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Review

Alkaline pH homeostasis in bacteria: New insights

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Abstract

The capacity of bacteria to survive and grow at alkaline pH values is of widespread importance in the epidemiology of pathogenic bacteria, in remediation and industrial settings, as well as in marine, plant-associated and extremely alkaline ecological niches. Alkali-tolerance and alkaliphily, in turn, strongly depend upon mechanisms for alkaline pH homeostasis, as shown in pH shift experiments and growth experiments in chemostats at different external pH values. Transcriptome and proteome analyses have recently complemented physiological and genetic studies, revealing numerous adaptations that contribute to alkaline pH homeostasis. These include elevated levels of transporters and enzymes that promote proton capture and retention (e.g., the ATP synthase and monovalent cation/proton antiporters), metabolic changes that lead to increased acid production, and changes in the cell surface layers that contribute to cytoplasmic proton retention. Targeted studies over the past decade have followed up the long-recognized importance of monovalent cations in active pH homeostasis. These studies show the centrality of monovalent cation/proton antiporters in this process while microbial genomics provides information about the constellation of such antiporters in individual strains. A comprehensive phylogenetic analysis of both eukaryotic and prokaryotic genome databases has identified orthologs from bacteria to humans that allow better understanding of the specific functions and physiological roles of the antiporters. Detailed information about the properties of multiple antiporters in individual strains is starting to explain how specific monovalent cation/proton antiporters play dominant roles in alkaline pH homeostasis in cells that have several additional antiporters catalyzing ostensibly similar reactions. New insights into the pH-dependent Na⁺/H⁺ antiporter NhaA that plays an important role in *Escherichia coli* have recently emerged from the determination of the structure of NhaA. This review highlights the approaches, major findings and unresolved problems in alkaline pH homeostasis, focusing on the small number of well-characterized alkali-tolerant and extremely alkaliphilic bacteria.

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Keywords: pH homeostasis; Na⁺(K⁺)/H⁺ antiporter; Alkaline pH homeostasis; NhaA; Mrp; MdfA; Tet(L)

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Abbreviations: ΔpH, transmembrane pH gradient; ΔΨ, transmembrane electrical gradient; AIB, α-aminoisobutyric acid; BCECF, 2',7'-bis-(2-carboxyethyl)-5-(and-6)-carboxyfluorescein; LB, Luria–Bertani medium; PMF, proton-motive force, a transmembrane electrochemical gradient of protons composed of the ΔpH (alkaline inside) and ΔΨ (negative inside); SCWP, secondary cell wall polymers; SMF, sodium-motive force; Tc, tetracycline; TMAO, trimethylamine *N*-oxide; TMS, trans membrane segment; MDR, multi-drug transporter; CPA, cation proton antiporters; Ec, *Escherichia coli*; Vc, *Vibrio cholerae*; Mj, *Methanococcus jannaschii*; Bs, *Bacillus subtilis*; Bh, *Bacillus halodurans* C-125; Bp, *Bacillus pseudofirmus* OF4; Sa, *Staphylococcus aureus*

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1. Introduction

Bacteria must maintain a cytoplasmic pH that is compatible with optimal functional and structural integrity of the cytoplasmic proteins that support growth. Most non-extremophilic bacteria grow over a broad range of external pH values, from 5.5–9.0, and maintain a cytoplasmic pH that lies within the narrow range of pH 7.4–7.8 [1–4]. Hence, they are able to acidify or alkalinize the cytoplasm relative to the external milieu. The consequences of being unable to do so are striking. *Caloramator fervidus* (previously named *Clostridium fervidus* [5]) possesses bioenergetic cycles that are entirely Na⁺-coupled and thus it lacks active H⁺ extrusion or uptake systems that can support pH homeostasis. As a result, it grows only within the narrow pH range of 6.3–7.7 that corresponds to an optimal cytoplasmic pH [6]. *Enterococcus* (formerly *Streptococcus*) *faecalis* can grow in the presence of iono-

phores that inhibit the H⁺ circulation required for pH homeostasis if the cells are grown in a rich medium that is K⁺-replete, low in Na⁺ and has a pH near pH 7.6 [7]. Acid pH homeostasis has been extensively studied and recently reviewed [8–12]. Here, we review bacterial alkaline pH homeostasis, which also features prominently in life cycles of pathogens [13,14], as well as ecologically and industrially important bacteria [15–17].

A shift to an alkaline environment, like a shift to an acid environment, is stressful for bacteria as shown by how *Escherichia coli* responds to alkali with SOS and heat shock-like responses [18–20]. It is important to distinguish between survival and growth in response to alkali. Survival involves no net increase in viable cells at the high test-pH and is usually accompanied by some loss of viability. Survival is monitored by counts of viable cells as colony-forming units on a neutral medium. Since survival is often monitored during stationary

phase, it should be noted that cultures that are not well buffered exhibit a marked rise in pH in many conventional growth media, e.g., LB [21]. Growth is accompanied by a logarithmic increase in viable cell count under the alkaline condition although the growth rate is reduced relative to the rate at the optimal neutral pH [20,22]. Turbidity (A_{600}) is frequently used to monitor growth, but viable counts are preferable here too because changes in shape as well as a non-viable cell component can introduce artifacts into the absorbance. The distinction between survival and growth at high pH raises the question of whether the underlying mechanisms are identical, overlapping or completely different.

Many natural habitats for microbes have high pH. Marine bacteria grow in alkaline marine environments at \sim pH 8.2 [23] and other bacteria spend part of their life cycle in marine environments, where they must survive or grow [24,25]. Within the human body, high pH (≥ 10) occurs at the pancreatic duct just below the pylorus [20,26]; thus, enteric flora must exhibit substantial alkali-resistance. Steady-state high pH is found consistently in regions of many insect guts [20,26–28]. A haloalkaliphilic archaeon *Natronococcus* is part of the termite gut flora [29], where a pH of 12 is reached [27]. Additional, extremely alkaline environments include saline alkaline soda lakes, non-saline alkaline lakes and man-made enrichments that are by-products of industrial activities such as the indigo dye production facilities from which the first alkaliphiles were isolated [30–34].

A large number of adaptive strategies are deployed for alkaline pH homeostasis and will be discussed in this review both in the context of the methods used to identify them and the context of the organism in which they are found. The strategies include: (i) increased metabolic acid production through amino acid deaminases and sugar fermentation; (ii) increased ATP synthase that couples H^+ entry to ATP generation; (iii) changes in cell surface properties; and (iv) increased expression and activity of monovalent cation/proton antiporters. Among these strategies, monovalent cation/proton antiporters play an essential and dominant role in alkaline pH homeostasis in many bacteria in addition to roles in Na^+ and volume homeostasis [3,35–41]. Similarly, Na^+/H^+ antiporters play essential roles in homeostasis of pH, Na^+ and volume in eukaryotic cells and organelles [37,42–45].

Bacteria have multiple monovalent cation/proton antiporters (e.g., at least four in *E. coli* [2,37], five in *B. subtilis* [46–49] and four in alkaliphilic *Bacillus* strains [40,50–52] see Fig. 1). Detailed information about individual Na^+/H^+ and $Na^+(K^+)/H^+$ antiporters is beginning to provide insights into the basis for the dominant role of specific antiporters in pH homeostasis of individual bacterial strains. This review will focus on several Gram-negative and Gram-positive bacteria in which studies of individual antiporters as well as genomic, global proteome and/or transcriptome information is available in relation to alkaline pH homeostasis. The methodological approaches used, the major findings and the major unresolved questions will be discussed.

2. Assays of pH homeostasis

The cytoplasmic pH of bacteria has usually been calculated from a pre-set external pH and the Δ pH across the cytoplasmic membrane of the cell. Most assays of the Δ pH, acid inside relative to outside, use labeled probes that are membrane-permeant weak bases. These weak bases distribute across the membrane according to the Δ pH and therefore accumulate within the more acidic cytoplasm that bacteria maintain when the external pH is high [53,54]. The weak base is either radiolabeled or is fluorescent, with fluorescence quenching occurring as the probe is internalized. Weak acids are similarly used to measure a Δ pH, alkaline inside relative to outside; it is best to do both a weak acid and weak base distribution assay when the Δ pH is small. ^{31}P -NMR has been used over a limited range of pH values to assess cytoplasmic pH in *E. coli*, but the conditions for such assays do not always replicate normal growth conditions because of the high cell density needed [3]. The cytoplasmic pH of bacteria has also been assayed using pH-sensitive fluorescent reporter proteins or dyes, such as green fluorescent proteins [12] or BCECF [55–57].

Measurements of the cytoplasmic pH of cells growing or surviving at different external pH values provide information about the limits of alkaline pH homeostasis when correlated with the viable count (at a permissive pH) of the bacterium. A rigorous assessment of the pH homeostatic capacity is further obtained from pH shift experiments, which are optimally conducted under carefully controlled pre- and post-shift pH values [58]. An alkaline shift in the external medium can be imposed on growing cells in pH-controlled batch cultures ([58] and Fig. 2A) or in pH stat (a chemostat in which pH is strictly controlled) [54,59,60]. Alternatively, cells are harvested at particular growth stages and equilibrated in pre-shift buffer until the cytoplasmic and external pH values are approximately the same; the shift is then initiated under specific conditions of energization and cation content ([51,61] and Fig. 2B). Survival data should be collected by pre- and post-shift viable counts (at a permissive pH) to be sure that survival is sufficient for meaningful cytoplasmic pH measurements. There are several parameters that can be varied in the protocols: (i) the size and range of the shift; (ii) the time over which the shift is imposed, i.e., sudden vs. gradual; (iii) the rate and duration of the measurements; (iv) the ion composition of the medium. Since, in general, the mechanism of alkaline pH homeostasis involves cation/ H^+ antiporters, the ion specificity of pH homeostasis can identify which antiporter might be involved and how.

3. Methods for identifying the proteins responsible for pH homeostasis

3.1. Global approaches

3.1.1. Proteomics

Two-dimensional gel electrophoresis of the proteome of bacteria grown at various pH values identifies changes in the level of individual proteins in response to a change in external pH. The proteins are identified by mass spectrometry. Findings

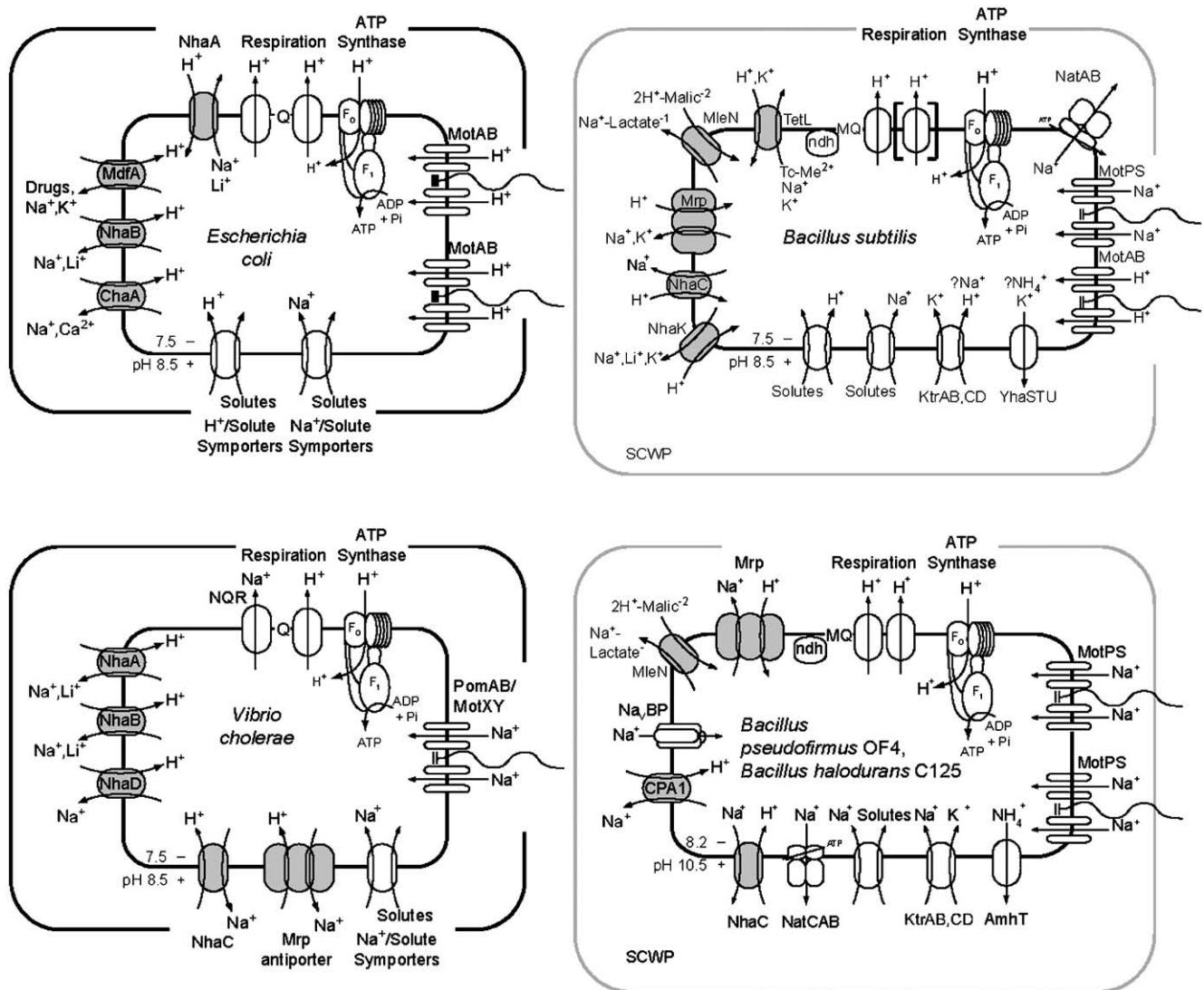


Fig. 1. Na^+/H^+ antiporters, the Na^+ cycle and selected H^+ - and K^+ -translocating proteins in membranes of four physiologically distinct bacteria. Transporters of the inner membrane of two Gram-negative neutralophilic examples, *E. coli* and *V. cholerae*, are schematically depicted within the inner membrane surrounded by the outer membrane. *E. coli*: a respiratory chain with Q representing ubiquinone or menaquinone, ATP synthase (F_0F_1) and flagellar motor that are all H^+ -coupled [236,237]; both H^+ - and Na^+ -coupled solute symporters [238,239]; and four monovalent cation/proton antiporters, one of which is also a multidrug/proton antiporter [37,122,144,238]. *V. cholerae*: a respiratory chain that extrudes both Na^+ and H^+ and H^+ -coupled ATP synthase; Na^+ -coupled solute symporters and polar flagellar motor [13,173] (whereas marine *Vibrio* species also have H^+ -coupled lateral flagella [240]); and at least five monovalent cation/proton antiporters; an annotated NhaP and MleN are not shown ([14,37,241], <http://www.membranetransport.org>). Transporters in Gram-positive neutralophilic *B. subtilis* and the alkaliphilic species *B. pseudofirmus* OF4 and *Bacillus halodurans* C-125 are shown in the cytoplasmic membrane surrounded by a peptidoglycan layer that has associated SCWP (secondary cell wall polymers). The *Bacillus* species have H^+ -extruding respiratory chains (MQ=menaquinone), H^+ -coupled ATP synthases [88,242,243], an ABC-type Na^+ efflux system [244,245], and a CPA:2-type K^+ -extruding system that has been shown to also extrude NH_4^+ in *B. pseudofirmus* OF4 [80,81]. *B. subtilis*: additionally has both H^+ - and Na^+ -coupled solute symporters and flagellar motors [214,246], five characterized monovalent cation/proton antiporters [47–49,193,247] and two Ktr-type K^+ -uptake systems whose ion-coupling properties are not established [136]. *B. pseudofirmus* OF4 has exclusively Na^+ -coupled solute symporters and flagellar rotors [88,214], two monovalent cation/proton antiporters thus far partially characterized [40] and a voltage-gated Na^+ channel that plays a role in supporting pH homeostasis as well as chemotaxis and motility [215]. The *B. halodurans* C-125 genome indicates the presence of MleN and a CPA1 type antiporter [52].

from such screens are well represented by the study of Stancik et al. [62] in *E. coli*. Under aerobic conditions (in LB medium, which contains tryptone, yeast extract and NaCl), the abundance of a variety of proteins is increased by high pH, including the periplasmic protein YceI, MalE and OmpA porins and the virulence-associated OmpX porin [62]. These changes presumably lead to metabolic patterns that are adaptive. For example, MalE would increase availability of

an acid-generating carbohydrate. Several enzymes of amino acid catabolism are also increased in abundance at alkaline pH, including TnaA (tryptophan deaminase) and TnaB (the tryptophan transporter) as well as serine deaminase (SdaA). Deaminases provide an acid-generating mechanism that is adaptive to alkaline challenge just as decarboxylases promote alkalization that is adaptive for an acid challenge [12,63,64], as predicted by early work by Gale and colleagues [65].

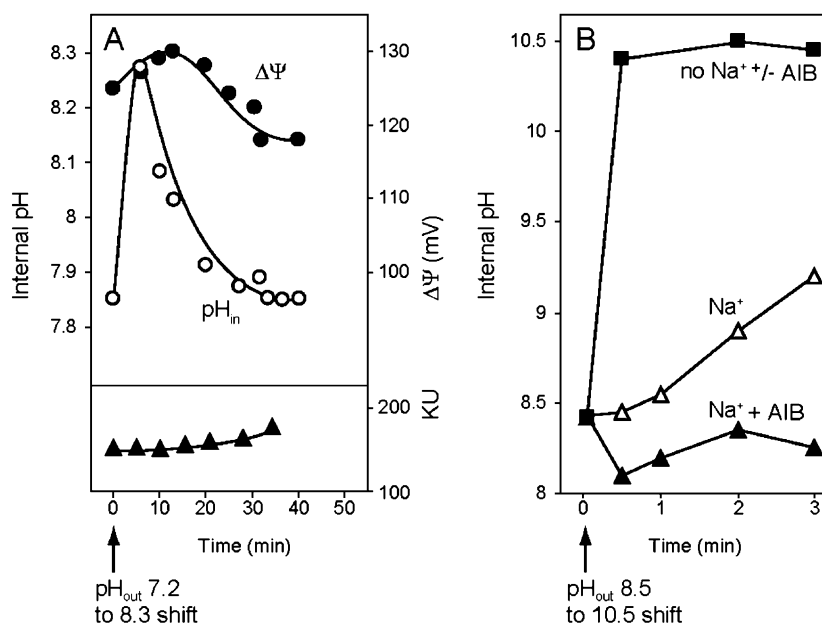


Fig. 2. Alkaline pH shift experiments with *E. coli* and alkaliphilic *B. pseudofirmus* RAB. (A) *E. coli* cells were grown logarithmically at pH 7.2 in minimal glycerol medium batch cultures before the pH of the medium was adjusted to 8.3 over 30 s. A rapid method of cytoplasmic pH and $\Delta\Psi$ determination in intact cells has made it possible to follow the time course and sequence of changes occurring following the pH shift [58]. Growth was also assessed (Klett units). The data are replotted from Fig. 3 in reference [58] with permission from the American Society for Microbiology. (B) Washed pH 10.5-grown cells of *B. pseudofirmus* RAB were equilibrated in potassium carbonate buffer at pH 8.5 and then rapidly shifted to pH 10.5 buffer with either 50 mM potassium carbonate (“no Na^+ ”, with or without added AIB), 50 mM sodium carbonate (“ Na^+ ”) or 50 mM sodium carbonate+500 μM AIB. The cytoplasmic pH was monitored after the shift. The figure is a modified version of Fig. 3 from reference [61], with permission from the American Society for Biochemistry and Molecular Biology.

Increased production of TnaA has also been implicated in a specific and remarkable defense against alkali when anaerobic *E. coli* uses TMAO as its terminal electron acceptor. Use of TMAO generates an alkaline reduction product and in apparent “anticipation” of that alkaline challenge, the Tor phospho-relay system that detects TMAO increases expression of *tnaA*. This results in a functional TnaA- and tryptophan-dependent defense against alkali [66]. It has not yet been shown whether deaminase activities lower cytoplasmic pH, external pH or both. Overall, the proteomic data are consistent with a model in which *E. coli* modulates multiple metabolic pathways of amino acid consumption so as to minimize the effect of an increase in the external pH [63]. A mechanism based on a change of external pH itself can only be realistic when the culture volume is small and/or the fluid layer surrounding the cells is unstirred so that its composition is not in equilibrium with that of the bulk external fluid. In general, the internal volume of cells is much smaller than the external volume. Hence, modest changes in intracellular acid or base production and/or in the protons pumped out of or taken up by the cytoplasm can have a significant impact on maintenance of a constant intracellular pH.

The proteome of a *B. subtilis* mutant disrupted in *Bs-mrpA* (a $\text{Na}^+(\text{K}^+)/\text{H}^+$ antiporter, see Section 5.3.4) and grown at high pH shows elevated levels of salt stress proteins as well as of the ATP synthase [67]. The latter is consistent with the alkali-adaptive value of ATP synthase inferred in *E. coli* ([20] and see Section 3.1.2.1). By contrast, ATP synthase genes are down-regulated during acid-adaptation of *H. pylori* [68] and in fermentative bacteria, the H^+ -pumping, hydrolytic mode of the ATPase is a critical piece of acid-adaptation [69–71].

3.1.2. Genomics-microarrays

In this approach, cDNA is synthesized labeled and hybridized to antisense genomic DNA arrays. Positive hybridization signals quantitatively reflect the level of gene expression and can be compared as a function of growth pH or before and after a shift in external pH.

3.1.2.1. *E. coli*. When grown in a buffered LBK medium (highly buffered LB in which KCl replaces the NaCl) at pH 7.0, *E. coli* cells exhibit a generation time of 18 min vs. pH 8.7, where the cells exhibit a generation time of 25 min [20]. Microarray studies show that among the genes repressed at the alkaline pH are genes encoding two respiratory chain complexes that pump protons outward during electron transport, *cyo* and *nuo*, as well as flagellar and chemotaxis genes [20]. Induced genes include those encoding the F_1F_0 -ATP synthase that imports protons during ATP synthesis, and the alternate *cyd*-encoded terminal oxidase, that generates a PMF without outward proton pumping. These observations indicate that alkali challenge, under growth conditions, changes the gene expression profile: robust oxidative phosphorylation together with the ATP synthase maximize proton retention and proton capture by the cell. This elegant study also shows that pH differentially regulates a large number of periplasmic and envelope proteins. Although the experimental conditions are not directly comparable, it is interesting to note an apparent difference in expression patterns in a microarray study conducted with *E. coli* in sea water (where the organism survives but does not grow) [25] vs. a glucose-containing laboratory medium [72]. It had earlier been found that the sea

water survival condition induces RpoS-controlled genes [73], a type of regulation usually involved in stationary phase ([74,75] and see Section 3.2.3). The microarray data further show that genes involved in cell division and nucleotide biosynthesis are repressed, consistent with cessation of growth. On the other hand, motility and chemotaxis genes as well as many metabolic genes are induced in sea water [72]; under these conditions, induction rather than repression of motility and chemotaxis may support the search by the surviving *E. coli* cells for a niche that can support growth.

3.1.2.2. *B. subtilis*. An alkaline shift protocol has been applied to *B. subtilis* [60]. In LB medium at pH of 6.2, *B. subtilis* initially grows rapidly and acidifies the medium. This is followed by a slower growth rate during which the medium is alkalinized to about pH 7.5. *B. subtilis* does not survive a sudden shift to pH ≥ 9.3 imposed just before the growth slow-down phase. A more modest shift, to pH 9.0, results in a growth arrest for 5 h during which is a significant loss of viable cells, while the medium acidifies gradually. Growth resumes at a slower rate when the outside pH is about 8.5, reaching the same final cell density as the nonshifted control.

Microarray studies have been conducted using the same shift protocol, comparing the pre-shift gene expression profile and the profile after a shift to pH 8.9 [60]. A group of the up-regulated genes is under the control of σ^w , one of the extracytoplasmic function family of σ factors (ECF) that commonly regulate groups of genes in response to extracytoplasmic stimuli [76]. It has been proposed that the induction of the σ^w regulon reflects an intersection of alkali-stress with cell wall stress and Na^+ -stress ([77] see also Section 7). Additional genes that are up-regulated after the alkaline shock include *Bsmrp* [60] genes that encode the Mrp $\text{Na}^+(\text{K}^+)/\text{H}^+$ antiporter ([41,49,78,79] and see Section 5.3.4). This is important since, as noted below (Section 3.1.4), the micro-array technique often does not detect significant up-regulation of genes encoding membrane transporters including antiporters of known involvement in alkali-stress. Another transporter gene locus that is up-regulated in the alkaline shock experiment is the *YhaSTU* locus that catalyzes K^+ efflux in *B. subtilis* and is regulated in a complex way by alkaline pH and Na^+ stress [80]. *YhaSTU* may extrude NH_4^+ (in addition to K^+), as does its alkaliphile homologue *AmhT* [81]. This could contribute to acidification of the medium (Fig. 1).

3.1.3. *lacZ* fusions

In this approach, in-frame promoter or genes encoding protein fusions are randomly introduced to the genome by transposition with the *lacZ* reporter gene [82]. The strains bearing the fusion are identified genetically and by the activity of the reporter gene *lacZ*. Depending on the fusion type, *LacZ* activity reflects transcriptional or translational regulation or both. In *E. coli*, a novel alkali-induced gene locus, *alk*, was identified by this approach [83]; its product was shown to be a membrane-associated redox modulator [62]. In a comparable gene fusion approach, plasmid-based libraries of promoter-reporter fusions (to a *luxCDABE* reporter) were introduced into

E. coli and used to detect promoters that were activated in sea water, including the *rpoS*-dependent promoters [73]. The promoter fusion approach has thus highlighted interplay of alkali-stress with both stationary phase stress and oxidative stress (see Section 7).

3.1.4. Caveats concerning use of the global approaches

The global approaches provide a valuable integrated picture of responses of genes and proteins to various growth conditions and stresses. However, there are several drawbacks that especially pertain to integral membrane proteins. The proteomic approach cannot appropriately resolve many of these proteins, which constitute around 20% of the bacterial proteome. The major reason is that integral membrane proteins are very hydrophobic proteins and often cross the membrane several times in a zig-zag fashion; their hydrophobicity commonly introduces artifacts in 2D as well as in ordinary gel electrophoresis [84,85] and some preparatory steps used for 2D samples (e.g., boiling) can induce their aggregation [84]. This is compounded by the fact that the proteins of greatest interest are generally present at very low levels in the membrane (e.g., the *Ec-NhaA* antiporter constitutes $<0.2\%$ of the *E. coli* membrane proteins [84]). Another problem in applying microarray approaches to membrane proteins such as transporters is that small changes in expression (e.g., 2- to 3-fold [46]) can have a significant physiological effect, but many microarray studies fail to capture such changes.

Another caveat is that neither proteome nor transcriptome studies can identify proteins that are regulated at the level of activity, a feature that is important in antiporters that participate in alkaline pH homeostasis (see Section 4.3). As yet, this property can be discerned only by combining classical genetics (including gene fusion approach) with biochemical approaches. It is not surprising therefore that very few antiporter genes or proteins have been identified in alkali-adaptation screens by any of the global approaches. Yet, their importance has unequivocally been shown by targeted studies of specific antiporter genes and proteins.

3.2. Focus on single genes and their products of potential interest for alkaline pH homeostasis

3.2.1. Identification of genes/proteins involved in alkaline pH homeostasis

For targeted identification of specific genes to be practical, a clue is needed as to which gene/protein might be involved in alkaline pH homeostasis. The global proteomic and genomic assays can provide such clues, but critical insights are often obtained from physiological studies. When Na^+/H^+ antiporters were described by West and Mitchell in intact *E. coli* cells [86], their potential for a role in adaptation to alkali pH was noted, initiating research on the Na^+/H^+ antiporters. The absolute requirement for Na^+ for pH homeostasis in alkaliphilic bacteria and diminished Na^+/H^+ antiporter activity in non-alkaliphilic mutants then provided strong support for an antiporter based pH homeostasis mechanism [39,87,88]. Study of a Li^+ resistant

mutant [89] led to the first cloning of an antiporter gene, *Ec-nhaA* of *E. coli* [90,91] and its regulator *Ec-nhaR* [92]. *Ec-NhaA* is the main antiporter of *E. coli* and many enterobacteria [37].

The genome project provided a flood of eukaryotic and prokaryotic putative antiporter genes of which products were grouped by Saier and colleagues in the monovalent cation proton antiporter superfamily (CPA) [http://tcdb.ucsd.edu/tcdb/\[93\]](http://tcdb.ucsd.edu/tcdb/[93]). A recent phylogenetic analysis of database information shows that *NhaA* belongs to CPA2, a sub family of the monovalent cation/H⁺ antiporters, and its orthologs exist in many prokaryotic genomes as well as eukaryotic genomes including that of humans [45]. The CPA1 family includes the main eukaryotic NHE Na⁺/H⁺ antiporters and has many prokaryotic orthologs such as Mj-NhaP1 ([45] and see Section 4.3.2).

The Mrp-type antiporters, classified as the CPA3 family, are the main antiporters of alkaliphilic bacilli [94]. They were discovered when a partial *mrp/sha* operon was found to complement a pH homeostasis-negative and Na⁺-sensitive phenotypes of an alkaliphilic *Bacillus halodurans* C-125 mutant [50]. Transpositional insertion libraries of *B. subtilis* for Na⁺- and alkali-sensitive mutants led to the identification of Bs-Tet(L) as a multifunctional (tetracycline-metal⁺)(Na⁺)(K⁺)/H⁺ antiporter with a role in pH homeostasis [47]. Once antiporter-deficient strains of *E. coli* were constructed, functional complementation screens have become a common route to clone new antiporters genes and study the encoded antiporters ([95–98] and see Section 3.2.2).

3.2.2. Changing the dosage of the candidate gene to probe its physiological role

The physiological role of a putative antiporter in alkaline pH homeostasis is best inferred from the effect of changes in the dosage of the gene on survival and growth at alkaline pH. For example, the *Ec-nhaA* carried on multicopy plasmid confers upon *E. coli* a markedly increased growth capacity in the presence of high Li⁺ [90] or Na⁺ at neutral pH or growth at alkaline pH in the presence of Na⁺ [95,98]. Deletion of *Ec-nhaA* produces a strain (NM81) that is sensitive to alkaline pH in the presence of Na⁺ [95]. This alkaline-sensitive strain was used to clone *Ec-NhaB*, a house-keeping antiporter in *E. coli* [99], leading to the generation of the double *nhaA* and *nhaB* deletion strain (EP432, [100]). *E. coli* EP432 was used to clone a third antiporter of *E. coli*, *Ec-chaA*, that has Na⁺(Ca²⁺)/H⁺ activity ([101,102], Fig. 1). *E. coli* strains NM81, EP432 and two triple mutants in all three antiporters strains, KNabc and TO114 [103,104], have been extensively used to identify Na⁺(K⁺)/H⁺ antiporter genes from diverse bacteria and to conduct studies of the properties of the antiporters encoded [37,105]. However, it should be stressed that for non-*E. coli* genes this is a heterologous system that cannot provide actual evidence for their roles in the native host. The effects of changing gene dosage in alkali-stressed pH homeostasis and the properties of the antiporters must be examined in the native host, in the correct context.

3.2.3. Studies of gene regulation provide further physiological insights: *Ec-NhaR*, *E. coli nhaA* regulator, as a case study

The pattern of transcriptional regulation of *Ec-nhaA* in *E. coli* offers a particularly interesting demonstration that the mechanisms regulating a single antiporter in response to alkalization can be totally different when supporting logarithmic growth than when supporting survival during stationary phase. The expression pattern of the reporter gene *lacZ* of a *nhaA-lacZ* fusion and Northern analyses of exponentially growing cells show that transcription of *Ec-nhaA* is specifically dependent on Na⁺ and increases at alkaline pH [106]. This regulatory system involves the positive regulator *Ec-NhaR* [92,107–109] of the LysR family of regulatory proteins [110–113], which is Na⁺- and Li⁺-specific and not modulated by ionic strength or osmolarity. Intracellular rather than extracellular Na⁺ is the signal for induction [109]. The binding site for *Ec-NhaR* on the *Ec-nhaA* was identified using purified *Ec-NhaR* in both gel retardation and foot printing assays [108]. *Ec-NhaR* binds the DNA under both inducing and non-inducing conditions. Na⁺ changes *Ec-NhaR* conformation so that its footprint on *Ec-nhaA* is altered in a Na⁺-specific and alkali-dependent fashion [108]. Two *Ec-nhaA* promoters, P1 and P2, were identified by primer extension analysis; only P1 is involved in the *Ec-NhaR*- and Na⁺-dependent regulation of *Ec-nhaA* [114].

In stationary phase, *Ec-nhaA* becomes part of the *rpoS* regulon and its expression is from the P2 promoter rather than from the P1 promoter used during logarithmic growth [114]. RpoS is a sigma factor that is recruited to the RNA polymerase to transcribe stationary phase specific genes, the *rpoS* regulon [74,75,115,116]. Accordingly, the reporter activity of P2-*lacZ* fusion is induced in stationary phase. Primer extension studies show that this induction is dependent neither on Na⁺ nor on *NhaR*, but on RpoS [114]. Most importantly, the *rpoS* dependent P2 is responsible for the survival of the cells in the stationary phase in the presence of Na⁺, at alkaline pH and at combined high Na⁺ and alkaline pH [114], the most stressful condition [117].

4. What properties underpin the ability of certain antiporters to support alkaline pH homeostasis?

Almost all bacteria have multiple monovalent cation/proton antiporters that form part of complex H⁺ and Na⁺ cycles in these cells (Fig. 1). As best studied in *E. coli* and *B. subtilis*, these antiporters do not contribute equally to alkaline pH homeostasis. This raises the question of what unique properties are needed for an antiporter to play a major role in this process. Properties of the antiporter that are considered include: the stoichiometry and kinetics of the exchange; the *K_D* of substrate binding; positive and negative effectors of the activity; structure–function relationships and the reaction mechanism. Many of the properties of antiporters have been studied in right-side-out [118] and everted membrane vesicle systems [119], often using expression of the antiporter from a plasmid in appropriate antiporter-deficient strains of *E. coli* [96]. The

use of these membrane preparations facilitates rapid screening of many conditions and properties.

However, there are two general *caveats* with respect to the properties determined in everted membrane vesicles. First, it is impossible to exclude the possibility that proteins, other than the test antiporter, are affecting the assay results. Second, if the membranes are from *E. coli* and the antiporter is from a different bacterial source, the properties may be altered by the phospholipid and heterologous protein complement of the membranes. A striking example of this is the change in cation coupling of the *Bacillus stearothermophilus* and *Bacillus caldotenax* L-glutamate transporters from $1\text{Na}^+ + 1\text{H}^+$ in the native membranes to $\geq 2\text{H}^+$ in *E. coli* membranes [120]. Therefore, it is of great importance to corroborate major findings using proteoliposomes, preferably made from the phospholipids of the natural bacterial host, into which purified antiporter is reconstituted. Among the antiporters involved in alkaline pH homeostasis, this was first accomplished with Ec-NhaA [84] and has since been achieved with several others [121,122].

4.1. Electrogenicity

4.1.1. Importance of electrogenicity for Na^+/H^+ antiport-dependent pH homeostasis

For pH homeostasis (maintenance of intracellular pH around 7.6) at external alkaline pH (>7.6), net cytoplasmic accumulation of H^+ must occur. When Na^+/H^+ antiporters are used for pH homeostasis, the antiport has to be electrogenic rather than electroneutral [39,98,123]. An electroneutral exchange of cytoplasmic Na^+ for external H^+ is a 1:1 exchange in which no net flux of charge is involved, so the $\Delta\Psi$ component of the PMF cannot contribute to energization of the exchange. Bacteria, like other living cells, exclude Na^+ from the cytoplasm by diverse extrusion mechanisms because it is cytotoxic [117]. Therefore, there is no outwardly directed Na^+ gradient to support electroneutral Na^+/H^+ antiport. Hence, an electroneutral antiport can equilibrate the ΔpH component of the PMF, but cannot catalyze a net H^+ accumulation needed to acidify the cytoplasm at alkaline pH. Electrogenic Na^+/H^+ antiporters exchange a larger number of entering H^+ than the number of exiting Na^+ during a turnover, so that net positive charge moves inward. Therefore, the $\Delta\Psi$ (negative inside relative to the outside) drives the inward H^+ movement and cytoplasmic acidification relative to the outside can be achieved.

4.1.2. Determination of coupling stoichiometries

As exemplified by Ec-NhaA and Ec-NhaB, the coupling stoichiometries of antiport must be determined in proteoliposomes containing the purified antiporter proteins [124,125]. Two approaches have been used: in a kinetic approach, the efflux of Na^+ and influx of H^+ are measured in parallel and the stoichiometry is calculated from the ratio of the fluxes under conditions in which the $\Delta\Psi$ and the ΔpH (or ΔpNa^+) created by the antiporter do not limit its rate [124]; in a steady-state thermodynamic approach, various Na^+ chemical gradients are

imposed across the proteoliposomal membrane under conditions in which the ΔpH is collapsed. The antiporter-dependent membrane potential that develops is measured, allowing calculation of stoichiometry assuming thermodynamic equilibrium and complete coupling [124]. Based on both methods, the coupling stoichiometry of Ec-NhaA is $2\text{H}^+/1\text{Na}^+$ [124]. The stoichiometry of Ec-NhaB is $1.5\text{H}^+/1\text{Na}^+$ [125]. These findings suggest that this small difference in electrogenicity underpins the ability of Ec-NhaA but not Ec-NhaB to support growth of *E. coli* at alkaline pH in the presence of Na^+ [100].

4.1.3. Assessing electrogenicity vs. electroneutrality without defining coupling stoichiometry

For other Na^+/H^+ and $\text{Na}^+(\text{K}^+)/\text{H}^+$ antiporters that function in alkaline pH homeostasis, precise coupling stoichiometries have not yet been determined. However, electrogenicity vs. electroneutrality of antiport activity is inferred from the effect of selectively abolishing the $\Delta\Psi$ or the ΔpH component of the PMF. $\Delta\Psi$ is abolished by permeant anions or cations. Commonly, the potassium ionophore valinomycin is used together with K^+ to abolish $\Delta\Psi$. ΔpH is usually dissipated by using low concentrations of nigericin; K^+ must be present and high concentrations of nigericin must be avoided because electrogenic exchange can occur under those conditions and abolish both components of the PMF [126,127]. Use of both nigericin and valinomycin together, in the presence of K^+ , or use of a protonophore (CCCP) abolishes both the $\Delta\Psi$ and ΔpH and serves as a negative control. Abolition of the $\Delta\Psi$ is accompanied by an increase in ΔpH and vice versa [1, 128]. Therefore, abolition of the $\Delta\Psi$ leads to stimulation of electroneutral antiporters while abolition of the ΔpH inhibits these antiporters. Electrogenic antiporters are inhibited by abolition of the $\Delta\Psi$ even though the ΔpH increases; antiport inhibition occurs because when ΔpH (directed outward in everted membrane vesicles) is the sole driving force, the charge that is transported by an electrogenic antiporter produces a $\Delta\Psi$, positive out, that limits the rate of the antiport in the everted system. Two more approaches are used to demonstrate an electrogenic antiport. One is to impose a gradient of the cationic substrate and use $\Delta\Psi$ probes to show that an antiporter- and cation-dependent electrical potential is generated [124,129,130]; it should be noted that chloride ion is permeant in *E. coli* membranes and should not be included in buffers where $\Delta\Psi$ is being measured [131]. The second approach is to drive the antiport with an imposed H^+ or a Na^+ chemical gradient and show that permeant ions increase the rate of the antiporter [132].

4.2. Substrate and cation specificity

Alkaline pH homeostasis can be achieved by antiporters other than the $\text{Na}^+(\text{Li}^+)/\text{H}^+$ specific antiporters. An example is the electroneutral Bs-MleN, a paralogue of the Na^+/H^+ antiporter Bs-NhaC [133]. Bs-MleN catalyzes $2\text{H}^+ - \text{Malate}^{-2}/\text{Na}^+ - \text{Lactate}^{-1}$ exchange (Fig. 1). When driven by inwardly directed malate and outwardly directed lactate gradients, the external malate enters with H^+ and the cytoplasmic lactate exits

with Na^+ . Therefore the exchange supports net cytoplasmic H^+ accumulation although a role in pH homeostasis of *B. subtilis* has not yet been tested [133]. In theory, an electroneutral K^+/H^+ antiporter could also support net H^+ uptake since bacteria usually maintain an outwardly directed K^+ gradient that could serve as the driving force [134–136]. However, since bacterial growth is inhibited when cytoplasmic K^+ is depleted, even a carefully regulated electroneutral K^+/H^+ antiporter might be a liability. On the other hand, when an electrogenic $\text{Na}^+(\text{K}^+)/\text{H}^+$ antiporter such as multifunctional Bs-Tet(L) and Bs-Mrp antiporter [47,49,121] use K^+ , the outwardly directed K^+ chemical gradient is an additional component to the driving force provided by the PMF. Involvement of K^+/H^+ antiport in alkaline pH homeostasis has been reported in several bacteria but the responsible genes and proteins have not been identified and characterized [137–139]. NhaK, a *B. subtilis* transporter belonging to the widely distributed CPA1 antiporter family [140], catalyzes K^+/H^+ antiport at alkaline pH [48]. This recent finding suggests that the K^+/H^+ antiport activity observed in other bacteria could depend upon other members of this transporter group, i.e., NhaK, NhaP and NhaG homologs [48].

4.3. pH regulation at the level of antiporter activity

The antiporters that are critically involved in alkaline pH homeostasis should be equipped with sensors of the external and/or cytoplasmic pH. They then must be able to transduce this environmental signal to produce a change in antiporter activity, so as to maintain pH homeostasis [98]. Failure to activate the antiporter will result in cytoplasmic alkalinization at alkaline extracellular pH [141]. Failure to shut off the antiporter could lead to over-acidification at lower pH values and energy depletion at higher pH values [132]. Indeed, one of the most characteristic properties of both eukaryotic and prokaryotic Na^+/H^+ antiporters is pH-dependent regulation of their activity at the protein level [37,43,142–147].

4.3.1. *E. coli* NhaA as a paradigm

Ec-NhaA is a paradigm for a prokaryotic, pH-regulated antiporter [144]. Thus, in proteoliposomes containing pure Ec-NhaA as the only protein, Ec-NhaA exhibits a pH response [84] that is identical to that in the native membrane [98]. It shuts off below pH 6.5. As the pH is shifted up to pH 8.5, the NhaA V_{max} increases over three orders of magnitude but does not increase further as pH is raised above pH 9. The pK of the pH response of Ec-NhaA is 7.6 that equals the “set point” for the intracellular pH [1]. At intracellular pH values above the “set point”, Ec-NhaA will be activated so as to restore pH homeostasis. At intracellular pH below the “set point”, it will be inactivated and it shuts off at pH 6.5, a mechanism that avoids over-acidification of the cytoplasm. When the set point of Ec-NhaA is altered by mutation, the result is a change in the pH profile of growth. For example, the G338S mutation of Ec-NhaA results in an antiporter that has lost pH control [132]. The protein is active throughout the range of pH 6.5–9. The G338S mutant is highly pH sensitive at alkaline pH, presumably because the excessive activity at high pH depletes

the cells of energy [132]. The pH-dependence of the Ec-NhaA protein with an H225R mutation is shifted towards acidic pH [141,148] while that of an E252C mutant is shifted towards alkaline pH [149]. These changes are accompanied by changes in the pH profiles of growth in the same direction [141,149].

Insights have been gained into how the Ec-NhaA “pH sensor” transduces the pH signal into a change in antiporter activity. There is only partial overlap between the amino acid residues that affect pH regulation and those that affect transport activity [144,150,151]. This implies that, as in eukaryotic Na^+/H^+ antiporters, there is a pH regulatory site that is distinct from the active site [43,144,152,153]. In Ec-NhaA many of the amino acids that are involved in pH regulation cluster at the N-terminus [154] and the loop between TMS VIII–IX [85,144,149] that, as revealed by cross-linking, are in close proximity [144,155]. Furthermore, alkali-dependent activation of Ec-NhaA is accompanied by a conformational change involving the N-terminus [154] and the loop between TMS VIII–IX [85,149].

4.3.2. Antiporter pH profiles: variations on the theme

Correlations between growth and the pH-dependence of antiporters have been noted in other bacteria. In *B. subtilis*, the $\text{Na}^+(\text{K}^+)/\text{H}^+$ antiport activity of Bs-Tet(L) exhibits a lower apparent K_m for Na^+ at pH 8.5 than 7.5 while the apparent K_m for the tetracycline–metal complex is higher at pH 8.5 than 7.5 [143]. These pH-dependent differences are adaptive, since Tc is most toxic in the lower pH range (because it enters cells by diffusion of the protonated form). On the other hand, Na^+ is more cytotoxic at the higher pH [117]. The NhaP Na^+/H^+ antiporter from *Methanococcus jannaschii*, a hyperthermophilic archaeon, exhibits no activity at pH ≥ 8.0 when expressed in *E. coli* and cannot complement antiporter-deficient *E. coli* strains at elevated pH in the presence of Na^+ . However, Mj-NhaP shows increasing activity as the pH is reduced to 6.0, correlating with the growth profile of its natural host [156]. NhaA from *Helicobacter pylori*, which is highly homologous to Ec-NhaA, lacks pH regulation and exhibits high activity from pH 6.5 to 8.5 when assayed in *E. coli* vesicles [157]. It will be interesting to study the still controversial pH homeostasis mechanism of this bacterium [158,159]. Notably, there are exceptions to these striking correlations between the pH profile of Ec-NhaA and other antiporters and the growth pH profiles of their natural hosts; this is consistent with existence of other participants in the adaptation to pH.

4.4. Kinetics

A fast antiporter such as Ec-NhaA (turn over of 89000 min^{-1} [84]) is advantageous for pH homeostasis because it can rapidly meet the challenge of alkali up to pH 9 once it is activated by an elevated cytoplasmic pH. *E. coli*, cells have been subjected to alkaline shock, during growth at pH 7.2 in a minimal medium with glycerol as carbon source, by imposition of a pH shift in the external pH to 8.3 over 30 s ([58,59], Fig. 2A). In the first sample taken after completion of the shift, the imposed pH gradient is observed, with the cytoplasmic pH at

the pre-shift value of 7.8. Rapid cytoplasmic alkalinization and an increase in $\Delta\Psi$ then occur. Within 4 min after the shift, the ΔpH is near zero and the cytoplasmic pH is almost as alkaline as the new external pH. During this period, growth is very slow but recovers when the ΔpH is re-established. More rapid and/or larger shifts in external pH (to $\text{pH} \leq 9$) result in longer and more complete growth arrest and a much slower growth rate upon recovery [58,59]. Hence, *E. coli* experiences and survives substantial cytoplasmic alkalinization after an alkaline shift. Little or no growth occurs while the cytoplasmic pH is above 8.0. Most likely, this cytoplasmic alkalinization is the pH signal that leads to tremendous activation of Ec-NhaA, which in turn establishes a cytoplasmic pH that supports substantial growth. Failure of one or more elements of the cell division mechanism may normally lead to growth arrest in *E. coli* at $\text{pH}_{\text{in}} \geq 8.3$ ([58] and see Section 3.1.2.1).

The extreme alkaliphile, *Bacillus pseudofirmus* OF4 (formerly *Bacillus firmus* OF4 [160]) can successfully respond to an even larger and more sudden alkaline shift (e.g., from pH 8.5 to 10.5) without a detectable alkalinization of the cytoplasm as long as Na^+ is available to the Bs-Mrp antiporter [61]. This is achieved by including a solute that is co-transported with Na^+ in the reaction mixture as shown for *B. pseudofirmus* RAB in Fig. 2B (and see Section 6.3). *B. pseudofirmus* RAB, an obligately alkaliphilic strain, is closely related to the facultatively alkaliphilic *B. pseudofirmus* OF4 that has been used more extensively in recent studies since its non-alkaliphilic mutants are viable [161,162]. In addition to the robust pH homeostasis upon a sudden alkaline shift, *B. pseudofirmus* OF4 differs from *E. coli* by growing when the cytoplasmic pH exceeds 8.5, i.e., when the external $\text{pH} > 10.5$ ([163] and see Section 6.1).

4.5. Meeting the challenge of the low external H^+ concentration

While a high H^+/Na^+ stoichiometry makes it possible for an antiporter to use the $\Delta\Psi$ to acidify the cytoplasm at alkaline pH, it does not solve the problem of capture of the H^+ by the antiporter when the external concentration of H^+ is very low. No features have yet been explicitly described could trap protons on the surface, comparable to those suggested for some membrane-embedded electron transport- and light-driven proton pumps [164–166]. Structural data such as those described below may reveal testable candidates.

4.6. New insights from the structure of Ec-NhaA

The crystal structure of NhaA, in the down-regulated conformation found at low pH, has recently been determined at 3.45 Å resolution [167] and (Fig. 3). The NhaA architecture is comprised of 12 TMS (Trans Membrane Segments). A negatively charged ion funnel formed of TMSs II, IX, IVc and V, opens to the cytoplasm and ends in the middle of the membrane at the putative ion binding site. There, a unique assembly of short helices of TMS IV and TMS XI are connected and crossed by extended chains. This

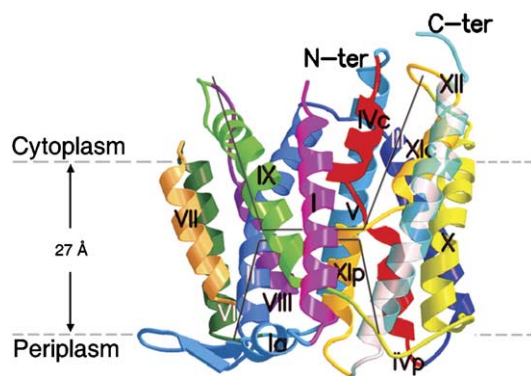


Fig. 3. Overall architecture of NhaA. A ribbon presentation of the NhaA molecule viewed parallel to the membrane (grey broken line) is shown. The 12 TMSs are labeled with Roman numerals. N and C indicate the N and C termini. Note the unique and novel fold formed by unwinding of helices VI and XI in the middle of the membrane. The remaining short helices at the periplasmic or cytoplasmic sides of the membrane are denoted by p or c, respectively. The cytoplasmic and periplasmic funnels are depicted by black line. For further details see [167].

TMS IV/ TMS XI assembly creates a delicately balanced electrostatic environment in the middle of the membrane. A shallow negatively-charged funnel formed by TMSs II, VIII and XIp opens to the periplasm [167] and (Fig. 3). The two funnels point to each other but are separated by a group of densely packed hydrophobic residues forming a periplasmic barrier. On the basis of the structure, it has been proposed that binding of the charged substrates causes electric imbalance allowing for a rapid alternating access mechanism. This ion exchange machinery is regulated by a conformational change elicited by a pH signal perceived at the pH sensor located at the cytoplasmic funnel entry and transduced to the TMSs IV/ XI assembly by helix IX [167] and (Fig. 3). This working model is strongly supported by genetic and biochemical data. At the N-terminus of helix IX and in its vicinity pH induced conformational changes occur [85,149,154] and many mutations affect the pH response of NhaA [144]. Many residues in TMS IV of the TMS IV/XI assembly are conserved in NhaA protein family and their mutations dramatically affect the apparent K_m for Na^+ and Li^+ [150,151]. Mutations in conserved residues in TMS XI of the assembly affect the pH response [144]. Furthermore the need for reorientation of helix IVc with respect to XIp is strongly supported by the double mutant F136C/S342C [151]. This mutant forms a disulfide bond that inhibits transport, while reducing the bond restores activity in a reversible manner [151]. Extensive cross-linking data are in accordance with the close contacts between the TMSs of the TMSs IV and XI assembly and their crucial role for activity and pH regulation [150,151]. The most informative example are the mutants G338S [132] and G338C [151] of the extended chain XI that completely remove the pH control and produce NhaA variants fully active in a pH-independent manner. In the inactive double mutant (T132C/G338C) both pH control and activity can be partially restored by re-establishing the physical interaction between the extended chains by chemical cross-linking [151].

The architecture of the Na^+/H^+ antiporter NhaA reveals the structural basis for the mechanism of Na^+/H^+ exchange and its unique regulation by pH. The physical separation between the “pH sensor” and the exchange machinery revealed by the structure entails long-range, pH induced conformational changes for pH activation. This observation is in agreement with biochemical data obtained from both prokaryotic [144] and eukaryotic [142,153] Na^+/H^+ antiporters. Furthermore, since the structure of NhaA represents a novel fold it contributes to the understanding of the architecture of membrane proteins in general.

5. Alkaline pH homeostasis in neutralophilic bacteria

5.1. pH homeostasis in *E. coli*, a combined effort of NhaA and MdfA antiporters

Even though Ec-NhaA is a paradigm of a pH-regulated antiporter that is suitable for pH homeostasis, Ec-*nhaA* deletion mutants can grow at alkaline pH as long as no Na^+ is added to the medium [98]. These observations suggest that either pH homeostasis is not required under low Na^+ conditions or that another system exists in *E. coli* that controls intracellular pH in the absence of Ec-NhaA. Very recently, this puzzle was solved. A deletion of *mdfA*, a multidrug transporter gene, results in modest alkali-sensitivity [122]. Furthermore, *mdfA* expression from a multi-copy plasmid enables *E. coli* to grow, albeit slowly and to a low yield, at pH values as high as 10 in a complex medium. The enhancing effect of Ec-*mdfA* over-expression on pH homeostasis is illustrated in an experiment which measures the response to a gradual alkaline shift from pH 7.5 to 8.3 or 9.3 of an Ec-*mdfA* deletion mutant, in comparison with a transformant of the same *E. coli* strain over-expressing Ec-*mdfA* from a plasmid (Table 1). Deletion of Ec-*mdfA* has a marginal effect on the response to the smaller shift but results in significantly reduced pH homeostasis in response to the larger pH shift. No significant capacity for cytoplasmic pH homeostasis is observed by over-expression of MdfA when tested without added Na^+ or K^+ , but upon a shift to pH 9.3 in the presence of either ion, multi-copy *mdfA* supports maintenance of a cytoplasmic pH of 7.8–7.9 ([122] and Table 1).

Table 1
E. coli MdfA confers a capacity for Na^+ - and K^+ -dependent pH homeostasis during an alkaline shift in the external pH^a

	Cytoplasmic pH after a gradual shift in pH _{out}					
	pH 7.0→8.3			pH 7.0→9.3		
	CholineCl	NaCl	KCl	CholineCl	NaCl	KCl
<i>E. coli</i> Δ <i>mdfA</i> mutant, control	7.9	7.8	7.8	9.3	9.3	9.3
<i>E. coli</i> Δ <i>mdfA</i> mutant, <i>mdfA</i> expressed from plasmid	7.6	7.6	7.6	9.3	7.9	7.8

^a Cytoplasmic pH was measured by the methylamine accumulation assay after cells were shifted from a growth medium at pH 7.0 to growth media at the final pH values indicated and had established logarithmic growth [122].

In both everted membrane vesicles and MdfA proteoliposomes, MdfA catalyzes Na^+/H^+ antiport and K^+/H^+ antiport activity in addition to the earlier described efflux of multiple drug substrates in exchange for H^+ [122]. The findings suggest that both Ec-NhaA and Ec-MdfA participate in pH homeostasis of *E. coli*. Ec-NhaA, with its very high and pH regulated activity (pH 7.6 “set point”), has a major function in the pH range up to pH 9 ([144] and Section 4.3.1). Ec-MdfA extends the pH tolerance up to pH 10 and also takes over when Ec-*nhaA* is deleted. These observations indicate that the presence of the right antiporter is sufficient to enable a non-alkaliphilic bacterium to grow, albeit sub-optimally, at an extremely high pH on a rich medium.

Why is Ec-NhaA unable to support *E. coli* growth at pH >9 while Ec-MdfA can? Possibly the contribution of Ec-NhaA is limited by its ability to capture external H^+ at the most alkaline edge of the *E. coli* pH range. Since Ec-MdfA can utilize K^+ as well as Na^+ in contrast to $\text{Na}^+(\text{Li}^+)$ -specific Ec-NhaA, Ec-MdfA might also have an advantage in the extremely alkaline range in being able to utilize the outwardly directed gradient of K^+ . We predict that an *E. coli* mutant deleted of both Ec-*nhaA* and Ec-*mdfA* will be a pH-conditional mutant, but such a strain is yet to be constructed. The existence of additional antiporters in *E. coli* cannot be excluded: two CPA1 type antiporters were deduced from the *E. coli* genome for which no associated Na^+/H^+ antiporter activity was demonstrated [168]. Many CPA1 type Na^+/H^+ antiporters are found in diverse bacteria [48,169–171] and, as noted, additional ones such as those in *E. coli* may function as K^+/H^+ antiporters [48].

5.2. The ecological importance of the antiporters in *E. coli* and *V. cholerae*

Enterobacteria such as *E. coli* and *Vibrio cholerae* are released into the alkaline sea water and estuaries during their life cycles. In these environments, they often survive rather than grow [24,25]. Measurements of survival of *E. coli* mutants show that expression of *nhaA* from the P2 promoter, via the *rpoS* regulon, is essential for the survival of *E. coli* in salinic environments at alkaline pH ([114] and Section 3.2.3).

The Na^+ cycle of *V. cholerae* plays a role (by a mechanism that is not yet understood) in the virulence of *V. cholerae* [13,172] and is expected to play a role in survival of this pathogen in salinic-alkaline environments throughout its life cycle. The genes that participate in Na^+ efflux of the *V. cholerae* Na^+ cycle include the *nqr* operon that encodes an electron transport-linked Na^+ pump, the NADH-quinone oxidoreductase complex (NQR) that is also found in marine *Vibrio* species such as *Vibrio alginolyticus* ([173,174], Fig. 1). In addition, *V. cholerae* possesses genes encoding three putative Na^+/H^+ antiporters that have been studied, Vc-NhaA, Vc-NhaB [14], Vc-NhaD [175]; four *nhaC* paralogues, a *mrp* operon, *nhaP* and *mleN* are also predicted from genomics (<http://www.membranetransport.org>) but have not yet been explored. Vc-*nhaA* and Vc-*nhaB* are homologous to the

respective *E. coli* genes and are also found in *V. alginolyticus* [176,177] and pathogenic marine *Vibrio parahaemolyticus* [103,178]. *Vc-nhaD* is homologous to that of *V. parahaemolyticus* [175,179,180].

In contrast to *E. coli*, where a mutant inactivated in the two antiporter genes, *nhaA* and *nhaB*, is more susceptible to NaCl than each of the respective single mutants [100], the inactivation of three putative antiporters (*Vc-nhaABD*) does not alter the exponential growth of *V. cholerae* in the presence of high Na⁺ concentrations and has only a slight effect on the bacterial survival in the stationary phase. In contrast, all mutants lacking *Vc-nhaA* show a Li⁺ sensitive phenotype during the exponential phase of growth and a marked decrease in survival of the triple mutant is observed in the stationary phase in the presence of Li⁺ [14]. The capacity of the NQR pump of *V. cholerae* to specifically extrude Na⁺ but not Li⁺ [13,174] may explain the higher contribution of the *V. cholerae* Na⁺/H⁺ antiporters to Li⁺ resistance as compared to Na⁺ resistance. In the absence of three of the Na⁺/H⁺ antiporters, the NQR pump can compensate for the Na⁺/H⁺ but not the Li⁺/H⁺ antiport activity, resulting in a Li⁺ sensitive but not a Na⁺ sensitive phenotype. Experiments with a quinone analogue that inhibits NQR support this idea [14]. Although the inhibitor has no effect on the growth of wild type *V. cholerae*, it greatly inhibits growth of both the *Vc-nhaA* and *Vc-nhaABD* mutants in the presence of elevated Na⁺ and pH [14]. These results suggest that NhaA is involved in the Na⁺ and H⁺ homeostasis of *V. cholerae* at alkaline conditions but its contribution is revealed only when the Na⁺ pump activity of NQR is inhibited.

5.3. *B. subtilis* has interlocking H⁺, Na⁺ and K⁺ cycles that support moderate alkali-resistance

5.3.1. Multi-functional antibiotic-resistance proteins Tet(L) and Tet(K) catalyzing (Tetracycline)(Na⁺)(K⁺)/H⁺ antiport

Tet(L) and Tet(K) genes are close homologues. Bs-Tet(L) is encoded in the *B. subtilis* chromosome and in plasmids of Gram-positive bacteria [181–183]. When expressed from a single chromosomal copy, the Bs-Tet(L) gene confers resistance only to very low levels of tetracycline (Tc), but when the gene is amplified under selective pressure or expressed from a plasmid, clinically relevant levels of antibiotic-resistance are observed [46,47,184,185]. Sa-Tet(K) is encoded in plasmids that can be incorporated in the chromosome of the human pathogen *Staphylococcus aureus*; there it supports a high level of antibiotic-resistance [186]. Bs-Tet(L) and Sa-Tet(K) form a branch of the 14-TMS member of the drug/H⁺ efflux antiporters of the Major Facilitator Superfamily of membrane transporters [187]. They both catalyze Tc–Me/H⁺ antiport. The Tc is complexed with a divalent metal ion (magnesium, manganese, cobalt are optimal) the Tc–Me complex bears a single net positive charge [143,188–190]. Bs-Tet(L) catalyzes Na⁺/H⁺ and K⁺/H⁺ antiport as well as Tc–Me/H⁺ antiport in both membrane vesicles from antiporter-deficient *E. coli* and in proteoliposomes reconstituted with purified Bs-Tet(L) [143,191]. Sa-Tet(K) also catalyzes these multiple antiport activities, as demonstrated in membrane vesicles [130,191].

Just like their monovalent cation substrates; neither Tc nor the divalent metal ion can be effluxed alone by Bs-Tet(L) or Sa-Tet(K). Competitive patterns of inhibition are evident between the three efflux substrates [191]. Data from site-directed mutagenesis experiments support the hypothesis that antiport occurs by a ping-pong mechanism using a single translocation pathway. However, the Tc–Me complex probably requires more contacts than Na⁺ or K⁺ for binding and positioning through the translocation pathway [191].

The Bs-Tet(L)-mediated antiport activities are electrogenic, with entry of a greater number of H⁺ than cytoplasmic Na⁺, K⁺ or Tc–Me exiting per turnover [121,130,143]; the precise stoichiometry has not yet been determined. An additional catalytic capacity of Bs-Tet(L) is observed when the external pH is elevated and the combined ΔΨ and external K⁺ concentration (relative to cytoplasmic or intravesicular [K⁺]) produces an inwardly directed electrochemical K⁺ gradient. Under these conditions, K⁺ and H⁺ may serve as coupling ions that are taken up in exchange for Na⁺ or Tc–Me [191].

5.3.2. Bs-Tet(L) has a major role in Na⁺(K⁺)/H⁺ antiport-dependent pH homeostasis in *B. subtilis*

Deletion of *Bs-tet(L)* in *B. subtilis* results in a tetracycline- and alkali-sensitive phenotype and reduced ability to grow on media with suboptimal [K⁺] at elevated pH ([46,192,193] and Section 3.2.1). When pH 7.5-equilibrated wild type cells are subjected to a shift in external pH to 8.5, they maintain a cytoplasmic pH of 7.5–7.6 in a Na⁺- or K⁺-dependent manner; *Bs-tet(L)* deletion strains completely lose this capacity (Table 2), but it is restored in transformants with either *Bs-tet(L)* or *Sa-tet(K)*-bearing plasmids [46]. The physiological functions of Bs-Tet(L) that are unrelated to the antibiotic are highly integrated into the physiology of *B. subtilis*, a conclusion supported by a transcription-level regulatory component that is mediated by Na⁺ and elevated pH [46] that is distinct from the post-translational regulation by Tc [194].

Table 2

Capacities of *Bacillus subtilis* and alkaliphilic *Bacillus pseudofirmus* OF4 for Na⁺- or K⁺-dependent pH homeostasis during a moderate or extreme alkaline shift in external pH^a

		Cytoplasmic pH, 10 min after a sudden shift				
		pH 7.5→8.5			pH 8.5→10.5	
		CholineCl	NaCl	KCl	Na ₂ CO ₃	K ₂ CO ₃
<i>B. subtilis</i>	Wild type	8.5	7.5	7.6	10.5	10.5
	Δ <i>tet(L)</i>	8.5	8.5	8.5	–	–
	Δ <i>mrpA</i>	8.5	7.6	7.8	–	–
<i>B. pseudofirmus</i> OF4	Wild type	8.5	7.5	8.4	8.2	10.5

^a For the pH 7.5→8.5 shift, washed logarithmic phase cells, grown on and energized with malate, were equilibrated at pH 7.5 in the presence of 100 mM cholineCl, KCl or NaCl before the external pH was rapidly raised to 8.5. For the pH 8.5→10.5 shift, comparable cells were equilibrated in 50 mM Na₂CO₃ or K₂CO₃ buffer at pH 8.5 before the external pH was rapidly raised to 10.5. In both sets of shift experiments, the cytoplasmic pH was determined after 10 min from a methylamine accumulation assay as described earlier [46,49].

5.3.3. Convergence of antibiotic/multi-drug resistance and alkali-resistance

The similar roles of the multi-drug efflux protein Ec-MdfA (see Section 5.1) and Tc efflux protein Bs-Tet(L) in Na⁺- and K⁺-dependent pH homeostasis and alkali-tolerance in their native hosts raises the questions of how and why this convergence of functions arises. Genomic evidence suggests that the 14-TMS Bs-Tet(L) and Sa-Tet(K) efflux antiporters evolved independently from the Gram-negative, 12-TMS Tn10-type Tet proteins [195,196]. They may have evolved in a pre-existing 12-TMS transporter by insertion of the two central TMS, VII and VIII [197]. This idea is supported by the loss of Tc–Me efflux capacity and retention of all the Na⁺(K⁺)/H⁺ and K⁺ uptake functions of Bs-Tet(L) by a synthetic “Bs-Tet(L)-12” which lacks those central TMS [198]. Interestingly, the conjoining of Tc–Me²⁺ efflux activity with Na⁺(K⁺)/H⁺ antiport activity may be an even more widespread phenomenon since a gene encoding a new member of the Tet(L)/Tet(K) group *tet38* has recently been found in the *S. aureus* chromosome [199] and the deduced product of another recently described Tc-resistance gene, *tet35*, from *Vibrio harveyi* exhibits 28% identity and even greater similarity to NhaC from alkaliphilic *B. pseudofirmus* RAB [200].

The particular combination, in one transporter, of drug–multidrug/H⁺ and Na⁺(K⁺)/H⁺ antiport has arisen multiple times, i.e. in Ec-MdfA and Bs-Tet(L) and perhaps Tet35. Apparently, this combination of antiport activities is readily achievable by protein evolution and bacteria are likely to be advantaged by having such a multifunctional antiporter. The Na⁺(K⁺)/H⁺ antiport could enhance the fitness of the host under conditions of alkaline pH, leading to persistence of the antiporter gene even when its drug substrate is absent [17]. Further study of the similar physiological roles of an Mdr, Ec-MdfA and of the single drug efflux protein, Tet(L), in the alkali-tolerance of *E. coli* and *B. subtilis*, respectively, will afford insights into general design principles of these multifunctional drug transport proteins and into the physiological networks in which they function.

5.3.4. Contributions of the Bs-Mrp antiporter and additional *B. subtilis* antiporters to pH homeostasis

The Bs-Mrp (also called Sha) antiporter is part of a large incompletely understood and unique antiporter family that is encoded by operons containing 6–7 *mrp* genes and has been designated CPA3 [41,140]. In contrast to the central, Na⁺-specific role of Mrp in pH homeostasis in alkaliphilic *Bacillus* species (see Section 6.2.1), in *B. subtilis*, Bs-Mrp hardly plays any role in Na⁺- and K⁺-dependent pH homeostasis (Table 2) while it plays a major role in Na⁺-resistance [49]. This critical role in Na⁺-resistance probably accounts for the appearance of Bs-*mrp* genes on a list of essential genes of *B. subtilis* compiles from assessments in LB medium, which contains added Na⁺ [201]; a Bs-*mrp* null mutant is viable on LBK medium, in which Na⁺ is replaced by K⁺ ([78] and Section 3.1.2.2). The basis for a more dominant role for Bs-Tet(L) in pH homeostasis and for Bs-Mrp in Na⁺-resistance is not yet clear. Nor is the basis for the apparent complexity of Mrp understood [41], but

perhaps Mrp proteins have more than a single Na⁺(K⁺)/H⁺ antiporter activity [202] and/or additional catalytic activities as suggested for MrpE [203] and supported by the involvement of Bs-MrpF in cholerae efflux [49,78].

B. subtilis has at least three additional Na⁺/H⁺ antiporters. The Bs-MleN antiporter couples this exchange to a malate uptake and lactate efflux and contributes to growth at low PMF, i.e., in the presence of inhibitory concentrations of uncouplers ([133] and Section 4.2). Bs-NhaC (YheL), another Na⁺/H⁺ antiporter, does not have a prominent role in wild-type pH homeostasis; its expression is increased in Bs-*tet*(L) deletion mutants, suggesting that it has a back-up role [193]. The newly characterized Bs-NhaK is a Na⁺(K⁺)(Li⁺)/H⁺ antiporter [48]. The precise contributions of these diverse antiporters to the physiology of *B. subtilis* are a puzzle to be solved. In the epiphytic niches of *B. subtilis*, there are variable pH-salt conditions, diverse plant metabolites that serve as growth substrates and numerous potential growth inhibitors [204]. They create different combinations of alkali, salinity and toxins that could account for the complexity of the Na⁺ cycle and its interactions with H⁺ and K⁺ cycles (Fig. 1).

6. Alkaliphilic bacilli: the Na⁺-cycle and other adaptations to extremely alkaline conditions

6.1. The centrality of active pH homeostasis in alkaliphily

Extremely alkaliphilic *Bacillus* species grow optimally at pH values of 10 and above [88,205]. Obligate alkaliphiles (*Bacillus alcalophilus*, *B. pseudofirmus* RAB) are better adapted to pH values above 11 than facultative strains [206] but facultative alkaliphiles are more useful for studies of the relationship of specific gene products to alkaliphily because non-alkaliphilic mutant strains can grow at pH 7–8.5. Among the most intensively studied facultative alkaliphiles (*B. pseudofirmus* OF4, *B. halodurans* C-125, and *Bacillus cohnii* YN2000), *B. pseudofirmus* OF4 is the most alkaliphilic, grows in semi-defined and defined media at extremely alkaline pH and is genetically accessible to targeted gene deletions [51,52,206,207]. *B. pseudofirmus* OF4 grows with a doubling time of 54 min at pH 7.5 vs. a doubling time of 38 min at external pH values from 8.5 to 10.6 in chemostat cultures containing semi-defined malate medium ([163], Fig. 4). At external pH values of 7.5, 8.5 and 9.5, the cytoplasmic pH remains close to 7.5, i.e., the alkaliphile produces a ΔpH, acid in, that rises from zero to 2.0 pH units (shown in parentheses in Fig. 4). Although the ΔpH at pH 10.6 is 2.3 units, larger than at pH 9.5, it does not keep pace with the external pH increase and the cytoplasmic pH rises to 8.3 at external pH of 10.6. The cytoplasmic pH and the doubling time increase almost in parallel above pH 10.6. Neither the ΔΨ nor the total PMF correlates similarly with the increasing doubling time in the high pH range. At the highest external pH examined, pH 11.4, the doubling time is more than 10 h and many of the cells form chains with multiple septa.

Major inferences can be drawn from the chemostat study about alkaliphily and pH homeostasis in extremophile *Bacillus*

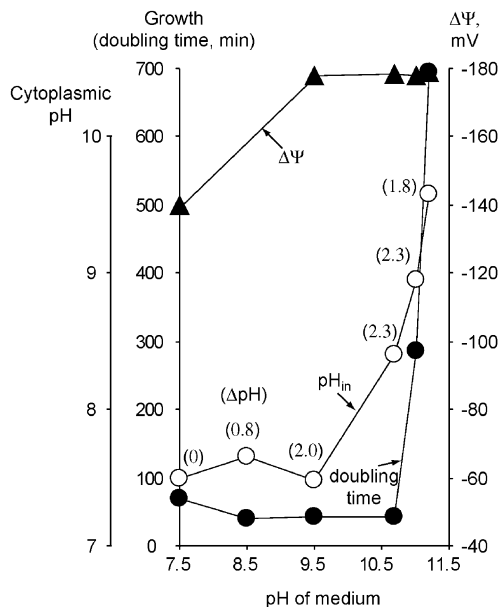


Fig. 4. Generation time and PMF parameters of *B. pseudofirmus* OF4 as a function of external pH in continuous cultures of malate-containing medium. Cells were grown in semi-defined malate containing medium maintained at the indicated pH values. tg, doubling time was calculated from the dilution rates in the chemostat and PMF parameters were determined as described in reference [88]; the data are replotted from Fig. 1 of this reference with permission from Blackwell Publishing Ltd.

growing under non-fermentative growth conditions. First, generation of a ΔpH , acid in, of 2–2.3 units is consistent with a remarkable pH homeostasis capacity that was also evident in pH shift experiments (Fig. 2B); it is not known what prevents further increases in the ΔpH at external pH of ≥ 10.6 . Second, the cells grow optimally even after the cytoplasmic pH increases significantly beyond 7.5. At external pH of 10.6 a doubling time of 38 min is still observed when the cells have a cytoplasmic pH of 8.3. We hypothesize that alkaliphiles have greater tolerance than non-alkaliphiles for elevated cytoplasmic pH (see the *E. coli* shift experiment in Fig. 2A and Section 4.4). This is consistent with an intriguing structural study showing that a cytoplasmic enzyme of alkaliphilic *B. alcalophilus* exhibits features not found in neutralophile homologues but are associated with extracellular enzymes that function at very high pH values [208]. Third, above a cytoplasmic pH of about 8.5, increasing cytoplasmic pH correlates well with a decreasing growth rate. As in *E. coli* cells ([58], Sections 4.4 and 3.1.2.1) elements of cell division may be more sensitive to alkalinization of the cytoplasm than other cellular processes. Cell wall stress could also be an important contributor to setting the upper limit for growth in the alkaliphiles.

Active mechanisms of pH homeostasis have a less exclusive role during alkaliphile growth on fermentable carbon sources. When wild type *B. pseudofirmus* OF4 cells are grown on and energized with glucose instead of malate during a pH 8.5 \rightarrow 10.5 shift (the same shift protocol as in Table 2), the cytoplasmic pH in the absence of Na^+ is 9.2 instead of 10.5 as observed in the presence of malate [209]. This suggests that an antiporter-independent capacity such as intracellular acid

production can partially replace the active transport-based mechanism.

6.2. The antiporters of the alkaliphile Na^+ cycle

In support of the essential role of Na^+/H^+ antiporters in alkaliphily, alkaliphile pH homeostasis is strictly dependent on the presence of Na^+ (Fig. 2B and Table 2, [40,87,210]). We hypothesize that the alkaliphiles use Na^+ -specific antiporters in order to avoid the risk of lowering their cytoplasmic K^+ concentration adversely during the robust, ongoing antiport that is required at high pH. Assays of the total, PMF-dependent $^{22}\text{Na}^+$ efflux capacity of whole cells of *B. subtilis* and *B. pseudofirmus* OF4 have been conducted under conditions that were at the upper pH range for *B. subtilis* and the lower end for the alkaliphile; the alkaliphile exhibits about 10-fold higher capacity for PMF-dependent Na^+ efflux (that is presumed to represent antiport activity) than the neutralophile [211].

6.2.1. Alkaliphile Mrp antiporter

Full *mrp* operons are widely distributed in bacterial genomes and usually contain 6–7 genes whose products (Mrp A, B, C, D, E, F, G) are all predicted to be hydrophobic membrane proteins [41]. The full *Bp-mrp* operon has been cloned and expressed in antiporter-deficient *E. coli* strains in which fluorescence-based assays demonstrate Na^+ -specific monovalent cation proton antiport [212]. Active Mrp antiporters have not yet been purified and studied in sufficient biochemical detail to indicate any adaptations that may help to meet the major challenge of adequate proton capture at pH ≥ 10 . Nevertheless, genetic data assign roles in alkaliphily and Na^+ resistance to *mrp*. A non-alkaliphilic mutant of *B. halodurans* C-125 no longer grows above pH 9.5, has a cytoplasmic pH of 10.4 when shifted to that external pH and exhibits no $\Delta\Psi$ -dependent Na^+/H^+ antiport in native vesicles although significant ΔpH -dependent antiport remains ([50,213] and Section 3.2.1). This mutant has a single point mutation in *Bh-mrpA* that could be corrected by cross-over from a partial clone of the *mrp* operon, leading to the restoration of normal pH homeostasis and alkaliphily. Deletions of *Bp-mrp* in *B. pseudofirmus* OF4 have thus far been lethal (Ito, M., unpublished data) but a *Bs-mrp* null mutant and in-frame deletion mutants of each *Bs-mrp* gene of *B. subtilis* are viable [78]. More robust Na^+ extrusion is probably necessary throughout the pH range of the alkaliphile than in *B. subtilis*, since all the ion-coupled solute symporters and flagellar motors of the alkaliphile are Na^+ coupled (Fig. 1) [88,214].

6.2.2. *Bp-NhaC* and additional antiporters of alkaliphilic *Bacillus*

Bp-NhaC plays a role in pH homeostasis at pH 10.5 under Na^+ -limiting conditions although its major role is in Na^+ extrusion at lower pH values [51]. Based on genomic data, *B. halodurans* C-125 [52] is predicted to have, in addition to *Bh-nhaC* and *Bh-mrp*, a homologue of the *Bs-mleN* gene that encodes the 2H-malate/Na-lactate antiporter ([133] and see Section 4.2) and a member of the CPA-1 family of antiporters

whose closest bacterial homologue is the NhaP Na⁺/H⁺ antiporters [169,171].

6.3. Re-entry routes for Na⁺ are required for optimal pH homeostasis

The essential role of Na⁺/H⁺ antiporters in alkaliphily also makes routes that recycle Na⁺ essential to provide substrate for ongoing antiporter activity. One such route is Na⁺ coupled transporters that increase Na⁺. During a pH 8.5 → 10.5 shift in external pH in the presence of 100 mM Na⁺, cells of *B. pseudofirmus* RAB exhibit a cytoplasmic pH that increases from just below 8.5, at 30 s after the shift, to pH 9.1 at 3 min (Fig. 2B). When the non-metabolizable amino acid AIB that is co-transported with Na⁺ is included in the shift buffer, pH homeostasis is greatly enhanced, with an initial reduction in the cytoplasmic pH to about 8.2, followed by a return to the initial pre-shift pH ([61], Fig. 2B).

The observed role of Na⁺/solute symporter in providing cytoplasmic Na⁺ for the antiporters raises the question of how alkaliphiles achieve significant (although not optimal) Na⁺/H⁺-dependent pH homeostasis when solutes that enter with Na⁺ are limiting or omitted. A Na⁺-channel that is activated at alkaline cytoplasmic pH was postulated [4,88]. Recently, this hypothesis has been supported by studies of two Na⁺ channels of *B. pseudofirmus* OF4, Bp-MotPS and Bp-Na_vBP [4,88,214,215]. MotPS is the Na⁺-translocating motility channel that was earlier suggested as a strong candidate for a role in Na⁺-entry in support of pH homeostasis [216]. Na_vBP, encoded by *ncbA*, is a member of a newly characterized family of novel voltage-gated, ion-selective Na⁺ channels in bacteria [215,217,218]. The effect of single and double deletions of Bp-*motPS* and the Bp-*ncbA* show that Bp-Na_vBP plays a significant role in supporting pH homeostasis while Bp-MotPS plays an ancillary role [215]. Bp-Na_vBP also has roles in motility and chemotaxis, which are observed only at high pH in the alkaliphile [163,214,215,219]. Consistently, the amplitude of channel-mediated current is increased and its activation is potentiated by high pH [215]. The detailed unraveling of interactions between Bp-Na_vBP and the chemotaxis apparatus and of the interplay between Na_vBP signaling and pH homeostasis will be an interesting new chapter of alkaliphile physiology that will have applications to the other salt- and alkaline-tolerant bacteria in which this family of channels has been identified [218].

6.4. Alkali-adaptive strategies of alkaliphiles apart from the Na⁺ cycle

6.4.1. A role for cell surface properties

Secondary cell wall polymers (SCWP) that are associated with the peptidoglycan layer of Gram-positive bacteria have been implicated in an ancillary role in *Bacillus* alkaliphily ([55,220,221] and Fig. 1). *B. halodurans* C-125 has teichuronic acid and teichuronopeptide as the major SCWP [222]. A mutant strain that does not produce teichuronic acids shows no defect in growth at pH 10 or 10.5, whereas a mutant lacking both teichuronic acids and teichuronopeptide grows poorly at

pH 10.5. At pH 10, growth is impaired but still substantial, correlating with a slightly higher cytoplasmic pH than wild type [55]. The major SCWP in *B. pseudofirmus* OF4 is an S-layer layer, encoded by the highly expressed *slpA* gene. At pH 7.5, deletion of *slpA* leads to significantly better growth in the mutant than wild type. A modest growth defect is observed at pH 10.5 that is exacerbated at pH 11. These observations suggest that the alkaliphile produces a polymer whose costly production reduces its growth rate at pH 7.5 but provides a small advantage in the event of a sudden alkaline shift [221]. In both *B. halodurans* C-125 and *B. pseudofirmus* OF4, SCWP are neither necessary nor sufficient substitutes for active antiporter-based pH homeostasis. The positive effect that these acidic SCWP do provide upon alkaline challenge is presumed to result from their ability to bind cations, perhaps even trapping them near the membrane surface. This may enhance the availability of H⁺ and Na⁺ for pH homeostasis and bioenergetic work [221,223,224].

Another cell surface feature that should be noted is the presence of high levels of cardiolipin as well as squalene in membranes of alkaliphilic *Bacillus*, with high pH-dependent increases in the cardiolipin [225]. Models have been advanced for a role of cardiolipin in trapping protons at the membrane surface [226] and for squalene in lowering H⁺ permeability of the lipid bilayer [227], roles that are yet to be probed experimentally.

6.4.2. Involvement of the oxidative phosphorylation machinery

The bioenergetic capacities of a panel of Bp-ATP synthase mutants in comparison with wild type *B. pseudofirmus* OF4 show that ATP synthesis capacity is a significant factor in overall pH homeostasis [228]. This is also inferred from microarray studies of *E. coli* and *B. subtilis* ([20,67] and see Section 3.1.2). The alkaliphile ATP synthase is further adapted for alkaliphily in having features that minimize loss of H⁺ to the bulk through the synthase when cells experience a sudden alkaline shift [228,229].

In contrast to the repression of H⁺-extruding terminal oxidase Cyo in *E. coli* growing at elevated pH ([20] and see Section 3.1.2), the H⁺-extruding Cta terminal oxidase of *B. pseudofirmus* OF4 is up-regulated at the alkaline edge of the pH range [230,231]. In fact, general properties of alkaliphilic *Bacillus* species include: alkali-dependent increases in level of H⁺-extruding respiratory chain components; and low redox potentials of alkaliphile respiratory chain components relative to neutralophile homologues [206,232–234]. Both these properties presumably relate to the importance of generating of a ΔΨ that is higher in alkaliphiles than in most bacteria (compare the *B. pseudofirmus* OF4 ΔΨ in Fig. 3 with that of *E. coli* in Fig. 2). This capacity is especially important at high pH, where the burden of ongoing electrogenic antiport and the higher energetic cost of ATP synthesis increase the demands upon ΔΨ generation [88].

7. The interaction between alkaline stress and other stresses

Alkaline stress has significant interplay with many other stresses. These overlaps complicate a linear association of a

protein with defense against alkali per se on the basis of its activation or induction upon alkaline challenge. Several examples follow. First, there is an overlap between alkaline stress and salt stress. It is due to the significant increase in the Na^+ cytotoxicity as the pH rises [117]. Moreover, the toxicity of Na^+ is highly dependent upon the status of the cytoplasmic K^+ concentration [22]. Second, alkaline pH has interplay with cell wall stress [77], because of the alkali-vulnerability of the subset of cell wall biosynthetic enzymes that are exposed on the outside of the membrane. Third and fourth, interplay of an alkaline challenge with that of high temperature arises because high temperature alters the H^+ permeability more than Na^+ permeability in many bacteria [235] and interplay with changes in osmolarity arises because some monovalent cation/proton antiporters have roles in osmoadaptation [98]. Fifth, interplay between alkaline stress and the stress of entry into stationary phase is strikingly illustrated by a switch in the regulatory mechanism of transcription of *Ec-nhaA* in *E. coli* ([114] and Section 3.2.3). Finally, the induction by alkaline conditions of an *E. coli* protein, Alx, that is involved in redox stress, adds another stress intersection to the list [62,83].

8. Closing remarks

The ability of bacteria to survive or to grow under alkaline conditions is of widespread ecological, industrial and epidemiological importance. The total repertoire and the overlap of proteins required for survival vs. growth are still being elucidated. They encompass proteins that contribute to structural, metabolic as well as bioenergetic adaptations. Active alkaline pH homeostasis that is centrally important in both survival and growth depends upon monovalent cation/proton antiporters, whose identification is generally achieved by combining genetic and biochemical approaches. Genomic and proteomic information is providing a wealth of information about the complexity of interlocking H^+ and Na^+ cycles that function in individual bacterial strains. An intricate interplay between alkaline and other stresses regulates the level of expression and activity of these antiporters. Studies of novel antiporters and channels involved in alkaliphily both of neutralophiles and extremely alkaliphilic species continue to reveal strategies that turn out to have counterparts across a broad range of important bacteria. Further biochemical studies are required to define the specific mechanisms that underpin the efficacy of the physiologically important antiporters and, in some cases, their capacity to catalyze multiple activities such as drug or multi-drug/ H^+ antiport as well as $\text{Na}^+(\text{K}^+)/\text{H}^+$ antiport. Given the insights provided by the novel NhaA structure, determination of additional antiporter structures will be needed to clarify structural principles related to the mechanism of Na^+/H^+ , its regulation by pH and capacity for cation capture.

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