Ethidium binding to *Salmonella enterica* ser. Typhimurium Cells and Salmon sperm DNA

Sandra Sakalauskaitė1, Valeryia Mikalayeva1,2, Rimantas Daugelavičius*

1Department of Biochemistry, Faculty of Natural Sciences, Vytautos Magnus University, Kaunas, Lithuania
2Institute of Cardiology, Lithuanian University of Health Sciences, Kaunas, Lithuania
*Correspondence: rimantas.daugelavičius@vdu.lt; Tel.: +370 37 327917 (R.D.)

**Abstract:** Bacterial resistance to antibiotics due to an increased efficiency of the efflux is a serious problem in clinics of infectious diseases. Knowledge of the factors affecting the activity of efflux pumps would help to find the solution. For this, fast and trustful methods for the efflux analysis are needed. Here we analyzed how the assay conditions affect the accumulation of efflux indicators ethidium (Et+) and tetraphenylphosphonium in *Salmonella enterica* ser. Typhimurium cells. An inhibitor phenylalanyl-arginyll-β-naphthylamide was applied to evaluate the input of RND family pumps into the total efflux. In parallel to spectrofluorimetric analysis, we used an electrochemical assessment of Et+ concentration. Results of our experiments indicated that Et+ fluorescence increases immediately after the penetration of this indicator into the cells. However, when cells bind a high amount of Et+, intensity of the fluorescence reaches the saturation level and stops reacting to the accumulated amount of this indicator. For this reason, electrochemical measurements provide more trustful information about the efficiency of efflux when cells accumulate high amounts of Et+. Measurements of Et+ interaction with the purified DNA demonstrated that affinity of this lipophilic cation to DNA depends on the medium composition. The capacity of DNA to bind Et+ considerably decreases in presence of Mg2+. Polymyxin B or when DNA is incubated in high ionic strength media.

**Keywords:** ethidium; tetraphenylphosphonium; multidrug resistance; outer membrane permeability; efflux inhibitor; phenylalanyl-arginyll-β-naphthylamide, Polymyxin B

1. Introduction

   Bacterial resistance to antimicrobials is a worldwide problem. The key part of the resistance are efflux pumps (EPs) [1]. AcrAB-ToIC complex in *Salmonella enterica* ser. Typhimurium belongs to resistance-nodulation-division (RND) family of the efflux transporters. EPs of this family are the most important in gram-negative bacteria because they extrude clinically relevant drugs [2]. It is necessary to understand peculiarities of the efflux to overcome this problem. Such knowledge would be essential for more efficient usage of the available antibacterials and for discovery of the new ones.

   Lipophilic cations ethidium (Et+) and tetraphenylphosphonium (TPP+) are well known EP substrates [3-6]. Transmembrane difference of electrical potential (Δψ, negative inside) drives the accumulation of lipophilic cations in the bacterial cytosol. Intracellular components bind low amount of TPP [7] and therefore this cation can be used to estimate the Δψ in bacteria and mitochondria [8]. The specific feature of Et+ is the affinity of this cation to double helix of nucleic acids. Intercalation of Et+ into the double stranded DNA or RNR considerably increases the fluorescence of this compound [9]. Therefore, measurements of Et+ fluorescence is one of the most popular methods for assay of EP activity in bacteria [10-13]. The intensity of fluorescence correlates with the amount of Et+ bound to DNA, and the latter is proportional to the intracellular concentration of this indicator. Measurements of Et+ fluorescence is a convenient method for
analysis of the competition between efflux substrates [14, 15], because it is possible to use several efflux indicators in the same sample. However, Et⁺ fluorescence in wells of the plates proceeds slower than electrochemically assayed binding of this indicator to bacteria [14]. Beside this, a gradual decrease of the fluorescence was observed in experiments with bacteria [13,14] although the cell bound Et⁺ was not destructed [13].

Here we investigated the fluorescence of Et⁺ in suspensions of S. Typhimurium cells at conditions close to ones used for the electrochemical monitoring of interaction of this indicator with the same bacteria. During the potentiometric analysis using Et⁺ selective electrode, samples of the bacterial suspension were taken from the vessels for immediate assays of the fluorescence of this indicator. In parallel to monitoring of Et⁺ interaction with the cells, at the same incubation conditions the electrochemical measurements of TPP⁺ accumulation in bacteria were performed. An efflux inhibitor phenylalanyl-arginyln-β-naphthylamide (PAβN) was used to analyze the role of RND-family pumps in extrusion of the indicators from intact and EDTA or Polymyxin B permeabilized cells. Results of experiments indicated that fluorescence of Et⁺ reaches the saturation level when millimolar concentrations of this cation accumulate the cytosol S. Typhimurium cells. Measurements of Et⁺ interaction with the purified DNA showed that the ability of DNA to bind Et⁺ depends on the composition of incubation medium. The presence of Mg²⁺ or Polymyxin B, or just increase of the ionic strength of the medium considerably decrease Et⁺ binding to DNA.

2. Results

2.1. Simultaneous measurements of Et⁺ and TPP⁺ interaction with S. Typhimurium cells

Results of our previous experiments indicated [14] that fluorescence of Et⁺ in S. Typhimurium suspensions correlate with the amount of this indicator bound to the cells. However, there were considerable differences in the kinetics and the amplitude of changes, registered by the spectrofluorimetric and the electrochemical techniques. This can be explained by the differences in cell incubation conditions during assays in vessels for the electrochemical measurements and in plate wells for the registration of Et⁺ fluorescence. To reduce the impact of incubation conditions, we combined electrochemical and spectrofluorimetric analyses: during the electrochemical monitoring of extracellular Et⁺ concentration, samples of S. Typhimurium suspension were taken to microtubes and the fluorescence was immediately measured. In parallel, the electrochemical monitoring of TPP⁺ concentration in the suspensions of cell incubated at the same conditions was performed (Figure 1).

Addition of S. Typhimurium cells to 100 mM Tris buffer had a very weak effect on Et⁺ fluorescence because of the outer membrane (OM) barrier and the efflux pumps. The wild type (wt) cells accumulated very low amounts of both lipophilic cations (Figure 1a, b), but concentrations of the indicators in the medium considerably decreased and the fluorescence of Et⁺ increased, when efflux pump mutant ΔtolC cells were added (Figure 1c, d). Chelator of the divalent cations EDTA permeabilized the OM and induced accumulation of the indicator cations by the cells of both strains, strongly increasing the fluorescence of Et⁺ in bacterial suspensions. The obtained results revealed that there was no considerable difference between the accumulated amounts of the indicators in S. Typhimurium wt cells, while ΔtolC mutant cells before permeabilization accumulated twice more Et⁺ than TPP⁺ (8 and 3.6 nmol respectively). The total amounts of accumulated indicators in EDTA-permeabilized cells were rather similar (Figure 1).

The most popular inhibitor of RND family pumps phenylalanyl-arginyln-β-naphthylamide (PAβN) increased the accumulation of Et⁺ and TPP⁺, and at the same time stimulated Et⁺ fluorescence. The maximal level of fluorescence was reached when PAβN concentration in wt cell suspension was 128 µM, but only 32 µM were needed for the maximal accumulation and fluorescence in the case of ΔtolC mutant cells (see Figure 1b, d). At higher PAβN concentrations the fluorescence of Et⁺ did not change, although the curves of electrochemical measurements indicated the leakage
of indicators to the medium (see Figure 1d). It should be noted, that Et⁺ leakage was electrochemically registered, when intensity of the fluorescence was rather stable and stayed at the maximum level (see Figure 1d).

Depending on the concentration, polycationic antibiotic Polymyxin B (PMB) permeabilizes the OM and depolarizes the plasma membrane (PM) [16]. Results of the potentiometric measurements demonstrated, that effects of PMB on Et⁺ accumulation in S. Typhimurium cells of both strains were dependent on the presence of PAβN: PMB was able to induce an additional accumulation of Et⁺ in the absence of PAβN (see Figure 1a, c), but a leakage of this indicator from cells was observed in the presence of the efflux inhibitor (see Figure 1b, d). In the absence of PAβN, addition of PMB considerably increased the fluorescence of Et⁺ (see Figure 1a, c).

![Figure 1](image_url)

**Figure 1.** Accumulation of Et⁺ and TPP⁺ ions in S. Typhimurium SL1344 wt (a, b) and ΔtolC mutant (c, d) cells during simultaneous potentiometric and spectrofluorimetric assays. The measurements were performed at 37 ºC in 100 mM Tris/HCl buffer, pH 8.0, containing 0.1 % glucose. The concentrated cell suspensions were added to obtain OD₆₀₀ of 1. EDTA was added to 1 mM, and PMB – to 50 mg/l. The final concentrations of PAβN (μM) are indicated in the figure (b and d). 75 μl samples of the cell suspensions were taken from the vessels to microtubes just before the indicated additions and intensities of the fluorescence were immediately measured.

In the presence of PAβN, Et⁺ fluorescence was at the maximal level and the addition of PMB had no effect on it (see Figure 1b, d). Correlation between the accumulated amount of Et⁺ and the intensity of fluorescence was lost when amount of the cells bound Et⁺ reached ~13 nmol (see Figure 1d). The fluorescence of Et⁺ did not changed when PAβN concentration in the medium increased from 32 to 128 μM, although the cell-accumulated amount of this cation decreased by 34 % (see Figure 1d).

These results indicate, that PMB induces an additional accumulation of Et⁺ when distribution of this ion between cells and the incubation medium is not in equilibrium. Permeabilization of the OM by EDTA and inhibition of the efflux by PAβN led to equilibrium distribution of TPP⁺ across the PM, enabling calculations of the ∆Ψ. In the case of *wt* cells, the maximal level of ∆Ψ, calculated according to the amount of
accumulated TPP⁺ in the presence of 64 μM PAβN, was 209±3 mV. In ΔtolC cells the highest amount of TPP⁺ was accumulated in the presence of 32 μM PAβN and the calculated ΔΨ at these conditions was 230±1 mV. Binding of Et⁺ to DNA does not allow to assay the intracellular concentration of unbound form of this indicator and, correspondingly, to calculate ΔΨ.

2.2. Et⁺ binding to S. Typhimurium cells

According to Rodrigues and colleagues [15], the possibility to use Et⁺ as an fluorescent efflux indicator depends on the concentration of this ion in the medium. Higher concentrations of Et⁺, exceeding the capacity of the efflux pumps, are expected to result in increased accumulations, which, if sufficiently high, can result in ethidium reaching DNA where it can readily intercalate. Choosing higher concentrations of Et⁺ in the medium, we added intact or heat-inactivated cells and monitored the bound amount of Et⁺ (see Figure 2). Additions of EDTA to Tris medium induced accumulation of Et⁺ by the cells, but the permeabilizing effect of this chelator was considerably weaker compared to 3 μM indicator containing medium (see Figure 1a, b). PMB induced a considerably stronger accumulation, but the amount of Et⁺ bound was not high compared to the cells, preliminary treated with Tris/EDTA (see Figure 2b). In the absence of EDTA, PMB was more efficient and the equilibrium distribution of Et⁺ across the cell envelope was achieved in two minutes after PMB addition (see Figure 2b). At lower concentrations, i.e. 24 μM, the PMB-induced accumulation of Et⁺ was not stable and a release of this indicator followed the accumulation (see Figure 2b). After the second PMB addition Et⁺ was only released to the medium. Because of the heat inactivated efflux and permeabilized the OM, after heating envelope of the cells demonstrated a very weak barrier to Et⁺. In spite of the absence cations in the cytosol accumulating ΔΨ, maximum amount of Et⁺ (190 nmol, when the initial concentration is 96 μM) was bound to the heat-inactivated cells (see Figure 2c). The accumulated amount of Et⁺ in heat-inactivated cells was stable. Starting the same initial concentration, the heat-inactivated cells were binding 24-50 % more Et⁺ compared to EDTA and PMB permeabilized cells (compare Figure 2b and c).
Figure 2. Binding of Et⁺ to intact and heat-inactivated S. Typhimurium SL1344 wt cells incubated with different concentrations of this cation. Concentrated intact (a, b) or heat-inactivated (c) cell suspensions were added to 100 mM Tris/HCl buffer, pH 8.0, containing 0.1 % glucose, to obtain OD₆₀₀ of 1. On Y-axes Et⁺ concentrations are presented in % of the initial ones. The initial concentrations of Et⁺ are indicated in the figure. PMB was added to the final concentration of 50 mg/l (a and c) or 50 and 100 mg/l (b), EDTA – to 1 mM. Numbers next to the curves indicate amount (nmol) of Et⁺ bound to the cell.

In order to get more information on the role of envelope barrier in Et⁺ binding to the cells, experiments with intact and Tris/EDTA-permeabilized cells, as well as purified salmon sperm DNA were performed. Cells or DNA were added to 100 or 400 mM Tris buffer containing 3 µM Et⁺. In both buffers intact cells did not bind Et⁺, and slow accumulation of this indicator was observed only after PMB addition. The concentration of Et⁺ in the medium immediately decreased after addition of Tris/EDTA permeabilized cells, but after the fast initial accumulation, a slow release of the indicator was observed,
more clearly expressed in 100 mM buffer (see Figure 3). PMB induced an additional release of the cell bound Et+. In general, the cells in 400 mM buffer accumulated less Et+ compared to ones, incubated in 100 mM Tris. The permeabilized cells in 400 mM buffer accumulated ~7 nmol of Et+, and ~1 nmol was released after PMB addition, but in 100 mM medium these values were around 10 nmol and 3 nmol, correspondingly. In both media after addition of PMB the intact cells accumulated higher amounts of Et+ compared to the preliminary permeabilized cells.

After DNA addition to the medium, a very fast decrease of Et+ concentration to the stable level was observed and in 100 mM Tris the bound amount of Et+ (11 nmol) was considerably higher than in 400 mM buffer (~6.5 nmol). In both media PMB released a considerable amount of bound Et+ and the left amount of the indicator bound in 100 mM buffer was even a bit lower than in 400 mM (around 4.6 and 5.6 nmol, correspondingly. The amount of Et+ bound to DNA after PMB addition was very close to the amount left in Tris/EDTA treated cells at the same conditions (see Figure 3).

**Figure 3.** Binding of Et+ to intact or Tris/EDTA-permeabilized *S. Typhimurium* SL1344 *wt* cells and salmon sperm DNA. The experiments were performed at 37 °C in 100 mM (a) or 400 mM (b) Tris/HCl buffer, pH 8.0, containing 0.1 % glucose. Concentrated suspensions of intact or Tris/EDTA-treated cells were added to obtain OD600 of 1. Red curves demonstrate addition of 100 μg of salmon sperm DNA. PMB was added to the final concentration of 50 mg/l. Numbers next to the curves indicate amount (nmol) of Et+ bound.

2.3. Binding of Et+ to DNA in solutions of various composition

To learn more, how Et+ binding to DNA depends on the composition of the medium, we extended potentiometric analysis of this process. During the first experiments we measured amount of Et+ bound to certain amount (100 μg) of DNA, increasing concentration of this indicator in 100 mM Tris buffer. We elucidated, that saturation level was reached in 60 μM and higher concentrations of this cation after binding of 50 nmol of Et+ to 100 μg of DNA (see Figure 4).
Figure 4. Dependence of the amount of Et⁺ bound to DNA on the concentration of this cation in the medium. 100 μg of salmon sperm DNA were added to various concentrations of Et⁺ containing 100 mM Tris/HCl buffer, pH 8. The experiment was performed at 37 ºC.

Continuing experiments with the purified DNA, we explored how changes in the medium composition could affect the binding of Et⁺ to DNA. In experiments with DNA solutions, the initial fluorescence of DNA-bound Et⁺ in 400 mM Tris buffer was ~20 % lower compared to 100 mM Tris/HCl. During the monitoring period the fluorescence gradually decreased and after 25-30 min it was ~20-30 % lower compared to the initial level (see Figure 5a, b). The presence of TPP⁺ or PAβN in DNA solutions had a weak effect on Et⁺ fluorescence. However, in the presence of PMB or Mg²⁺ (also Ca²⁺, data not shown) the levels of fluorescence were considerably lower and very similar in both concentrations of Tris (see Figure 5a and b).

In our previous experiments with S. Typhimurium cells [14], the strongest decrease of Et⁺ fluorescence was observed after addition of PMB to the suspension of S. Typhimurium cells in 400 mM Tris/HCl buffer. Electrochemical Et⁺ measurements showed that binding of this indicator to DNA is a fast process and the amount bound is rather stable. In the medium with 1.2 μM Et⁺ (see Figure 5c), the initial intensity of the fluorescence correlated well with the amount of this cation bound to DNA (Fig. 5a and b). In 400 mM Tris/HCl buffer DNA bound lower amount of Et⁺ than in 100 mM Tris, and PMB or Mg²⁺ in the medium decreased the bound amount. Mg²⁺ or PMB in 100 mM Tris/HCl buffer induced a slight time-dependent release of bound Et⁺ (see Figure 5c). Addition of PMB to DNA solution in 100 mM Tris buffer at the end of experiment, immediately decreased the amount of bound Et⁺. However, effect of PMB addition to DNA solution in 400 mM Tris was weak, as well as addition of this polycationic antibiotic to DNA solution in 100 mM Tris, already containing Mg²⁺ or PMB (see Figure 5c). This demonstrates that ability of PMB to displace Et⁺ depends on the amount of this indicator bound.
Figure 5. Influence of the medium composition on the binding of Et⁺ to DNA. Experiments were performed at 37 ºC in 100 mM (a, c, d) or 400 mM (b, c, d) Tris/HCl buffer, pH 8, at 37 ºC. Intensity of the initial Et⁺ fluorescence in 100 mM Tris/HCl buffer was taken as 100% (a, b). The concentration of salmon sperm DNA was 20 mg/l, Et⁺ - 1.2 μM (a-c) or 60 μM (d). Concentrations of PAβN, Mg²⁺, TPP⁺ and PMB were 250 μM, 10 mM, 125 μM and 100 mg/l, correspondingly. In c, additional 100 mg/l of PMB were added at the end of experiment. However, the binding of Et⁺ to DNA was partly reversible process when initial concentration of this indicator was 60 µM, and a time-dependent release of the initially bound Et⁺ was observed (Fig. 5d). In the presence of PMB in 400 mM buffer, the potential of Et⁺-selective electrode decreased to values, lower than the initial ones, before addition of DNA. These results suggest that some lipophilic cationic compounds were released from salmon sperm DNA in result of interaction with PMB.

3. Discussion

Antibiotics and other to bacteria noxious compounds are pumped out of the cells by energy-dependent transporters, mainly by the proton motive force-driven efflux pumps [1]. The energy-dependent processes in bacteria are sensitive to the conditions of incubation. Different efficiencies and kinetics of the efflux from S. Typhimurium wt cells registered at various laboratories [4,12,14] can be explained by different methods of registration used for the assay and, correspondingly, different cell incubation conditions and procedures used during evaluation of the efflux.

Here we potentiometrically and fluorimetrically analyzed the efflux in S. Typhimurium cells determining the bound amounts of Et⁺ and TPP⁺. In liquid cultures of gram-negative bacteria, the electrochemical real-time monitoring of the efflux using Et⁺ and/or TPP⁺ as the indicators, PAβN - as the efflux inhibitor, and PMB - as the permeabilizer of cell envelope, is a convenient method. Bacterial suspensions are thermostated and constantly stirred during the experiments, reagents can be added, and the samples taken for complementary analysis without any interruption of the registration and any changes of the incubation conditions.

In slightly alkaline media, the proton motive force of bacteria consists in most of ΔΨ [8]. Despite of a high ΔΨ, S. Typhimurium wt cells with the intact OM in 100 mM Tris/HCl buffer accumulate neither TPP⁺, nor Et⁺ ions. However, ΔtolC mutant cells...
slowly bind these indicators in the absence of the OM permeabilizing compounds, but the presence of EDTA or PMB drastically increases rates of influx (see Figure 1). In the absence of TolC, AcrB or other energy-dependent components of RND and ABC family transporters extrude the lipophilic cations to the periplasm, instead of the incubation medium, as wt cells do. More efficient accumulation of Et\textsuperscript{+} by ΔtolC cells before permeabilization of the OM can be explained by higher selectivity to TPP\textsuperscript{+} of EPs, expressed in the absence of TolC. At low concentrations blocking the activity of RND family pumps, PAβN increases the accumulation of indicator cations and stimulates Et\textsuperscript{+} fluorescence. The maximum accumulation of the indicators and the maximum fluorescence of Et\textsuperscript{+} in ΔtolC mutant cells was achieved at lower PAβN concentration than for wt cells (see Figure 1b, d). This difference clearly indicates that inhibition of the efflux PAβN must cross the OM, and this crossing is more efficient in the case of mutant. The same tendency was observed also for ΔAcrB cells [17].

The altered extrusion of Et\textsuperscript{+} ions from cells, lacking the major EP complex MexAB–OprM, was observed in experiments with 

*Pseudomonas aeruginosa* cells [13,18]. Xu et al. [18] observed that intensity of fluorescence of cells with an inactive MexAB-OprM pump decreased below the fluorescence in buffer solution, when Et\textsuperscript{+} accumulation in the cells reached some “critical” values. This was explained by degradation of Et\textsuperscript{+} in MexAB-OprM mutant cells or the efflux through the assembly of unknown efflux transporters [18]. The authors were considering ethidium bromide as non-dissociated neutral molecule, entering the viable bacterial cells through passive diffusion. Possibility to measure Et\textsuperscript{+} by ion-selective electrode indicates that ethidium bromide is dissociated, and near Nernstian behavior of the electrode shows that the degree of dissociation is close to 1. Babayan and Nikaido [13] extracted the cell-accumulated Et\textsuperscript{+} and presented evidences that this indicator is not destroyed by the cells after accumulation. It was concluded that self-quenching is the main reason of the decreased intensity of fluorescence. Our observations are in agreement with the results from Nikaido group, that the accumulated Et\textsuperscript{+} is not destructed by the cells. The self-quenching could be due to the reduction of affinity of DNA to Et\textsuperscript{+} after the binding of higher amount of this indicator. It is possible that PMB also more efficiently displace this indicator from the binding sites when the high amount of Et\textsuperscript{+} is bound.

Results of our experiments with pure DNA (see Figure 5) suggest that structural and/or functional changes inside the cells during the incubation period alter the DNA bound amount of Et\textsuperscript{+}: When binding equilibrium is reached in DNA solutions containing excess of Et\textsuperscript{+}, one Et\textsuperscript{+} cation is bound for every five nucleotides in DNA, and one per ten nucleotides in RNA molecules [19]. There could be several reasons of fluorescence quenching. According to Hayashi and Harada [20], intercalation of Et\textsuperscript{+} lengthens and unwinds DNA, and the isotherm of intercalation shows negative cooperativity between adjoining Et\textsuperscript{+} molecules. On the other hand, a concomitant drop in the intensity of fluorescence without a change in the amount of Et\textsuperscript{+} bound to DNA was observed with increasing amount of the unbound Et\textsuperscript{+} [21]. In our experiments comparison of Et\textsuperscript{+} binding to purified DNA at low (1.2 μM) and high (60 μM) concentrations of this ion in the medium revealed, that the release of accumulated Et\textsuperscript{+} to the incubation medium could be the main reason of the decrease in fluorescence at high concentrations of this ion (see Figure 5).

1 mg dry weight of *Escherichia coli* cells contains around 100 nmol of DNA base [22]. Considering that size S. Typhimurium chromosome (4,857 kbp [23]) is like the one of *E. coli* (4,639 kbp [24]), in our experiments 5 ml of S. Typhimurium suspension contained ~150 nmol DNA bases. Maximum intensity of the fluorescence was achieved when amount of the bound Et\textsuperscript{+} was around 20 nmol. It means, that Et\textsuperscript{+}/DNA ratio is less than 0.2, even when we consider also the RNA bases (about 20 nmol in 0.03 mg dry weight cells [22]).

We start to register the increase in fluorescence of Et\textsuperscript{+} simultaneously with the potentiometrically observed influx of this indicator into the cells. It is also indicated that primary binding occurs up to a bound Et\textsuperscript{+} to DNA nucleotide ratio of 0.20-0.25 and if the
Et/DNA ratio is below 0.14 there are no changes in the spectrum [25]. However, we have not detected any threshold for increase in Et+ fluorescence after this indicator binding to DNA. It is possible, that the increase in Et+ fluorescence starts before it intercalates into DNA. Et+ binding to DNA is the ionic strength, as well as medium composition dependent, and the presence of Mg2+ in the medium decrease the binding of Et+ to DNA and the fluorescence of this compound (see Figure 5). The decrease of Et+ fluorescence during the incubation with DNA was observed also in the absence of PMB or Mg2+ (see Figure 5 a, b). This could be because of DNA-ethidium complexes become better accessible to water which is highly efficient fluorescence quencher [26], and removal of water increases the fluorescence.

There are evidences of additional non-intercalative, less fluorescence-enhancing sites, involving electrostatic binding to nucleic acids. The enhancement of Et+ fluorescence observed due to electrostatic binding to DNA or to hydrophobic solvents is attributed to a reduction in the excited state proton transfer rate [19]. However, this secondary binding occurs only at low ionic concentrations (i.e., 0.01 M), and when binding at the primary site is saturated. Low ionic strength conditions are not typical to cytosol of viable bacteria, although it could happen after depolarization of the cells, when Et+ fluorescence in bacterial suspensions is measured in low ionic strength media [11]. However, we must be careful in interpreting these results. Our results indicate (see Figure 2, also [14]) that heat killed cells bind considerably more Et+ than intact cells, despite very strong accumulation of the cations in the cytosol by membrane potential. These data are in agreement with [27], that fluorescence of metabolically inhibited cells never exceeded more than 50% of the dead cells values. Most probably, after cell death the complex nature of bacterial chromosome [28] is lost and relative amount of Et+ accessible double-stranded DNA considerably increases.

According to Rodrigues and colleagues [16], at concentrations lower than 3 μg/ml (7.5 μM) Et+ does not bind to DNA, but already is a substrate of efflux pumps, and such concentrations should be used for studies of the efflux. Results of our experiments indicate that an increase of fluorescence is observed when Et+ concentration in the medium is lower, only 3 μM, and we have not found any concentration threshold for binding of this indicator to DNA. On the other hand, there are ideas that EtBr becomes strongly fluorescent when it gets into the periplasm of Gram-negative bacteria or into the cytoplasm of Gram-positive ones [28].

When cells accumulate high amount of Et+ and fluorescence reaches the maximum level, it stops to correlate with the intracellular concentration of this cation and Et+ loses the role of efflux indicator. At increased, cells depolarizing concentrations, PAβN and PMB cause the release of accumulated Et+. Leakage of TPP+ indicates the total depolarization of the PM, but a considerable amount of Et+ remains inside the cells, most probably, because of the binding to nucleic acids. Our results indicate that the decrease of intracellular concentration of this cation not immediately leads to the decrease of the fluorescence. It looks, that the release of bound Et+ is rather slow process [14]. The fast release of Et+ after additions of PAβN and PMB agrees with [28] that if Et+ is not intercalated between nucleic bases of DNA, it is a subject to extrusion or leakage after depolarization of the plasma membrane, as in our case. When it is intercalated, the binding constant is sufficiently high to keep Et+ from access to the efflux pump systems of the bacterium [19]. The dependence of Polymyxin B induced Et+ binding to the cells on the presence of PAβN in the medium and the kinetics of Et+ binding/release indicate a complex nature of this process. Depending on the concentration used, PMB initially increases permeability of the OM, only, and does not affect the PM of Gram-negative cells [16]. At higher concentrations PMB damages the PM and switches off energy-dependent processes, including efflux. When the cell suspension does not contain PAβN and Et+ fluorescence is not at its maximum, we see the additional Et+ accumulation after PMB addition (see Figure 1a and c). Permeabilization of the OM and switching off the efflux by PMB facilitates Et+ entry into the cytosol and binding to DNA, although depolarization of the PM repeats the ΔΨ-dependent Et+ accumulation-driving component. However, our previous experi-
ments [14] demonstrated, that such Et⁺ accumulation in S. Typhimurium wt cells is only temporary: PMB causes the leakage of Et⁺ after the equilibrium is reached. This means that decreasing fluorescence of Et⁺ after PMB addition is not only because of the self-quenching, but also caused by displacement of Et⁺ by PMB at its binding sites. In the presence of EDTA, effect PMB on Et⁺ binding is weaker, most probably, because of the reduced binding of PMB to the OM due to EDTA-induced release of LPS [29] (compare curves in Figure 2a and 2b).

The presence of PAβN in the cell incubation medium considerably changes the run of events (see Figure 1b and d): depolarizing the PM and causing leakage of TPP⁺, PMB induces release of more than 50% of accumulated Et⁺. How PAβN can so drastically change PMB effects on Et⁺ accumulation? Both compounds have high affinity to LPS, but PMB added after PAβN displaces the latter at the binding sites (Sakalauskaite et al., in preparation). The increased concentration of the free PAβN, also subsequent added PMB, induce a release of accumulated Et⁺. Rather stable maximum intensity of the fluorescence in situation when 50 % of intracellular Et⁺ leaked out supports the idea, that the release of DNA-intercalated Et⁺ is a slow process.

4. Materials and Methods

4.1 Bacteria cultivation and preparation for experiments

Salmonella enterica ser. Typhimurium SL1344 cells of wild type (WT) and ΔtolC mutant strains were obtained from Prof. Séamus Fanning (Institute of Food and Health, University College Dublin, Ireland). Overnight cultures of these cells were grown in Luria-Bertani broth, containing 0.5 % NaCl (Sigma-Aldrich, Munich, Germany), diluted 1:50 in fresh medium, and the incubation was continued until the OD₆₀₀ reached 1.0. The cells were collected by centrifugation at 4 °C for 10 min at 3000 x g (HeraeusTM MegafugeTM 16R, Thermo Scientific, Germany). The pelleted cells were re-suspended in 100 mM Tris-hydroxyaminomethane (Tris)/HCl (Roth, Karlsruhe, Germany), pH 8.0, to obtain ~2×10¹¹ cells/ml. The concentrated cell suspensions were kept on ice until used, but not longer than 4 h. For permeabilization of the outer membrane (OM) during the measurements, ethylene diamine tetraacetic acid (EDTA; Sharlau, Barcelona, Spain), pH 8.0, was added to the final concentration of 1 mM. To permeabilize the OM before measurements, the cells were at 37 °C 10 min incubated in 100 mmol/L Tris/HCl containing 10 mmol/L ethylene diamine tetraacetic acid (EDTA; Sharlau, Barcelona, Spain), pH 8.0, then pelleted and re-suspended as described above. Heat treatment of the cells was performed incubating 1 mL of the concentrated suspension in a 1.5-ml Eppendorf tube for 10 min in a boiling water bath.

4.2 Potentiometric measurements

TPP⁺ and Et⁺ concentrations in the incubation media were potentiometrically monitored using selective electrodes as described previously [14,17]. While assembling TPP⁺ and Et⁺-selective electrodes the sensors were filled with 0.1 mM TPP⁺ chloride (Fluka, St. Gallen, Switzerland) or Et⁺ bromide (Acros Organics, New Jersey, USA) solutions in 100 mM NaCl and connected to internal Ag/AgCl half-cell electrodes. In between measurements the sensors were stored dry at room temperature.

The thermostated and magnetically stirred glass vessels were filled with 5 ml of 100 or 400 mM Tris/HCl, pH 8.0, containing 0.1 % glucose. After calibration of the electrodes the concentrated cell suspension was added to obtain an OD₆₀₀ of 1, or shared salmon sperm DNA stock solution was added to the final concentration of 20 mg/l. We used the electrode potential-amplifying system with an ultralow-input bias current operational amplifier AD549JH (Analog Devices, Norwood, MA, USA). The data acquisition system PowerLab 8/35 (AD Instruments, Oxford, UK) was used to connect the amplifying system to a computer. The agar salt bridges were used for indirect connection of the Ag/AgCl reference electrodes (Thermo Inc.; Orion model 9001) to cell suspensions or
DNA solutions in the vessels. The measurements were performed simultaneously in 2–4 reaction vessels. The ΔΨ values were calculated as described previously [30], assuming that OD600 1 correspond to 8.3x10^8 cells/ml, 2.3x10^9 cells correspond to 1 mg of dry mass, and the intracellular water volume of S. enterica is 1.1 ml/g of dry mass. The representative sets of curves from 3-5 independent series of measurements are presented in figures.

4.3 Fluorescence measurements

Single tube measurements were performed using microtubes with 75 µl of samples taken from vessels for electrochemical measurements. Intensity of Et⁺ fluorescence was measured by Modulus™ Single Tube Reader (Turner BioSystems, Inc., USA) using green filter set (excitation 525 nm, emission 580-640 nm).

For evaluation of Et⁺ interaction with sheared salmon sperm DNA (Eppendorf AG, Hamburg, Germany), stock solutions of PAβN hydrochloride, polymyxin B (PMB) sulphate (7730 U of PMB base/mg; Sigma-Aldrich, Munich, Germany), TPP⁺ chloride, or MgCl₂ (Roth, Karlsruhe, Germany) were added to the incubation buffer and mixed. Then salmon sperm DNA solution was added to the final concentration of 20 mg/l, and Et⁺ to the corresponding concentration. The samples were mixed and transferred into 96-well flat-bottom black plates, 100 µl per well. Relative intensity of the fluorescence (excitation 535 nm, emission 612 nm) was monitored in “TECAN GENios Pro™” (Männedorf, Switzerland) plate reader, thermostatting the plate at 37 °C. The plate was shaken 5 s before each registration point.

5. Conclusions

Inhibiting activity of efflux, phenylalanine-arginine-β-naphtylamide (PAβN), the most popular inhibitor of RND family pumps, increases the accumulation of Et⁺ and TPP⁺ in the cells and stimulates Et⁺ fluorescence. Lower concentrations of PAβN are needed to reach the maximal level of accumulation of lipophilic cations in the case of ΔtolC mutant than wt cells. Electrochemical measurements indicate that the main reason of decreasing Et⁺ fluorescence at high concentrations of this ion is the release of accumulated Et⁺ to the incubation medium. However, at maximum level of Et⁺ fluorescence, PAβN as well as PMB cause the leakage of Et⁺ ions from the cytosol depolarizing their plasma membrane, but this leakage is not considerably affecting Et⁺ fluorescence. Et⁺ binding to DNA is medium ionic strength, as well as the composition dependent, and the presence of PMB or Mg²⁺ in the medium decrease it. The intensity of fluorescence reaches the saturation level and stops reacting to the intracellular concentration of this indicator when cells accumulate high amount of Et⁺.

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