO release from 10 µM DEA/NO and 1 µM GTN was measured in the absence and presence of 1 mM dithiothreitol (DTT), 1 mM chloral hydrate, 500 µM LPA-H2, and 10 µM PX-12 as indicated in the text and figure legends. DEA/NO, GTN, DTT, LPA-H2, chloral hydrate, and PX-12 were transiently applied to the cells using a gravity-based perfusion system. To test for the reversibility of ALDH2 inactivation, cells were incubated with 1 µM GTN for 10 minutes or 1 µM GTN and 1 mM DTT for 1 hour in the absence and presence of 10 µg/ml cycloheximide. After varying time intervals of 1, 3, or 6 hours, C301S/C303S-ALDH2-catalyzed NO formation was determined in the presence of 1 µM GTN. Measurements were performed using an Axiosvert 200 M fluorescence microscope (Zeiss, Gottingen, Germany). C-geNOp was excited at 430 nm. Emitted light was collected at 480 nm and 530 nm with a 20× objective and a charge-coupled device camera (Retiga 1350B; QImaging, Surrey, Canada). The fluorescence microscope was controlled using VisiView software (Visitron Systems GmbH, Puchheim, Germany). From the changes in fluorescence intensity of C-geNOp over time, values of ΔI/Δt [%] were calculated with the equation ΔI/Δt [%] = (I − I0)/I0 · 100%, where I is the measured fluorescence intensity over time and I0 the fluorescence intensity of C-geNOp of cells before treatment. On the basis of a previously determined calibration (Eroglu et al., 2016), these fluorescence changes were converted to concentrations of NO using the empirical equation [NO] = 35 × ΔI/Δt/(19.9 − ΔI/Δt). For the sake of clarity, the NO concentrations calculated in this way were plotted on the vertical axes of Figs. 1–5; however, please note that since calibrations were not performed for each experiment, the absolute NO concentrations reported here must be considered crude approximations, in contrast to the relative concentrations within each set of experiments.

**Immunoblotting.** Expression of C301S/C303S-ALDH2 in the absence and presence of cycloheximide was determined by immunoblotting. Cells were harvested and homogenized by sonication (3x5s) in 10 mM Tris buffer, pH 7.4, containing 125 mM KCl, 5 mM ECTA, 2 mM MgCl2, and Complete Protease Inhibitor Cocktail. Protein concentrations were determined with the Pierce BCA Protein Assay Kit using bovine serum albumin as standard. Denatured samples (10 µg) were separated by SDS-PAGE on 10% gels and transferred electrophoretically to nitrocellulose membranes. After blocking with...
Fig. 1. C-geNOp-determined time course of NO concentration in C301S/C303S-ALDH2 overexpressing VSMCs in the presence and absence of GTN, DTT, and chloral hydrate. (A) C301S/C303S-ALDH2 overexpressing VSMCs were perfused with 1 μM GTN and 1 mM DTT as indicated, and the NO concentration was monitored by fluorescence microscopy. See Materials and Methods for further details. The curve shown is the average of 13 traces of three individual experiments (n = 3). The individual traces are shown in Supplemental Fig. S7. (B) Effect of chloral hydrate on NO generation from GTN. Perfusion was carried out with 1 μM GTN and 1 mM chloral hydrate as indicated (n = 3, 16 control traces, 14 curves with chloral hydrate). See Materials and Methods for further details.

![Diagram](image1.png)

5% nonfat dry milk in phosphate-buffered saline containing 0.1% (v/v) Tween-20 for 1 hour, membranes were incubated overnight at 4°C with a primary polyclonal antibody to human ALDH2 (1:20 000; kindly provided by Dr. Henry Weiner) or to β-actin (1:200 000; Sigma). Thereafter, membranes were washed three times and incubated for 1 hour with a horseradish peroxidase-conjugated anti-rabbit (ALDH2) or anti-mouse (β-actin) IgG secondary antibody (1:5000). Immunoreactive bands were visualized by chemiluminescence using ECL detection reagent (Biozym Scientific, Hessisch Oldendorf, Germany) and quantified densitometrically using the Fusion SL system (Peglab, Erlangen, Germany).

Animals and Tissue Preparation. Thoracic aortas were harvested from unsexed Sprague-Dawley rats (bought from Charles River, Sulzdorf, Germany) that were housed at the local animal facility in approved cages. Animals were fed standard chow (Altromin 3023; obtained from Königshofer Futtermittel, Ebergassing, Austria) and received water ad libitum. Animals were euthanized in a box that was gradually filled with CO2 until no more vital signs (cessation of respiration and circulation) were noticed. Subsequently, the thorax of the animals was opened, the thoracic aorta was removed and placed in chilled buffer. Before assessment of vessel function, the endothelium was removed by gently rubbing the intimal surface with a wooden stick. All animal experiments were performed in compliance with the Austrian law on experimentation with laboratory animals (last amendment 2012).

Isometric Tension Vasomotor Studies. For isometric tension measurements, vessel rings were suspended in 5-mL organ baths containing oxygenated Krebs-Henseleit buffer (118.4 mM NaCl, 25 mM NaHCO3, 4.7 mM KCl, 1.2 mM KH2PO4, 2.5 mM CaCl2, 1.2 mM MgCl2, 11 mM d-glucose, pH 7.4) as previously described in detail (Neubauer et al., 2015). After equilibration for 60 minutes, maximal contractile activity was determined with a depolarizing solution containing 100 mM K+ and 1 mM DTT as indicated, and the NO formation was re-established, this time in a continuous fashion that was terminated or restored by arresting or resuming GTN perfusion. When GTN was perfused in combination with DTT, NO formation was re-established, this time in a continuous fashion that was terminated or restored by arresting or resuming GTN perfusion. NO formation occurred in subsequent bouts of GTN perfusion. When GTN was perfused in combination with DTT, NO formation was re-established, this time in a continuous fashion that was terminated or restored by arresting or resuming GTN perfusion. These results confirm previous observations reported by us (Opelt et al., 2016) and are in line with the hypothesis that: 1) NO is formed from a reaction of GTN with ALDH2, 2) the catalytically active cysteine residue of ALDH2 (C302) is oxidized in the process, and 3) rereduction of C302 by a reductant like DTT is required to sustain turnover.

A couple of observations require closer examination. First and foremost, although in the absence of DTT the NO level rapidly decreased after the initial burst, it did not return to zero. A low level of NO, amounting to ~2.0 nM, persisted. This residual NO formation was dependent on the presence of GTN. The veracity of the observed difference between NO levels during and between perfusion was ascertained by paired t tests of the NO levels before and after initiation/cessation of perfusion (see Supplemental Material). We previously

Statistical Procedures. For reasons of clarity, all time courses, except Fig. 4, are shown without error bars. The variability of single curves can be envisioned from the complete set of single curves that is presented for Fig. 1A (NO time courses) and Fig. 6A (relaxation studies). In Fig. 4 the error bars represent the S.E. In Fig. 5, data are presented ± S.D. The veracity of the low level of NO persisting after the burst phase of the first perfusion was estimated by a paired t test (see Supplemental Material). The time-dependent changes in Figs. 3B and 5 were tested by one-way analysis of variance followed by Dunnett’s post hoc test with the initial values as control. Observed NO time courses were fitted nonlinearly to the equations specified in the main text and derived in the Supplemental Material. The appropriateness of the equations was tested by fits to simulated curves (see Supplemental Material). The fitting parameter corresponding to low-level steady-state NO formation (v_{catal}) was tested against 0 with a Mann-Whitney test.

Results

NO Formation from C301S/C303S-ALDH2 Expressing VSMCs upon Perfusion with GTN. As illustrated in Fig. 1A, perfusion of C301S/C303S-ALDH2 overexpressing VSMCs with 1 μM GTN caused a burst of NO that lasted approximately 3 minutes. No burst of NO formation occurred in subsequent bouts of GTN perfusion. When GTN was perfused in combination with DTT, NO formation was re-established, this time in a continuous fashion that was terminated or restored by arresting or resuming GTN perfusion. These results confirm previous observations reported by us (Opelt et al., 2016) and are in line with the hypothesis that: 1) NO is formed from a reaction of GTN with ALDH2, 2) the catalytically active cysteine residue of ALDH2 (C302) is oxidized in the process, and 3) rereduction of C302 by a reductant like DTT is required to sustain turnover.

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demonstrated that the ALDH-2 inhibitor daidzin prevented NO formation in ALDH-2 overexpressing VSMCs (Opelt et al., 2016). To investigate the source of low-level NO generation observed here, we perfused chloral hydrate after the initial burst. As illustrated in Fig. 1B, this resulted in complete inhibition of NO generation, demonstrating the involvement of ALDH2 in low-level NO generation. When chloral hydrate was already present at the start of GTN perfusion, formation of NO was not observed (Supplemental Fig. S10). These results suggest that even in the absence of exogenous reductants, an endogenous compound is present in VSMCs that sustains a low level of steady-state NO generation by ALDH2 (please note that the term steady-state refers to ALDH2, not to the NO concentration; the enzyme steady-state is established long before NO reaches a constant level) (Supplemental Fig. S3).

In addition, there appears to be a discrepancy between the time courses observed in the presence and absence of DTT. Specifically, the addition of DTT caused a decrease in the initial rate of NO formation upon GTN addition, which is unexpected if DTT functions as an additional reductant of oxidized ALDH2 (see Supplemental Fig. S1 and accompanying discussion). This effect is not due to partial inactivation of ALDH2 during the series of GTN perfusions preceding the addition of DTT since it was also observed when DTT was already present at start of the experiment (Fig. 2A). It has been reported that in a purely chemical system, thiols, including DTT, consume NO, which results in diminished apparent rates of NO formation from NO donors such as DEA/NO (Kolesnik et al., 2013). To determine whether this phenomenon occurs in the present system, we perfused C301S/C303S-ALDH2 overexpressing VSMCs with 10 μM DEA/NO in the presence and absence of 1 mM DTT and observed marked diminution of apparent NO release in the presence of DTT (Fig. 2B). Although it cannot be excluded that DTT is affecting the observed NO time course in other ways as well, we conclude that the apparent decrease in the initial rate of NO formation in the presence of DTT is caused by consumption of NO by DTT.

LPA-H2 was reported to support ALDH2-catalyzed NO formation from GTN, although less efficiently than DTT (Chen and Stamler, 2006; Wenzel et al., 2007; Beretta et al., 2008). In agreement with those studies, LPA-H2 caused steady-state formation of NO at a significantly lower level than DTT (Fig. 3A).

We previously reported (Beretta et al., 2008) that, in addition to the inhibition of NO formation by oxidation of C302 that can be reversed by DTT or LPA-H2, ALDH2 is also inhibited irreversibly in the course of GTN biotransformation in a relatively slow reaction that is accelerated upon mutation of C301 and C303 (Wenzl et al., 2011). Likewise, in the present study, NO formation decreased with time under low-steady-state conditions in the absence of reductants (Figs. 1 and 3), as well as in the presence of DTT (Fig. 3B). Plotting the amplitude of the NO concentration against the perfusion number yielded a linear decrease of ∼12.3% per perfusion (Supplemental Fig. S9). This gradual loss of activity was not rescued by the addition of ascorbate or N-acetylcysteine to the perfusion solution (not shown).

Kinetic Analysis: Identification of the Relevant Reaction Steps. The model we adopted for the time course of the NO signal upon perfusion with GTN is described by reactions 1–3, where R stands for an unspecified reductant and X represents any unspecified reaction product in addition to disappearance of NO with the perfusion solution after diffusion out of the cell:

\[
\text{GTN} + \text{ALDH2}_{\text{red}} \rightarrow \text{NO} + \text{ALDH2}_{\text{ox}} \quad \text{(Reaction 1)}
\]

\[
\text{R}_{\text{red}} + \text{ALDH2}_{\text{ox}} \rightarrow \text{R}_{\text{ox}} + \text{ALDH2}_{\text{red}} \quad \text{(Reaction 2)}
\]
The equation we derived for Reactions 1–3 (eq. 1) yielded excellent fits to the averaged time courses both in the absence and presence of DTT (Fig. 4). The decay of the NO signal in the presence of DTT upon termination of GTN perfusion was fitted to a single exponential (eq. 2). In principle, the fitting parameters to eq. 1 will yield values for the rate and the amplitude of the fast initial reaction (burst phase) between GTN and ALDH2red (k1 and [ALDH2red0], Reaction 1), the subsequent slow turnover rate (vcat, Reactions 1 + 2), and the rate of NO decay (k2, Reaction 3). A detailed analysis is afforded in the Supplemental Figs. S4 and S5; Supplemental Table S3.

\[ [\text{NO}] = \frac{v_{\text{cat}}}{k_2} \cdot [\text{ALDH2}^{\text{red}}_0] \exp(-k_1 \cdot t) - \left( \frac{v_{\text{cat}}}{k_2} + \frac{k_1 [\text{ALDH2}^{\text{red}}_0]}{k_2 - k_1} \right) \exp(-k_2 \cdot t) \]

with

\[ v_{\text{cat}} = v_{\text{cat}} \left(1 - e^{-k_1 \cdot t}\right) \]

\[ [\text{NO}]_0 = \Delta [\text{NO}] \cdot e^{-k_1 \cdot t} + [\text{NO}]_0 \]

As an alternative to the approach represented in Fig. 4, we fitted all single curves separately and averaged the obtained fitting parameters, with similar, but not identical, results (see Supplemental Fig. S6). This alternative approach also allowed us to perform significance testing on the low-level steady-state turnover rates. A Mann-Whitney test of the obtained fit values \( v_{\text{cat}} \) against 0 yielded \( P = 0.018 \). Taken together, we estimated an apparent rate constant of 1.5–2.5 minutes\(^{-1}\) for the burst phase in the absence of DTT, which corresponds to a second-order rate constant for the reaction between ALDH2\(^{\text{red}}_0\) and GTN of \(-3 \times 10^4 \text{ M}^{-1} \cdot \text{s}^{-1}\). For the subsequent low steady-state turnover rate (\( v_{\text{cat}} \)), we derived a value of 1.1 nM/min, corresponding to a turnover number of 0.03–0.06 minutes\(^{-1}\). For the reaction in the presence of DTT a clear burst phase could not be discerned, whereas \( v_{\text{cat}} \) increased to 10–20 nM/min. This result suggests that DTT increases the steady-state turnover rate by 10- to 20-fold. Alternatively, one can estimate the stimulation by DTT by comparing the steady-state NO levels observed during the second and subsequent perfusions (1.4–1.9 and 9.3–13.8 nM in the absence and presence of DTT, respectively, see Fig. 1A), which yields a stimulation factor of 6 to 9. Because of the consumption of NO by DTT (Fig. 2), this may still be an underestimation by \(-43\%\). For the decay of the NO signal after termination of GTN perfusion, which we ascribe to diffusion of GTN out of the cell (see Supplemental Material), we obtained a rate constant of 2.0–2.3 minutes\(^{-1}\).

**Regeneration of ALDH2 Activity After GTN Perfusion.** To study the reversibility of ALDH2 inactivation, we varied the time interval between consecutive perfusions with GTN. As illustrated in Fig. 5A, a second perfusion after 1 hour...
resulted in strongly diminished NO formation, whereas NO generation was restored after 3 and 6 hours. This effect did not involve de novo protein synthesis, as it was not affected by cycloheximide (Fig. 5B); ALDH2 expression was comparable under all conditions (Fig. 5C). These results strongly suggest that an endogenous compound present in VSMCs slowly regenerates ALDH2red, confirming the model described by Reactions 1–3. Preincubation with GTN and DTT for 1 hour also resulted in virtually complete inhibition of NO formation from subsequently added GTN, but in this case activity did not recover after 3 or 6 hours (Fig. 5D), indicating irreversible inactivation of the enzyme under high-turnover conditions.

**Relaxation Studies.** To compare ALDH2-catalyzed NO formation in VSMCs with GTN-induced vascular relaxation, we recorded the time-dependent changes in contractile force of endothelium-denuded rat aortic rings exposed to 0.1 and 1 μM GTN and DEA/NO (Fig. 6). At a concentration of 0.1 μM, DEA/NO and GTN relaxed the vessels to similar extent (maximal relaxations of 67.5% ± 3.3% and 72.0% ± 4.7% for GTN and DEA/NO, respectively), but the maximal effect was reached about twice as quickly with DEA/NO (after ∼8.2 minutes compared with 16.8 minutes with GTN, Fig. 6A). Moreover, the effect of GTN lasted significantly longer than that of DEA/NO. The apparent half-lives, measured from the time the maximal effect was reached, were ~9.8 and ~13.2 minutes for DEA/NO and GTN, respectively. As a result, after ~11.3 minutes, relaxation in the presence of 0.1 μM GTN became more pronounced than the response to 0.1 μM DEA/NO. Similar observations were made in the presence of 1 μM of the compounds. Maximal effects increased to 95.0% ± 3.1% and 82.9% ± 5.0% relaxation for DEA/NO and GTN, respectively (Fig. 6A). Again, relaxation increased and decreased faster in the presence of 1 μM DEA/NO than in the presence of 1 μM GTN. Relaxation in response to 1 μM GTN was still quite significant even after 1 hour. Addition of 1 mM chloral hydrate 200 seconds after the addition of GTN (0.1 or 1 μM) immediately blocked relaxation, demonstrating the involvement of ALDH in GTN biotransformation (Fig. 6B).

**Discussion**

Since biotransformation of GTN by ALDH2 results in mechanism-based oxidation of the enzyme, a reductant is

![Fig. 5. Reversibility of the loss of NO generation from GTN by C301S/C303S-ALDH2 overexpressing VSMCs.](image-url)
required to support continuous turnover. Reactivation of ALDH2 occurs in the presence of the artificial thiol DTT, whereas GSH, the most abundant intracellular thiol, and cysteine are ineffective (Chen and Stamler, 2006). These findings led to the assumption that VSMCs contain an endogenous reducing agent with an efficiency similar to that of DTT, but to date this putative reductant has not been identified.

**Sustained Formation of Nanomolar GTN-Derived NO in the Absence of Added Reductants.** To clarify the mechanism of ALDH2 reactivation in vascular smooth muscle, we used the novel fluorescent NO sensor C-geNOp (Eroglu et al., 2016) to study the kinetics of ALDH2-catalyzed GTN bioactivation in isolated VSMCs. As expected from the results obtained with purified ALDH2 (Beretta et al., 2008; Wenzl et al., 2011), perfusion of the cells with GTN led to a short burst of NO that reflected a single turnover of the enzyme in the absence of an exogenously added reductant.

In the presence of added DTT, maximal NO formation persisted, showing that oxidized ALDH2 is efficiently reduced by the thiol; however, although purified ALDH2 becomes fully inactivated in the absence of DTT, the enzyme expressed in VSMCs continued producing small amounts of NO from GTN, even without an added reductant. This observation suggests that VSMCs contain an endogenous reducing agent that causes minor but significant reactivation of oxidized ALDH2.

**Sustained Relaxation of Aortic Rings Exposed to GTN in the Absence of Added Reductants.** The organ bath experiments revealed long-lasting relaxation of aortic rings in the presence of GTN at two different concentrations (0.1 and 1 μM), exceeding the duration of DEA/NO-induced relaxation. This observation is not compatible with a short burst of NO, lasting ~3 minutes in VSMCs in the absence of added reductants. This raises the question as to whether the low-level continuous formation of NO, observed in VSMCs in the absence of DTT, is sufficient to cause long-lasting relaxation of GTN-exposed blood vessels.

Garthwaite and coworkers determined the EC50 values of 0.9 and 0.5 nM NO for cGMP formation by the α1β1 and α2β1 heterodimers of soluble guanylate cyclase (sGC), respectively (Griffiths et al., 2003); they also estimated the physiologic concentration of NO to be in the subnanomolar range and biologic activity of NO to be achieved at concentrations as low as 10 pM (Hall and Garthwaite, 2009). Moreover, blood vessels were reported to contain a large fraction (~95%) of sGC NO receptors (Mergia et al., 2006), and an increase in cGMP levels to less than 5% of the maximal effect caused by NO donors was sufficient for maximal relaxation (Kollau et al., 2005). We estimated a low-level continuous concentration of NO in the perfusion studies of ~2 nM. Considering the lower rates of NO formation by wild-type ALDH2 compared with the double mutant used in the present study, the corresponding concentration with wild-type ALDH2 might be ~0.7 nM. Moreover, overexpression of ALDH-2 may have resulted in increased NO production. On the other hand, the concentration of NO in isolated aortas exposed to 1 μM GTN was probably much greater than the ~2 nM observed in the perfusion studies since a major sink (i.e., perfusion of NO out of the system) was absent in the relaxation studies (see Supplemental Material). Thus, the estimated concentration of GTN-derived NO generated by partially reactivated ALDH2 in VSMCs is in the physiologic range and expected to cause significant sGC activation. Indeed, the inhibition of relaxation by chloral hydrate clearly demonstrates the involvement of ALDH-2 in relaxation by GTN.

By contrast, the NO originating from 0.1 μM DEA/NO, which was roughly equieffective to 0.1 μM GTN in terms of vascular relaxation on a time scale of minutes, is rapidly exhausted owing to the relatively short half-life of DEA/NO, autoxidation of released NO, and unspecified first-order intracellular processes of NO consumption. Thus, compared with GTN-induced relaxation, relaxation by 0.1 μM DEA/NO is short-lived.
Based on these results, we propose that vascular relaxation to GTN is mediated by low steady-state generation of NO that is maintained by reactivation of oxidatively inactivated ALDH2 by a rather inefficient endogenous reductant. The identity of this reductant is unknown, but in view of the abandoned requirement of high potency, LPA-H2 appears to be a promising candidate (Wenzel et al., 2007). We also considered thioredoxin, which is essentially involved in the cellular regeneration of various reduced sulphhydryl proteins, but the specific thioredoxin inhibitor PX-12 (Kirkpatrick et al., 1998) did not affect NO formation by the ALDH2 double mutant overexpressed in VSMCs (10 μM, 24 hours; n = 7, Supplemental Fig. S12).

Development of Nitrate Tolerance. Previous studies indicate that loss of ALDH2 activity essentially contributes to the development of vascular GTN tolerance (Fung, 2004; Sydow et al., 2004; Chen and Stamler, 2006; Beretta et al., 2008; Mayer and Beretta, 2008; Münzel et al., 2011). Specifically, oxidation of ALDH2 that occurs upon reaction with GTN appears to explain both the requirement of an added reductant for GTN biotransformation and the partial reversal of nitrate tolerance by thiols, antioxidants, and various other reducing agents. Since oxidation of ALDH2 occurs in a single turnover, resulting in immediate inhibition of NO formation and decrease of the NO concentration to virtually zero within minutes, it has been suggested that continuous GTN turnover leads to gradual depletion of an unknown endogenous reductant and consequent loss of vascular GTN sensitivity (Fung, 2004; Sydow et al., 2004; Chen and Stamler, 2006; Mayer and Beretta, 2008; Münzel et al., 2011). Involvement of thiol depletion in the development of nitrate tolerance was originally proposed by Needleman and Johnson (1973) and has been discussed controversially ever since because nitrate tolerance is not consistently and not completely reversed by supplementation with thiols (Fung, 2004; Mayer and Beretta, 2008). It is conceivable that gradual depletion of the reductant that supports low level NO formation from GTN by ALDH2 contributes to the loss of GTN sensitivity.

We have previously reported irreversible inactivation of ALDH2 that takes place to a minor extent concurrently with the catalytic cycle (Beretta et al., 2008). Thus, with every turnover, a small fraction of the enzyme is inactivated even in the presence of DTT (see Fig. 5B), and enzyme activity does not recover within 6 hours after washout of GTN under these conditions (compare with Fig. 5D). Our earlier findings with the alternative ALDH2 substrate pentaerythrityl tetranitrate (PETN) suggest that the observed irreversible inactivation of ALDH2 is pharmacologically relevant (Griesberger et al., 2011). As expected, bioactivation of PETN was accompanied by turnover-dependent inactivation of ALDH2 that was prevented and reversed by DTT; however, unlike GTN, PETN did not cause irreversible ALDH2 inactivation. Together with animal and clinical studies consistently showing that PETN does not cause tolerance in vivo (Fink and Bassenge, 1997; Jurt et al., 2001; Mullenhaim et al., 2001; Gori et al., 2003), irreversible ALDH2 inactivation appears to be the essential cause for the development of vascular tolerance to GTN.

Summary and General Conclusions. In summary, we demonstrate that ALDH2 produces low nanomolar concentrations of NO from GTN in VSMC without exogenously added reductants and suggests that this is sufficient for sustained relaxation of GTN-exposed blood vessels. According to our findings, the search for a highly efficient endogenous reductant with DTT-like potency may be futile. LPA-H2 or a similar thiol with relatively low efficiency may fulfill the requirement of partial ALDH2 reactivation. Since biotransformation of GTN is not a physiologic function of ALDH2, there is no compelling reason to assume that the ALDH2/GTN reaction takes place at maximal rates in vivo. GTN-induced vasodilation appears to be the result of a minor side reaction (<10% relative yield) (Wenzel et al., 2011) of the organic nitrate with ALDH2 in combination with the high sGC binding affinity of NO and the high vasodilatory potency of cGMP. Vascular tolerance to GTN may be the consequence of two interdependent processes, depletion of the intracellular reductant that supports low level NO formation, and, probably more relevant, slow mechanism-based irreversible inactivation of ALDH2.

References

Authorship Contributions

Conducted experiments: Opelt, Wolkart.

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