The Saccharomyces cerevisiae quinone oxidoreductase Lot6p: stability, inhibition and cooperativity


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The Saccharomyces cerevisiae quinone oxidoreductase
Lot6p: stability, inhibition and cooperativity

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Dear Prof. Nielsen,

We would like you to consider our paper entitled "The *Saccharomyces cerevisiae* quinone oxidoreductase Lot6p: inhibition, stability and cooperativity" for publication in *FEMS Yeast Research*.

We believe that this paper is suitable for publication because:

1. It documents the steady state enzyme kinetics of Lot6p with NADH as well as with NADPH (which has been the focus of previous studies).
2. It establishes assays to measure the thermal stability of Lot6p and shows that several compounds can stabilize Lot6p in a concentration dependent manner.
3. It shows that the same group of compounds act as inhibitors of Lot6p - and that some of these compounds act with negative cooperativity, inferring information transmission is possible between the active sites. To our knowledge, this is the first report of negative cooperativity in Lot6p (or the functionally related "nitroreductase" enzymes in bacteria).
4. It presents molecular models which suggest a plausible molecular mechanism for this information transmission.
5. It presents an experimental test of this mechanism by altering a key residue in the proposed communication pathway which reduces the degree of cooperativity observed.

We hope that you will agree that this paper is suitable for publication and look forward to hearing from you in the near future.

Yours sincerely,

David J Timson
$h = 0.75$
The *Saccharomyces cerevisiae* quinone oxidoreductase Lot6p:

stability, inhibition and cooperativity

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Running title: Lot6p: stability, inhibition and cooperativity
Abstract

Lot6p (EC 1.5.1.39; Ylr011wp) is the sole quinone oxidoreductase in the budding yeast, *Saccharomyces cerevisiae*. Using hexahistidine tagged recombinant Lot6p, we demonstrated that the enzyme can function with either NADH or NADPH as an electron donor; no cooperativity was observed with these substrates. The NQO1 inhibitor curcumin, the NQO2 inhibitor resveratrol, the bacterial nitroreductase inhibitor nicotinamide and the phosphate mimic vanadate all stabilise the enzyme as judged by thermal scanning fluorimetry. All except vanadate have no observable effect on the chemical crosslinking of the two subunits of the Lot6p dimer. These compounds all inhibit Lot6p’s oxidoreductase activity and all except nicotinamide exhibit negative cooperativity. Molecular modelling suggests that curcumin, resveratrol and nicotinamide all bind over the isoalloxazine ring of the FMN cofactor in Lot6p. However, resveratrol and curcumin stretch further from the cofactor and contact an α-helix that links the two active sites. Mutation of Gly-142, which forms part of this helix, to serine does not greatly affect the kinetics or stability of the enzyme. However, this variant shows less cooperativity towards resveratrol and vanadate. This suggests a plausible hypothesis for the transmission of information between the subunits and, thus, the molecular mechanism of negative cooperativity in Lot6p.

**Keywords:** Quinone oxidoreductase; resveratrol; negative cooperativity; nitroreductase; FMN-containing enzyme
Introduction

Many species contain relatively non-specific quinone oxidoreductases. The best characterised of these is the human NAD(P)H quinone oxidoreductase 1 (NQO1, DT-diaphorase, EC 1.6.5.2). This enzyme has attracted considerable interest because of its role in vitamin K metabolism, its potential as an anti-cancer drug target and its ability to stabilise the tumour suppressor protein p53 (Anwar et al., 2003; Gong et al., 2007; Nolan et al., 2007; Gong et al., 2008; Dinkova-Kostova & Talalay, 2010; Tie et al., 2011). NQO1 is a dimeric enzyme with an FAD cofactor tightly bound to both subunits (Li et al., 1995). This enzyme is a target for the anticoagulant drug dicoumarol (Hosoda et al., 1974; Rase et al., 1976). In humans, there is a second, structurally similar enzyme, NQO2 (EC 1.10.99.2) (Long & Jaiswal, 2000; Vella et al., 2005). The functions and substrates of NQO2 are currently less clear, although it is known that this enzyme is inhibited by resveratrol (Buryanovskyy et al., 2004). Bacteria have a range of quinone oxidoreductases including the NQO1-like MdaB, the tetrameric oxidoreductase WrbA, the azoreductase AzoR and the so-called nitroreductases (e.g. NemA, NfsA, NfsB, YdjA) (Zenno et al., 1996a; Zenno et al., 1996b; Kobori et al., 2001; Lovering et al., 2001; Adams & Jia, 2006; Ito et al., 2006; Patridge & Ferry, 2006; Andrade et al., 2007; Carey et al., 2007; Choi et al., 2008; Hong et al., 2008; Prosser et al., 2010; Mercier et al., 2013). The name of the last group arises due to their ability to catalyse the reduction of a range of nitrogenous compounds, for example nitrobenzene, trinitrotoluene, chloramphenicol, and the pro-drug 5-(aziridin-1-yl)-2,4-dinitrobenzamide (CB1954) (Anlezark et al., 1992; Caballero et al., 2005; Smith et al., 2007; Yanto et al., 2010). In the budding yeast, Saccharomyces cerevisiae, there is one such enzyme, Lot6p (EC 1.5.1.39; Ylr011wp) (Zhang et al., 2001). This enzyme resembles the bacterial azo- and nitroreductases in
that it is dimeric and uses FMN, not FAD, as its tightly bound cofactor (Liger et al., 2004). Despite this difference it still catalyses similar reactions to NQO1 and MdaB (Liger et al., 2004; Sollner et al., 2007).

All quinone oxidoreductases characterised to date have a substituted enzyme (“ping-pong”) mechanism in which a reducing agent (usually NADH or NADPH), transfers a hydride to the FAD or FMN cofactor. The oxidised NAD(P)⁺ then leaves the active site, enabling the second substrate to enter and be reduced by the cofactor (Hosoda et al., 1974; Sollner et al., 2007; Sollner et al., 2009b). The range of possible substrates is broad. In addition to a wide range of quinones and quinone derivatives, human NQO1 has been shown in vitro to catalyse the reduction of aromatic nitrogen compounds and iron(III) ions (Sarlauskas et al., 1997; Anusevicius et al., 2002; Miseviciene et al., 2006; Newsome et al., 2007; Onyenwoke & Wiegel, 2007).

Lot6p was first identified as the protein encoded by one of a diverse group of low temperature (LOT) sensitive genes. This group also included fructose 1,6-bisphosphate aldolase (FBA1 or LOT1), a ribosomal subunit (RPL2B or LOT2), a nucleolar RNA processing enzyme (NOP1 or LOT3), a gene of unknown function (LOT5) (Zhang et al., 2001). The structure of Lot6p has been solved and revealed a typical flavodoxin fold (Liger et al., 2004). The enzyme has been shown to catalyse the NADPH-dependent reduction of a range of biologically relevant quinones including 1,4-benzoquinone, 1,4-naphthoquinone and duroquinone (Sollner et al., 2007) in addition to ferricyanide, azo dyes and nitrocompounds (Liger et al., 2004). The activity with quinones is important in minimising toxicity by these compounds: a strain deleted for LOT6 is less viable than the wild-type in the presence of micromolar
concentrations of 1,4-benzoquinone, whereas one which overexpressed the protein was more viable (Sollner et al., 2007). Like NQO1, Lot6p has roles beyond its enzymatic activity. It interacts directly with the 20S proteasome. Reduction of the FMN in 20S-associated Lot6p results in the recruitment of the transcription factor Yap4p and protects Yap4p from proteosomal degradation (Sollner et al., 2009c). Lot6p also induces apoptosis: strains deleted for LOT6 are less likely to undergo apoptosis when challenged with hydrogen peroxide (Sollner et al., 2009a).

In contrast to mammalian NQO1 and NQO2, relatively little data has been collected on the biochemical properties of Lot6p. Here we extend previous enzymological and biochemical work by describing Lot6p’s kinetics, inhibition and stability to thermal denaturation. We describe, for the first time in detail, the enzyme’s kinetics with NADH as a reducing agent and quantify its inhibition by the organic compounds curcumin, resveratrol and nicotinamide. These compounds were selected for their ability to bind to the active sites of NQO1, NQO2 and bacterial nitroreductases respectively (Buryanovskyy et al., 2004; Tsvetkov et al., 2005; LinWu et al., 2012). Vanadate ions were also studied as a phosphate analogue. In some cases, but not all, these compounds acted with negative cooperativity. Models of Lot6p in complex with these compounds were built in order to help explain their different behaviour. These models informed site-directed mutagenesis studies which further illuminated cooperativity in Lot6p.

**Materials and Methods**

*Expression and purification of wild type and mutant Lot6p*
The coding sequence for LOT6 was amplified by PCR using a single colony from *Saccharomyces cerevisiae* strain BY4741 (MATa; his3Δ1; leu2Δ0; met15Δ0; uralΔ0; EUROSCARF Consortium, Germany) as a template. The product was inserted into pET-46 Ek/LIC (Merck, Nottingham, UK) and competent *E. coli* cells (NovaBlue GigaSingles, Merck) were transformed with the annealed Ek/LIC vector and insert. Colonies resulting from this transformation were picked and grown shaking at 37 °C overnight in 5 ml of Luria-Burtani (LB) broth supplemented with ampicillin (100 µg ml⁻¹), plasmid DNA was isolated using a miniprep kit (Yorkshire Biosciences, York, UK) and the presence of inserts verified by restriction digestion and PCR. Competent *E. coli* HMS174(DE3) were transformed with the plasmid DNA and a single colony picked and grown shaking overnight at 37 °C in 100 ml of LB supplemented with ampicillin (100 µg ml⁻¹); DNA was isolated using a midiprep kit (Sigma, Poole, UK) and sequenced by GATC Biotech, (Konstanz, Germany).

Another colony was picked and grown overnight shaking at 37 °C in 5 ml of LB broth supplemented with 100 µg ml⁻¹ ampicillin. The culture was diluted into 1 l of the same medium and grown at 37 °C for 4 h until mid-log phase was reached (A₆₀₀~0.6). The culture was induced by the addition of IPTG to a final concentration of 1.7 mM, and grown for a further 4 h at 37 °C. The cells were then collected by centrifugation at 4200 g for 15 min and the pellets resuspended in buffer R (50mM HEPES-OH pH 7.4; 150mM NaCl; 10% v/v glycerol). Sonication on ice (30 s pulses at 100 W with 30 s breaks for cooling) broke the cells and insoluble material was removed by centrifugation (24,000 g for 15 min). The supernatant was poured into a column containing 1 ml cold nickel agarose resin (His-select, Sigma, Poole, UK) which had been pre-equilibrated with 25 ml of buffer A (50 mM HEPES-OH pH 7.4; 500 mM...
NaCl; 10% v/v glycerol), and allowed to pass through under gravity. The column was then washed with 50 ml of buffer A. Lot6p was eluted by application of two 2 ml volumes of buffer A supplemented with imidazole (250 mM). The fraction(s) containing Lot6p was identified by 10% SDS-PAGE and were dialysed overnight at 4 °C against 1 l of buffer R supplemented with 2 mM dithiothreitol (DTT).

Site-directed mutagenesis was carried out by the QuikChange method (Wang & Malcolm, 1999) and the mutation verified by DNA sequencing of the complete LOT6 coding sequence (GATC Biotech). The mutant enzyme was expressed and purified using the same procedures as for the wild type.

Protein concentrations were determined using the method of Bradford (Bradford, 1976) using BSA as a standard and the purified Lot6p stored in 20 µl and 50 µl aliquots frozen at -80 °C.

**Enzyme kinetic analysis of Lot6p**

Lot6p activity was measured at 30 °C in 50 mM HEPES, pH 7.3 using either NADH or NADPH as the electron donor and DCPIP as the electron acceptor. Reaction rates were determined by measuring the rate of decrease in absorbance at 600 nm resulting from the reduction of DCPIP (70 µM). The enzyme concentration ranged from 1.25 nM to 10 nM. Absorbance readings were taken every 5 s and all reactions were carried out in triplicate in 96-well plates. A calibration curve of $A_{600}$ against [DCPIP] was constructed in triplicate in a 96-well plate to determine a value for $\varepsilon L$ in the Beer Lambert Law, $A=\varepsilon Lc$ were $A$ is absorbance, $c$ is concentration, $\varepsilon$ is the molar extinction coefficient of DCPIP and $L$ is the pathlength. The resulting equation of the
line was used to convert all rates in terms of change in absorbance at 600 nm, to rates in terms of change in concentration of DCPIP.

To ensure initial, enzyme-catalyzed rates were obtained, rates for the non-enzymatic, direct reduction of DCPIP by both NADH and NADPH were determined at each concentration of electron donor with 70 µM DCPIP. These background rates were then subtracted from the rates obtained from the linear section at the beginning of each progress curve with enzyme included.

$K_{m,app}$ and $V_{max,app}$ values for each electron donor were determined, at a constant DCPIP (70 µM) by plotting enzyme-catalyzed rate ($v$) divided by the enzyme concentration ([E]) against the corresponding NADH or NADPH concentration. The data were fitted to a modified form of the Michaelis-Menten equation (1) using non-linear curve fitting (Marquardt, 1963) as implemented in Graphpad Prism 5.0 (GraphPad Software Inc, CA, USA.). All points were weighted equally.

$$\frac{v}{[E]} = \frac{k_{cat,app}[S]}{K_{m,app}+[S]} \quad (1)$$

where $k_{cat,app}$ is the apparent turnover number (equal to the apparent maximal rate, $V_{max,app}$, divided by the enzyme concentration), $K_{m,app}$ is the apparent Michaelis-Menten constant, [E] is the concentration of Lot6p dimer and [S] is the concentration of either NADH or NADPH.

**Inhibition kinetics**

The effect of potential inhibitors (resveratrol, nicotinamide, curcumin and vanadate ions) on the enzyme-catalyzed rate measured at three concentrations of NADH and three concentrations of NADPH with a constant DCPIP concentration (70 µM).
Dilution series of resveratrol and curcumin were prepared such that the final volume of DMSO in each reaction mixture was 0.5% (v/v). Nicotinamide and sodium metavanadate were dissolved in 50 mM HEPES pH 7.3. Dixon plots (Dixon, 1953) for each concentration of electron donor were constructed and the apparent inhibition constant, \( K_{i,app} \), was obtained by determining the inhibitor concentration at which the lines corresponding to each NAD(P)H concentration intercepted.

A concentration range for each inhibitor was chosen based on these \( K_{i,app} \) values and an inhibitor titration was carried out using a concentration approximately equal to \( K_{m,app} \) for NADH (250 µM) or NADPH (100 µM) and DCPIP (70 µM). Linearized Hill plots \((-\log_{10}(v/(v_0-v))\) against \(-\log_{10}[\text{inhibitor}]\), where \( v \) is the rate of the inhibited reaction and \( v_0 \) is the rate in the absence of inhibitor) were constructed for each inhibitor; the gradient of these is the Hill coefficient, \( h \) (Hill, 1910).

**Crosslinking and limited proteolysis**

Increasing concentrations of the chemical cross-linkers \( \text{bis} \text{ulfsu} \text{cinimidy} \text{ls} \text{uberate (BS\textsuperscript{3}) and \( N\text{-} (3\text{-dimethylaminopropyl})-N'\text{-ethylcarbodiimide hydrochloride (EDC) were} \} \) added to a constant concentration of Lot6p (35 µM dimer) which had been preincubated at 30 °C for 5 min. The reaction was allowed to proceed for 30 min after which time it was stopped by the addition of an equal volume of SDS loading buffer (120 mM TrisHCl pH 6.8, 4% (w/v) SDS, 20% (v/v) glycerol, 5% (w/v) bromophenol blue, 1% (w/v) DTT). Samples were denatured by heating at 95 °C for 5 min and analysed by 10% SDS PAGE. An optimum concentration of each crosslinker was chosen and the reaction repeated in the presence of each inhibitor.
Differential scanning fluorimetry

Lot6p was diluted in 50 mM HEPES, pH 7.3 to final concentration of 0.25 µM dimer in a final volume of 20 µl. The samples were loaded, in triplicate, into a Rotor-Gene Q cycler (Qiagen) and the high resolution melt protocol was used. Lot6p was subjected to an increase in temperature from 25 °C to 95 °C in steps of 1 K (no gain optimisation) with excitation at 460 nm and emission measured at 510 nm throughout, thereby exploiting the fluorescence of its cofactor, FMN (Forneris et al., 2009). The melting temperature \( T_m \) of the enzyme was determined from the first derivative of the melting curve, using the inbuilt analysis software. A dilution series of each ligand (except curcumin which fluoresces in the same region as FMN (Chignell et al., 1994)) was prepared such that the final volume of DMSO in the reaction mixture was 0.5% v/v. Reactions were prepared in triplicate and kept on ice until loading into the instrument. The melting temperature for each concentration of ligand was plotted against the corresponding concentration of ligand and the data were fitted to equation (2) using non-linear curve fitting in GraphPad Prism.

\[
\Delta T_m = \Delta T_{m,\text{max}} [\text{ligand}]/(K_{D,\text{app}} + [\text{ligand}]) \quad (2)
\]

where \( \Delta T_{m,\text{max}} \) is the maximum, limiting change in melting temperature \( T_m \), and \( K_{D,\text{app}} \) is the apparent dissociation constant for ligand and Lot6p.

Molecular modelling

Models of Lot6p with ligands bound were based on the experimentally determined x-ray crystal structure (PDB: 1T0I) (Liger et al., 2004). This structure was aligned with that of NQO2 with resveratrol bound (PDB: 1SG0 (Buryanovskyy et al., 2004)) using PyMol (http://www.pymol.org). Lot6p and the resveratrol molecules were saved into a single .pdb file which was then energy minimised and computationally
solvated using YASARA (Krieger et al., 2009) to generate the final model.

Nicotinamide was modelled using the structure of NQO1 bound to NADP⁺ (Li et al., 1995). In this case, the structural alignment between the proteins was poor and so the structures were aligned using three atoms (C8, N3 and N10) in the isoalloxazine ring of the FAD or FMN cofactor. NADP⁺ was then inserted into the Lot6p structure and this was used as a template to insert a nicotimamide molecule (from the RCSB PDB Ligand Expo; http://ligand-expo.rcsb.org/) which overlapped the nicotinamide moiety of NADP⁺. The resulting structure was then energy minimised in YASARA to create the final model. Since there are no currently available structures of curcumin bound to a quinone oxidoreductase, this molecule was fitted into the active site by aligning the O1, C9 and C14 atoms of resveratrol with equivalent atoms in one ring of curcumin (from Ligand Expo) and assuming that the molecule adopts a similar orientation in Lot6p’s active site. This initial model was then energy minimised in YASARA to generate the final model. The three final models are available as supplementary data to this paper.

Results

Recombinant expression and dimerisation of Lot6p

Lot6p can be expressed in, and purified from, E. coli with a typical yield of approximately 12 mg purified protein per litre of initial culture (Fig 1a). Crosslinking with both EDC and BS³ showed that Lot6p is a dimer (Fig. 1b) and the amount of crosslinking observed was unchanged in the presence of resveratrol, nicotinamide and curcumin; however, vanadate ions reduced the amount of crosslinked product observed (Fig. 1c).
Steady state enzyme kinetics: no cooperativity with either NADH or NADPH

Lot6p can use both NADH and NADPH as electron donors; the $K_{m,\text{app}}$ for NADH was 273±35 µM and for NAD(P)H it was 131±22 µM. The $k_{\text{cat,app}}$ values for NADH and NADPH were 297±11 s$^{-1}$ and 449±29 s$^{-1}$ respectively (Fig. 2). Lot6p exhibits Michaelis-Menten kinetics (i.e. no detectable cooperativity) towards both electron donors with a Hill coefficient of 1.05±0.08 and 0.99±0.07 for NADH and NADPH respectively (Fig. 2).

Quinone oxidoreductase inhibitors stabilise Lot6p

The melting temperature of Lot6p, as judged by TSF, was 56.9±0.4 °C in HEPES buffer. Lot6p is slightly stabilised by phosphate ions; the $T_m$ increased to 58.9±0.4 °C in 50 mM phosphate buffer (Supplementary Fig. S1). Resveratrol and nicotinamide stabilised Lot6p in a saturatable, concentration-dependent manner. The apparent dissociation constants with resveratrol and nicotinamide were 0.091±0.026 mM and 24.5±3.4 mM respectively (Fig. 3).

Inhibition of Lot6p is negatively cooperative with some compounds

The oxidoreductase activity of Lot6p was inhibited by curcumin, resveratrol, nicotinamide and vanadate. Dixon plots for all four compounds with NADH and NADPH as the reducing agent intersected as expected for competitive inhibition (Dixon, 1953). For each inhibitor except vanadate, the $K_{i,\text{app}}$ values obtained with NADH and NADPH were similar (Fig. 4; Table 1). Of the organic compounds, nicotinamide has the highest $K_{i,\text{app}}$ followed by curcumin and resveratrol which was the most effective inhibitor tested (Fig. 4; Table 1). Lot6p exhibits negative cooperativity with the inhibitors resveratrol, curcumin, and vanadate when inhibiting
with respect to both NADH and NADPH; in contrast, nicotinamide does not induce negative cooperativity in Lot6p with either electron donor (Fig. 5; Table 2).

Molecular modelling of Lot6p and inhibitors
Nicotinamide, resveratrol and curcumin were all predicted to bind in the enzyme’s active site, lying across the partly exposed surface of the isoalloxazine ring of the FMN cofactor, consistent with their role as competitive inhibitors (Fig. 6, left column). The two larger molecules, resveratrol and curcumin, extend beyond the active site into a cleft between the two polypeptide chains of the Lot6p dimer. Interestingly, resveratrol is predicted to make contact with an α-helix (residues Ser-130 to Leu-143) which links the two active sites of the dimeric enzyme (Fig. 6, right column). Curcumin contacts the equivalent α-helix in the other subunit. The smaller molecule, nicotinamide, does not contact this helix. Vanadate could, potentially, bind at a variety of sites including those for the three phosphate groups in NADPH. In the absence of any structures of quinone oxidoreductases bound to vanadate, no modelling was attempted with this inhibitor.

Mutation of a key residue reduces the degree of negative cooperativity
From the molecular models, we noticed that resveratrol contacts the α-helix which links the two active sites close to a glycine residue (Gly-142). Given the role of glycine in protein flexibility and the importance of such mobility in phenomena such as cooperativity (Goodey & Benkovic, 2008), we reasoned that altering this residue might affect cooperativity towards the inhibitors. Alteration of Gly-142 to serine results in an active enzyme, which is dimeric, has Michaelis-Menten kinetics with NADH and is inhibited by resveratrol and vanadate (Supplementary Fig S2, S3, S4;
Table 3). The apparent inhibition constants for these compounds with respect to NADH were increased approximately two-fold (Table 3). The enzyme is slightly more stable towards thermal denaturation than the wild type ($T_m$ values of 60.9±0.2 °C and 61.5±0.6 °C in phosphate and hepes buffer respectively; Supplementary Fig. S5). This is consistent with a less flexible overall structure. Resveratrol and nicotinamide both stabilise the G142S variant protein, with apparent dissociation constants similar to those of the wild-type (Table 3, Supplementary Fig. S6). However, the degree of negative cooperativity towards resveratrol and vanadate was less than that seen with the wild type; for both compounds, the Hill coefficient rose towards one (Table 3, Fig. 7).

Discussion

Previous studies of Lot6p have concentrated on its activity with NADPH as an electron donor (Sollner et al., 2007). Here, it is demonstrated that NADH functions as an electron donor, albeit with a higher apparent Michaelis constant and lower turnover number. Lot6p’s location in the cytoplasm (Sollner et al., 2007) means that it is more likely to encounter NADPH since the NADPH:NADH concentration ratio in the cytoplasm is generally greater than one (Jacobson & Kaplan, 1957). However, it has been demonstrated that, in *S. cerevisiae*, environmental changes (e.g. altered nutrient sources) can alter this ratio (Satrustegui et al., 1983; Nissen et al., 2001; Celton et al., 2012; Ask et al., 2013). Therefore, both reducing agents are likely to be important in vivo. Despite the dimeric nature of the enzyme, and the involvement of residues from both polypeptides in the two active sites, no cooperativity with respect to NADH or NADPH was detected in our experiments.
Through thermal scanning fluorimetry experiments, we established that resveratrol,
nicotinamide and vanadate ions all bind to, and stabilise, Lot6p. Furthermore, these
molecules (and curcumin) competitively inhibit the enzyme with respect to NAD(P)H
– all except nicotinamide with negative cooperativity. Negative cooperativity towards
inhibitors has previously been observed with mammalian NQO1 when inhibited with
dicoumarol – although the molecular mechanism and physiological significance
remains obscure (Rase et al., 1976). The molecular models (Fig. 6) suggest a
potential mechanism for inter-active site communication and also why nicotinamide is
unable to initiate this. Both resveratrol and curcumin are predicted to contact an
extended α-helix at the interface between the two polypeptides of the dimer. This
helix links the two active sites and so it is plausible that alterations to its conformation
induced by an inhibitor binding at one active site may affect the conformation of the
second active site. Nicotinamide is predicted to bind parallel to the FMN cofactor and
does not make contact with this helix; therefore it would be unable to initiate
information transmission between the active sites. Given that, like all models, these
should be treated with some caution, we tested the hypothesis by mutating a glycine
residue in this helix. The resulting protein had similar kinetic properties and stability
to the wild type protein, but showed less negative cooperativity (i.e. higher value of $h$)
towards resveratrol and vanadate. This result supports the hypothesis that this helix
plays a key role in transmitting information between the active sites in Lot6p. It also
confirms that the non-Michaelian kinetics observed with inhibitors arises from
genuine inter-active site cooperativity and not from intra-molecular aggregation or
some other artefact.
The physiological importance of the negative cooperativity observed in Lot6p is not yet clear. In many signalling systems, negative cooperativity functions to extend the concentration range over which the system is sensitive to concentration changes (Ferrell, 2009). Therefore it is possible that some, as yet unidentified, naturally occurring molecule in the yeast cytoplasm regulates Lot6p’s activity and perhaps also its role in apoptosis or in the proteasome.

Acknowledgements

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For Peer Review


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Figure legends

Figure 1: **Expression and dimerization of recombinant Lot6p.** (A) SDS-PAGE (10%) showing the progress of a typical expression and purification of Lot6p. (B) Crosslinking of recombinant Lot6p with EDC (0, 4, 8, 16, 32, 64 mM) and BS₃ (0, 50, 100, 200, 400, 800 µM). The effects of solvent (DMSO 1 %(v/v)) and ligands (resveratrol 0.7 mM, curcumin 40 µM, nicotinamide 160 mM, vanadate 18 mM) on crosslinking by EDC (32 mM) and BS₃ (800 µM). In (A), (B) and (C) lane M represents the molecular mass markers (with masses shown to the left of the gel in kDa).

Figure 2: **Lot6p kinetics with electron donors.** Initial rates of DCPIP (70 µM) reduction were measured and plotted as a function of either NADH or NADPH concentration (top graphs). These data were replotted as linear Hill plots (bottom graphs). Each point represents the mean of three independent determinations and the error bars the standard error of these means.

Figure 3: **Ligands stabilise Lot6p against thermal denaturation.** Increasing concentrations of resveratrol and nicotinamide were mixed with Lot6p (0.25 µM) in Hepes-OH buffer (pH 7.3) and the melting temperature determined by TSF. The first derivative curves of the fluorescence against temperature (top) were used to determine Tₘ values, which were plotted against ligand concentration (bottom). Each point represents the mean of three values and error bars the standard errors of these means.

Figure 4: **Ligands inhibit the oxidoreductase activity of Lot6p.** Initial rates of DCPIP reduction were measured at different NADH (left column) and NADPH (right)
concentrations in the presence of increasing concentrations of ligand. See Materials and Methods for the conditions of these experiments. Dixon plots (1/v against [Inhibitor]) were constructed and the $K_{i,app}$ estimated for each inhibitor. Each point represents the mean of three separate determinations and the error bars the standard errors of these means.

Figure 5: Some inhibitors exhibit negative cooperativity. Linear Hill plots with (A) NADH and (B) NADPH as the reducing agent were constructed in order to determine the Hill coefficient, $h$. See Materials and Methods for the conditions of these experiments. Each point represents the mean of three separate determinations and the error bars the standard errors of these means.

Figure 6: Molecular modelling predicts how the inhibitors bind to Lot6p. In the left hand column a close up of the active site with inhibitor bound is shown. In the right hand column the inhibitor is shown bound to one active site and, in the case of curcumin and resveratrol, contacting an α-helix (orange) which connects the two active sites of the enzyme. (Note that curcumin and resveratrol contact the equivalent helix in different subunits.) FMN is shown in yellow.

Figure 7: The G142S variant of Lot6p has reduced negative cooperativity towards some inhibitors. Linear Hill plots showing the effect of resveratrol and vanadate on the NADH mediated reduction of DCPIP. See Materials and Methods for the conditions of these experiments. Each point represents the mean of three separate determinations and the error bars the standard errors of these means.
### Tables

**Table 1:**

Inhibition of Lot6p.

<table>
<thead>
<tr>
<th>Inhibitor</th>
<th>$K_{i,app}$/mM with NADH</th>
<th>$K_{i,app}$/mM with NADPH</th>
</tr>
</thead>
<tbody>
<tr>
<td>Resveratrol</td>
<td>0.067±0.003</td>
<td>0.063±0.033</td>
</tr>
<tr>
<td>Curcumin</td>
<td>5.8±1.0</td>
<td>5.0±2.5</td>
</tr>
<tr>
<td>Nicotinamide</td>
<td>20.0±4.8</td>
<td>11.9±4.5</td>
</tr>
<tr>
<td>Vanadate</td>
<td>21.1±4.3</td>
<td>3.4±0.2</td>
</tr>
</tbody>
</table>

Values are the mean of the intersection points in the Dixon plot and the errors are the standard deviation of these means.
Table 2:

Hill coefficients ($h$) for the inhibition of Lot6p.

<table>
<thead>
<tr>
<th>Inhibitor</th>
<th>Hill coefficient ($h$) with NADH</th>
<th>Hill coefficient ($h$) with NADPH</th>
</tr>
</thead>
<tbody>
<tr>
<td>Resveratrol</td>
<td>0.75±0.11</td>
<td>0.83 ±0.05</td>
</tr>
<tr>
<td>Curcumin</td>
<td>0.60±0.09</td>
<td>0.61 ±0.10</td>
</tr>
<tr>
<td>Nicotinamide</td>
<td>1.02±0.07</td>
<td>1.03 ±0.05</td>
</tr>
<tr>
<td>Vanadate</td>
<td>0.44±0.05</td>
<td>0.71 ±0.01</td>
</tr>
</tbody>
</table>

Values were determined by constructing three separate linearised Hill plots, each based on triplicate data. The values quoted are the means of the three estimates of $h$ and the errors the standard deviations of these means.
Table 3

Properties of the G142S variant of Lot6p

<table>
<thead>
<tr>
<th>Property</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$K_{m,\text{app}}$ (NADH)</td>
<td>1200±110 µM</td>
</tr>
<tr>
<td>$k_{\text{cat,app}}$ (NADH)</td>
<td>160±5 s⁻¹</td>
</tr>
<tr>
<td>$h$ (NADH)</td>
<td>1.08±0.03</td>
</tr>
<tr>
<td>$K_{i,\text{app}}$ (Resveratrol)</td>
<td>0.059±0.030 mM</td>
</tr>
<tr>
<td>$K_{i,\text{app}}$ (Vanadate)</td>
<td>46±16 mM</td>
</tr>
<tr>
<td>$K_{D,\text{app}}$ (Resveratrol)</td>
<td>0.22±0.07 mM</td>
</tr>
<tr>
<td>$K_{D,\text{app}}$ (Nicotinamide)</td>
<td>20.8±2.3 mM</td>
</tr>
<tr>
<td>$h$ (Resveratrol)</td>
<td>0.94±0.04</td>
</tr>
<tr>
<td>$h$ (Vanadate)</td>
<td>0.82±0.08</td>
</tr>
</tbody>
</table>
**Supplementary Figure S1:** Phosphate ions slightly stabilise Lot6p. Melting point determination of Lot6p (0.25 µM) by TSF in Hepes-OH (pH 7.3) and phosphate buffer (pH 7.4).

**Supplementary Figure S2:** The G142S variant of Lot6p forms dimers. Upper panels: Crosslinking of the G142S variant with EDC (0, 4, 8, 16, 32, 64 mM) and BS$_3$ (0, 50, 100, 200, 400, 800 µM). Lower panels: The effects of solvent (DMSO 1 % (v/v)) and ligands (resveratrol 0.7 mM, curcumin 40 µM, nicotinamide 160 mM, vanadate 18 mM) on crosslinking by EDC (32 mM) and BS$_3$ (800 µM). In all gels, lane M represents the molecular mass markers (with masses shown to the left of the gel in kDa).

**Supplementary Figure S3:** The G142S variant of Lot6p has Michaelis-Menten kinetics. Initial rates of DCPIP (70 µM) reduction were measured and plotted as a function of NADH (upper panels). The data were replotted as a linear Hill plot (lower panels). Each point represents the mean of three independent determinations and the error bars the standard error of these means.

**Supplementary Figure S4:** The G142S variant of Lot6p is inhibited by resveratrol and vanadate. Initial rates of DCPIP reduction were measured at different NADH concentrations in the presence of increasing concentrations of inhibitor. See Materials and Methods for the conditions of these experiments. Dixon plots (1/v against [Inhibitor]) were constructed and the $K_{i,app}$ estimated for each inhibitor. Each point represents the mean of three separate determinations and the error bars the standard errors of these means.
Supplementary Figure S5: The G142S variant of Lot6p has similar thermal stability to the wild type. Melting point determination of G142S (0.25 µM) by TSF in Hepes-OH (pH 7.3) and phosphate buffer (pH 7.4).

Supplementary Figure S6: The G142S variant of Lot6p is stabilised by resveratrol and nicotinamide. Increasing concentrations of resveratrol and nicotinamide were mixed with G142S (0.25 µM) in Hepes-OH buffer (pH 7.3) and the melting temperature determined by TSF. The first derivative curves of the fluorescence against temperature were used to determine $T_m$ values, which were plotted against ligand concentration. Each point represents the mean of three values and error bars the standard errors of these means.