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PAPER

The interplay between vacuolar and siderophore-mediated iron storage in *Aspergillus fumigatus*[†]

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Iron is an essential element for all eukaryotes but its excess has deleterious effects. *Aspergillus fumigatus* produces extracellular siderophores for iron uptake and the intracellular siderophore ferricrocin (FC) for distribution and storage of iron. Iron excess has previously been shown to increase the content of ferric FC and the expression of the putative vacuolar iron importer CccA (AFUA_4G12530), indicating a role of both the vacuole and FC in iron detoxification. In this study, we show that CccA-deficiency decreases iron resistance in particular in combination with derepressed iron uptake, while overproduction of CccA increases iron resistance. Green fluorescence protein-tagging confirmed localization of CccA in the vacuolar membrane. In contrast to CccA-deficiency, inactivation of FC biosynthesis did not affect iron resistance, which indicates that vacuolar rather than FC-mediated iron storage is the major iron detoxifying mechanism. After uptake, extracellular siderophore backbones are hydrolyzed and recycled. Lack of FC, CccA, and in particular lack of both increased the cellular content of iron chelated by siderophore breakdown products. These data indicate that the transfer of iron from extracellular siderophores to the metabolism, FC or the vacuole precedes recycling of siderophore breakdown products. Furthermore, this study indicates that CccA does not play an exclusive role in vacuolar iron storage for nutritional reuse.

Introduction

Due to its crucial role in a wide variety of cellular oxidation–reduction reactions, iron is an essential element for all eukaryotes. Nevertheless, iron excess can cause oxidative stress *via* the Haber–Weiss/Fenton chemistry.¹ Therefore, cellular iron homeostasis has to maintain a delicate balance between sufficient iron supply and prevention of iron toxicity. As microorganisms lack iron excretory mechanisms, the maintenance of iron homeostasis results from a tight regulation of iron acquisition and iron storage.

Aspergillus fumigatus is a ubiquitous saprophytic mold, but at the same time the most common cause of airborne fungal infections in immunocompromised patients.² During both saprophytic growth and infection, this opportunistic pathogen has to cope with rapidly changing environments, including

high variations of iron availability. For the cellular import, *A. fumigatus* employs low-affinity iron acquisition and two high-affinity iron uptake systems, reductive iron assimilation and siderophore (low molecular mass, ferric iron chelators)-assisted iron uptake.³ In order to mobilize extracellular iron *A. fumigatus* excretes two different siderophores, fusarinine C (FSC) and triacetyl-fusarinine C (TAFC).^{4,5} After the cellular uptake of the ferri-forms of FSC and TAFC *via* specific siderophore-iron transporters, these siderophores are degraded for iron release.^{6,7} Extracellular siderophores are found in most fungal and bacterial species as well as in some plants.⁶ In contrast to plants and bacteria, most fungi also possess intracellular siderophores. *A. fumigatus* produces two intracellular siderophores, ferricrocin (FC) for hyphal iron storage and distribution, and hydroxyferricrocin (HFC) for conidial iron storage.^{4,5} In recent years, individual components of the *A. fumigatus* siderophore and reductive iron assimilatory systems have been characterized at the molecular level,⁶ *e.g.* the ornithine hydroxylase SidA that is essential for biosynthesis of both intra- and extracellular siderophores, the non-ribosomal peptide synthetase SidC that is indispensable for biosynthesis of intra- but not extracellular siderophores, and the iron permease FtrA that is essential for reductive iron assimilation. Due to its role in intracellular iron storage and distribution, FC plays a

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crucial role in germination and conidiation.⁴ Moreover, FC has also been shown to be essential for sexual development in *Aspergillus nidulans*.⁸ Biosynthesis of extra- and intracellular siderophores proved to be crucial for full virulence of *A. fumigatus* in various aspergillosis models,^{3,4,9,10} and for intracellular survival after phagocytosis by murine alveolar macrophages.^{11,12}

Inactivation of the transcription factor SreA, which represses excessive iron uptake *via* the downregulation of siderophore-mediated and reductive iron assimilation, results in toxic accumulation of iron under iron-replete conditions and subsequent growth retardation.^{6,13} Excessive iron uptake has been shown to elevate the cellular FC content, which indicates an additional role of FC in iron detoxification. Furthermore, iron excess leads to an upregulation of the transcript level of AFUA_4G12530, termed CccA,¹³ which displays significant similarity to *Saccharomyces cerevisiae* Ccc1 and *Arabidopsis thaliana* Vit1. Ccc1 mediates vacuolar import of iron and manganese to prevent toxic levels of these metals in the cytosol.^{14,15} Vit1 has also been shown to transport iron into the vacuole.¹⁶ Both *S. cerevisiae* and *A. thaliana* possess vacuolar iron export systems for the reuse of vacuolar-stored iron.^{17–20}

Apart from *S. cerevisiae* and *A. thaliana*, which both lack siderophores, the role of vacuolar iron storage has not been functionally characterized in any other species. In the present study we characterized the role of *A. fumigatus* CccA in

vacuolar iron storage and analyzed the interplay between siderophore-mediated and vacuolar iron storage.

Results

A. fumigatus encodes two homologs of *S. cerevisiae* Ccc1

BLASTP homology searches (<http://blast.ncbi.nlm.nih.gov/Blast.cgi>, http://www.ncbi.nlm.nih.gov/sutils/genom_table.cgi?organism=fungi) and phylogenetic analysis identified homologs of the *S. cerevisiae* vacuolar iron importer Ccc1 in all fungal genomes available with the exception of *Microsporidia*, *Pleosporales* and the *Exobasidiomycete* *Malassezia globosa*, *i.e.*, in 68 of 76 fungal species (Fig. 1 and data not shown). Interestingly, Ccc1 homologs are not only found in fungi and plants but also in bacteria such as *Pseudomonas aeruginosa* and *Streptococcus sanguinis*. In contrast to most other fungal species, *Trichocomaceae* (including *Neosartorya*, *Aspergillus*, and *Penicillium* species) and *Hypocreales* (including *Gibberella*, *Nectria*, and *Metarhizium* species) encode two paralogs (Fig. 1). An amino acid sequence alignment of the Ccc1 homologs from *A. fumigatus*, *Penicillium chrysogenum*, *S. cerevisiae* and *A. thaliana* is shown in Fig. S1 (ESI[†]). All these proteins are about 300 amino acids in length and display 12% identity (56 amino acid residues) in the 228 amino acid core region.

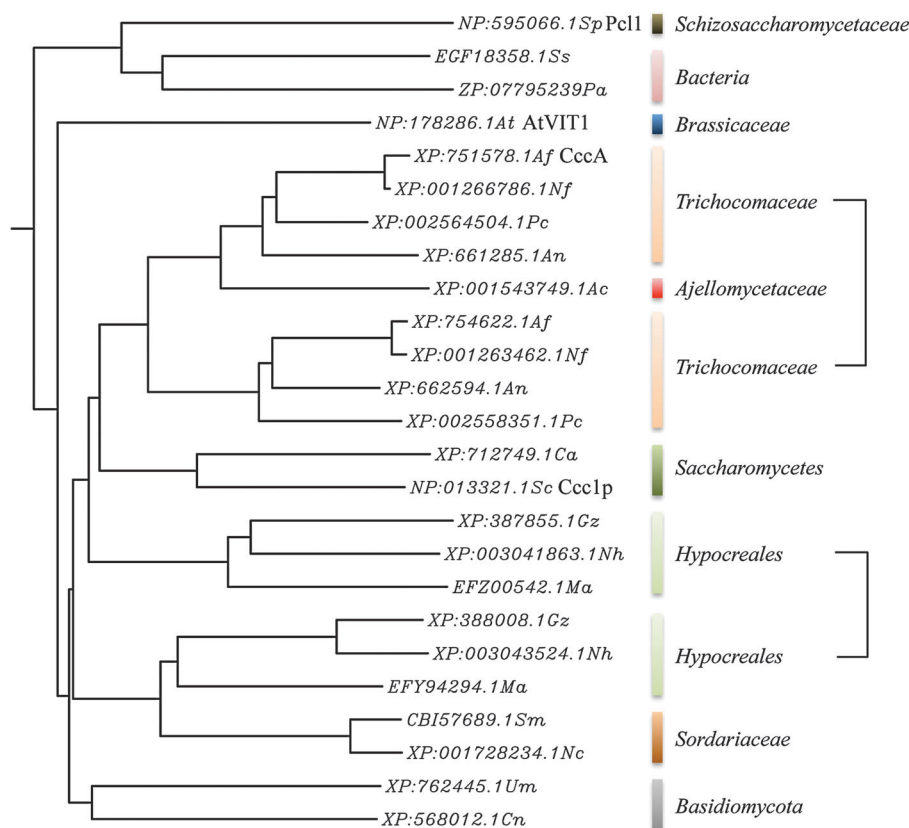


Fig. 1 Phylogenetic analysis (rooted neighbour joining tree) of CccA homologs from selected fungi, bacteria and plants. The *Trichocomaceae* and *Hypocreales* paralogs are marked in brackets. Species abbreviations: Af, *A. fumigatus*; An, *A. nidulans*; At, *A. thaliana*; Ca, *Candida albicans*; Cn, *Cryptococcus neoformans*; Gz, *Gibberella zeae*; Ma, *Metarhizium anisopliae*; Nc, *Neurospora crassa*; Nf, *Neosartorya fischeri*; Nh, *Nectria haematococca*; Pa, *P. aeruginosa*; Pc, *Penicillium chrysogenum*; Ss, *S. sanguinis*; Sc, *S. cerevisiae*; Sm, *Sordaria macrospora*; Sp, *Schizosaccharomyces pombe*; Um, *Ustilago maydis*.

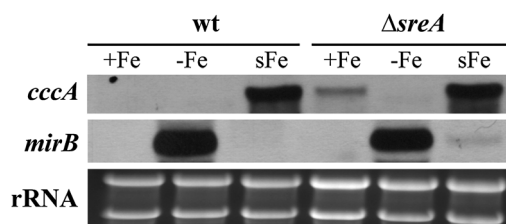


Fig. 2 Expression of *cccA* is induced by iron excess. For Northern analysis of *cccA*, *A. fumigatus* wt and Δ *sreA* strains were grown for 24 h in liquid flask cultures under iron-depleted (–Fe), iron-replete conditions (+Fe; 30 μ M FeSO_4) or shifted from –Fe to 0.1 mM FeSO_4 for 1 h (sFe) to force excessive iron uptake. Ethidium bromide-stained rRNA is shown as a control for loading and quality of RNA. As a control for the cellular iron status, Northern analysis was also performed with the siderophore transporter-encoding *mirB* gene, the expression of which is repressed by iron.¹³

Expression of CccA is induced by iron

In *S. cerevisiae*, Ccc1 expression is transcriptionally induced by iron.²¹ As shown previously¹³ and confirmed in this study, the transcript level of AFUA_4G12530, termed CccA, is upregulated in the *A. fumigatus* wild type strain ATCC46645 (wt) during a shift from iron starvation to iron sufficiency (Fig. 2). Furthermore, derepression of iron uptake under iron-replete conditions due to deficiency in the iron uptake repressor SreA increased the *cccA* steady-state transcript level. As a control for the cellular iron status, Northern analysis was also performed with the siderophore transporter-encoding *mirB* gene, the expression of which is in contrast to *cccA* repressed by iron.¹³ These data indicate the function of CccA in the iron metabolism.

As observed previously,¹³ transcripts of *cccA* paralog AFUA_3G09970, termed *cccB*, could not be detected in Northern analysis, neither in the wt nor in the SreA-deficient strain (data not shown), which suggests a minor role of this gene in iron metabolism. As indicated by phylogenetic analysis (Fig. 1), BLAST searches revealed that the CccA and CccB orthologs are more closely related to each other than to the other homologs (Table S3, ESI†). Interestingly, the CccB orthologs from *A. fumigatus* and *P. chrysogenum* display an extended C-terminus, an insertion of two amino acid residues and a deletion of one amino acid in the core region (Fig. S1, ESI†).

CccA but not siderophore biosynthesis is important for iron resistance

In order to functionally characterize CccA, the *cccA* coding region was replaced by the pyrithiamine resistance gene *ptrA* in the *A. fumigatus* wt, yielding strain Δ *cccA*, as described in Materials and methods and Fig. S1 (ESI†). Additionally, CccA-deficiency was combined with deficiency in siderophore biosynthesis and reductive iron assimilation (Δ *cccA* Δ *sidA* Δ *ftrA*), FC biosynthesis (Δ *cccA* Δ *sidC*), the iron uptake repressor SreA (Δ *cccA* Δ *sreA*), as well as SreA and siderophore biosynthesis (Δ *cccA* Δ *sidA* Δ *sreA*). Radial growth of wt and mutant strains was compared on plates containing different iron concentrations (Fig. 3). As shown previously,^{3,4} deficiency in siderophore biosynthesis (Δ *sidA*) blocked growth in the presence of the iron

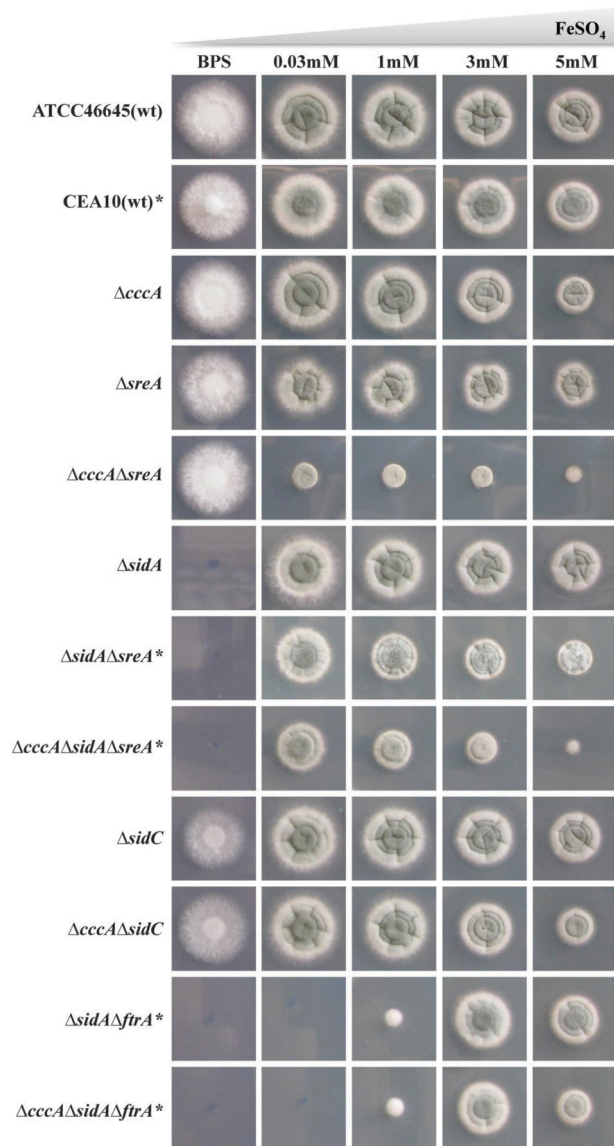


Fig. 3 CccA-deficiency decreases iron resistance. 10^4 conidia of the respective fungal strain were point-inoculated on AMM plates containing the indicated iron concentration and incubated for 48 h at 37 °C. The BPS medium contained 0.2 mM of the ferrous iron chelator bathophenanthroline disulfonic acid, which inhibits reductive iron assimilation. CEA10 genetic background strains are marked with *, the other strains have an ATCC46645 background. The fungal strains are listed and described in Table 1.

chelator bathophenanthroline disulfonic acid (BPS). FC-deficiency (compare wt/ Δ *sidC*) decreased growth in the presence of BPS, and deficiency in both siderophore-mediated and reductive iron assimilation (compare wt/ Δ *sidA* Δ *ftrA*) blocked growth up to an iron concentration of 30 μ M. Genetically unidentified low-affinity iron uptake enabled partial growth of Δ *sidA* Δ *ftrA* at an iron concentration of 1 mM and wt-like growth at \geq 3 mM. SreA-deficiency decreased growth under conditions of high iron availability, which was best visible at 5 mM, due to toxicity of excessive iron uptake (compare wt/ Δ *sreA*).¹³ In all SreA-functional genetic backgrounds, CccA-deficiency decreased growth under high iron conditions, which was best

seen at 5 mM iron (compare wt/ $\Delta cccA$, $\Delta sidC/\Delta cccA\Delta sidC$, $\Delta sidA\Delta firA/\Delta cccA\Delta sidA\Delta firA$). Combined deficiency in both SreA and CccA reduced growth already at iron concentrations $\geq 30 \mu\text{M}$ (compare $\Delta sreA/\Delta cccA\Delta sreA$, $\Delta sidA\Delta sreA/\Delta cccA\Delta sidA\Delta sreA$). Taken together, CccA-deficiency led to decreased iron resistance, especially in strains with derepressed iron uptake due to lack of SreA. These data indicate a crucial role of CccA in iron detoxification. Consistently, CccA-deficiency did not affect growth in any genetic background under iron limitation (Fig. 3).

In further agreement with the function of CccA in iron detoxification, expression of *cccA* under the control of the *P. chrysogenum xylP* promoter, which is xylose-inducible,^{22,23} increased iron resistance in wt (strain wt $cccA^{oe}$) and $\Delta cccA$ (strain $\Delta cccAcccA^{oe}$) strains under inducing but not repressing conditions (Fig. 4A).

In *Ustilago maydis*, simultaneous deficiency in the SreA ortholog Urbs1 and siderophore biosynthesis appears to be lethal, indicating an essential role of intracellular siderophores in iron detoxification.²⁴ In *A. fumigatus*, inactivation of neither the entire siderophore production nor FC biosynthesis decreased iron resistance in any background tested (compare wt/ $\Delta sidA$, wt/ $\Delta sidC$). Moreover, the simultaneous deficiency in SreA and siderophore biosynthesis was not lethal and did not result in increased iron sensitivity ($\Delta sreA\Delta sidA$). Even simultaneous deficiency in SreA, siderophore biosynthesis and CccA was not lethal ($\Delta cccA\Delta sreA\Delta sidA$). These data demonstrate that in *A. fumigatus* siderophore biosynthesis is not essential for iron detoxification. Notably, the mutants described here have two different genetic backgrounds (ATCC46645 and CEA10) but in the comparisons above only mutants with the same genetic background have been included. Moreover, iron metabolism and siderophore metabolism are highly similar in these two genetic backgrounds.^{3,25}

It is generally believed that iron leads to damage through the generation of oxygen radicals.¹ To investigate the basis of iron susceptibility caused by CccA-deficiency we analyzed the impact of CccA-deficiency in wt and $\Delta sidC$ genetic backgrounds on resistance to oxidative stress caused by 2 mM hydrogen peroxide under iron starvation, iron sufficiency (30 μM) and iron excess (3 mM) in radial growth assays. Remarkably, CccA-deficiency did not decrease resistance to hydrogen peroxide under any of the tested conditions indicating that iron toxicity may not be solely due to oxidative damage (data not shown). In addition, neither hydrogen peroxide-caused oxidative stress nor inactivation of various components of the oxidative damage response including the master regulator of oxidative stress response, Yap1, affected the iron susceptibility of the *S. cerevisiae* $\Delta ccc1$ mutant.²⁶ Moreover, iron toxicity mediated by Ccc1-deficiency occurs in this system under both aerobic and anaerobic conditions.²⁶

CccA is located in the vacuolar membrane

To investigate the localization of CccA, we generated an *A. fumigatus* strain expressing CccA N-terminally tagged with enhanced green fluorescence protein (GFP) under the control of the constitutive *cgxA*-XAH promoter in $\Delta cccA$ (strain $\Delta cccAcccA^{gfp}$) as described in Materials and methods and Fig. S1 (ESI[†]).²⁷ Epifluorescence microscopy revealed that GFP-CccA co-localizes to the vacuolar membrane with the FM4-64 marker (Fig. 4B) as previously shown for N-terminal GFP fusions of the CccA homologs Ccc1 in *S. cerevisiae* and Vit 1 in *A. thaliana*.^{15,16} The GFP-CccA fusion partially cured the iron sensitivity of $\Delta cccA$ (Fig. 4A) indicating at least partial correct biochemical function and localization. Lack of full complementation might be due to decreased transport

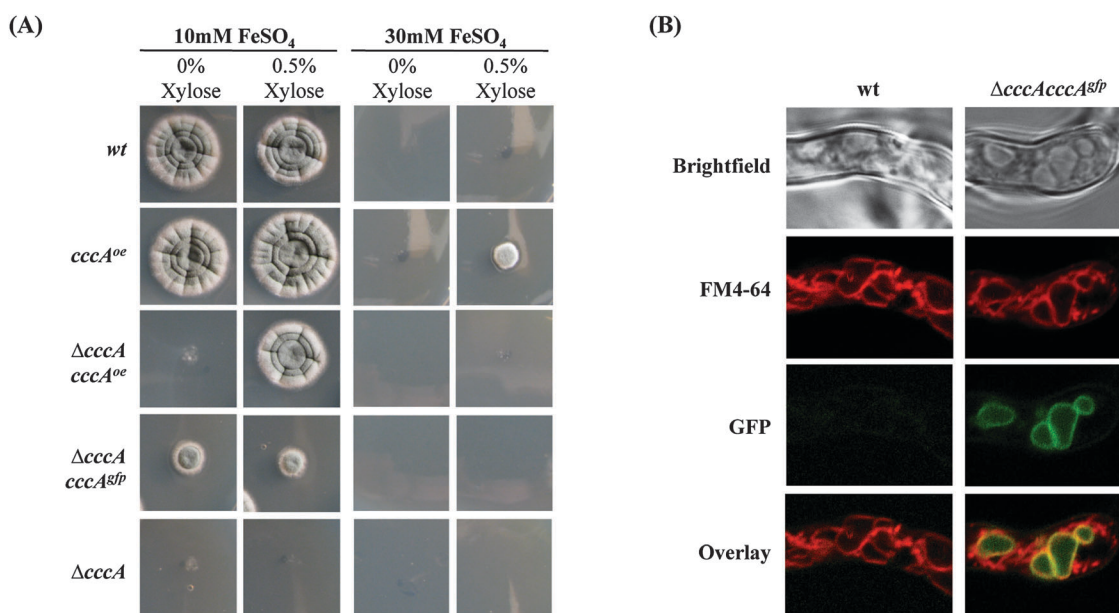


Fig. 4 Xylose-induced expression of *cccA* increases iron resistance in wt and $\Delta cccA$ backgrounds (A) and GFP-tagged CccA localizes to the vacuolar membrane (B). (A) 10^4 conidia of the respective fungal strain were point-inoculated on AMM plates containing FeSO₄ at a final concentration of 10 mM and 30 mM, respectively, with or without 0.5% xylose and incubated for 48 h at 37 °C. (B) The GFP fluorescence of $\Delta cccAcccA^{gfp}$ was visualized using confocal microscopy. Cellular membranes were stained by preincubation with 8 μM FM4-64 for 1 h.

activity caused by the GFP-fusion or inappropriate expression as a result of the use of a heterologous promoter. The latter can be excluded as Northern analysis indicated an even higher transcript level of GFP-tagged *cccA* compared to *cccA* (data not shown). Notably, C-terminal GFP-tagged CccA under the native promoter was not functional (data not shown).

Taken together, the localization of CccA in the vacuolar membrane suggests that CccA transports iron into the vacuole as shown previously for its homologs in *S. cerevisiae* and *A. thaliana*.^{15,16}

Vacuolar iron storage increases the degradation of siderophores

After cellular uptake, ferric forms of TAFC and FSC are degraded *via* esterase-mediated hydrolysis, the iron is transferred to the metabolism or FC and the FSC/TAFC degradation products are recycled.⁷ To analyse a possible interplay between vacuolar iron storage and intracellular siderophores, wt, $\Delta cccA$, $\Delta sidC$, and $\Delta cccA\Delta sidC$ strains were grown in liquid cultures for 20 h under iron starvation to activate siderophore production. Subsequently, 100 μ M iron were added to the growth media and the strains were further incubated for 4 h, which resulted in uptake of iron chelated by FSC and TAFC. Mycelia were harvested and cell extracts were analyzed by reversed-phase HPLC and mass spectrometry (Fig. 5). After the iron shift, the cell extracts of all strains were orange-coloured (Fig. 5A), which is typical for abundant iron resources chelated by siderophores or siderophore breakdown products.⁷ Interestingly, the orange coloration of the cell extract was more intense in the case of $\Delta cccA$ compared to wt and $\Delta cccA\Delta sidC$ compared to $\Delta sidC$. The latter indicated that the orange coloration was not exclusively caused by ferric FC as $\Delta sidC$ strains lack FC. Reversed-phase HPLC (Fig. 5) combined with mass spectrometry identified desferri-FC as a main compound in the wt cell extract during iron starvation. After the addition of iron, the desferri-FC was converted to FC and various FSC/TAFC degradation products accumulated. Compared to wt, CccA-deficiency did not affect the desferri-FC content during iron starvation but resulted in increased accumulation of FSC/TAFC degradation products, in particular *N*⁵-anhydromevalonyl-*N*⁵-hydroxyornithine (fusarinine). SidC-deficiency blocked FC production and increased accumulation of FSC/TAFC degradation products. The highest levels of FSC/TAFC degradation products were observed in $\Delta cccA\Delta sidC$, which lacks both vacuolar iron storage and FC-mediated intracellular iron chelation. The orange coloration of the cell extract and the detection of ferric FSC/TAFC breakdown products (Fig. 5) underline that the intracellular siderophore breakdown products indeed chelate iron *in vivo*. FSC and TAFC are cyclic (*N*⁵-anhydromevalonyl-*N*⁵-hydroxyornithine)₃ and (*N*²-acetyl-*N*⁵-anhydromevalonyl-*N*⁵-hydroxyornithine)₃, respectively, connected by head to tail ester bonds.²⁸ The detection of *N*⁵-anhydromevalonyl-*N*⁵-hydroxyornithine-*N*²-acetyl-*N*⁵-anhydromevalonyl-*N*⁵-hydroxyornithine demonstrates that TAFC is degraded by hydrolysis and deacetylation and therefore, the detected *N*⁵-anhydromevalonyl-*N*⁵-hydroxyornithine is most likely not only derived from degradation of FSC but also TAFC.

Taken together, the increased accumulation of FSC/TAFC degradation products due to lack of vacuolar iron storage or intracellular iron chelation by FC and in particular both indicates that the transfer of iron from extracellular siderophores to the metabolism, FC or the vacuole precedes recycling of siderophore breakdown products. In other words, the absence of FC and/or vacuolar iron storage decreases the release of iron from iron-chelating siderophore breakdown products. As chelation decreases iron toxicity, this mechanism might represent another iron detoxification strategy.

CccA-deficiency does not affect the putative nutritional use of vacuolar iron

Both *S. cerevisiae* and *A. thaliana* are able to recycle vacuolar-stored iron and deficiency in the CccA homolog blocks the use of the vacuole as an iron storage compartment in both organisms.^{15,17–20} The current study indicates that CccA is a major vacuolar iron transporter in *A. fumigatus*.

To study the possible recycling of vacuolar iron in *A. fumigatus*, we compared in a first approach wt and $\Delta cccA$ strains during a shift from iron sufficiency (30 μ M TAFC) to iron starvation (Fig. 6). The two strains did not significantly differ in timing and the extent of siderophore synthesis (Fig. 6A) or biomass production (Fig. 6B). As reuse of vacuolar iron is expected to delay iron starvation and consequently siderophore production as a symptom of iron starvation in wt compared to $\Delta cccA$, these data indicate that CccA does not play an exclusive role in the use of the vacuole as a nutritional iron source.

In a second approach, $\Delta sidA\Delta ftrA$ and $\Delta cccA\Delta sidA\Delta ftrA$ strains, which lack both reductive and siderophore-mediated iron assimilation and therefore high-affinity iron uptake, were compared to exclude compensation of possible defects by uptake of extracellular iron (Fig. 6C). The two mutants were iron-saturated by incubation for 16 h at 37 °C on plates containing either TAFC or FC separated from the growth media by dialysis membranes. Subsequently, the mycelia-containing dialysis membranes were transferred onto iron-depleted minimal media, from which they are unable to acquire iron (compare Fig. 3). After incubation for five days, CccA-deficiency did not decrease growth but even supported increased conidiation. Again, these data make an exclusive role of CccA in nutritional vacuolar iron storage unlikely.

In the third approach, $\Delta sidA\Delta ftrA$ and $\Delta cccA\Delta sidA\Delta ftrA$ strains were iron-saturated by culturing in liquid media for 18 h with TAFC or FC as an iron source. Subsequently, mycelia were washed and transferred into iron-depleted media and the iron status was monitored in a time course by Northern analysis of *mirB* and *cycA* expression (Fig. 6D), which are repressed and induced by iron, respectively.^{13,29} Interestingly, *mirB* transcription was induced significantly later and, consistently, *cycA* expression was repressed later in FC- compared to TAFC-precultured mycelia. These data suggest that FC is a better iron source compared to TAFC due to either better uptake or better intracellular iron supply. Nevertheless, the *cycA* and *mirB* expression patterns did not reveal significant differences in the two strains (Fig. 6D), which again makes an exclusive role of CccA in vacuolar iron storage for nutritional use unlikely.

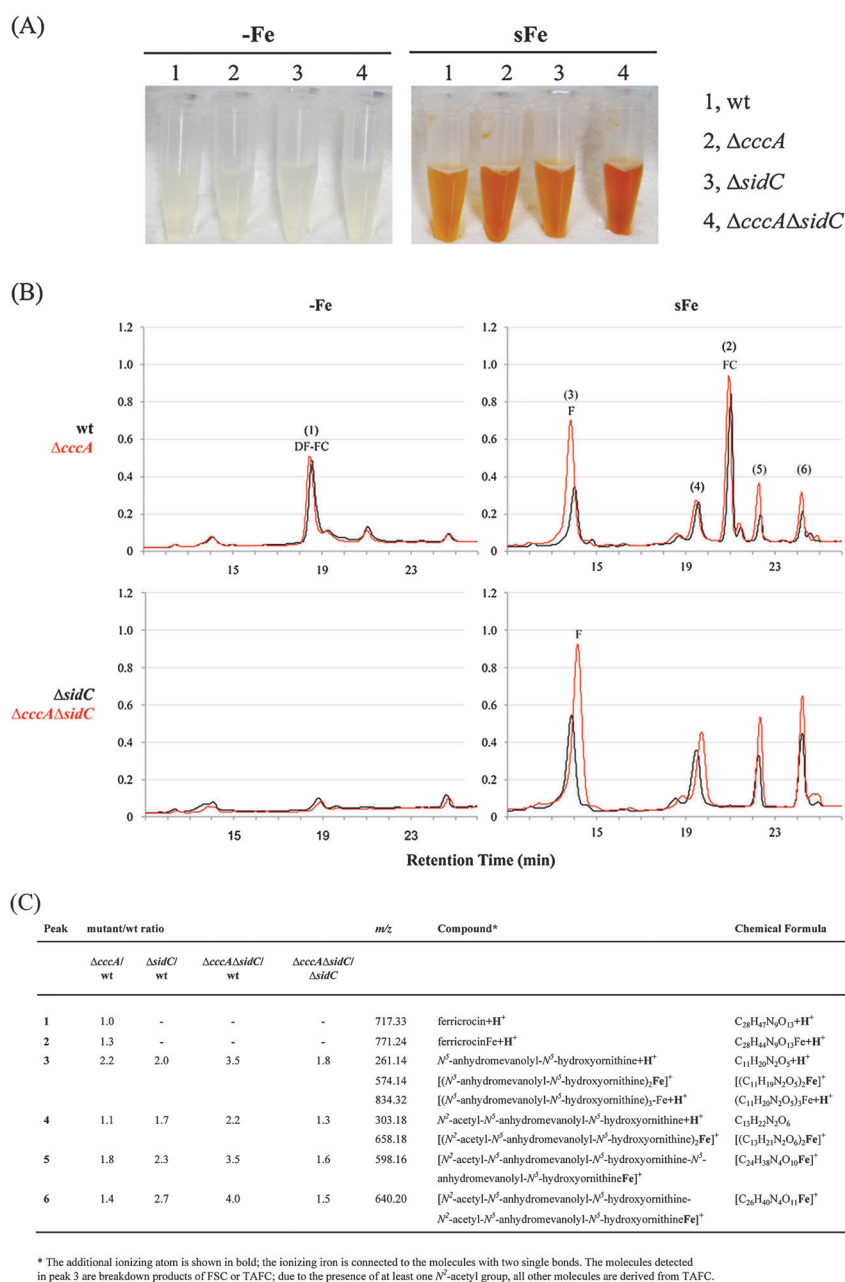


Fig. 5 Deficiency in FC, CccA, and in particular both increases the cellular content in FSC/TAFC breakdown products. *A. fumigatus* wt, $\Delta cccA$, $\Delta sidC$, and $\Delta cccA\Delta sidC$ were grown in liquid cultures for 20 h at 37 °C under iron starvation (–Fe). Subsequently, FeSO₄ was added to a final concentration of 0.1 mM and cultures were incubated for another 4 h (sFe). (A) Pictures of cell extracts (50 mg dry weight in 1 ml) before and after the iron shift. The orange coloring indicates the presence of siderophores and/or their degradation products.⁷ (B) Representative reversed-phase HPLC analysis (absorption at 210 nm in milliabsorption units) of the intracellular siderophore content comparing $\Delta cccA$ (in red) and wt (in black) as well as $\Delta cccA\Delta sidC$ (in red) and $\Delta sidC$ (in black) during –Fe and sFe, respectively. (C) The detected compounds were analyzed by mass spectrometry and the abundance of the sFe compounds was normalized to that of the wt. The data represent the mean from three experiments; the SD did not exceed 15%.

Taken together, these data indicate that vacuolar iron-import by CccA does not seem to play an exclusive role for nutritional reuse, which contrasts the situation in *S. cerevisiae*.¹⁵ Possibly *A. fumigatus* lacks recycling of vacuolar iron in general. *S. cerevisiae* extracts vacuolar iron via vacuolar membrane-localized paralogs of proteins involved in reductive iron assimilation, termed Fet5 and Fth1.¹⁷ In agreement with a possible lack of recycling of vacuolar iron, *A. fumigatus* does

not possess paralogs to the proteins involved in reductive iron assimilation.⁶ However, iron might be exported from the vacuole in *A. fumigatus* by other mechanisms and the data could be explained by the presence of CccA-independent vacuolar iron importers. One potential alternative vacuolar iron importer could be the CccB. However, expression of the encoding gene could not be detected, even in strains lacking CccA (data not shown).

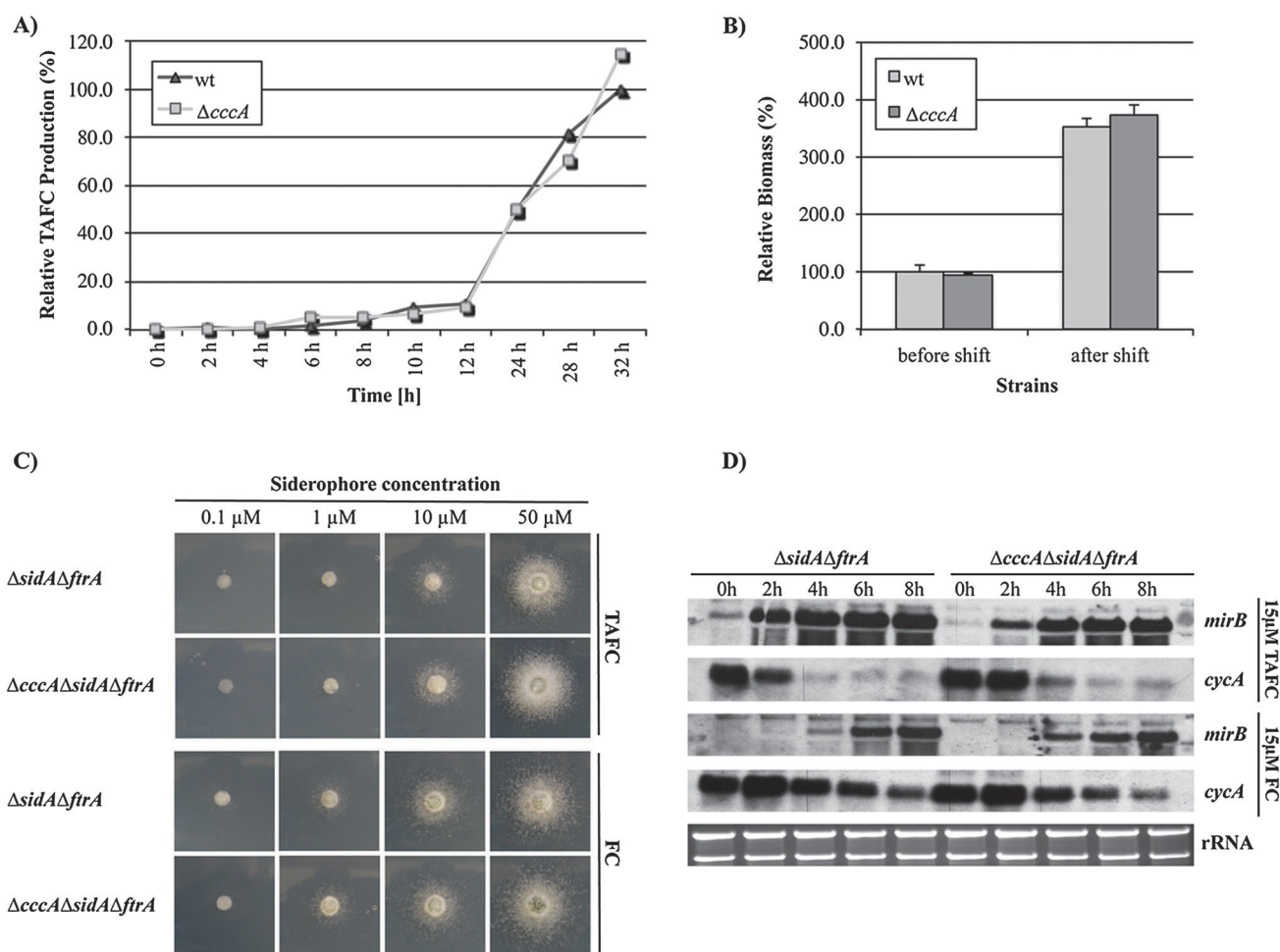


Fig. 6 CccA-deficiency does not affect the potential reuse of vacuolar iron. CccA-deficiency does not influence production of TAFC (A) and biomass (B) of 19 h-liquid iron-replete cultures (AMM + 30 μ M TAFC as an iron source) transferred to iron-free AMM. TAFC and biomass production is displayed normalized to wt. (C) CccA-deficiency does not influence radial growth after a shift from iron sufficiency to iron starvation in a Δ *sidA* Δ *ftrA* background. 10^4 conidia were point-inoculated on plates containing the indicated concentration of TAFC or FC for 18 h separated from the growth media by a dialysis membrane. Subsequently, the mycelia-containing dialysis membranes were transferred onto plates lacking iron and incubated for 5 days. (D) During a shift from iron sufficiency to iron starvation, CccA-deficiency does not influence the expression pattern of *mirB* and *cycA* in a Δ *sidA* Δ *ftrA* background, the transcription of which is activated and repressed by iron starvation, respectively. For Northern analysis, RNA was isolated from mycelia of 20 h-liquid cultures containing 15 μ M TAFC or FC before (0 h) or after transfer (2–8 h) into iron-lacking media. Ethidium bromide-stained rRNA is shown as a control for loading and quality of RNA.

Materials and methods

Growth conditions and siderophore analysis

Generally, *A. fumigatus* strains were grown at 37 °C in *Aspergillus* minimal medium (AMM) according to Pontecorvo *et al.*³⁰ containing 1% (wt/vol) glucose as a carbon source, 20 mM glutamine as a nitrogen source, and 30 μ M FeSO₄. For iron-depleted conditions, iron was omitted. Production of conidia was performed on AMM plates containing 30 μ M FeSO₄; for strains lacking high affinity iron uptake systems (Δ *sidA* Δ *ftrA*, Δ *cccA* Δ *sidA* Δ *ftrA*) 4 mM FeSO₄ was supplemented. The *xylP*^P promoter was induced by adding 0.5% xylose to AMM. Siderophore production was analyzed by reversed phase HPLC as described previously.³¹

Mass-spectrometric analysis

Determination of the molecular mass of the samples obtained by RP-HPLC was carried out using a LTQ Velos ion trap

mass spectrometer (Thermo Fisher Scientific) equipped with an electrospray source (ESI-MS, electrospray ionization mass spectrometry). Samples were dissolved in 50% aqueous methanol containing 0.1% formic acid, and infused directly into the ion source using the syringe pump. The electrospray voltage was set at 4.0 kV and the heated capillary was held at 270 °C.

cccA-disruption (Δ *cccA*), *cccA*-overexpression (*cccA*^{oe}) and *gfp-cccA* fusion (*cccA*^{gfp})

All strains used in this study are listed and described in Table 1. For generating the *cccA*-deletion strain, the bipartite marker technique was used.³² Therefore, *A. fumigatus* was co-transformed with two DNA fragments, each containing overlapping but incomplete fragments of the pyrithiamine resistance-conferring *ptrA* gene fused to 0.9-kb 5'- and 1.2-kb c-flanking regions of *cccA*, respectively. The *cccA* 5'-flanking region was PCR-amplified from genomic DNA using primers oAfcccA-1 and oAfcccA-4r.

Table 1 Fungal strains used in this study

Strains	Genotype	Ref.
wt ATCC46645		American type culture collection
wt CEA10		37
$\Delta cccA$	ATCC46645; $\Delta cccA::ptrA$	This study
$\Delta sreA$	ATCC46645; $\Delta sreA::hph$	13
$\Delta sidA$	ATCC46645; $\Delta sidA::hph$	3
$\Delta sidC$	ATCC46645; $\Delta sidC::hph$	4
$\Delta cccA\Delta sreA$	ATCC46645; $\Delta cccA::ptrA$; $\Delta sreA::hph$	This study
$\Delta cccA\Delta sidC$	ATCC46645; $\Delta cccA::ptrA$; $\Delta sidC::hph$	This study
$\Delta sidA\Delta ptrA$	CEA17; $\Delta sidA::pyrG$; $\Delta ptrA::hph$	3
$\Delta cccA\Delta sidA\Delta ptrA$	CEA17; $\Delta cccA::ptrA$; $\Delta sidA::pyrG$; $\Delta ptrA::hph$	This study
$\Delta sidA\Delta sreA$	CEA17; $\Delta sidA::pyrG$; $\Delta sreA::hph$	This study
$\Delta cccA\Delta sidA\Delta sreA$	CEA17; $\Delta cccA::ptrA$; $\Delta sidA::pyrG$; $\Delta sreA::hph$	This study
$cccA^{oe}$	ATCC46645; $pxylP^{ccA}$; $ptrA$	This study
$\Delta cccAcccA^{oe}$	ATCC46645; $\Delta cccA::ptrA$; $pxylP^{ccA}$; hph	This study
$\Delta cccAcccA^{gfp}$	ATCC46645; $\Delta cccA::ptrA$; $pgfpcccA$; hph	This study

For amplification of the 3'-flanking region primers oAfcccA-3 and oAfcccA-2r were employed. Subsequent to gel-purification, these fragments were digested with *Stu*I (5'-flanking region) and *Hind*III (3'-flanking region), respectively. The *ptrA* selection marker was released from the plasmid pSK275 by digestion with *Sma*I and *Hind*III and ligated with the 5'- and 3'-flanking region, respectively. The transformation construct A (2266 bp, fusion of the *cccA* 5'-flanking region and the *ptrA* split marker) was amplified from the ligation product using primers oAfcccA-5 and optrA-2r. For amplification of the transformation construct B (2267 bp, fusion of the *cccA* 3'-flanking region and the supplementary *ptrA* split marker) primers occcA-6r and optrA-1 were employed. For transformation of *A. fumigatus* strains, both constructs A and B were used simultaneously. In the $\Delta cccA$ mutant, the region -150 to 1155 bp relative to the *cccA* sequence was replaced by the *ptrA* resistance cassette.

For the reconstitution of $\Delta cccA$ and overexpression of *cccA* in wt and $\Delta cccA$, strains were transformed with the *cccA* coding sequence plus 110 bp of its 3' untranslated region under the control of the xylose-inducible promoter of the *P. chrysogenum* endoxylanase *xyIP* gene.²² Therefore, a 1.6 kb fragment of the *xyIP* promoter (*xyIP*^P) was amplified using pXyluidA-P as a template with synthetic oligos XylP-NotI and XylP-NcoI, thus introducing a *Nco*I site into the *xyIP* start codon and a *Not*I site at 1650 bp.²² The fragment was then digested and ligated to the *Nco*I/*Not*I-cleaved pGEM5zf(+) vector (Promega) resulting in *pxylIP*^P. Next, a 2.2 kb fragment of *cccA* was amplified using XylPcccA1 and oAfcccA-6r, introducing a *Pci*I site into the *cccA* start codon. The fragment was then cleaved and ligated to *Nco*I-cleaved *pxylIP*^P, giving rise to *pxylIP*^P*cccA*.

In order to determine the subcellular localization of CccA, the plasmid pgfpcccA was generated. Therefore the *cccA* gene was amplified using the oligonucleotides oAfcccA-BglIII.f and oAfcccA-NotI.r. After digestion, the fragment was subcloned into the *Bgl*III/*Not*I digested plasmid pCAME-703-*AohapX*-full replacing *AohapX*.²⁷

For selection, *A. fumigatus* strains harbouring *pxylIP*^P*cccA* and pgfpcccA *A. fumigatus* wt were cotransformed with the plasmid pSK275 carrying the pyrithiamine resistance gene and mutant strains with pAN7-1 carrying the hygromycin resistance gene, respectively. Generally, 0.1 μ g ml⁻¹ pyrithiamine (Sigma) or 0.2 mg ml⁻¹ hygromycin B (Sigma) was used for

selection. Screening of transformants was performed by Southern blot analysis (Fig. S1, ESI†). The hybridization probes for Southern blot analysis of $\Delta cccA$, $\Delta cccAcccA^{gfp}$ and $\Delta cccAcccA^{oe}$ strains were generated by PCR using the primers listed in Table S1 (ESI†). For extraction of genomic DNA, mycelia were homogenized and DNA was isolated according to Sambrook *et al.*³³ For general DNA propagations *Escherichia coli* DH5 α strain was used as the host.

Northern analysis

RNA isolation was performed using TRI Reagent[®] (Sigma). For Northern analysis, 10 μ g of total RNA was electrophoresed on 1.2% agarose-2.2 M formaldehyde gels and blotted onto Hybond N membranes (Amersham). The hybridization probes used in this study were generated by PCR. Hybridization probes and primers used are listed in Table S1 (ESI†).

Fluorescence microscopy

Confocal microscopy images were taken on a confocal laser scanning microscope (Axiovert 100 LSM510 [Carl Zeiss] or SP5 [Leica]) equipped with a 63 \times oil immersion objective (numerical aperture, 1.40). Images were processed using LSM Image Browser (version 4.2) software, ImageJ (version 1.44o), and Adobe Photoshop CS3.

Phylogenetic analysis

A BLASTP search was carried out with the PSI-BLAST algorithm with a NCBI protein database containing 76 different fungal species to determine CccA homologs in different fungi.³⁴ A multiple sequence alignment was done with ClustalW³⁵ using selected sequences with *E* values higher than 2×10^{-4} in the PSI-BLAST search. Based on this alignment, a rooted tree was generated (Fig. 1). The alignment was generated using ClustalW and BOXSHADE 3.31.³⁶

Abbreviations

BPS	Bathophenanthroline disulfonic acid
GFP	Enhanced green fluorescence protein
FC	Ferricrocin
FSC	Fusarinine C
TAFC	Triacetylfulvarinine C
wt	Wild type

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