The Golgi α-1,6 mannosyltransferase KlOch1p of Kluyveromyces lactis is required for Ca²⁺/calmodulin-based signaling and for proper mitochondrial functionality

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Abstract

Background: Protein N-glycosylation is a relevant metabolic pathway in eukaryotes and plays key roles in cell processes. In yeasts, outer chain branching is initiated in the Golgi apparatus by the alpha-1,6-mannosyltransferase Och1p.

Results: Here we report that, in Kluyveromyces lactis, this glycosyltransferase is also required to maintain functional mitochondria and calcium homeostasis. Cells carrying a mutation in KlOCH1 gene showed altered mitochondrial morphology, increased accumulation of ROS and reduced expression of calcium signalling genes such as calmodulin and calcineurin. Intracellular calcium concentration was also reduced in the mutant cells with respect to the wild type counterparts.

Phenotypes that occur in cells lacking the alpha-1,6-mannosyltransferase, including oxidative stress and impaired mitochondria functionality, were suppressed by increased dosage of KlCmd1p. This, in turn, acts through the action of calcineurin.

Conclusions: Proper functioning of the alpha-1,6-mannosyltransferase in the N-glycosylation pathway of K. lactis is required for maintaining normal calcium homeostasis; this is necessary for physiological mitochondria dynamics and functionality.

Background

In eukaryotic cells, transmembrane and secreted proteins undergo several modifications during their maturation; N-Glycosylation is one of these modifications, and contributes to the functional conformation and to the selection of the final destination of these proteins [1]. Yeast and mammals share the initial steps of N-glycosylation in the ER, even if the pathways diverge between mammals and yeast in the Golgi apparatus. Indeed, in this cellular compartment, the extension of the oligosaccharide chain involves many specific glycosyltransferases. This specificity generates the observed diversity of glycan structures between different species and cell types [2]. In the yeast Saccharomyces cerevisiae the glycosyltransferases are consti-
tuted by mannosyltransferases, which lead to the formation of two main types of mannan outer chain. Many proteins of the cell wall and periplasm receive a large mannan structure that contains a long -1,6-linked backbone of about 50 mannoses with short -1,2 and -1,3 side chains. In contrast, the proteins of the internal organelles display a smaller core-type structure, with only a few mannoses [3]. The structure of the mannan outer chain has been investigated by the study of mnn (mannan defective) mutants isolated by Ballou and co-workers [4]. The analysis of the partial N-glycan structures in these mutants has allowed the ordering of the steps of mannan synthesis. The work of Munro suggests a model for the complete N-glycosylation pathway in S. cerevisiae [3,5]. In this model, the formation of the mannan outer chain is initiated by the Och1p protein, a type II membrane -1,6-mannosyltransferase that defines an early Golgi compartment [6-8]. Upon arrival in the Golgi, all N-glycan cores receive a single α-1,6-mannose added by Och1p. The formation of the long α-1,6-linked backbone is generated by two enzyme complexes: the M-Pol I and M-Pol II complexes.

*Kluyveromyces lactis* is a yeast species related to *S. cerevisiae*; its outer mannan chains differ from those of *S. cerevisiae* by having terminal N-acetylglucosamine and no mannose phosphate [9]. Previous studies from our laboratory reported that the Kloch1-1 mutation of *Kluyveromyces lactis* yeasts was mapped at 738 bp of the KLOCH1 gene and we found that the corresponding wild type allele encodes the functional homologue of the *S. cerevisiae* α1,6-mannosyltransferase [10]. Quantitative analysis of cell-wall components indicated a noticeable increase of chitin and β1,6-glucans and a severe decrease of mannoproteins in the Kloch1-1 cells as compared to the wild-type counterparts. In addition, fine-structure determination of the β1,6-glucan polymer showed that, in the Kloch1-1 strain, the β1,6-glucans are shorter and have more branches than in the wild-type strain. Moreover, the Kloch1-1 cells showed a significantly improved capability of secreting heterologous proteins.

Recent reports provide evidences that defects in protein N-glycosylation can cause mitochondrial deficiencies. An ER-derived oxidative stress from misfolded proteins in an eru29 mutant of *S.cerevisiae* has been shown to lead to UPR activation, which cause the accumulation of ROS from both the ER and mitochondria, and resulted in cell death [11]. It has also been reported that dosage attenuation of Dol-P-dependent GlcNAc-1-P transferase, ALG7, in haploid cells led to the loss of mtDNA and respiratory deficiency [12]. Also Eos1, a protein involved in the N-glycosylation, was required for tolerance to oxidative stress [13]. However, in most of the cases the mechanisms linking glycosylation and mitochondrial dysfunctions need to be elucidated. In this work we found that altered mitochondria functionality in cells deprived of the Golgi α-1,6-mannosyltransferase is caused by a defective calcium homeostasis.

**Results**

**Altered mitochondrial biogenesis in Kloch1-1 cells**

The *K. lactis* KLOCH1 gene encodes the α-1,6-mannosyltransferase localized in the Golgi apparatus and involved in the glycosylation process and in cell wall morphogenesis. The Kloch1-1 mutant strain showed enhanced heterologous protein production [10]. However, the Kloch1-1 cells displayed a higher generation time in rich fermentable medium in comparison with wild type cells, and in the presence of respiratory carbon source, the biomass yield was strongly reduced. This prompted us to investigate if the lack of the α 1,6-mannosyltransferase activity in *K. lactis* cells could be associated with altered mitochondrial functionality. To this end cells were incubated with DASPMI, a fluorescent probe that is taken up by mitochondria as a function of membrane electrochemical potential. We found indeed a different distribution of the dye in the two strains: in the Kloch1-1 background the cells showed a punctuate pattern and some ring structures instead of the regular tubular network of the wild type counterpart (Figure 1A). In order to analyze possible alterations of the mitochondria structures, we transformed the mutant and wild type strains with a plasmid carrying the GFP fused to the mitochondrial signal sequence from the subunit 9 of the F0-ATPase from *Neurospora crassa*; this construct has been demonstrated to correctly deliver functional GFP into yeast mitochondria [14]. The fluorescence microscope observation of the mitochondrial matrix (Figure 1B) showed, for the mutant cells, an hyperbranched mitochondrial net not revealed by the DASPMI staining, whereas the mitochondrial morphology of the wild-type cells resulted identical to the one observed with DASPMI staining. To further study the altered mitochondrial morphology, the mutant strain and its isogenic wild type counterpart were analyzed by electron microscopy. In ultrathin sections of the wild type cells (Figure 1C) the mitochondria appeared as tubular structures, with normal morphology and typical cell peripheral distribution. The Kloch1-1 cells showed instead mitochondria that were either stretched and without crests in the middle part either round swollen at the ends, in agreement with the altered morphology observed by DASPMI staining.

Mitochondrial dysfunction is often associated with accumulation of reactive oxygen species (ROS), we therefore evaluated the amount of ROS by using the fluorescent dye 123 dihydrorhodamine (DHR). This compound accumulates inside the cells and it is oxidized to the corresponding fluorescent chromophore by ROS. The fluorescence microscope observation revealed that 28% of Kloch1-1 cells accumulated ROS as compared to 5% of the parental
strain; these values increased up to 48% and 18% in the mutant and wild type cells respectively after challenge with a generator of oxidative stress, such as the hydrogen peroxide (Figure 2A). To further investigate the oxidative stress taking place in the mutant, \textit{K. lactis} cells were challenged for 2 and 5 h with a cytotoxic concentration of 20 mM hydrogen peroxide. A strong reduction of the survival rate was observed after a 2 hours treatment in cells lacking the $\alpha$1,6-mannosyltransferase activity (Figure 2B), and after 5 hours only 41% of mutant cells survived versus 93% of the parental ones. In addition, the mutant strain was not able to grow in the presence of 4 mM hydrogen peroxide (Figure 3A).

\textbf{Isolation of KlCMD1 as extragenic suppressor of oxidative stress occurring in Kloch1-1 cells}

In order to highlight genetic interactions underlying the phenotypes observed, we performed a screen to identify multicopy suppressors able to relieve the growth defect of cells lacking $\alpha$1,6-mannosyltransferase activity on YPD supplemented with 4 mM hydrogen peroxide (Figure 3A). Three of the plasmids, isolated from the corresponding clones that survived the selection procedure, resulted to be identical and were further analyzed. Sequencing analysis revealed that the \textit{K. lactis} DNA fragment present in the plasmid contained an ORF of 441 bp with 1000 bp upstream of the putative ATG start codon and 100 bp downstream of the putative stop codon. The protein encoded by this ORF resulted to be \textit{KlCMD1}, the homologue of the \textit{S. cerevisiae} calmodulin gene [15]. The gene was able to restore the growth defect of Kloch1-1 cells on 4 mM H$_2$O$_2$ also when cloned in a centromeric plasmid (CpKICMD1).

\begin{figure}[h]
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\includegraphics[width=\textwidth]{figure1.png}
\caption{\textbf{Morphological and functional analysis of mitochondria in cells lacking $\alpha$1,6-mannosyltransferase activity.} (A) DASPMI staining of Kloch1-1 cells and the parental strain. (B) GFP fluorescence of mitochondrial matrix of wt and mutant strains harbouring p426SD11 mtGFP vector. (C) Mitochondrial analysis by electron microscopy of Kloch1-1 cells and the parental strain. Cultures were grown to exponential phase into YPD medium. n, nucleus; m, mitochondrion; cw, cell wall; bar 2 $\mu$m.}
\end{figure}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure2.png}
\caption{\textbf{Oxidative stress in Kloch1-1 cells.} (A) Estimation of ROS accumulation in the indicated strains by DHR-staining after growth to exponential phase in SD medium. Measurements were also obtained after exposure to H$_2$O$_2$ for 2 h. (B) Cell viability after H$_2$O$_2$ exposure: parental strain (black bars) and cells lacking $\alpha$1,6-mannosyltransferase activity (white bars), grown to exponential phase on YPD medium, were challenged with 20 mM H$_2$O$_2$ for 2 or 5 h. The viability was evaluated plating the samples on YPD and was expressed as the CFU percentage of the corresponding untreated cultures. The values of both panels were the mean of three independent experiments and showed an SD < 10%.}
\end{figure}
We wondered if transcriptional variations of *KlCMD1* gene could occur in *Kloch1-1* mutant cells as compared to the parental strain. In these cells we effectively observed a strong reduction of the mRNA of the calmodulin protein as revealed by northern blotting analysis (Figure 3B).

The presence of *KlCMD1* on a centromeric plasmid in the mutant strain was able to reduce the oxidative stress of these cells as revealed by the DHR staining. The amount of positively stained cells of the transformants, in fact, was significantly reduced in comparison to that of the mutant cells: 8% versus 28% respectively (Figure 4A). However, the *Kloch1-1* cells transformed with CpKlCMD1 did not show an increase in the survival capabilities with respect to the mutant cells, when challenged for 2 hours with 20 mM (additional file 1).

The mitochondrial functionality of the *Kloch1-1* cells carrying either the calmodulin on a centromeric (CpKlCMD1) or on a multicopy plasmid (MpKlCMD1) was studied by DASPMI staining (Figure 4B). The *Kloch1-1* cells transformed with CpKlCMD1 showed a functional hyperbranched mitochondrial net in contrast to the dots phenotype, typical of the mutant (see Figure 1A). These data indicate that the mitochondrial net that was visualised by mitoGFP in *Kloch1-1* cells (see Figure 1B) became able to establish a membrane electrochemical potential just by adding a few copies of calmodulin in these cells.
In addition, when the mutant cells were transformed with MpKlCMD1, the DASPMI staining revealed a wild type-like mitochondrial network, even if in some cases ring structures were still present (Figure 4B).

In addition to the generation of cellular energy, mitochondria also play an important role in regulating calcium homeostasis [16]. Based on this and on the suppression of the mutant phenotypes by calmodulin, we looked for possible calcium homeostasis alterations in Kloch1-1 cells. We thus investigated if cells lacking α 1,6-mannosyltransferase activity could grow in presence of EGTA, a cationic chelator. As reported in the panel A of the figure 5, the growth of mutant cells was strongly inhibited when YPD plates were supplemented with this chelating agent. This inhibition was suppressed by adding in the growth medium, 20 mM CaCl₂ together with the EGTA, indicating that an altered calcium homeostasis occurs in the mutant strain.

Intracellular calcium determinations, carried out using Fura-2AM, indeed showed that the calcium content of Kloch1-1 cells was significantly reduced with respect to the cation concentration present in the wild-type cells determined in the same conditions (Figure 5B). Measurements employing cells deleted for the Golgi Ca²⁺-ATPase, KlPmr1p, previously reported to have increased cytosolic calcium content, were used as additional calibration control.

**Increased calcineurin activity is required in Kloch1-1 cells for normal mitochondrial morphology and cell wall structure**

Since calmodulin acts as a mediator of calcium signals in eukaryotic cells mainly through the Ca²⁺/calmodulin-dependent phosphatase calcineurin, we investigated the possible role of this phosphatase in the mitochondrial phenotypes of Kloch1-1 mutant cells.

Calcineurin requires both the regulatory and the catalytic subunits for full activity, and the KCNB1 and KCNA1 genes respectively were isolated (see Materials and Methods). The reduced expression of KCMD1 observed in the mutant strain prompted us to first analyze whether a similar behaviour could be observed also for the calcineurin subunits. The Kloch1-1 cells effectively showed a decrease of KCNB1 transcript in comparison to the wild type cells, and a similar result was observed also for the catalytic subunit (Figure 6A and 6B). In order to obtain the expression balance of both regulatory and catalytic subunits of calcineurin, we transformed Kloch1-1 strain with the plasmids harbouring KCNB1 (pKlCNB1) and KCNA1 (pKlCNA1) genes. Surprisingly, the effect of the increased dosage of the regulatory subunit alone in the Kloch1-1 cells on the EGTA sensitivity of the mutant strain was identical to that of the overexpression of both subunits (Figure 6C). Similar results were also obtained when the amount of ROS was analyzed: 15% of the Kloch1-1 cells transformed with pKlCNB1 alone or together with pKlCNA1 were positive to the staining with DHR as compared to 28% of the mutant cells. In this case the suppression was not complete as compared to the almost complete recovery observed in Kloch1-1 cells by increasing the dosage of calmodulin gene.

We then looked up to the mitochondrial structures and functionality by DASPMI staining of the transformants. Mutant cells carrying pKlCNB1 vector alone showed a partial recovery of the tubular phenotype. We observed a reduced amount of dots and the appearance of tubules, whereas the same cells transformed also with the pKlCN1A plasmid showed the typical tubular wild type-like network, although with a peripheral distribution (Figure 7A). By electron microscopy analysis we observed only a partial recovery of tubular mitochondria in Kloch1-1...
cells harbouring only the pKlCNB1 vector (Figure 7B). However, when in the same strain we co-overexpressed also the catalytic subunit of calcineurin, the relieve was almost complete: mitochondria appeared very similar to the wild type ones and mitochondrial cristae were visible, in agreement with the structures observed by fluorescence microscopy (Figure 7B).

**Figure 6**

Analysis of calcineurin in Kloch1-1 cells. (A) Northern Blotting analysis of KlCNB1 WT and Kloch1-1 cells. The same amount of total RNA (40 μg) from the strains was loaded on each lane; the ethidium bromide-stained gel of the autoradiogram is shown in the bottom part of the panel and the mRNA loading was normalized using the 26S rRNA bands. Quantification, by densitometric analysis, of the radiolabeled signal on the blot is shown in the right part of the panel. The hybridization signal for wild type strain was set as 1. (B) RT-PCR semi-quantitative analysis of KlCNA1 gene. Exponentially growing wild-type and Kloch1-1 cells were collected and RNA was isolated. Reverse transcription and PCR reactions with the specific primers described in materials and methods was performed. Shown is the electrophoresis in a 2% agarose gel of 10 μl of each PCR reaction with 5 μl (1×) and 10 μl (2×) of cDNA as template. RT-PCR of the KlUBC6 gene was performed as an internal control. RNA extraction was performed twice and the results shown are representative of four independent RT-PCR experiments. (C) Growth of indicated yeast strains onto solid medium supplemented with 20 mM EGTA.

**Figure 7**

Phenotypical analysis of Kloch1-1 overexpressing KlCNB1 or both subunits of calcineurin. (A) DASPMI staining of the Kloch1-1 cells harbouring pKlCNB1 plasmid alone or with the pKlCNA1 plasmid. (B) Ultra-thin sections of above strains. n, nucleus; m, mitochondrion; cw, cell wall; bar 2 μm. (C) Serial dilutions of cultures from the same strains onto YPD agar plates supplemented with the cell wall interfering agent calcofluor white (CFW). The growth at 28°C was monitored after 3 d; three independent transformants have been checked, obtaining identical results.
We previously reported that the Kloch1-1 mutant had increased cell wall thickness and some dark-stained rims were present within the amorphous layer in comparison with wild-type cells [10]. Notably, in cells depleted of αL,6 mannosyltransferase activity but transformed with the pKICNB1 and pKICNA1 plasmids, the thickness of the cell wall resembled that of the wild-type parent (Figure 7B). In agreement with the electron microscopy observations, we found that the ability of the mutant cells to grow in presence of the cell wall-perturbing agent calcofluor white was completely restored only when the cells were transformed with both subunits of calcineurin (Figure 7C). Identical results were also obtained when congo red, another molecule interfering with the cell wall, was used (data not shown). This phenotype has to be ascribed to the increased activation of the calcineurin signaling pathway that generates a rescue of a normal biogenesis of cell wall.

Discussion
Protein N-glycosylation is one of the fundamental metabolic pathways in cell fate. Deciphering how N-glycosylation controls specific metabolic and signaling events has become important to unraveling the underlying basis of cellular behavior. In yeasts, outer chain branching is initiated in the Golgi apparatus by the α-1,6-mannosyltransferase Och1p. Here, we reported that altered mitochondrial functionality and oxidative stress take place in K. lactis cells carrying a mutation in KIOCH1 gene. The mutant cells showed also a reduction in the calcium content and in the expression of genes related to calcium signalling. Although a direct link between N-glycosylation defects and mitochondrial functionality has not been reported, underglycosylation of proteins destined for the mitochondria could interfere with, or abolish, the import of these proteins into the organelle. Indeed, a 45 kDa protein has been identified in rat liver inner mitochondrial proteins into the mitochondria, could interfere with, or abolish, the import of these proteins. Underglycosylation of proteins destined for the mitochondria and mitochondrial functionality has not been reported, place in K. lactis cells carrying a mutation in KIOCH1 gene. The mutant cells showed also a reduction in the calcium content and in the expression of genes related to calcium signalling.

S. cerevisiae calmodulin, a Ca²⁺ binding protein, regulates many cell processes both depending or not upon the intervention of Ca²⁺ ions. Among those Ca²⁺-dependent is the organization of the actin cytoskeleton; moreover mutations in CMD1 resulted coletal, suggestive of functional interactions, with the inactivation of genes encoding components of the glycosylation pathways like ANP1, CWH8 and MNN10 [22]; mutations in such genes also result in altered morphology of actin cytoskeleton. We should also take into account that, although och1 deletion mutant of S.cerevisiae in the BY4741 background was sensitive to EGTA, the mutant strain was not altered in the expression of calmodulin and calcineurin genes (Zanni et al., unpublished results). However, we can not exclude the possibility that a reduction in the activity of the calcium signalling proteins can occur in the OCH1 deleted cells.

Mitochondrial plasticity and functionality strongly depend upon the interactions between mitochondria and cytoskeleton. Several shape-related proteins have been described in S.cerevisiae, localized on the mitochondria surface and reported to interact with actin [23,24]; however the individual role and underlying mechanisms are still unsolved.
Another unanswered question is how do cells change the mitochondrial shape upon cell signals. In the case of calcium signalling, a relevant player could well be Gem1p, a member of the Miro GTPase family [25]. Gem1p is also localized on the outer mitochondrial membrane with its GTPase domain and, most notably, its EF-hand calcium binding domain exposed in the cytosol.

We are tempted to speculate that the altered calcium availability we observed in Kloch1-1 cells could be originated by a defective calcium membrane channel Mid1/Ch. In fact, it has been demonstrated that S. cerevisiae Mid1 requires a full glycosylation to correctly localize and assemble at the level of the plasma membrane [26].

In mammalian cells Ca^{2+} influx through voltage-dependent Ca^{2+} channels (VDCCs) causes a rapid halt in mitochondrial movement and induces mitochondrial fission. VDCC-associated Ca^{2+} signaling stimulates phosphorylation of dynamin-related protein 1 (Drp1) at serine 600 via activation of Ca^{2+}/calmodulin-dependent protein kinase Iα (CaMKIα). In neurons and HeLa cells, phosphorylation of Drp1 at serine 600 and dephosphorylation at serine 637, both calcineurin-dependent, are associated with increase in Drp1 translocation to mitochondria [27,28]. Nevertheless, one cannot exclude that defective Kloch1 gene induce reduced glycosylation of other proteins relevant for calcium handling; scrutiny of such picture will deserve future work.

**Conclusions**

A proper functioning of outer chain-extension of manno-proteins in K. lactis is required for correct calcium homeostasis. The impairment of Kloch1 results in a low calcium/calmodulin based signaling and altered mitochondria morphology and functionality. The reported data strongly indicate a novel link between relevant cell processes taking place in separate compartments.

**Methods**

**Yeast Strains and Growth Conditions**

The strains used in this study were MW278-20C (MAT a, ade2, leu2, uraA), CPV3 (MAT a, ade2, leu2, uraA, Kloch1-1) and CPK1 (MAT a, ade2, leu2, uraA, KlPMR1::Kan R). Yeast strains were grown in YPD medium (1% yeast extract, 1% peptone, 2% glucose) or SD minimal medium (2% glucose, 0.67% yeast nitrogen base without amino acids) with the appropriate auxotrophic requirements. Fivefold serial dilution from concentrated suspensions of exponentially growing cells (5 × 10^{6} cell/ml) were spotted onto synthetic YPD agar plates supplemented or not with 4 mM H_{2}O_{2}, 20 mM EGTA, 200 μg/ml Congo red or 50 μg/ml CFW and the plates were incubated at 30°C for 48 h.

**Plasmids Construction**

Construction of the pCXJ3-U and pCXJ6-L plasmids: the kanamycin-resistance encoding gene (kan) was excised by PstI digestion from pCXJ3 and pCXJ6 [29] plasmids, giving multicopy vectors with the selectable marker URA3 or LEU2, respectively.

Construction of the pCXJ3-K plasmid: the URA3 gene was excised by BglII digestion from pCXJ3 plasmid, obtaining multicopy vector with the selectable marker KAN.

Construction of the CpKlCMD1 and MpKlCMD1 plasmids: the 1545 bp fragment containing the full ORF (444 bp of KlCMD1 plus 1000 bp upstream and 100 bp downstream was amplified by PCR, using primers modified with the recognition site for the restriction endonuclease BamHI. The PCR fragment encoding the KlCmd1p was cloned into the pGEM-T-Easy vector (Promega) according to the manufacturer’s instructions, giving pGEM-KlCMD1 and the gene correctness was confirmed by DNA sequencing (MWG Biotech, Martinsried, Germany). The fragment was excised by BamHI digestion from pGEM-KlCMD1 and was ligated into the centromeric (pCXJ20) or multicopy (pCXJ6-L) vectors, linearized by the same endonuclease, to obtain the CpKlCMD1 and MpKlCMD1 plasmids respectively.

Construction of the pKlCNB1 and pKlCNA1 plasmids: the KlcNB1 and KlcNA1 genes were PCR amplified from K. lactis DNA genome using the primers 5'-CGGGATCC-GGCCAGAGAGCAGGTCAAC-3' and 5'-CGGGATCCGCTG-CCITCATCTGATCCGCGCCG-3', 5'-CGGGATCCGCCGC TCCATACCATACGCGCGC-3' respectively (the BamHI restriction site underlined). The PCR products were ligated into the pGEM-T-Easy vector (Promega) giving the pGEM-KlCNB1 and pGEM-KlCNA1 plasmids.

After sequencing (MWG Biotech, Ebersberg, Germany) the fragments were successively cloned in BamHI-digested pCXJ3-U and pCXJ6-K plasmids, obtaining pKlCNB1 and pKlCNA1 vectors, respectively.

**Yeast Transformation and Selection of Suppressor Genes**

The Kloch1-1 strain was transformed to saturation with the yeast genomic library constructed in the pKepl multicopy vector (kindly provided by Wesołowski-Louvel) by electroporation [30]. All the Ura+ transformants were replicated on to YPD medium supplemented with 4 mM H_{2}O_{2}. The plasmids isolated from the Ura+/H_{2}O_{2}^8 transformants were used to transform the Kloch1-1 strain. Plasmids capable of restoring the H_{2}O_{2}^8 phenotype to the Kloch1-1 after retransformation were analyzed. Restriction enzymes analysis of the genomic fragments from the isolated plasmids showed that one of these plasmids carrying
an insert of about 8000 bp was able to restore the H₂O₂ phenotype. The plasmid was then sequenced (MWG Biotech). The 1600 bp fragment contained the full ORF (444 bp) of KlCMD1 plus 1000 bp upstream and 100 bp downstream.

**Stress Condition and Viability**

Yeast cells were grown aerobically at 28°C in liquid medium for 24 h and were challenged with hydrogen peroxide. This was directly added to the growth medium to the final concentration of 20 mM. Untreated cultures were incubated in parallel over the same periods. Viability was determined by colony counts on YPD plates after 2 and 5 h of incubation at 28°C and was expressed as the percentage of the corresponding control cultures. The values are the mean of three independent experiments with a SD < 15%.

**Measurement of Intracellular Oxidation Levels**

The oxidant-sensitive probe dihydrorhodamine 123 (Sigma) was used to measure intracellular oxidation levels in yeast according to [31].

Cells were concentrated by centrifugation and resuspended in 10 μl of fresh medium. Five microliters of cells were loaded onto slides and observed immediately under epifluorescence microscopy (excitation at 488 nm and emission at 530 nm). At least 300 cells per sample were scored manually as fluorescent or nonfluorescent.

**Fluorescence Microscopy**

Cells grown in 2% glucose medium were harvested in exponential growth phase (6 × 10⁷ cells/ml), washed with water and then incubated for 30 min in the presence of 5 μM of DASPMI [32]. Epifluorescence microscopy was carried out with a Zeiss Axiospho microscope fitted with a 10× immersion objective and a standard FITC filter set.

**Electron Microscopy**

Cells were prepared as described in [21].

**Northern Blot Analysis**

Total RNA of K. lactis strains was extracted by the hot phenol method [33]. The RNAs were quantified by absorption (OD₂₆₀) and separated by denaturing agarose electrophoresis. After electrophoresis the RNAs were transferred to nylon membranes and hybridized with 32P-labeled random primed probes (Roche, Lewes, East Sussex, United Kingdom). All the probes were PCR amplified from the K. lactis DNA genome. The 1300-bp PCR product of KlCMD1 was obtained using primers 5'-CGGGATCCCGTACCCTGATAGCTCTACC-3' and 5'-CGGGATCCGTTGGAATTTGAGCGATGG-3'. The fragment of 410 bp containing the KlCNB1 gene was obtained with primers 5'-GAATTGAATGGGAGCAGCA-3' and 5'-CTTGAAAATCAACATCTCCG-3'. The densitometric analysis was done with an image analyzer (Phoretix 1D; Non Linear Dynamics Ltd.).

**Semi quantitative RT-PCR**

Total RNA extracted as before was subjected to TURBO™ DNase treatment according to manufacturer's instructions (Ambion, USA). Reverse transcription was performed using Promega Reverse Transcription System with 1 μg total RNA to yield 20 μl cDNA. PCRs were then performed to determine the linear range of amplification for the genes that would allow a semi-quantitative assessment of expression levels. The primers used for KlCNB1 were 5'-GTGAATGCACTCTGCTGATC-3' and 5'-CAGCTGATGATATGGTCCCTTCTC-3', while for KlUBC6 were 5'-ATTACGTGTATTACCGGATCCA-3' and 5'-GCCTCTGGATGATAATC-3'. The optimal parameters determined for each PCR were 95°C, 30 s; 52°C, 30 s; 72°C, 30 s; and 20 cycles for both the KlCNB1 and KlUBC6 genes. The primers used were designed to yield small amplicons (KlCNB1, 241 bp; KlUBC6, 178 bp) to improve the efficiency and reproducibility of the PCR. 10 μl of each PCR reaction with 5 μl (1×) and 10 μl (2×) of cDNA diluted 1:5 as template were separated on a 2% agarose gel, stained with ethidium bromide and photographed.

**Ca²⁺ Measurements**

A suspension of 50 μl freshly prepared spheroplasts were diluted in 1 ml of spheroplast buffer (SB) containing 1 M sorbitol, 50 mM Tris buffer, pH 7.5, 10 mM Mg²⁺ and left into a polylysine-coated plate in agitation at 4°C overnight. Plates were washed twice with 1 ml of SB. Spheroplasts were incubated for 60 min at 37°C in standard reaction medium (125 mM sucrose, 65 mM KCl, 10 mM HEPES, pH 7.2 and 500 μM ethanol) in the presence of 0.1 mg/ml bovine serum albumin (Sigma) and 10 μM Fura-2AM (Molecular Probes, Eugene, OR). To measure fluorescence changes, a Hamamatsu Argus 50 computerized analysis system was used, recording every 6 sec the ratio between the values of light intensity at 340 and 380 nm stimulation. Spheroplasts were prepared according to [32].

**Authors’ contributions**

EZ, DJ, designed, carried out the experiments, analyzed data and drafted the manuscript. FF performed the genetic screening, analyzed data and commented on the manuscript. CF performed the calcium measurements, analyzed data and commented on the manuscript. AR participated to the electron microscope analysis. PM performed the electron microscope analysis, analyzed data and commented on the manuscript. CP designed experiments, analyzed data and commented on the manuscript. DU

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supervised the project and corrected the final manuscript. All authors read and approved the final manuscript.

Additional material

Additional file 1
Cell viability after H2O2 challenge. Indicated strains, grown to exponential phase on YPD medium, were challenged with 20 mM H2O2 for 2 h. The viability was evaluated plating the samples on YPD and was expressed as the CFU percentage of the corresponding untreated controls. The values were the mean of three independent experiments and showed an SD < 10%.

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