Tolerance to alkaline ambient pH in *Aspergillus nidulans* depends on the activity of ENA proteins

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Tolerance of microorganisms to abiotic stress is enabled by regulatory mechanisms that coordinate the expression and activity of resistance genes. Alkalinity and high salt concentrations are major environmental physicochemical stresses. Here, we analyzed the roles of sodium-extrusion family (ENA) transporters EnaA, EnaB and EnaC in the response to these stress conditions in the filamentous fungus *Aspergillus nidulans*. While enac has a minor role, enaB is a key element for tolerance to Na⁺ and Li⁺ toxicity. Adaptation to alkaline pH requires the concerted action of enaB with EnaA. Accordingly, expression of enaA and enaB was induced by Na⁺, Li⁺ and pH 8. These expression patterns are altered in a sltAΔ background and completely inhibited in a mutant expressing non-functional PacC protein (*palH72*). However, a constitutively active PacC form was not sufficient to restore maximum enaA expression. In agreement with their predicted role as membrane ATPases, EnaA localized to the plasma membrane while EnaB accumulated at structures resembling the endoplasmic reticulum. Overall, results suggest different PacC- and SltA-dependent roles for EnaB in pH and salt homeostasis, acting in coordination with EnaA at pH 8 but independently under salt stress.

Ion homeostasis is an essential biochemical process for cell life influencing a wide range of cellular functions from osmotic regulation to enzyme activity. In addition, certain ions, such as sodium, calcium, and lithium become toxic at a high cytoplasmic concentration. Therefore, microorganisms have developed specific transport mechanisms to maintain an adequate intra and extracellular distribution of certain ions. Examples are the maintenance of low cytoplasmic Na⁺/K⁺ ratio and the extremely low cytoplasmic concentrations of Li⁺ (< 10⁻⁷ M) and Ca²⁺ (approx. 10⁻⁷ M)²–⁴. Modulation of cytoplasmic cation contents is mainly achieved by active transmembrane transporters that facilitate efflux and influx of ions through the plasma membrane or by an efficient storage of ions in intracellular compartments such as vacuoles. In addition, the activity of plasma-membrane transporters generates an electrochemical gradient across the membrane which stores energy that can be used by other transport mechanisms acting as antiporters and/or symporters. In this respect, the transmembrane electrochemical proton gradient has a major influence on the transmembrane potential and on the functionality of ion transport across the plasma membrane in both directions. But, most importantly, proton concentration determines the pH value and is the basis of a major abiotic stress/stimulus for the cell.

Filamentous fungi are able to grow over a wide pH range with almost constant intracellular pH and transmembrane potential of approximately – 200 mV⁶. A change in the extracellular pH, either towards acidification or to alkalization, is an important stimulus to a cell or organism. This might cause alterations in the electrochemical gradient and its maintenance, with direct impact on the bidirectional transport of solutes and other compounds. Changes in extracellular pH also influence the activity of secreted enzymes and metabolites, most of which have an optimal pH for maximal activity. Major adaptation processes occur, mainly altering the expression pattern of genes that are important for ion homeostasis and whose products are pH regulated⁷.

Regarding transport mechanisms, intracellular levels of Na⁺/K⁺ are maintained at neutral and acidic pH by Na⁺/H⁺ and K⁺/H⁺ antiporters. In *S. cerevisiae* intracellular low sodium maintenance relies on the Nha1p Na⁺/H⁺ antiporter which is able to extrude Na⁺, K⁺, and Li⁺. These antiporters depend on the transmembrane ΔpH, being less functional when the external pH is higher than the cytoplasmic pH, resulting in a decreased pH gradient. The intracellular pH value of *Aspergillus spp.* is approximately 7.6⁶, and when the external pH turns alkaline,
translators are required that can act independently of an external proton gradient. In most cases, P-type Ena ATPases mediate Na⁺/K⁺ transport under alkaline conditions\(^5\).

ENA ATPases have been identified in bryophytes, protozoa and fungi\(^12\text{--}^{16}\). Furthermore, it has been shown that fungal Ena-like ATPases display also transport activities for Li⁺ and Rb⁺ in addition to Na⁺ and K⁺\(^5\). Most extensive studies are available for *Saccharomyces cerevisiae* Ena ATPases. However, this Saccharomyceete is an acidophilic organism and first results from human pathogenic *Candida* species and the plant pathogen *Ustilago maydis* point to major differences in the activity and regulation of these transporters\(^7\). It is worth noting the fact that pathogenic microorganisms might have adapted to the high intracellular potassium ion concentrations of their hosts, being a cause of evolutionary specialization. In the case of *Aspergillus nidulans*, EnaA (AN6642) was first reported by Han and colleagues\(^18\) in connection with the high osmolarity glycerol pathway (HOG). These authors observed a transcriptional up-regulation pattern of a plasma membrane ATPase similar to the members of the HOG pathway, dependent on the presence of high NaCl concentration. However, induction of *enaA* was not observed when heat or oxidative stress was applied\(^19\) or to be subjected to the control of the BZIP transcription factor AtfA, a key element in the coordination of environmental stress response including oxidative stress\(^19\).

Three major regulatory pathways mediate tolerance of *A. nidulans* to alkalinity. Ambient pH response is mediated by the well-known Pal/Pac pathway\(^20\). The signalling pathway comprises six so called Pal proteins (PalA, PalB, PalC, PalF, PalH, and PalI) and transduces the alkaline-pH signal into the cell to its final acceptor, the zinc-finger transcription factor PacC. At neutral-to-alkaline ambient pH conditions PacC is activated through two proteolytic steps, being the first modification dependent on the Pal pathway. The final 27 kDa processed form of PacC acts as an activator of alkaline-expressed genes and as a repressor of acid-expressed genes\(^20\). The role of CrzA and SLT transcription factors in cation homeostasis and alkaline pH regulation has been determined\(^16,22,23\). In *S. cerevisiae*, expression of *ena* genes requires the integrity of the PacC/Pal homologous RIM pathway\(^5\). ScCrz1p, the CrzA orthologue, is one major regulator of ScEna1p\(^5,25\). To date, in *A. nidulans*, regulation of *ena* genes has not been addressed in detail. Expression of *enaA* is not regulated by CrzA and SLT was shown to play a minor effect on *enaA* transcription\(^5\).

In this study, we focus on the functional analysis of the three ScEna1p orthologues of *Aspergillus nidulans*, EnaA, EnaB and EnaC\(^16,18\). Tolerance to extracellular excess of sodium and lithium at low or high pH has been evaluated in strains carrying all possible combinations of null *ENA* alleles. Transcriptional regulation of *ena* genes and the subcellular localization of EnaA and EnaB has been explored in growth conditions combining excess of Na⁺/Li⁺ and alkalinity, as well as the involvement of ambient pH regulatory pathway mediated by PacC and the cation stress responsive pathway mediated by SltA as possible regulatory mechanisms. Results suggest a key role of EnaB in cation homeostasis and a concerted action with EnaA in the response to alkaline pH.

### Results

**Genes coding for ENA-like proteins in *A. nidulans*.** As previously noted by Spielvogel and collaborators\(^16\), the genome of *A. nidulans* contains three genes coding for putative ENA proteins which were designated *enaA* (AN6642), *enaB* (AN1628) and *enaC* (AN10982). A multiple alignment of An-ENA sequences including ScEna1p is shown in Supplementary Fig. 1. EnaA, EnaB and EnaC show high conservation with ScEna1p (YDR040C) sequence (65%, 67% and 59%, respectively). Nevertheless, the three An-ENA proteins clustered in one of the eight clades that allocate the 22 ATPases identified in *A. nidulans* (Supplementary Fig. 2A)\(^5\). Conserved domains (Pfam domains) are found among ENA homologues denoting their roles as P-type ATPases, such as (I) cation-ATPase-N (pfam00690), (II) E1–E2 ATPase (pfam00122), (III) haloacid dehalogenase-like hydrolase (HAD) (pfam00509 and (IV) cation-ATPase (pfam00689) (Supplementary Fig. 2B). A more detailed analysis of *A. nidulans* ENA protein sequences compared to ScEna1p showed the presence of several small motifs characteristic of this family of transporters, such as the Actuator domain (TGES\(^{16}\)), the Mg\(^{2+}\) binding motif (DGVN\(^{71}\)) and sequences belonging to the catalytic site of P-type ATPases: the signature sequences DKTG\(^{70}\), TGD\(^{70}\) and DPPR\(^{62}\) and residues of the nucleotide binding domain (F\(^{337}\), K\(^{442}\), K\(^{561}\)) (indicated coordinates as in ScEna1p, Supplementary Fig. 1)\(^5,22\). Distribution of these functional motifs was also conserved among the An-ENA ATPases (Supplementary Fig. 1). As for ScEna1p P-type ATPases, An-ENA transporters were predicted to harbour 10 putative transmembrane spanning regions (TMHMM and HMHMM predictions) with N- and C-terminal regions being directed to the cytoplasmic side if located at the plasma membrane (Fig. 1A).
Additional in silico searches showed the presence of at least one and up to four ENA-like ATPase coding genes in Saccharomyces and filamentous fungal genomes (Fig. 1B). The phylogram, based on the amino acid sequence of putative and described \( Na^+ \), \( K^+ \), \( H^+ \) ATPases, revealed that EnaA, EnaB, and EnaC were clustered in two different clades. While the EnaA and EnaB groups clustered, EnaC was included into a separate group of ATPases. EnaC shared similarities with HwEna1 and HwEna2 of Hortaea werneckii, which have been described to constitute a new family of ENA-like ATPases, although their functions are not completely determined yet\(^2\)

Of interest is the fact that Ena-P-type ATPases from Saccharomyces constitute a monophyletic clade in this phylogenetic analysis.

In summary, fungal genomes encode several ENA-like ATPases (2–4) showing significant conservation in their sequences and composition of functional domains. However, specific transport activities as well as regulatory mechanisms or alternative functions must be determined experimentally and not only inferred from in silico predictions.

**Functional analyses of AnENA transporters.** To determine a functional role for An-ENA orthologues, single, double and triple knock-out of \( enaA \), enaB, and \( enaC \) were constructed by gene replacement and subsequent meiotic recombination to obtain all possible mutant combinations (see list of strains in Supplementary Table 1). Single, double and triple null \( ena \) mutant colonies grew on solid standard *Aspergillus* minimal medium (AMM) showing no appreciable alterations on radial growth or conidiation (Fig. 2A, row A). Subsequent meiotic recombination to obtain all possible mutant combinations (see list of strains in Supplementary Table 1) showed no alterations on growth traits (Fig. 2A, rows B and C, respectively). When each single knocks-out mutants were grown in liquid medium (AMM) showing no appreciable alterations on radial growth or conidiation (Fig. 2A, see AMM, row A).

From Northern analyses\(^1\)\(^6\)\(^8\), and RNA sequencing\(^10\), we confirmed that the expression of both \( enaA \) and \( enaB \) was tested (Fig. 2A, rows B and C, respectively). When each the three single knock-out mutants were grown in liquid medium (AMM) showing no appreciable alterations on radial growth or conidiation (Fig. 2A, see AMM, row A). Subsequent meiotic recombination to obtain all possible mutant combinations (see list of strains in Supplementary Table 1) showed no alterations on growth traits (Fig. 2A, rows B and C, respectively). When each single knocks-out mutants were grown in liquid medium (AMM) showing no appreciable alterations on radial growth or conidiation (Fig. 2A, see AMM, row A).

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Northern blot in Fig. 3A, and its quantification in 3B, shows that \( enaB \) transcript was also detected under non alkaline growth conditions (Fig. 2A, rows E and F). Besides the expected growth inhibition of strains lacking both EnaA and EnaB activities, a negative effect of combining alkalinity and high \( Na^+ \) concentration was not observed in the other null \( ena \) mutant strains. When alkaline pH and elevated \( Li^+ \) concentration were combined, growth of the wild-type strain was largely impaired. In this growth condition, the null \( enaC \) strain showed a wild-type phenotype and growth of the wild-type strain was strongly inhibited. There was a positive effect was observed after addition of large amounts of \( Na^+ \) or \( Li^+ \), and the mixture of high levels of either \( Na^+ \) or \( Li^+ \) with alkalinity. In agreement with previous results,雙\( enaA \) and \( enaB \), and the triple mutant. In contrast, the double null \( enaA \) and \( enaB \) mutant showed a level of tolerance to sodium comparable to that of wild-type strain. These results identified EnaB as the principal element in tolerance to \( Na^+ \) and \( Li^+ \) stress.

Since alkalinity is a signal inducing expression of \( enaA \) gene (see references\(^16\)\(^8\)\(^,\) and below), tolerance of \( ena \) null mutant strains to alkaline pH was also analyzed (Fig. 2A, row D). Single null \( ena \) strains grew as well as the wild-type strain at pH 8. However, double deletion of \( enaA \) and \( enaB \) resulted in strong sensitivity. Other double null mutant combinations displayed normal colonial growth. As expected, the triple null mutant displayed a pH 8-sensitive phenotype comparable to that displayed by the double \( enaA \) and \( enaB \) strain. These results indicate that activities of EnaA and EnaB are jointly required for tolerance to ambient alkalinity.

Expression of null \( enaB \) strain to high concentrations of \( Na^+ \) and \( Li^+ \) was also tested at alkaline pH (Fig. 2A, rows E and F). Besides the expected growth inhibition of strains lacking both EnaA and EnaB activities, a negative effect of combining alkalinity and high \( Na^+ \) concentration was not observed in the other null \( ena \) mutant strains. When alkaline pH and elevated \( Li^+ \) concentration were combined, growth of the wild-type strain was largely impaired. In this growth condition, the null \( enaC \) strain showed a wild-type phenotype and growth of the wild-type strain was strongly inhibited. There was a positive effect was observed after addition of large amounts of \( Na^+ \) or \( Li^+ \), and the mixture of high levels of either \( Na^+ \) or \( Li^+ \) with alkalinity. In agreement with previous results, double \( enaA \) and \( enaB \), and the triple mutant. In contrast, the double null \( enaA \) and \( enaB \) mutant showed a level of tolerance to sodium comparable to that of wild-type strain. These results identified EnaB as the principal element in tolerance to \( Na^+ \) and \( Li^+ \) stress.

Since alkalinity is a signal inducing expression of \( enaA \) gene (see references\(^16\)\(^8\)\(^,\) and below), tolerance of \( ena \) null mutant strains to alkaline pH was also analyzed (Fig. 2A, row D). Single null \( enaB \) strains grew as well as the wild-type strain at pH 8. However, double deletion of \( enaA \) and \( enaB \) resulted in strong sensitivity. Other double null mutant combinations displayed normal colonial growth. As expected, the triple null mutant displayed a pH 8-sensitive phenotype comparable to that displayed by the double \( enaA \) and \( enaB \) strain. These results indicate that activities of EnaA and EnaB are jointly required for tolerance to ambient alkalinity.

Transcriptional regulation of \( ena \) genes. Series of Northern blot analyses were conducted to analyze the regulation of expression of \( ena \) genes in total RNAs isolated from mycelia grown at alkaline pH, in excess of \( Na^+ \) or \( Li^+ \), and the mixture of high levels of either \( Na^+ \) or \( Li^+ \) with alkalinity. In agreement with previous results from Northern analyses\(^16\)\(^8\)\(^,\) and RNA sequencing\(^10\), we confirmed that the expression of both \( enaA \) and \( enaB \) is regulated by these stresses. Under non stressing conditions (C, control condition), expression of \( enaA \) was detected and a positive effect was observed after addition of large amounts of \( Na^+ \) or \( Li^+ \), or by alkalinizing the cultures (Fig. 3A). Addition of 1.0 M \( Na^+ \) or 0.3 M \( Li^+ \) resulted in a moderate increase of \( enaA \) mRNA levels, sixfold and fivefold change, respectively, but a stronger effect, 19-fold change, was observed in response to alkaline pH, with a noticeable peak of expression at 30 min (Fig. 3B). Yet, when elevated \( Na^+ \) concentrations were combined with pH 8, transcript levels continued increasing throughout the experiment (60 min) showing a 27-fold change with respect to initial values. However, this synergic response was not observed when combining alkalinity and 0.3 M \( Li^+ \). Despite the fact that reads mapping at the \( enaA \) locus in our RNA-seq experiments\(^15\) suggest the existence of alternative splicing events at the three introns predicted in the coding region (i.e. the processed form seems to be more abundant in the wild-type at pH 8 than under standard culture conditions), Northern hybridization did not indicate the generation of transcripts of different sizes (Fig. 3A).

Northern blot in Fig. 3A, and its quantification in 3B, shows that \( enaB \) transcript was also detected under non stressing conditions. Variation of transcript abundance of \( enaB \) and \( enaA \) genes seemed to follow a comparable...
Figure 2. Functional analysis of null ena mutants. (A) Spores of single-, double- and triple-null strains were point-inoculated and images of colonies were taken after 48 h of incubation at 37 °C in solid AMM (row 1), AMM supplemented with 1 M NaCl (Na+; row 2) or 0.3 M LiCl (Li+; row 3), AMM adjusted to pH 8 with 50 mM Tris–HCl (alkaline pH; row 4), and AMM that combined the Tris–HCl buffer with either 1 M NaCl (pH 8 + Na+; row 5) or 0.3 M LiCl (pH 8 + Li+; row 6). (B) Biomass production was determined by measuring the dry weight of cells grown in liquid AMM for 24 h at 37 °C with and without elevated concentration of cations (1.0 M NaCl and 0.3 M LiCl) or alkaline pH. Data is presented as percentages and normalized by designating the growth of each strain in standard AMM (no stress agents) as 100%. Graphs show the means of three replicates per strain and condition, and error bars indicate standard deviation. Asterisks indicate a significant variation in biomass production (p < 0.05) as compared to the reference wild-type strain. N = 3.
trend, but with lower intensity in the case of \(enaB\), probably due to higher levels of expression under non-stressing conditions, compared to \(enaA\). Even so, differences were observed principally regarding \(Na^+\) stress, which induced a continuous increase in \(enaB\) expression through 1 h of treatment. As previously noted for \(enaA\), combination of high \(Na^+\) and alkaline pH resulted in a higher expression than singly implemented stress conditions. It is worth mentioning that \(enaB\) mRNA levels remained almost unchanged after addition of 0.3 M Li\(^+\), which contrasts with the severity of the growth defect shown by the null \(enaB\) strain under these stress conditions but on solid medium.

In the case of \(enaC\), Northern blot analyses detected two possible transcripts for this gene (indicated with arrows, right side Fig. 3A), being the high mobility band slightly induced by the presence of alkaline pH. The size of the two hybridization signals did not coincide with those of \(enaA\) or \(enaB\) transcripts, and thus, a cross-hybridization phenomenon was excluded. The presence of two transcripts may be a consequence of possible alternative splicing events at introns 1 and 3 of the coding region, as shown by our RNA-seq data. Furthermore, the last intron of \(enaC\) is not processed, adding 13 codons to the sequence but without altering the reading frame. The corresponding set of 13 amino acids is part of the fifth transmembrane domain. The corrected \(enaC\) sequence, including the additional 13 amino acids, was used in the analysis shown in Fig. 1 and Supplementary Fig. 1 (see also below in Fig. 4A). However, these 39 nucleotides do not explain the difference in size observed between both \(enaC\) transcripts. Considering that multiple RNA-seq reads map to the intergenic region for \(enaC\) and \(An7665\) (not shown), and that this region would include an additional target site for PacC and two sites for
SltA, it is tempting to suggest that the presence of two transcription start sites could account for the existence of two *enaC* transcripts.

MsnA is a transcription factor involved in saline stress response\(^1\). Figure 3A shows expression levels of *msnA* following a similar pattern to that described before for *enaB*. High Na\(^+\) elevated the expression of *msnA* (Fig. 3A,B). In contrast, high Li\(^+\) had no effect or even we detected a negative effect of this cation by preventing activation of *msnA* expression in the presence of alkaline pH. These results suggest a common transcriptional regulatory mechanism acting on *msnA* and *enaB* genes, and probably on *enaA*.

To further investigate a possible influence of Ena proteins in their own transcriptional regulation, expression of the remaining *ena* genes was studied in single knock-out mutants *enaA\(\Delta\) and *enaB\(\Delta\). Figure 3C shows that the temporal profile of *enaB*, *enaC* and *msnA* did not significantly change in the absence of *enaA*. Likewise, in the absence of *enaB* (Fig. 3D), no major differences were observed in *enaC* and *msnA* gene regulation, whereas the response of *enaA* to the combination of alkalinity and Li\(^+\) resulted in a stronger early (maybe compensatory) response (30 min) compared to samples from cells where *enaB* is present.

Taken together, these results indicate that separately Na\(^+\), Li\(^+\) and alkaline pH may induce *enaA* and *enaB* expression at different levels. However, a combination of high Na\(^+\) and alkalinity caused the greatest levels of expression for these genes, suggesting a major need of these ATPases under both stresses. Interestingly, the presence of lithium prevented the synergic effect of alkaline pH, indicating the presence of differential regulatory mechanisms acting under each cation stress.
Regulation of *ena* genes by PacC and SltA transcription factors. Since Na\(^+\) and Li\(^+\) cations, alkalinity and its combination differently modulate *enaA* and *enaB* expression, the role in *ena* expression of two transcription factors mediating response to ambient pH and cation stresses, PacC and SltA, was monitored\(^ {16,20}\).

To study the role of PacC on the regulation of *ena* genes, we analyzed their expression levels in two opposed ambient-pH mutant backgrounds. A mutant with a null *pahH72* mutation prevents proteolytic processing of the primary translational form of PacC, PacC72kDa, disabling functionality\(^ {30}\). On the other hand, the *pacC*14 mutation, constitutively generates the active 27kDa version of PacC, at any ambient pH\(^ {10}\). Northern analyses showed the strict requirement of proper PacC signalling for *enaA* and *enaB* expression (Fig. 4B). Transcripts of *enaA* and *enaB* were not detected in the null *pahH72* background under any condition (Fig. 4B). In contrast, detection of putative *enaC* transcripts and expression of *msnA* remained essentially as described in a wild-type PacC background (compare with Fig. 3A).

The constitutive activation of PacC did not result in a pH-independent expression of *ena* genes. Expression of *enaA* in the *pacC*14 mutant required addition of Na\(^+\) or ambient pH alkalinisation, but the expected synergic effect by combining both stresses was not observed (Fig. 4C). Excess of extracellular Li\(^+\) had a partial effect and *enaA* expression was high at 30 min but strongly inhibited at 60 min. A similar effect was observed in samples treated with Li\(^+\) and alkaline pH. In contrast, the *enaB* transcript showed almost constant levels under most conditions tested, but a strong transcriptional response after alkaline stress induction was still observed at 60 min. A negative effect of Li\(^+\) on *enaB* expression was observed at 60 min, with or without media alkalinisation, and also, the mixture of Na\(^+\) and alkaline pH did not induce the expected positive effect. Transcript levels of *enaC* and *msnA* followed similar patterns to those described in the wild-type or *pahH72* backgrounds. These results suggest the existence of two additional regulatory mechanisms with opposite effects: a positive alkaline-pH dependent effect and a negative high-Na\(^+\) modulation.

To determine the role of SltA on their regulation, the expression profiles of *ena* genes were analyzed in a *sltA* mutant strain under the above stress conditions. Figure 4D shows that the expression pattern of *enaA* in the null *sltA* strain was similar to that described in the wild-type background. Absence of *sltA* resulted in higher basal *enaB* transcript levels which were increased at alkaline pH but, as mentioned above, a negative effect was also detected when high concentrations of Na\(^+\) or Li\(^+\) were added to cultures. The expression pattern of *msnA* nearly paralleled that of *enaB*.

In agreement with possible direct roles of PacC and SltA on the regulation of *enaA* and *enaB* genes, we found putative target sites for these transcription factors in their promoters. Searches confirmed the presence of five consensus PacC (5′-GGCCAG-3′) and three SltA (5′-AGGCA-3′) sites in *enaA* promoter (1 kb upstream of ATG). Six PacC and one SltA target sequences in *enaB* promoter and two PacC and four SltA sites in *enaC* promoter (Fig. 4A). Number and locations of PacC sites in *enaA* and *enaB* promoters are in accordance with a direct role for this TF, as has been demonstrated before for other genes (see discussion). Scattering of SltA sites might be a cause for the reduced role of this TF in their regulation.

Expression profiles of EnaA and EnaB proteins under stress conditions. To further analyze EnaA and EnaB function, we constructed strains expressing C-terminally GFP-tagged versions. These strains showed a wild-type phenotype in all conditions tested (not shown), indicating that the chimaeras did not cause any dys-function to the ENA system. Both EnaA-GFP and EnaB-GFP were detected at very low levels in mycelia grown in AMM without Na\(^+\)/Li\(^+\) or alkaline pH stresses, in line with Northern blot results. Immunodetection experiments further confirmed barely detected EnaA-GFP (with expected size of 150 kDa) in protein extracts from mycelia grown for 19 h and 24 h in standard AMM (CA in Fig. 5A,B, respectively). In the case of EnaB-GFP, and EnaA-GFP were expressed early in alkaline medium subjected to alkaline pH stress (Fig. 5). These dispersed high-molecular weight species of fusion proteins might reflect post-translationally modified forms. EnaA-GFP and EnaB-GFP were expressed in AMM without Na\(^+\)/Li\(^+\) or alkaline pH stresses, in line with Northern blot results. Immunodetection experiments suggest that compared to pH 8, an increase in the concentration of both proteins cause for the reduced role of this TF in their regulation.

Subcellular localization of EnaA and EnaB. EnaA and EnaB are expected to be transmembrane proteins. In agreement with previous protein expression results, cells from *enaA*-gfp and *enaB*-gfp transformants showed no fluorescence in standard microscopy medium (WMM; not shown). For fluorescence observation, cells were exposed for 60 min to alkaline pH (by addition of 100 mM Na\(_2\)HPO\(_4\) to WMM). The images depicted in Fig. 6A show that, under alkaline conditions, EnaA-GFP displayed preferential plasma membrane localization (PM). A regular distribution of fluorescence was observed at PM of apical and distal hylphal compartments. Thus, response to alkaline pH seems to involve transport to, and regular distribution of EnaA ATPase to all parts of a hypha avoiding a preferential accumulation at the cell tip, where major exocytic and endocytic processes have been shown to occur\(^ {31–33}\). Fluorescence of EnaA was present at the apical region (t), at branching points (b) and also between compartments, where septa were formed (s), as well as in spots dispersed within the cytosol. At earlier times of induction (15–30 min exposure to pH 8, images at the bottom of panel A), EnaA-GFP fluorescence was visible along the apical compartment but location at PM was not uniform, indicating the existence of
clusters of transporters probably associated to specific regions at the PM such as lipid rafts (indicated with white arrowheads and dotted lines).

After induction of the expression of the EnaB-GFP chimera by alkaline pH, microscopy showed a preferential location of EnaB in intracellular, elongated and round structures (Fig. 6B). Elongated structures accumulated at the tip of the cell while round ones distributed at basal locations of hyphal compartment. Noticeably, EnaB-GFP localizations resemble those illuminated by Sec63-GFP, a protein associated to ER and nuclear periphery (possible nuclei, black arrowheads, Fig. 6C). Presence of EnaB at the periphery of nuclei is shown in Supplementary Fig. 3. Thus, EnaA and EnaB have distinct locations, being EnaA located at the PM and EnaB probably in ER as well as additional intracellular membranous compartments such as nuclei, Golgi and transport vesicles.

Alkaline pH sensitivity of loss of function mutants in the pH regulatory system is not a result of reduced expression of enaA gene. The strong dependency of ena expression on the Pal/PacC pathway and its preferential PM localization, led us to speculate that EnaA might be one of the major factors in determining alkaline pH tolerance in the absence of a functional Pal/PacC pathway. Thus, we analyzed whether a PacC independent expression of ena could rescue the alkaline-sensitive phenotype caused by a loss-of-function mutation in the Pal signalling pathway such as palA1.

To express enaA in a pH-independent manner, we transformed a wild-type strain with a plasmid carrying an insert in which the coding region of enaA was under the control of a truncated version of the gpdA promoter, gpdAmini, which leads to constitutive but moderate expression levels. Of selected transformants, MAD3042 carried a single copy and MAD3043 two copies of the construct integrated at the pyroA locus. Both strains displayed a wild-type phenotype for salt or alkaline pH tolerance, although, as expected, different expression levels of enaA independent of pH condition were detected (Fig. 7A). The amount of enaA mRNA increased consistently with the copy number and in the single copy gpdAmini::enaA strain enaA mRNA levels were similar to those produced by
the endogenous ena locus at alkaline pH. Subsequently, both strains were independently crossed with a palA1 mutant strain (MAD1865). Progeny carrying one or two copies of gpdAmini::enaA and the mutation palA1 were selected and analyzed (strains I25 (single copy) and I4 (double copy)). Using Northern analysis, we confirmed a constitutive expression of enaA independent of inducing conditions and of a functional Pal/PacC pathway in both strains (Fig. 7B). However, phenotypic analysis could not detect any effect of expressing enaA in a palA1 mutant background regarding salt or pH sensitivity (Fig. 7C). Indeed, addition of neomycin (2 mg/mL) to the culture medium showed hyper-tolerance of palA1 strains expressing gpdAmini-driven enaA to this compound, as has been traditionally described for this group of mutants. Therefore, PacC-independent expression of enaA does not suppress the sensitivity to alkaline pH, supporting the broad regulatory activity of a functional Pal/PacC pathway and the complexity of alkaline pH response by A. nidulans.

Discussion

In this work we investigated, through a combination of genetics and cell biology techniques, the roles of the three putative sodium ATPases of the ENA family identified in A. nidulans. Sequence similarities, domain distribution and the phylogenetic association (Fig. 1) indicate that A. nidulans Ena-like ATPases probably fulfil the same biological function described for several filamentous fungal and yeast ENA homologues. The action of Ena P-type ATPases in fungi has been commonly related to potassium, lithium and sodium ion export from the cytoplasm, especially at alkaline and high salinity conditions. However, phylogenic comparison supports the existence of genetic variability of Ena P-type ATPases, which are believed to have evolved from an ancestral K+ ATPase by gene duplication in adaptation to salt and/or alkaline conditions. Thus, it is not surprising that evolved functional variants cannot be directly inferred by sequence similarity analyses. This is exemplified by unsuccessful attempts to suppress growth defects in S. cerevisiae ENA mutants by expression of an Ena P-type ATPase from Neurospora crassa or from Hortaea werneckii. Of the three Ena proteins found in A. nidulans, EnaA and EnaB share higher sequence conservation than with EnaC, suggesting a different phylogenetic origin or divergent evolutionary paths.

In contrast to previously cited ENA functions in other organisms, phenotypic analyses of A. nidulans null enaA strains clearly established that a functional EnaA ATPase is not essential for Na+ or for Li+ detoxification,
but it is necessary at alkaline pH (Fig. 2). A different scenario was found when studying EnaB function. EnaB is necessary for growth in high salinity (Li\(^+\) and Na\(^+\)) and alkalinity environments. Strong sensitivity to alkalinity is observed in the absence of both EnaA and EnaB proteins, suggesting the existence of overlapping functions of these ATPases in adaptation to alkalinity. At alkaline conditions the importance of EnaB in Na\(^+\) and Li\(^+\) tolerance is more prominent, being indispensable in the latter. Thus, the described properties of EnaB are more similar to the function that has been observed for typical Ena-like P-type ATPases. The function of EnaC remains, as yet, undefined. In our phylogenetic analyses, EnaC clustered together with HwENA1 and HwENA2 of H. werneckii; however, a requirement of EnaC function in adaptation to alkaline condition or salinity was not observed in this work. This suggests that EnaC undergoes specific spatial and temporal regulations, is activated by other cations or may even have become a pseudogene during Ena evolution in A. nidulans.

Figure 7. Constitutive expression of enaA under the control of the gpdA\(^{\text{mini}}\) promoter. (A) Expression, in a wild-type background and under inducing (+) or non-inducing (−) conditions, of enaA driven by the native or the gpdA\(^{\text{mini}}\) promoters. Strains that integrated one or two copies of the gpdA\(^{\text{mini}}\):enaA plasmid were analyzed. Ribosomal RNA is shown as a loading control. (B) Same experiment as in (A) but carried out with samples corresponding to the palA1 mutant background. (C) Phenotypes of wild-type (left block of images) or palA1 (right) strains bearing zero (−), one (1 ×) or two (2 ×) copies of the gpdA\(^{\text{mini}}\):enaA plasmid after 48 h of culture at 37 °C on AMM (row 1), AMM with LiCl (0.3 M; row 2), KCl (1 M; row 3), NaCl (1 M; row 4) or neomycin (2 mg/mL; row 6), and AMM adjusted to pH 8 (row 5).
The above-described differential expression patterns of *A. nidulans* ENA proteins correlate with what has been observed in other fungal species. Indeed, *ena* expression patterns in response to high external ion concentrations and pH differ significantly among orthologues and paralogues of different fungal species. In salt tolerant species such as the Dothideomycete *H. werneckii* (HwENA1 and HwENA2) or the Saccharomycete *Zygosaccharomyces rouxii* (ZENA1), expression of Ena P-type ATPases is induced only at elevated sodium ion concentrations, (5 M and 3 M, respectively) (ZENA1 and ZENA2). In Debaryomyces (formerly Schwanniomyces) occidentalis, ENA1 and ENA2 transcript levels are very low at neutral pH and 0.2 M NaCl concentration. On the other hand, in *S. cerevisiae*, which is more salt sensitive, ENA1 expression is strongly induced by addition to the medium of 0.4 M Na+, 0.05–0.1 M Li+, as well as by alkaline pH. UmENA1 of *Ustilago maydis* has been characterized as a specific Na+ and K+ efflux ATPase and is strongly induced by 0.5 M NaCl, whereas UmENA2 is only marginally induced.

The present work highlighted that *A. nidulans* *ena* genes are differentially regulated by Na+, Li+ and alkaline pH. We determined that *enaA* is weakly expressed in standard culture conditions, AMM with pH close to neutral (~6.8), and is induced by 1.0 M NaCl, 0.3 M Li+ and by alkaline pH (8.0) (Fig. 3A). Furthermore, at alkaline pH, ENA expression is strongly elevated. A similar effect has also been described for UmENA1 and UmENA2 and HwENA1 and HwENA2. The highest expression levels for this group of Ena P-type ATPases are achieved when alkalinity is combined with high-Na+. However, these high gene expression levels do not correlate with an essentiality of the gene function, as observed in the phenotypic analysis of the null *enaA* strain. *enaB* followed a comparable expression profile to that of *enaA*, although in general with lower concentrations than those of *enaA* (except for basal conditions). Still, differences were observed, especially regarding to Na+-induction time, which was later in the case of *enaB*. A hypothetical coordination of an early response mediated by *enaA* and a late response mediated by *enaB* would require both ATPases to efflux Na+ at alkaline pH. It is interesting that, despite the essentiality of EnaB on Li+ tolerance at neutral and at high pH conditions (Fig. 2), Li+ did not induce such a remarkable increase in *enaB* expression as did Na+ or alkaline pH.

Our work also focused on the transcriptional regulation of Ena-like ATPases. In *S. cerevisiae*, in addition to the Pal/PacC (Rim101) pathway, the calcineurin/Crz1 and high osmolarity glycerol (HOG) osmo-responsive pathway are involved in the control of ENA1 expression together with a regulation by nutrients via Snf1 and Mig1.2 (reviewed in). Our previous work demonstrated that *enaA* expression is independent of Crz1. *AtfA* of *A. nidulans* was characterized as a candidate transcription factor acting downstream of Hog1/Sak1 MAPK and playing a role in oxidative and osmotic stress response. Expression of *enaA* has been shown to be independent of AtfA, but *enaB* and *enaC* displayed AtfA-dependent expression especially in response to oxidative stress. This observation establishes a putative link between ENA expression and the response to oxidative stress that should be assessed more deeply in the future.

Elevated transcript levels of *enaA* and *enaB* at alkaline pH, together with the presence of several putative binding sites for PacC (5'-GCCAAG-3') at their promoters, supported a role of the Pal/PacC regulatory pathway and we showed that PacC is an essential positive regulator of *enaA* and *enaB* expression. In the absence of the Pal signalling pathway (palH72) that finally activates the PacC transcription factor, neither *enaA* nor *enaB* transcripts were detected. In the constitutive PacC background (pacC), different transcript levels were obtained to those observed under inducing conditions in the wild type background. This result contrasts with previous reports on the mechanisms PacC-dependent gene regulation (e.g. *ipnA*). The absence of a full upregulated expression of *enaA* and *enaB* in the pacC14 background reveals the need of at least one additional regulatory pathway to influence *enaA* and *enaB* expression in response to salinity. A minor positive regulatory role for the salt activated transcription factor SlmA on expression of *ena* genes has been shown in this and previous works. This in agreement with our RNA-seq results, in which both *enaA* and *enaB* (but not *enaC*) were among the most intensely upregulated genes at pH 8 in a wild-type background (Top45 and Top2 genes with the highest log2FC values, respectively), but none of them was significantly downregulated in the slmA mutant under these stress conditions.

One possible factor causing the different cation and pH sensitivity of *enaA* and *enaB* null strains could be the different subcellular localization of the proteins. *ENA* as *S. cerevisiae* Enap1 localizes in the plasma membrane, whereas ENA2 localizes in an intracellular membranes system that resembles the ER and Golgi network. However, since Li+ is very toxic at low concentrations, compartmentalization into it by Enap2 could greatly increase Li+ tolerance. Results described in previous reports demonstrated that *Ustilago maydis* UmEna2p and *Neurospora crassa* NcEna2 localize in the proximity of the ER and other endomembranes.

We further considered that, since a functional *enaA* or *enaB* was necessary for alkaline tolerance, constitutive expression of *enaA* could be sufficient to rescue the alkaline sensitive phenotype of a Pal/PacC loss-of-function mutation. A strain constitutively expressing *enaA* in a palA1 genetic background maintained the *palA1* phenotype, however, confirming that other Pal/PacC-controlled elements are required. This is in agreement with the envisaged role of PacC as a regulator of multiple pathways in response to environmental pH changes. In this context, next-generation sequencing and analysis tools should help unveil further details of this complex process.

**Methods**

**Fungal strains and growth conditions.** *Aspergillus* minimal media (AMM) adjusted to pH 6.8, containing 2% glucose and 71 mM sodium nitrate as main carbon and nitrogen sources, respectively, was prepared as described previously. *A. nidulans* strains used in this study are listed in Supplementary Table 1. Strain MAD1427 was used for the systematic deletions. Meiotic crosses and analysis of the progeny followed standard procedures as described previously. Strain BD575 was obtained from the progeny of a cross between MAD2446 and BD488, and strains BD604 and BD612 were obtained from the crossing between BD486 and BD575. Progeny was verified by PCR analysis using oligonucleotides listed in Supplementary Table 2.
Phenotypic analyses of mutant strains were done on solid AMM supplemented with a range of compounds to generate cation or pH stress. Sodium and lithium stress were induced by addition of 1.0 M NaCl and 0.3 M LiCl, respectively. Alkaline stress was induced by adjusting media pH to 8 with either 50 mM Tris–HCl pH 8 or 100 mM Na2HPO4, always with similar results. Specifically, the use of 100 mM Na2HPO4 on solid medium was avoided to prevent precipitation when combining with lithium chloride. Radial extension of mutant strains was always compared to that of the reference wild-type strain. Strains were incubated at 37 °C for 2 days and at least 3 replicates per strain and condition were analyzed.

For investigating the effects of alkali-cations and alkaline pH on biomass production, 1 × 10⁶ spores of the mutant strains of interest were inoculated in liquid medium (30 mL of medium in 100 mL flask) with or without addition of the stress agent (1.0 M NaCl, 0.3 M LiCl or 50 mM Tris–HCl pH 8). After cultures were incubated for 24 h at 37 °C in an orbital incubator at 200 rpm, mycelia were collected, dried at 100 °C and weighed. Three biological replicates were performed per strain and condition. For data normalization, growth of each strain at liquid AMM without addition of the stress agent was designated as 100%. Data were presented as percentages. Statistical significance of differences observed in biomass production between the wild-type and mutant strains was evaluated using Two-tailed Student’s t-test for unpaired samples.

Construction of deletion cassettes for enaA, enaB and enaC. Gene deletion was achieved by using deletion cassettes comprising *Aspergillus fumigatus* pyrG (AN6642/enaA; AN10982/enaC), riboB (AN1628/enaB), or pyroA (AN10982/enaC) genes, respectively, as selectable markers flanked by 700–1,500 bp up- and downstream of the targeted *A. nidulans* open reading frame. The selectable markers were amplified with primer pairs gsp2–gsp3* (listed in Supplementary Table 2), which contained tails (20 bp) specific to each terminal of both the 5′ and 3′ ends of the targeted gene, and using the cloned selectable marker as a template present on plasmids (see Supplementary Table 3). Fragments corresponding to the 5′-UTR and 3′-UTR of the targeted genes were amplified using genomic DNA of *A. nidulans* as template and oligonucleotide pairs gsp1–gsp2 or gsp3–gsp4, respectively. The final cassette was generated by fusing the three fragments by Fusion PCR using primers gsp1–gsp4. Cassette were purified and directly used to transform *A. nidulans* protoplasts (Supplementary Table 1) following the protocol described previously. The double-null mutants were generated by sequential transformations of strains deficient in heterologous recombination with the required fusion cassettes. Prototrophic transformants were isolated and deletion of the targeted open reading frame confirmed by Southern-blot after genomic DNA extraction.

GFP tagging of EnaA and EnaB. Construction of *A. nidulans* strains expressing C-terminally GFP-tagged proteins followed the 3-way PCR methodology described previously, ~1.5 kb from the most 3′ end of the target gene and the 3′ UTR region were amplified with primers gsp5–gsp6 and gsp3–gsp4, respectively (Supplementary Table 2), and fused in a 3-way PCR with a fragment, amplified with primers gsp6–gsp3*, containing the gfp gene together with the selectable pyrG gene of *A. fumigatus*. The fusion cassette was transformed into recipient MAD1427 strain that carries pyrG89 mutation. Uracl and uridine prototrophs were selected, purified to homokaryosis and then analyzed by Southern-blot for verification of a single integration event of the gene replacement cassette. Functional replacement was verified by phenotypic analysis using wild-type and deletion phenotypes as references.

Constitutive expression of enaA. The enaA coding sequence was amplified using primers EnaAup and EnaAdown, which introduced EcoRI restrictions sites for further cloning into plasmid p1660. This plasmid carries a truncated version of gpdA promoter, gpdAmini, allowing constitutive expression of target gene, and a truncated version of pyrA gene, restricting integration at the pyroA locus by complementation of pyroA mutation (Table Supplementary 3). Selection of positive transformants is based on recovery of pyrimidine prototrophy (see also). The resulting plasmid was named pGPDA-EnaA and the integrity of the enaA open reading frame was confirmed by sequencing. Protoplasts of strain MAD1739 were transformed using pGPDA-EnaA and positive transformants and copy number of integration events at the pyroA locus were confirmed by Southern blot analysis.

RNA extraction and gene expression analyses. For RNA extraction, 1 × 10⁶ spores of wild-type and single null mutant strains were inoculated in fermenters and cultured for 16 h at 37 °C and 200 rpm. Mycelia were harvested and transferred to flasks with fresh medium containing 1.0 M NaCl, 0.3 M LiCl or pH 8.1, with further incubation of the culture as specified in the figures. Mycelia samples were collected by filtration, frozen in liquid nitrogen and grounded.

Total RNA was isolated following manufacturer’s protocol by adding 1 mL TRIzol reagent (Fluka, Sigma-Aldrich Quimica SL) to 100 mg of grounded samples. Total RNA concentration was calculated using a Nanodrop 2000c system (Thermo Fisher Scientific, Waltham, MA). Northern blot assays were carried out as described previously. 10 µg of total RNA per sample was loaded in 1.2% agarose gels and transferred to positively charged nylon filters (Roche). Equal loading of total RNA was evaluated by methylene blue staining of rRNA. Transcripts of *A. nidulans* ena-like genes were detected using specific PCR-amplified probes as described previously, mRNA levels of msnA gene (AN1652) were detected using a probe of 1,995 bp amplified using primers described in Table S2 and which covered 100% of the ORF. Labelling of DNA probes was performed using Roche radioactive labelling kit, detected using a PhosphorImager screen (Molecular Dynamics) and developed using a FLA-5100 Reader (Fujifilm). Band intensity quantification was performed using Multi-Gauge V3.0 software (Fujifilm).
Protein isolation and western blot. Sodium, lithium and alkalinity-shock effects on protein concentration was studied using western-blot technique. Total protein extracts were isolated from mycelium of GFP-tagged strains cultivated for 16 h at 37°C in appropriately supplemented Covel's minimal medium34 plus the indicated time after addition of the stress agent. Mycelia were harvested by filtration, frozen in dry ice and lyophilized for 16 h. Protein extraction from lyophilized samples was performed by alkaline-lysis extraction procedure described previously88. Proteins were resolved in 8% SDS-polyacrylamide gels and transferred to nitrocellulose membrane using TransBlot Turbo System (Bio-Rad). GFP-tagged versions of EnaA and EnaB were detected using mouse anti-GFP monoclonal antibody (1:5,000 dilution; Roche). As loading control, gamma-actin was detected using mouse anti-actin C4 antibody (1:5,000 dilution; MP Biomedicals). As secondary antibody peroxidase-conjugated goat anti-mouse IgG immunoglobulin (1:4,000 dilution; Jackson) was used.

Western blots were developed using ECL kit (GE Heathcare) and images were taken using a Luminescent Image Analyzer LAS-3000 (Fujifilm) and processed with Multi-Gauge V3.0 software (Fujifilm).

Fluorescence microscopy. Aspergillus nidulans conidiospores were inoculated in uncoated glass-bottom µ-dishes (Ibidi GmbH) containing 2.5 mL watch minimal medium (WMM57) supplemented with 25 mM NaH₂PO₄, 5 mM ammonium (+)-tartrate, and 0.5% glucose. After 16 h at 25°C medium was replaced with freshly prepared WMM supplemented with 100 mM Na₂HPO₄. Samples were cultured for an additional 1.5 h at 37°C prior to observation of fluorescence (see text for details of enaA and enaB induction).

Different Interference Contrast (Nornarski optics) and fluorescence images were acquired from in vivo cultures with a Leica DMI-6000b inverted microscope coupled to an ORCA-ER digital camera (Hamamatsu Photonics) and equipped with a 63 Plan Apochromat 1.4 N.A. oil immersion objective (Leica) and a GFP filter (excitation 470 nm; emission 525 nm). Images were acquired using Metamorph (Molecular Dynamics) software and processed using ImageJ free-software (https://imagej.nih.gov/ij/).

Bioinformatics. Multiple sequence alignments were made using BLAST and Clustal Omega open source program (https://www.ebi.ac.uk/Tools/msa/clustalomega/). Jalview application was used for alignment visualization and domain analyses. Transmembrane domains of Ena-like proteins were predicted using Hidden Markov Models (HMM) in the Institute Pasteur Mobyle server (https://mobyle.pasteur.fr/). Phylogenetic trees were generated using Mega Version 7.0 software58. Maximum-Likelihood (80 replicates) and Neighbor-Joining (10,000 replicates) methods rendered the same organization of the clades. The Maximum-likelihood tree was edited using iTOL59.

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References
1. Findon, H. et al. Analysis of a novel calcium auxotrophy in Aspergillus nidulans. Fungal Genet. Biol. 47, 647–655 (2010).
2. Lopez, F. et al. The yeast inositol monophosphatase is a lithium- and sodium-sensitive enzyme encoded by a non-essential gene pair. Mol. Microbiol. 31, 1255–1264 (1999).
3. Serrano, R. Salt tolerance in plants and microorganisms: toxicity targets and defense responses. Int. Rev. Cytol. 165, 1–52 (1996).
4. Mulet, J. M. et al. A novel mechanism of ion homeostasis and salt tolerance in yeast: the Hal4 and Hal5 protein kinases modulate the Trk1–Trk2 potassium transporter. Mol. Cell Biol. 19, 3328–3337 (1999).
5. Ariño, I., Ramos, I. & Sychrová, H. Alkali metal cation transport and homeostasis in yeasts. Microbiol. Mol. Biol. Rev. 74, 95–120 (2010).
6. Portillo, F. Regulation of plasma membrane H(+)-ATPase in fungi and plants. Biochim. Biophys. Acta 1469, 31–42 (2000).
7. Portillo, F., Eraso, P. & Serrano, R. The plasma membrane H⁺-ATPase of fungi and plants. In Biomembranes: A Multi-Volume Treatise ATPases (ed. Lee, A. G.) 225–240 (JAI Press, London, 1996).
8. Bagar, T., Altenbach, K., Read, N. D. & Bencina, M. Live-cell imaging and measurement of intracellular pH in filamentous fungi using a genetically encoded ratiometric probe. Eukaryot. Cell 8, 703–712 (2009).
9. Miller, A. J., Vogg, G. & Sanders, D. Cytosolic calcium homeostasis in fungi: roles of plasma membrane transport and intracellular sequestration of calcium. Proc. Natl. Acad. Sci. U.S.A. 87, 9548–9552 (1990).
10. Picazo, J. et al. Defining the transcriptional responses of Aspergillus nidulans to cation/alcaline pH stress and the role of the transcription factor SlaA. Microb. Genomics https://doi.org/10.1099/mgen.0.000415 (2020).
11. Ruiz, A. & Ariño, J. Function and regulation of the Saccharomyces cerevisiae ENA sodium ATPase system. Eukaryot. Cell 6, 2175–2183 (2007).
12. Benito, B., Garcíadelblás, B. & Rodríguez-Navarro, A. Molecular cloning of the calcium and sodium ATPases in Neurospora crassa. Mol. Microbiol. 35, 1079–1088 (2000).
13. Benito, B., García-delblás, B. & Rodríguez-Navarro, A. Potassium- or sodium-efflux ATPase, a key enzyme in the evolution of fungi. Microbiology 148, 933–941 (2002).
14. Ramos, J., Ariño, J. & Sychrová, H. Alkali-metal-cation influx and efflux systems in nonconventional yeast species. FEMS Microbiol. Lett. 317, 1–8 (2011).
15. Rodríguez-Navarro, A., Benito, B. Sodium or potassium efflux ATPase a fungal, bryophyte, and protozoal ATPase. Biochim. Biophys. Acta 1798, 1841–1853 (2010).
16. Spielvogel, A. et al. Two zinc transcription factors, CrzA and SltA, are involved in cation homeostasis and detoxification in Aspergillus nidulans. Biochem. J. 414, 419–429 (2008).
17. Benito, B., García-delblás, B., Fraile-Escanciano, A. & Rodríguez-Navarro, A. Potassium and sodium uptake systems in fungi. The transporter diversity of Magnaporthe oryzae. Fungal Genet. Biol. 48, 812–822 (2011).
18. Han, K. H. & Prade, R. A. Osmotic stress-coupled maintenance of polar growth in Aspergillus nidulans. Mol. Microbiol. 43, 1065–1078 (2002).
19. Miskei, M., Karányi, Z. & Pócsi, I. Annotation of stress-response proteins in the aspergilli. Fungal Genet. Biol. 46(Suppl 1), S105–S120 (2009).
20. Peñalva, M. A., Tilburn, J., Bignell, E. & Arst, H. N. Jr. Ambient pH gene regulation in fungi: making connections. Trends Microbiol. 16, 291–300 (2008).

21. Arst, H. N. Jr. & Peñalva, M. A. Recognizing gene regulation by ambient pH. Fungal Genet. Biol. 40, 1–3 (2003).

22. Hernández-Ortiz, P. & Espeso, E. A. Phospho-regulation and nucleocytoplasmic trafficking of CrzA in response to calcium and alkaline-pH stress in Aspergillus nidulans. Mol. Microbiol. 89, 532–551 (2013).

23. Mendizabal, I., Pascual-Aluiri, A., Serrano, R. & de Larrinoa, I. F. A second component of the SltA-dependent cation tolerance pathway in Aspergillus nidulans. Fungal Genet. Biol. 82, 116–128 (2015).

24. Mendizabal, I., Pascual-Aluiri, A., Serrano, R. & de Larrinoa, I. F. Promoter sequences regulated by the calcineurin-activated transcription factor Crz1 in the yeast ENA1 gene. Mol. Genet. Genomics 265, 801–811 (2001).

25. Petrezselyova, S. et al. Regulation of the Na+/K+-ATPase Ena1 expression by calcineurin/Crz1 under high pH stress: a quantitative study. PLoS ONE 11, e0158424 (2016).

26. Harris, S. D. et al. Morphology and development in Aspergillus nidulans: a complex puzzle. Fungal Genet. Biol. 46(Suppl 1), S82–S92 (2009).

27. Palmgren, M. G. & Nissen, P. P-type ATPases. Annu. Rev. Biophys. 40, 243–266 (2011).

28. Gorjan, A. & Plemenitas, A. Identification and characterization of ENA ATPases HwENA1 and HwENA2 from the halophilic black yeast Hortaea werneckii. FEMS Microbiol. Lett. 265, 41–50 (2006).

29. Herrvías-Aguilar, A., Galindo, A. & Peñalva, M. A. Receptor-independent ambient pH signaling by ubiquitin attachment to fungal arrestin-like PaF. J. Biol. Chem. 285, 18095–18102 (2010).

30. Orejas, M. et al. Activation of the Aspergillus PaC transcription factor in response to alkaline ambient pH requires proteinolysis of the carboxy-terminal moiety. Genes Dev. 9, 1622–1632 (1995).

31. Araújo-Bazán, L., Peñalva, M. A. & Espeso, E. A. Preferential localization of the endocytic internalization machinery to hyphal tips underlies polarization of the actin cytoskeleton in Aspergillus nidulans. Mol. Microbiol. 67, 891–905 (2008).

32. Pantazopoulou, A. & Peñalva, M. A. Organization and dynamics of the Aspergillus nidulans Golgi during apical extension and mitosis. Mol. Biol. Cell 20, 4335–4347 (2009).

33. Taheri-Talesh, N. et al. The tip growth apparatus of Aspergillus nidulans. Mol. Biol. Cell 19, 1439–1449 (2008).

34. Marinka-Iñarrairaegui, A., Pantazopoulou, A., Espeso, E. A. & Peñalva, M. A. The Aspergillus nidulans peripheral ER: disorganization by ER stress and persistence during mitosis. PLoS ONE 8, e7154 (2013).

35. Punt, P. J., Kuyvenhoven, A. & van den Hondel, C. A. A mini-promoter lacZ gene fusion for the analysis of fungal transcription control sequences. Gene 158, 119–123 (1995).

36. Diez, E. et al. Activation of the Aspergillus PacC zinc finger transcription factor requires two proteolytic steps. EMBO J. 21, 1350–1359 (2002).

37. Cockx, M. X., Brownlee, A. G. & Arst, H. N. Jr. Regulation of gene expression by pH of the growth medium in Aspergillus nidulans. Mol. Gen. Genet. 203, 346–353 (1986).

38. Balázs, A. et al. AtfA bZIP-type transcription factor regulates oxidative and osmotic stress responses in Aspergillus nidulans. Mol. Genet. Genomics 283, 289–303 (2010).

39. Batuñes, M. A. et al. The Nha1 antiporter of Saccharomyces cerevisiae mediates sodium and potassium efflux. Microbiology 144(Pt 1), 2749–2758 (1998).

40. Benito, B., García-deBlas, B., Pérez-Martin, J. & Rodríguez-Navarro, A. Growth at high pH and sodium and potassium tolerance in media above the cytoplasmic pH depend on ENA ATPases in Ustilago maydis. Eukaryot. Cell 8, 821–829 (2009).

41. Watanabe, Y. et al. Characterization of the Na+–ATPase gene (ZEN1A) from the salt-tolerant yeast Zygosaccharomyces rouxii. J. Biont. Bioeng. 88, 136–142 (1999).

42. Batuñes, M. A. & Rodríguez-Navarro, A. P-type ATPases mediate sodium and potassium effluxes in Schwannomyces occidentalis. J. Biol. Chem. 273, 1640–1646 (1998).

43. Sychrová, H. Yeast as a model organism to study transport and homeostasis of alkali metal cations. Physiol. Res. 53(Suppl 1), S91–S98 (2004).

44. Mendoza-Martínez, A. E., Lara-Rojas, F., Sanchez, O. & Aguirre, J. N.A. mediates a redox regulation of the antioxidant response, carbon utilization and development in Aspergillus nidulans. Front. Microbiol. 8, 516 (2017).

45. Espeso, E. A., Tilburn, J., Arst, H. N. Jr. & Penvala, M. A. pH regulation is a major determinant in expression of a fungal penicillin biosynthetic gene. EMBO J. 12, 3947–3956 (1993).

46. Marques, M. C. et al. A functional Rim101 complex is required for proper accumulation of the ENa +/ATPase protein in response to salt stress in Saccharomyces cerevisiae. FEMS Yeast Res. 15, for017 (2015).

47. Käfer, E. Origins of translocations in Aspergillus nidulans. Genetics 152, 217–232 (1995).

48. Todd, R. B., Davis, M. A. & Hynes, M. J. Genetic manipulation of Aspergillus nidulans: meiotic progenesis for genetic analysis and strain construction. Nat. Protoc. 2, 811–821 (2007).

49. Yang, L. et al. Rapid production of gene replacement constructs and generation of a green fluorescent protein-tagged centromeric marker in Aspergillus nidulans. Eukaryot. Cell 3, 1359–1362 (2004).

50. Tilburn, J. et al. Transformation by integration in Aspergillus nidulans. Gene 26, 205–212 (1983).

51. Naya, T. et al. A versatile and efficient gene-targeting system for Aspergillus nidulans. Genetics 172, 1557–1566 (2006).

52. Sambrook, J., Fritsch, E. F. & Maniatis, T. Molecular Cloning: A Laboratory Manual 2nd edn. (Cold Spring Harbor Laboratory, New York, 1989).

53. Ettebeeste, O. et al. Basic zipper-type transcription factor FlbB controls asexual development in Aspergillus nidulans. Eukaryot. Cell 7, 38–48 (2008).

54. Herrero-Garcia, E. et al. Tip-to-nucleus migration dynamics of the axenic development regulator FlbB in vegetative cells. Mol. Microbiol. https://doi.org/10.1111/mmi.13156 (2015).

55. Garzia, A. et al. Aspergillus nidulans FlbB is an upstream developmental activator of conidiation functionally associated with the putative transcription factor FlbB. Mol. Microbiol. 71, 172–184 (2009).

56. Cove, D. J. The induction and repression of nitrate reductase in the fungus Aspergillus nidulans. Biochem. Biophys. Acta 113, 51–56 (1966).

57. Peñalva, M. A. Tracing the endocytic pathway of Aspergillus nidulans with FM4-64. Fungal Genet. Biol. 42, 963–975 (2005).

58. Kumar, S., Stecher, G. & Tamura, K. MEGA7: molecular evolutionary genetics analysis version 7.0 for bigger datasets. Mol. Biol. Evol. 33, 1870–1874 (2016).

59. Letunic, I. & Bork, P. Interactive Tree Of Life (iTOL) v4: recent updates and new developments. Nucleic Acids Res. 47, W256–W259 (2019).

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Author contributions
E.A.E. and U.U. proposed this research line. E.A.E. conceived and supervised the experiments, which were carried out by A.M.-I. and A.S. E.A.E. and O.E. performed bioinformatic analyses. E.A.E. and A.M.-I. wrote the original draft of the manuscript. All authors contributed to the improvement of the text and figures.

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Competing interests
The authors declare no competing interests.

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