Functional comparison of plasma-membrane Na⁺/H⁺ antiporters from two pathogenic Candida species

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Abstract

Background: The virulence of Candida species depends on many environmental conditions. Extracellular pH and concentration of alkali metal cations belong among important factors. Nevertheless, the contribution of transporters mediating the exchange of alkali metal cations for protons across the plasma membrane to the cell salt tolerance and other physiological properties of various Candida species has not been studied so far.

Results: The tolerance/sensitivity of four pathogenic Candida species to alkali metal cations was tested and the role of one of the cation transporters in that tolerance (presumed to be the plasma-membrane Na⁺/H⁺ antiporter) was studied. The genes encoding these antiporters in the most and least salt sensitive species, C. dubliniensis and C. parapsilosis respectively, were identified, cloned and functionally expressed in the plasma membranes of Saccharomyces cerevisiae cells lacking their own cation exporters. Both CpCnh1 and CdCnh1 antiporters had broad substrate specificity and transported Na⁺, K⁺, Li⁺, and Rb⁺. Their activity in S. cerevisiae cells differed; CpCnh1p provided cells with a much higher salt tolerance than the CdCnh1 antiporter. The observed difference in activity was confirmed by direct measurements of sodium and potassium efflux mediated by these antiporters.

Conclusion: We have cloned two genes encoding putative Na⁺/H⁺ antiporters in C. parapsilosis and C. dubliniensis, and characterized the transport properties of encoded proteins. Our results show that the activity of plasma-membrane Na⁺/H⁺ antiporters is one of the factors determining the tolerance of pathogenic Candida species to high external concentrations of alkali metal cations.

Background

The family of Candida species, normally a harmless human commensal of the gastrointestinal and genitourinary tract, can become a human pathogen under certain circumstances. Mainly in HIV and immunocompromised patients, Candida cause a wide range of infections and are the most prevalent pathogenic yeast. One key feature of these fungi is their ability to grow in three different morphologies: yeast, pseudohyphae and true hyphae [1]. This reversible switching from one form to another is dependent on environmental conditions like temperature, pH, nutritional status and external/internal concentration of cations [2].

In general, pathogenic Candida species are osmotolerant yeasts and can grow, with the exception of Candida dubliniensis [3], at relatively high NaCl concentrations, although the presence of salt was shown to negatively...
influence several virulence traits of *Candida albicans* [4]. Recent experiments also suggest a relationship between the formation of *C. albicans* hyphae and the intracellular concentration of potassium [5]. Nevertheless, the regulation of intracellular potassium and sodium concentrations in *Candida* species has not been studied in detail.

Yeast species in general have several transport systems in their plasma membranes at their disposal to maintain homeostasis in alkali metal cations, i.e. a high ratio between potassium (which is the main intracellular cation) and toxic sodium concentrations [6-8]. Among these transport systems, Na⁺/H⁺ antiporters play an important role. Most yeasts that have so far been studied (e.g. *Saccharomyces cerevisiae, Debaryomyces hansenii*) possess only one type of this antiporter in their plasma membranes, which efficiently transports both sodium and potassium cations from the cells, as well as their analogues lithium and rubidium [7,9]. A few yeast species (*Yarrowia lipolytica, Schizosaccharomyces pombe*) have two antiporters of this family at their disposal, one of them with a substrate preference for sodium and lithium, the other preferring potassium and rubidium [10,11].

The Na⁺/H⁺ antiporter that has been studied the most so far is from *S. cerevisiae*, encoded by the *NHA1* gene, and has 12 predicted trans-membrane domains and a very long hydrophilic C-terminus [7]. Beside its function in removing toxic Na⁺ from cells and maintaining potassium homeostasis, it is involved in several other cellular functions such as regulating intracellular pH [12,13], cell volume [7], plasma membrane potential [14] and the cell cycle [15,16], and it participates in the cell response to osmotic shock [7,17]. The Nha1p orthologs from *C. albicans* and *Candida tropicalis* CaCnh1p and CtCnh1p, respectively, were functionally characterized upon heterologous expression in *S. cerevisiae*. Both showed the same broad substrate specificity as ScNha1p [18,19]. The deletion of *CNH1* in *C. albicans* results in cell sensitivity to high external potassium concentrations [20] and under some conditions causes slight changes in cell morphology [21].

In this work we compared the tolerance of four different pathogenic *Candida* species to alkali metal cations, performed a search for Nha1/Cnh1 antiporter-encoding orthologs in their genomes, and characterized the transport properties of the Na⁺/H⁺ antiporters from the most and least tolerant species, *C. parapsilosis* and *C. dubliniensis* respectively.
Results

Candida species differ in their halotolerance

According to the literature [3], C. dubliniensis is relatively sodium sensitive, whereas C. parapsilosis was shown to tolerate high NaCl concentrations [22]. In order to estimate the tolerance of Candida species to different alkali metal cations, the growth of four Candida species and a S. cerevisiae wild type (as a non-osmotolerant control) in the presence of increasing concentrations of various salts was estimated. In the absence of salts, S. cerevisiae cells grew more slowly than all four Candida species, and also the growth of C. parapsilosis was not as robust as with the other three Candida species (Figure 1). All yeast species grew equally well in the presence of lower salt concentrations (cf. Methods) but as the amount of alkali-metal-ion on the plates increased, important differences were observed. Of the tested Candida species, C. dubliniensis had the lowest tolerance to all of the tested salts, it is not able to grow when the salt concentrations are above 1600 mM NaCl, 2300 mM KCl or 200 mM LiCl. Nevertheless, C. dubliniensis is more sodium, potassium and rubidium tolerant than S. cerevisiae, but is much more sensitive to toxic lithium cations (Figure 1). Also, the lithium sensitivity of C. glabrata is higher than that of S. cerevisiae but C. glabrata cells can grow in much higher concentrations of the other salts than S. cerevisiae cells (up to 2300 mM NaCl, 2400 mM KCl, 1800 mM RbCl; Figure 1). C. albicans and C. parapsilosis are the most halotolerant species, and C. parapsilosis seems to grow even better in the presence of high salt concentrations than C. albicans (Figure 1). To summarize, the four Candida species have not only different sensitivity to NaCl, as published previously [3,22] but they differ in their tolerance to alkali metal cations in general. C. albicans and C. parapsilosis are highly halotolerant and C. dubliniensis is halosensitive.

Comparison of Candida CNH1 genes and encoded antiporters

A search in databases revealed the existence of open reading frames homologous to the CaCNH1 gene in the genomes of C. dubliniensis, C. glabrata and C. parapsilosis. In these species, just one homologous sequence was found, suggesting that their plasma-membrane Na+/H+ antiporters have a broad substrate specificity to alkali metal cations, similar to those of C. albicans and C. tropicalis. We named the identified orthologous genes according to their species of origin, CdCNH1, CgCNH1 and CpCNH1. The CdCNH1 gene is 2454 nt (818 aa) long, CgCNH1 has 2835 nt (945 aa), and the CpCNH1 gene is composed of 2955 nt (985 aa). Neither of them have any introns.

The predicted protein structures of these three antiporters were compared with the two Candida antiporters that have already been characterized, CaCNH1 and CtCNH1. Comparison of the protein length and predicted structure of the Cnh1 proteins from five Candida species and S. cerevisiae revealed that CpCNH1 is the longest and C. dubliniensis antiporter the shortest member of the Candida Na+/H+-antiporters’ subfamily (Table 1). For all Candida proteins, the Kyte-Doolittle method predicted a similar structure to ScNha1p with highly conserved N-termini and 12 trans-membrane sections (Tables 1 & 2). On the other hand, they differ in the length and composition of their hydrophilic C-termini, as do the antiporters from non-Candida yeast species [23]. The most significant is the difference in length (approx. 180 aa) between the C-termini of CgCNH1p (555 aa), CaCNH1p (366 aa) and CdCNH1p (388 aa). Of the analyzed proteins, CaCNH1p and CdCNH1p show the highest sequence identity in all parts of the protein, though the hydrophobic trans-membrane domains and connecting loops are highly conserved (approx. 90% identity) in all Candida antiporters except for CgCNH1p (Table 2). C. glabrata Cnh1p is more similar to S. cerevisiae Nha1p than to the antiporters from other Candida species in all the features analyzed, which corresponds to the phylogenetic relationships among these yeast species [24].

Though the highest divergence from identity was found in the C-termini of all the compared antiporters, the existence of six conserved C-terminal regions described previously [18] was also confirmed in C. dubliniensis, C. parapsilosis and C. glabrata species (not shown). A new, approximately 25 aa-long conserved region (K/R)(L/I)SR(S/T)(L/A)SRRS(Y/F)Y(K/R)KDDP(H/N)(K/;

Table 1: Comparison of deduced secondary structures of plasma-membrane Na+/H+ antiporters from Candida species and S. cerevisiae

| Yeast         | Antiporter | Number of amino acid residues | N-terminus | Tms + loops | C-terminus | Whole protein |
|---------------|------------|-----------------------------|------------|-------------|------------|---------------|
| C. albicans   | CaCNH1p    | 11                          | 419        | 366         | 796        |
| C. dubliniensis| CdCNH1p   | 11                          | 419        | 388         | 818        |
| C. glabrata   | CgCNH1p    | 12                          | 418        | 515         | 945        |
| C. parapsilosis| CpCNH1p   | 11                          | 419        | 555         | 985        |
| C. tropicalis | CdNha1p    | 11                          | 419        | 545         | 975        |
| S. cerevisiae | ScNha1p    | 12                          | 418        | 555         | 985        |
Both the *C. dubliniensis* and *C. parapsilosis* species belong to the group of yeasts in which the CTG codon encodes a serine and not a leucine, as in other yeast species (i.e. *S. cerevisiae*) [25]. One CTG codon exists at aa position 621 in *CpCNH1*. This serine 621 is localized in the antiporter’s hydrophilic C-terminus and not in the membrane part of the protein. It is localized in a small weakly conserved area where at a similar position *CdCnh1p* (aa 568) and *CtCnh1p* (aa 644) also have a serine and *CaCnh1p* (aa 552) has an isoleucine.

**Table 2: Identity (%) of *Candida* and *S. cerevisiae* alkali-metal-cation antiporters**

| Antipporter | CaCnh1p | CdCnh1p | CpCnh1p | CpCnh1p | GtCnh1p | GtCnh1p |
|-------------|---------|---------|---------|---------|---------|---------|
| CaCnh1p     | -       | 84.2/98.6/67.5 | 46.0/69.9/20.5 | 46.3/88.3/39.6 | 69.0/93.1/41.0 | 44.0/68.7/23.5 |
| CdCnh1p     | 46.2/69.4/22.2 | -       | 62.7/88.1/35.3 | 67.5/93.1/43.8 | 46.7/69.5/19.8 |
| CpCnh1p     | 42.7/70.1/21.4 | 39.7/69.4/19.6 | -       | 39.7/69.4/19.6 | 57.0/86.8/31.8 |
| CtCnh1p     | 57.1/86.8/34.2 | 38.8/68.7/20.6 | 69.0/93.1/41.0 | -       | 41.0/69.4/19.0 |
| ScNha1p     | 69.4 | 86.8 | 93.1 | 93.1 | - | 68.7 |

RJKVYAHHR (in CaCnh1p aa 639–664) preceding conserved region no. 5 [18]; was found in the four *Candida* species, except for *C. glabrata*.

The functional expression of all the constructs were first tested in drop experiments, which showed that 1) the presence of the constructs did not influence the growth rate of cells in standard media, i.e. the heterologous expression of these membrane proteins was not toxic for *S. cerevisiae*, and 2) the expression of both GFP-tagged and non-tagged *CdCnh1* and *CpCnh1* proteins brought about the same ability to grow on 800 mM NaCl or 1800 mM KCl, as did the positive controls with *ScNha1* and *CaCnh1* proteins, whereas the cells without antiporters were not able to grow (not shown). This result also confirmed that the C-terminal GFP-tagging did not influence the activity of the antiporters. In order to estimate the substrate specificity and transport capacity of the antiporters, BW31a cells expressing the four antiporters or transformed with an empty vector were spotted on a series of YNB plates containing increasing NaCl, KCl, LiCl and RbCl concentrations. Cells expressing *CpCnh1p* were able to grow in the highest concentrations of salts, as did cells expressing *CaCnh1p*. Both these *Candida* antiporters conferred a slightly higher tolerance to the cells than equivalent expression of the native *S. cerevisiae* antiporter, Nha1p (Figure 2). The tolerance of cells expressing *CdCnh1p* to high external potassium and rubidium was almost the same as for cells expressing *CpCnh1p* and *CaCnh1p* (Figure 2), but their tolerance to toxic cations was significantly lower, only 1000 mM NaCl, and there was no increase in LiCl tolerance compared to cells with the empty vector (30 mM LiCl in both cases).

The proton-antiport mechanism of these Cnh1 proteins was verified in a series of drop tests on plates with various pH values (Table 3). As was previously thought, the cells expressing antiporters showed the highest salt tolerance (at least for three of their four substrates) when grown at lower external pH, i.e. in conditions where the proton gradient across the plasma membrane is the highest. Surprisingly, the expression of both the *S. cerevisiae* and *C. dubliniensis* antiporters did not increase the cell tolerance to lithium cations at pH 3.5, suggesting that Li⁺ was not recognized as their substrate under these conditions. On the other hand, all four antiporters were partially active, even at an externally neutral pH 7.0, as their presence ena-
bled the cells to support higher salt concentrations. This detailed study confirmed again that \( \text{CdCnh1p} \) has the least ability to improve the salt tolerance of cells (Table 3).

Localization of \( \text{CdCnh1} \) and \( \text{CpCnh1} \) antiporters in \( \text{S. cerevisiae} \) cells
As mentioned above, C-terminal GFP-tagging of the \( \text{CdCnh1} \) and \( \text{CpCnh1} \) proteins did not affect their functionality. Both antiporters improved the cell salt tolerance to a similar degree as the non-tagged versions, and fluorescence microscopy localized them to the plasma membrane of \( \text{S. cerevisiae} \) BW31a cells (Figure 3) and not to the membranes of intracellular organelles. This result indicates a high probability of the same localization in their organisms of origin, as was previously shown for \( \text{C. albicans} \) Cnh1p [20].

Western blot analysis of antiporters' amount in cells
To verify whether the use of the same vector and promoter for expression ensures similar levels of the four antiporters in BW31a cells, the GFP-tagged proteins were visualized on western blots. Fig. 4 shows that 1) the size of \( \text{Sc} \) and \( \text{Cp} \) antiporters was alike, similarly as the size of \( \text{Ca} \) and \( \text{Cd} \) transporters (and in agreement with the size deduced

Table 3: pH dependence of salt tolerance of BW31a cells expressing various antiporters.

| Antiporter | NaCl (mM) | KCl (mM) | LiCl (mM) | pH_{out} |
|------------|-----------|-----------|-----------|----------|
|            | 3.5       | 5.5       | 7.0       |          |
|            | 3.5       | 5.5       | 7.0       |          |
|            | 3.5       | 5.5       | 7.0       |          |
| -          | <500      | 200       | 75        |          |
| ScNha1p    | 1300      | 800       | 150       | 1000     |
| CaCnh1p    | 1300      | 1000      | 200       | 2000     |
| CdCnh1p    | 1100      | 800       | 200       | 2000     |
| CpCnh1p    | 1300      | 1000      | 200       | 2000     |

Maximum salt concentrations [mM] allowing growth of \( \text{S. cerevisiae} \) BW31a cells (transformed with the YEp352 plasmid either empty or harbouring genes encoding antiporters) on YNB media with various pH values.
from the gene sequence, cf. Table 1 and 2) the quantity of antiporters in extracts of exponentially growing cells was similar, the highest amount apparently being observed for CdCnh1p and the lowest one for CpCnh1 antiporter. The analysis was repeated three times with the same result. The fluorescence microscopy and western blot analysis confirmed that the observed low activity of CdCnh1p and the high activity of CpCnh1 antiporter did not reflect different protein levels in the cells.

**Efflux of Na\(^+\) and K\(^+\) from BW31a cells expressing various antiporters**

To confirm the results from drop test experiments and to determine the activity and efflux rate of CdCnh1p and CpCnh1p, the efflux of K\(^+\) and Na\(^+\) was directly measured. The loss of K\(^+\) from BW31a cells expressing *C. dubliniensis* or *C. parapsilosis* antiporters was measured directly; to measure Na\(^+\) efflux, cell preloading with 100 mM NaCl was necessary (cf. Methods). Cells expressing the CaCnh1 antiporter served as positive, and cells transformed with an empty vector as negative controls, respectively. The initial internal concentration of K\(^+\) in exponentially growing cells was almost the same in all strains, in the representative experiment about 549.5 ± 25.7 nmol (mg dry wt\(^-1\)). After preloading, cells contained 110.5 ± 6.8 nmol (mg dry wt\(^-1\)) Na\(^+\). As shown in Figure 5 and Table 4, CpCnh1p exported Na\(^+\) and K\(^+\) much more efficiently than CdCnh1p. Within 60 minutes, cells with CpCnh1p lost 80% of their internal sodium and 32% of their potassium compared to cells with CdCnh1p, which over the same period only lost 38% of their sodium and 20% of their potassium. The *C. albicans* antiporter was the most effective, exporting 82% of its host cells' sodium and 55% of their potassium in 60 min, which agrees with previously published results [19]. The Na\(^+\) efflux curves for CaCnh1p and CpCnh1p are very similar and almost exponential; most of the sodium is exported in about 40 minutes. The efflux of sodium via CdCnh1p is linear and slow. The sodium efflux via all three antiporters is faster than their potassium efflux, though the initial intracellular concentration of K\(^+\) is much higher than that of Na\(^+\), approx. 300 vs. 50 mM. These results suggest that the *Candida* antiporters have, at least upon heterologous expression in *S. cerevisiae*, a higher affinity for sodium than for potassium cations.

**Discussion and Conclusion**

The variable sensitivity of *Candida* species to NaCl has been observed and exploited previously, mainly in connection with the original niche of the species, e.g. the high NaCl tolerance of *C. parapsilosis* strains isolated from sea water or hypersaline brines [22,27] or in clinical microbiological tests aiming to distinguish among *Candida* species (NaCl sensitivity of *C. dubliniensis*[3]). In this work, we performed a series of tests to characterize the tolerance of four different pathogenic *Candida* species to different alkali metal cations, and addressed the question of whether the observed differences in salt tolerance could be based on the transport activity and/or specificity of *Candidas’* plasma-membrane Na\(^+\)/H\(^+\) antiporters. Our results showed clearly that, besides differences in their tolerance to sodium, the four species tested also differ in their sensitivity to highly toxic lithium and their tolerance of high external concentrations of non-toxic potassium.
cations (K\(^+\), Li\(^+\), Na\(^+\), Rb\(^+\)), but transport them with varying substrate specificity. They recognize at least four different ion exporters revealed that both antiporters have broad transport capacities exceed that of Na\(^+\)/H\(^+\) antiporters are not the most important factor in this activity [7] and that a single amino-acid exchange in one of the transmembrane domains can significantly influence both the substrate specificity and transport capacity of the antiporter [28,29].

Though the C. parapsilosis Cnh1 antipporter mediates a high and efficient sodium efflux from S. cerevisiae cells (similarly as the CaCNH1p, Figure 5 and Table 4), its physiological role in C. parapsilosis remains to be established. A recent study [20] showed that the Cnh1 antipporter is mainly important in potassium homeostasis in C. albicans cells and its role in Na\(^+\) detoxification is rather marginal.

The differences observed in alkali-metal-cation tolerance between C. dubliniensis and C. parapsilosis/C. albicans species were also found upon testing the alkali-metal-cation tolerance of S. cerevisiae cells expressing the antipporters of these three species. Efflux measurements confirmed that the differing tolerances of S. cerevisiae cells were based on the differing transport activities of the Candida antipporters. Altogether, our results suggest that the activity of plasma-membrane Na\(^+\)/H\(^+\) antipporters is one of the factors determining the tolerance to high external concentrations of alkali metal cations in pathogenic Candida species.

**Methods**

**Yeast strains, media and growth conditions**

To determine the salt tolerance of various Candida species, C. albicans SC5314, C. glabrata ATCC2001, C. dubliniensis CD36 and C. parapsilosis CBS604 were used, together with S. cerevisiae S288c as a control. The CdCNH1 and CpCNH1 genes were isolated from C. dubliniensis CD36 and C. parapsilosis CBS604, and heterologously expressed in S. cerevisiae BW31a (ena1-4Δ nha1Δ, W303 derivative, [8]). Yeast cells were grown in YPD or YNB-NH\(_4\) media supplemented with 2% glucose at 30°C. Salts were added to the media prior to and auxotrophic supplements after autoclaving.

**DNA manipulations, plasmid construction and DNA sequencing**

For DNA manipulations, standard protocols [30] were used. The CdCNH1 and CpCNH1 gene were amplified by
PCR with platinum Pfx polymerase with proofreading activity (Invitrogen) using their isolated genomic DNA [31] as a template.

Plasmids for the heterologous expression of these antiporters in S. cerevisiae were constructed by homologous recombination in BW31a cells. The oligonucleotides used are listed in Table 5. Two types of plasmids were constructed. CdCNH1 and CpCNH1 coding sequences were cloned behind the ScNHA1 promoter either in multicopy YEp352 or in pGRU1, enabling C-terminal GFP tagging [7]. All constructs were analysed by sequencing in an ABI PRISM 3100 DNA sequencer using the BigDye Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems).

Salt tolerance determination

The cell tolerance to various alkali metal cations was determined by drop test experiments. 3 μl of serial 10-fold dilutions of saturated cell cultures were spotted on solid media. For Candida species, YPD – 3% agar plates supplemented with 10 mM KCl or 10 mM RbCl to prevent Na+ or K+ reuptake respectively) was used. For the determination of Na+ efflux, the cells were preloaded for 60 min with 100 mM NaCl in YNB adjusted to pH 7.0 with NH₄OH. The K⁺ efflux was measured directly in the harvested cells. Samples were taken from cell incubation cultures at regular time intervals over a period of 60 min and the cellular cation content was determined by atomic absorption spectroscopy [7]. Each efflux experiment was repeated at least three times and representative results are shown.

Microscopy analysis

Exponential phase cells (grown in YNB at 30°C, OD₆₀₀ ≈ 0.15) expressing the CdCNH1 or CpCNH1 gene tagged with the GFP sequence were viewed with an Olympus AX70 microscope using a U-MWB cube with a 450–480 nm excitation filter and 515 nm barrier filter. The micrographs were recorded with a DP70 digital camera using the program DP Controller. For whole-cell pictures, Nomarski optics was used.

Immunoblotting

Exponentially growing (OD₆₀₀ ≈ 0.15) BW31a cells expressing GFP-tagged antiporters were harvested and concentrated by centrifugation to OD₆₀₀ ≈ 3.0. The proteins were extracted according to [32] with some modifications. After resuspension of cell pellet in 150 μl freshly prepared 1.85 M NaOH with 7.5% β-mercaptoethanol and incubation for 15 min on ice, 150 μl of cold 50% trichloroacetic acid were added. After incubation on ice for 20 min the collection of precipitates by centrifugation at 20,000 × g for 20 min followed. The pellet was resuspended in 190 μl of cold 50% trichloroacetic acid and incubation for 30 min at 37°C the samples were centrifuged at 20,000 × g for 30 min. Supernatant (7.5 μl) were directly loaded on 8% glycin gel and separated by polyacrylamid gel electrophoresis (PAGE). Separated proteins were transferred via electroblotting on nitrocellulose membrane. To detect the GFP-tagged proteins on membranes, rabbit polyclonal anti-GFP antibody (Santa Cruz Biotech., diluted 1:200), secondary goat anti-rabbit IgG antibody with conjugated peroxidase (BioRad, diluted 1:10,000) and ECL detection kit (Pierce) were used.

Table 5: Oligonucleotides used for amplification of C. dubliniensis and C. parapsilosis CNH1 genes. Sections homologous to CNH1 genes are underlined.

| Primer                  | Sequence (5’-3’)                                                                 |
|------------------------|---------------------------------------------------------------------------------|
| CdCNH1-F               | TTTTTTGTACATTATAAAAAAAAAAAATCCCTGACTTATGCTAGATCTTATGGCTGGATCGTTAGAA             |
| CdCNH1-R               | ACCAGCTGTAAACGGACGCTGAAGCTCAGATGGCTGGATCGTTAGATCTTATGGCTGGATCGTTAGAA           |
| CpCNH1-F               | TTTTTTGTACATTATAAAAAAAAAAAATCCCTGACTTATGCTAGATCTTATGGCTGGATCGTTAGAA           |
| CpCNH1-R               | ACCAGCTGTAAACGGACGCTGAAGCTCAGATGGCTGGATCGTTAGATCTTATGGCTGGATCGTTAGAA           |
| CpCNH1-GFP-R           | CATTCTTAAAGCTCCGGAGCTTGCATGCGTTGCAGTTGCAGTGAATGCTGCAAGCTTGCAGTTGCAGTTGCAGTTGACG |
**Authors’ contributions**

HS was involved in the design phase, YK provided the experimental data, and both authors drafted the manuscript, read and approved its final version.

**Acknowledgements**

This work was supported by the EU grant MRTN-CT-2004-512481 - CanTrain and Czech grants LC531 and AV0Z5010509. We thank Olga Zimmernannova for helpful discussions, Gary Moran, Karl Kuchler and Jozef Nosek for providing Candida strains, and Jaroslav Horak for the help with Western blots.

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