Expression and Purification of Transmembrane Protein MerE from Mercury-Resistant *Bacillus cereus*

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Mercury-resistant (Hg⁰) bacteria were isolated from heavy metal polluted wastewater and soil collected near to tanneries of district Kasur, Pakistan. Bacterial isolates AZ-1, AZ-2 and AZ-3 showed resistance up to 40 µg/ml against mercuric chloride (HgCl₂). 16S rDNA ribotyping and phylogenetic analysis were performed for the characterization of selected isolates as *Bacillus* sp. AZ-1 (KT270477), *Bacillus cereus* AZ-2 (KT270478) and *Bacillus cereus* AZ-3 (KT270479). Phylogenetic relationship on the basis of merA nucleotide sequence confirmed 51–100% homology with the corresponding region of the merA gene of already reported mercury-resistant Gram-positive bacteria. The merE gene involved in the transportation of elemental mercury (Hg⁰) via cell membrane was cloned for the first time into pHLV vector and transformed in overexpressed C43(DE3) E. coli cells. The recombinant plasmid (pHLMerE) was expressed and the native MerE protein was obtained after thrombin cleavage by size exclusion chromatography (SEC). The purification of fusion/recombinant and native protein MerE by Ni-NTA column, dialysis and fast protein liquid chromatography (FPLC/SEC) involved unfolding/refolding techniques. A small-scale reservoir of wastewater containing 30 µg/ml of HgCl₂ was designed to check the detoxification ability of selected strains. It resulted in 83% detoxification of mercury by *B. cereus* AZ-2 and *B. cereus* AZ-3, and 76% detoxification by *Bacillus* sp. AZ-1 respectively (p < 0.05).

Keywords: 16S rRNA, Ni-NTA chromatography, pHLMerE, thrombin, Hg-detoxification

Introduction

Mercury pollution is a global problem due to its toxic effects on humans and animals. Anthropogenic sources and activities, e.g., the release of untreated industrial wastewater from chlor-alkali plants, metal mining and the incineration of coal, are increasing mercury pollution day by day in the environment [1]. It is evident that the cytotoxic and neurotoxic effects to humans and animals are caused by inorganic (ionic) and organic forms of mercury [2]. In Pakistan, the maximum release of mercury (kg/year) is estimated as 1,625.11, 2,500, 21,120, 1,071.79, 5,779, and 4,637.55 through extraction processes, cement industries, chlor-alkali industries, intentional uses of mercury in consumer products, waste deposition/land filling and wastewater treatment plants, respectively, indicating 36,898.77 kg of total cumulative release of mercury per year [3].

*Bacillus* sp. RC607, identified for the first time as having chromosomal resistance against mercury, was isolated from a contaminated site, Boston Harbor, USA [4]. In later studies, strain RC607-like sequences were found in 74 *Bacillus* spp. isolated from Minimata Bay sediment, Japan, and also in several Russian environmental bacterial isolates. It was suggested that the possible global distribution of the RC607 mer operon had occurred, as *Bacillus mer* operon-bearing plasmids were found in several strains analyzed in the Russian study [5, 6].

Due to the presence of mercury in the environment and its toxic effects, microorganisms have evolved mercury...
detoxification systems to deal with mercury-containing compounds. Bacteria have the ability to detoxify the toxic form of mercury (Hg2+) into the non-toxic elemental form (Hg0) through a cytoplasmic enzyme, mercuric reductase, which is encoded by the merA gene of mer operon [7]. A lot of research analysis was done on this cytoplasmic enzyme, a member of flavin-containing, NADPH-dependent dithiol oxidoreductase that has been isolated from Gram-positive as well as Gram-negative bacteria [8]. Gram-positive and some Gram-negative bacteria also possess organomercurial lyase encoded by merB gene of mer operon which cleaves the C–Hg bond of many mercury-containing compounds. The enzyme organomercurial lyase confers broad spectrum resistance against mercury compounds which results in the elemental mercury (Hg0) being diffused out of the cell [9].

Bacterial genes involved in mercury resistance are well-organized and well-characterized into genetic elements known as mer operons. The Gram-negative bacterial mer operons contain a broad-spectrum structural gene, merE, that encodes a CH2Hg+ and Hg2+ detoxification system. [10]. The merE gene, which immediately follows the merD at the downstream end of the bacterial mer operon, is frequently found in many narrow and broad-spectrum mer operons [11]. Kiyono et al., [12] have reported the transcription and translation of merE gene into a membrane protein with molecular mass of ~8 kDa in Gram-negative bacteria. Moreover, MerE protein is a methylmercury membrane transporter. This system is involved in the transportation of mercury ions across the cell membrane [12].

For structural and biochemical studies, active and properly folded forms of soluble and transmembrane proteins are essential during expression and purification procedures. For rapid screening of soluble and transmembrane proteins, fast, high-yield production has been developed by recombinant expression systems. Among bacterial, insect, yeast, mammalian and plant protein expression systems, the bacterial expression system is less expensive and fast but has two major problems. The first is that the protein-folding machinery is the least complex and the second is the formation of inclusion bodies (IBs) which are insoluble aggregates of recombinant proteins [13].

Refolding/activity of proteins from IBs is affected by several factors such as solubilization of IBs by denaturants, removal of the denaturant, and assistance of refolding by small molecule additives. The methodologies being used nowadays for refolding aggregated proteins can be divided into two major categories: chromatographic, i.e., solvent exchange by size-exclusion chromatography (SEC), on-column refolding and chaperone-assisted refolding [14] and non-chromatographic for rapid dilution of denaturants from denatured proteins, i.e., dialysis, dilution, chemical additives (NH4)2SO4, amino acids (ArgHCl), glycerol and cyclodextrins [15–17].

In this study, screening, molecular identification and phylogenetic analysis of Hg-resistant bacteria were described on the basis of 16S rDNA ribotyping and merA gene sequence. Following that, recombinant plasmid construction and expression and partial purification by Ni-NTA column of the transmembrane protein MerE involved in the mercury detoxification system were performed. Furthermore, the native MerE protein was purified by size exclusion chromatography (FPLC). Finally, the detoxification potential of Hg-resistant bacteria from Hg-contaminated wastewater was also checked.

**Materials and Methods**

**Chemicals and Expression Vectors**

For DNA restriction, the enzymes HindIII and SacI were purchased from New England Biolabs (www.neb.com) unless otherwise noted, and the oligonucleotides were synthesized by Integrated DNA Technologies (www.idtdna.com). Ligation Kit was purchased from life sciences (www.lifescience.roche.com). For bacterial genomic DNA, a GeneJET Purification Kit and E. coli cells (DH5a Competent) were obtained from Thermo Fisher© (www.thermofisher.com). Bacterial competent cells (E. coli) C43(DE3) were obtained from Lucigen© (www.lucigen.com). For plasmid preparations, small scale a Gel Extraction Kit (Miniprep Kit) and Ni-NTA resin were obtained from Qiagen© (www.qiagen.com). The PCR reagents were purchased from Agilent Technologies© (www.genomics.agilent.com). Sephacryl S-200 column for size exclusion chromatography was purchased from Pharmacia LKB (USA).

**Isolation and Screening of Hg2+ Bacteria**

Mercury-resistant bacteria were isolated from wastewater and contaminated soil samples obtained from the tanneries near Kasur district in Pakistan. The physico-chemical properties such as temperature, soluble salts, organic matter, Hg2+ concentration and soluble N, P, and K, of all samples were checked. To this end, serial dilutions of 102, 103, and 104 were prepared from 1% of the initial water and soil samples. For obtaining the isolated colonies of Hg-resistant bacteria, 100 μl from the 103 and 1004 dilutions were spread on Lauria Bertani (LB) agar plates supplemented with HgCl2 of different concentrations ranging from 1–200 μM. The plates were incubated at 37°C for 24 h. After incubation, the isolated and Hg-resistant colonies were obtained and these were re-streaked on new LB agar plates without HgCl2 for obtaining purified colonies and stored as glycerol stocks at ~80°C.
PCR-Amplification of 16S rRNA and merA Genes

The bacterial isolates AZ-1, AZ-2, and AZ-3 showing high resistance against HgCl₂ were characterized by 16S rDNA ribotyping. Bacterial strain ZA-15 was used as a negative control. The bacterial genomic DNA was extracted by GeneJET Genomic DNA Kit. The universal primers used for the amplification of 16S rRNA gene were: 16S-F (5’ AGAGTTTGATCCTGGCTCAG3’) and 16S-R (5’ AAGGAGGTGATCCAGCAGCGA3’) [18] and for merA gene; merAF : 5’TGCGTGGAACCTGGCTAA3’ and merAR : 5’TTTACCAGCACAGCAAGATA3’ using thermal cycler. The amplification conditions for both genes consisted of initial denaturation at 95°C (5 min) and then 30 cycles of 95°C (1 min), 55°C (1 min), and 72°C (1.5 min), plus an additional cycle at 72°C (5 min) for chain elongation. The amplified products were sent to the Macrogen sequencing core facility in Korea and the obtained sequence results were checked by NCBI nucleotide BLAST and finally submitted to GenBank.

Phylogenetic Relationship

The multiple sequence alignment through ClustalW was used to check phylogeny among bacterial species of the Bacillus genera on the basis of 16S rRNA and merA genes through the neighboring-joining method using MEGA 5 software. The percentage of homology among different clades was checked with a bootstrap test at a value of 1,000 replicas as shown next to the branches. The tree is drawn to scale which represents the nucleotide change. The units of branch lengths and the evolutionary distances are the same and used to infer the phylogenetic tree.

Construction and Expression of Recombinant Plasmid pHLMerE

The merE gene of mercury-resistant isolate Bacillus cereus AZ-2 was amplified using the following primers; F-MerE [5’ ATCTAT AAGCTTATGAAAAACATAATAAAAAGTTCAGGTTGT 3’] and R-MerE [5’ ATATATGAGCTCTATAGGACCTGAC TTCTCTCT GCCATGCAGCA3’]. Bacillus megaterium MB1 was used as a positive control for both merA and merE genes [9]. The amplified DNA of 270 bp was cloned in HindIII-Sacl-cleaved pHLV expression vector. The designed recombinant plasmid was transformed into DH5α competent cells. The pHLMerE (3,842 bp) was confirmed by restriction with the same enzymes and visualized on agarose gel electrophoresis. The nucleotide sequence of merE gene was also confirmed by a DNA sequencing facility provided by the University of California, San Diego (UCSD), USA. The recombinant plasmid pHLMerE was then transformed into E. coli competent cells C43(DE3) as described in our previous study [19].

Five μl of clone C43(DE3):pHLMerE from glycerol stock was inoculated in 5 ml of LB medium containing 50 mg/l of carbenicillin and incubated for 5 h at 37°C. After incubation, 1 ml was taken from 5 ml culture and inoculated in 50 ml of M9 minimal medium (Na₂HPO₄ 7.0 g, KH₂PO₄ 3.0 g, NaCl 0.5g, CaCl₂ 0.1 mM, MgSO₄ 1 mM, thiamine 50 mg, D-glucose10 g, (NH₄)₂SO₄ 1 g and 50 mg/l of carbenicillin for 1 L) and kept in shaking incubator at 37°C for overnight with 300 rpm. An overnight 50-ml culture was poured into 450 ml of the same M9 medium and allowed to grow at the same growth conditions. When the optical density of the culture reached at O.D₆₀₀ ~ 0.6, isopropyl-β-D-thiogalactoside (IPTG) with final concentration of 1 mM was added for the production of His-tag_trpΔLE_thrombin site_MerE fusion protein and shaking was continued in the same incubator for another 7 h. The cell pellet harboring recombinant plasmid was obtained by centrifugation at 7,000 × g for 30 min at 4°C.

Purification of the Recombinant Protein by Ni²⁺-NTA Column

The purification of the recombinant protein by Ni²⁺-NTA column was performed by using a method as described in our previous studies with some modifications [20–22]. The cell pellet was re-suspended in 30 ml lysis buffer consisting of 50 mM Tris hydrochloride, 15% glycerol (v/v), 1 mM NaCl, with pH 8.0 and incubated at room temperature for 10 min. The collected cell lysate was disrupted by sonication for 5 min on ice (5 sec ON and 10 sec OFF) and spun down at 17,000 × g for 30 min at 4°C. At this stage, purified inclusion bodies contained presumably 90% of MerE recombinant protein. The inclusion bodies were solubilized in the binding buffer consisting of 20 mM Tris hydrochloride, 500 mM NaCl, 6 M GdnCl, and 5 mM imidazole at pH 8.0 by tip sonication. The cell lysate was centrifuged again at 19,000 × g for 1 h to remove protein and lipid-associated cell debris. The Ni²⁺-NTA histidine-binding resin was pre-equilibrated with binding buffer and then the supernatant was loaded onto the column. The resin was extensively washed by washing buffer consisting of 20 mM Tris hydrochloride, 500 mM NaCl, 6 M GdnCl, and 50 mM imidazole with pH 8.0. The polyhistidine-tagged fusion protein was eluted with elution buffer consisting of 20 mM Tris hydrochloride, 500 mM NaCl, 6 M GdnCl, and 500 mM imidazole with pH 8.0. The eluted polyhistidine-tagged fusion protein MerE was monitored by SDS-PAGE and then concentrated to 30 ml by using a YM10 filter membrane containing Amicon stirred concentrator cells. The eluate was dialyzed against ddH₂O in a 10 kDa MWCO dialysis membrane with four water changes until the protein precipitated out of the solution and then lyophilized.

Thrombin Cleavage of the Fusion Protein

The cleavage of the fusion partner by thrombin was performed by the following method described by Das et al. [23] with some modifications. The lyophilized polypeptide was re-suspended in 30 ml of binding buffer consisting of 20 mM Tris HCl, 500 mM NaCl, 1% SDS, 10 mM imidazole, and 0.1% TCEP with pH 8.0, and loaded on to a Ni²⁺-NTA column which was pre-equilibrated by 20 bed volumes of thrombin cleavage buffer consisting of 20 mM HEPES, 250 mM NaCl, 0.1% hexadecylphosphocholine, pH 8.0. The fusion partner trpΔLE was cut-off by thrombin (1000 units) by incubating for 2 h at room temperature. The column was then washed with 10 bed volumes of washing buffer consisting of 20 mM Tris HCl, 0.5% dodecylphosphocholine, 50 mM NaCl, and 20 mM imidazole with pH 7.3). The native MerE and fusion
partner were eluted with 3 bed volumes of elution buffer consisting of 20 mM HEPES, 0.5% dodecylphosphocholine, 500 mM imidazole, and 50 mM NaCl with pH 7.3. Approximately 2–3 mg of MerE protein was obtained from each 10 ml elute and lyophilized. The cleaved polypeptide was washed and precipitated by using a 3,500-kDa MWCO dialysis membrane against ddH2O and then lyophilized.

**Purification of MerE by FPLC**

The purification of MerE protein by FPLC was performed as described by Amin and Latif [22]. The cleaved and lyophilized MerE protein was solubilized in 5 ml phosphate SDS buffer containing 100 mM Na2HPO4, 20 mM DTT, 4 mM SDS, 1 mM EDTA, and 1 mM NaN3 with pH 8.2. The aggregated protein was disturbed by extensive bath sonication which presumably incorporated SDS into micelles for maintaining stable tertiary structure. For the separation of leader peptide + His-tag from the pure MerE protein, a Pharmacia FPLC system was used containing a Sephacryl S-200 column. The column was pre-equilibrated by phosphate-SDS buffer and 5 ml of sample was loaded. Fractions containing protein MerE were pooled and concentrated down to 30 ml using a 3,500-kDa MWCO membrane of Amicon stirred cells. The pure MerE protein was then dialyzed against ddH2O with 30 ml using a 3,500-kDa MWCO dialysis membrane against ddH2O and then lyophilized.

**Detoxification of Hg²⁺ by Selected Bacteria at a Lab Scale**

Selected bacterial strains (AZ-1, AZ-2, AZ-3, and ZA-15) were inoculated in four flasks containing 30 ml of wastewater containing 30 μg/ml of HgCl₂ separately. The detoxification efficiency of mercury was determined by inoculating 1.5 ml of overnight free cell culture (O.D. 2.0 at 600 nm) in four flasks (4th flask was taken as control) in triplicates and incubated at 37°C for 24 h at 120 ×g of agitation. After incubation, cultures were centrifuged at 14,000 ×g for 15 min and the supernatant was separated. The pH of supernatant for each strain was adjusted to 0.4 by adding concentrated H2SO4 and then transferred to a separating funnel. To the cooled solution, 4 ml of 6N acetic acid and 2.5 ml chloroform were added and vigorously shaken for 1 min. The phases were allowed to separate completely before the chloroform layer was taken out and discarded. Freshly prepared 5 ml of 0.001% dithizone solution was added in the remaining solution of each strain and again shaken vigorously for 1 min. The layers were allowed to separate and cotton was placed in the tips of separating funnels to elute the dithizone-mercury complex. In the absence of mercury (Hg²⁺) no significant color was extracted into dithizone, while in the presence of Hg²⁺, an orange color was observed in the organic layer, and the color became more pronounced with higher concentration of Hg²⁺. The optical density (O.D) of water–free chloroform extracts was determined at 500 nm against reagent (dithizone) blank to estimate the detoxification of Hg²⁺ [24, 25].

**Statistical Analysis**

All statistical analyses i.e., mean, standard deviation and analysis of variance (ANOVA) of this lab scale mercury detoxification experiment were performed by using SPSS V.20 software.

**Results and Discussion**

The interaction between heavy metals and heavy metal-resistant microorganisms has been determined by bacterial transformation and the conversion of metallic ion forms to elemental forms by reduction [26]. Some of these microorganisms have shown promising activity to decrease mercury in contaminated environments. In the present study, three bacterial strains out of 30 were selected on the basis of resistance to higher HgCl₂ concentrations (MIC: up to 1–200 μM) and two strains, ZA-15 and MB1 were used as negative and positive controls respectively (Table 1).

The physicochemical parameters of selected samples at the time of collection were checked as temperature 24°C, pH 8.2, soluble salts 0.7 g/kg, organic matter 8.4 g/kg, Hg²⁺ concentration 10–20 μg/ml, soluble N, P and K 53.5, 3.5, and 45 mg/kg. In literature, 100 μM, 50 ppm and 100 μg/ml were found to be the MIC for mercury in Bacillus spp. [27–29].

Mercury-resistant bacterial isolates (AZ-1, AZ-2, and AZ-3) were characterized by 16S rDNA ribotyping (≈ 1.5 kb) and identified as Bacillus sp. (KT270477), Bacillus cereus (KT270479), and Bacillus cereus (KT270479), respectively. Other close matches to Bacillus sp. (KT270477) included Bacillus anthracis HQ200405 and Bacillus sp. JN593078 and GU566355. The bacterial isolates also showed similarity (98%) among themselves. In the same clade, B. cereus AZ-1 and AZ-2 showed 98% similarity with already reported

<table>
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<th>Isolates</th>
<th>Growth against HgCl₂ (μg/ml)</th>
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<td>10</td>
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<td>AZ-1</td>
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<td>ZA-15 (-ve control)</td>
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<td>MB1 (+ve control)</td>
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B. thuringiensis KP306751, Bacillus sp. JQ691603 and B. cereus KP407139 and LK392517 (Fig. 1).

Phylogenetic analysis of selected bacterial strains on the basis of merA gene showed 100% homology with already reported sequences of merA genes in B. cereus NG034957, B. megaterium NG035056, Bacillus sp. LC015492, Paenibacillus sp. LC015492, 98% in B. macroides Y09906, 89% in Bacillus sp. RC607 A13RF77 and Clostridium butyricum AB024961 and 88% in B. cereus AB066362 and also with themselves (Fig. 2).

The gene construct containing merE was designed by modifying previously reported pHLLVpu plasmid constructed by pHHLV_ML vector which is derivative of the commonly used vector pET-3a [30]. The vector consisted of 9-His-tag tryptophan-leader-Vpu fusion on plasmid encoding carbenicillin resistance. In modified vector, Vpu gene was replaced by merE gene by designing specific primers as shown in Fig. 3A. In the gene construct, His-tag and Trp leader peptide sequences were inserted for the ease of purification and the formation of inclusion bodies respectively. The recombinant plasmid was transformed into competent DH5α and amplified in LB medium. The plasmid was then transformed into E. coli C43(DE3) cells for producing high yield of MerE recombinant protein. The fusion protein was found non-toxic to C43(DE3) cells and expressed at levels up to approximately 20% of total cellular protein.

After expression, the separation and purification of MerE were performed. Briefly, the inclusion bodies containing recombinant/fusion protein were separated from the total cell lysate by centrifugation. The fusion protein was partially purified by nickel affinity chromatography and dialyzed to decrease gradually the denaturant concentration for efficient protein refolding. The data in Fig. 4 illustrated the expression and isolation of inclusion bodies of full-length recombinant MerE by Ni²⁺-NTA column. During dialysis, the protein refolding activity of chemically denatured protein was increased sufficiently due to the decrease in denaturant concentration. The protein concentrations remained almost constant during the refolding process. The rate of refolding to the native (active) structure was also increased by increasing dialyzing

**Fig. 1.** Evolutionary relationships of different Bacillus sp. on the basis of 16S rDNA ribotyping.

**Fig. 2.** Evolutionary relationship of different Bacillus sp. on the basis of merA gene was inferred using the Neighbor-Joining method with the sum of branch length 0.53623375. The branch lengths and the evolutionary distances used to infer the phylogenetic tree are in the same units with the bootstrap test value 500. The analysis involved 17 nucleotide sequences and codon positions included were 1st+2nd+3rd+Noncoding. A total of 1279 positions were in the final dataset with no gaps and missing data. Evolutionary analyses were performed by MEGA 5 software [42].
time along with the decrease in concentration of denaturant [15, 31].

After elution by nickel column, the targeted fusion protein was solubilized in thrombin cleavage buffer and again bound to the affinity column for cleaving native MerE from fusion partner (TrpΔLE) and dialyzed. Lane 2 in Fig. 5A showed the cleavage of fusion protein and lane 3 contained the impurified fraction of MerE while lanes 4–7 showed the purified fractions of native MerE after size-exclusion chromatography (FPLC). Fig. 5B showed the peaks of MerE and TrpΔLE in FPLC equipped with a Sephacyrl S-200 HR column. The fully denatured proteins by GndCl or urea, usually have a high tendency to aggregate due to the exposure of buried hydrophobic residues. The gel filtration column used for SEC/FPLC restricts the aggregation by physically isolating the molecules.
and allows the successful recovery of biologically active (refolded) proteins. Using this approach, Neely et al. [32] successfully obtained refolded/active $\beta_{1b}$ subunit of a calcium channel. In a similar way, SEC refolding was successfully used to obtain biologically active forms of the $\alpha_5$ subunit of the 20S proteasome from human [33]. In another example, Ouellette et al. [34] used denatured inclusion bodies of human interleukin (IL)-7 to load onto a Superdex 200 column and obtained its biologically active (refolded) form.

This was a first study for reporting the plasmid construction, expression and purification of MerE protein from Gram-positive Bacillus cereus AZ-2. Previously, Sone et al. [35] had reported the expression and preliminary purification by Nickel agarose column of MerE protein from Gram-negative bacteria. Park SH, Opella SJ [36] and Lu et al. [37] had also performed NMR structural studies of purified samples of MerE from Gram-negative bacteria. The gene and protein sequences of MerE of both types of bacteria were totally different as shown in Fig. 6. In Gram-positive bacteria, the polypeptide consisted of 89 amino acids with internal methionine residues at different sites while in Gram-negative bacteria, 78 amino acids containing peptide had only single methionine at N-terminal. Due to the presence of internal methionine residues, the gene construction, expression and cleavage of recombinant protein for obtaining the native sequence of MerE varied

![Fig. 4. SDS-PAGE of MerE fusion protein.](image)


![Fig. 5. (A) SDS-PAGE of native protein MerE and fusion partner TrpΔLE after cleavage by thrombin. L: ladder (kDa). C: cleavage by thrombin. F1-F5: fractions of native MerE from Sephacryl S-200 column. (B) SEC chromatogram showing peaks of MerE and TrpΔLE.](image)

![Fig. 6. Topological alignment of all amino acids of MerE separated according to the predicted hydrophobic elements generated by TMHMM software.](image)
Expression and Purification of MerE

It was reported that certain species such as cyanobacteria, *Klebsiella aerogenes*, *K. aerogenes* NCTC418 and *Pseudomonas putida* were involved in bioreduction of toxic mercury compounds under laboratory conditions [38, 39]. Nakamura K, Hagimine M, Sakai M and Furukawa K [40] reported the decrease of mercury from various mercury compounds like mercuric chloride (HgCl\(_2\)) 88.9%, methyl mercuric chloride (MeHg) 95.4%, ethylmercuric chloride (EtHg) 83.8%, thimerosal 91.9%, fluorescein mercuric acetate (FMA) 74.6%, phenylmercuric acetate (PMA) 5.7%, and p-chloromercuric benzoate (p-CMB) 92.3% by *Pseudoalteromonas haloplanktis* strain M-1. The biosorption capability of the immobilized *B. cereus* cells was described by Sinha A, Pant KK, Khare SK [41] as 104.1 mg/g Hg\(^{2+}\) at 30 °C and pH 7.0, biomass concentration of 0.02 g/l and contact time of 72 h. A lab scale experiment was designed to analyze the detoxification potential of mercury by selected bacterial strains. Results indicated 23 (76%), 25 (83%), 25 (83%) and 27 (90%) μg/ml detoxification of Hg\(^{2+}\) out of 30 μg/ml (100%) by selected bacterial strains as *Bacillus* sp. AZ-1, *B. cereus* AZ-2, *B. cereus* AZ-3 and *E. cloacae* ZA-15 respectively (Fig. 7). In conclusion, phylogenetically similar and mer operon-harboring mercury-resistant bacterial strains *B. cereus* AZ-2, *B. cereus* AZ-3 can be used for the detoxification of mercury from industrial effluent.

**Acknowledgments**

Aatif Amin cloned the plasmid, expressed and purified the MerE protein under the supervision of Stanley J. Opella. Arslan Sarwar and Mushtaq A. Saleem helped to execute mercury detoxification experiments and then also wrote the draft. Zakia Latif as a my supervisor, proofread the manuscript.

**Conflict of interest**

The authors declare that they have no conflict of interest.

**References**


