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Journal of Biomedical Science 2010, 17:84 doi:10.1186/1423-0127-17-84

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ISSN	1423-0127
Article type	Research
Submission date	13 June 2010
Acceptance date	9 November 2010
Publication date	9 November 2010
Article URL	http://www.jbiomedsci.com/content/17/1/84

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A role of *ygfZ* in the *Escherichia coli* response to plumbagin challenge

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Abstract

Plumbagin is found in many herbal plants and inhibits the growth of various bacteria. Escherichia coli strains are relatively resistant to this drug. The mechanism of resistance is not clear. Previous findings showed that plumbagin treatment triggered up-regulation of many genes in E. coli including ahpC, mdaB, nfnB, nfo, sodA, yggX and ygfZ. By analyzing minimal inhibition concentration and inhibition zones of plumbagin in various gene-disruption mutants, ygfZ and sodA were found critical for the bacteria to resist plumbagin toxicity. We also found that the roles of YgfZ and SodA in detoxifying plumbagin are independent of each other. This is because of the fact that ectopically expressed SodA reduced the superoxide stress but not restore the resistance of bacteria when encountering plumbagin at the absence of ygfZ. On the other hand, an ectopically expressed YgfZ was unable to complement and failed to rescue the plumbagin resistance when sodA was perturbed. Furthermore, mutagenesis analysis showed that residue Cys228 within YgfZ fingerprint region was critical for the resistance of E. coli to plumbagin. By solvent extraction and HPLC analysis to follow the fate of the chemical, it was found that plumbagin vanished apparently from the culture of YgfZ-expressing E. coli. A less toxic form, methylated plumbagin, which may represent one of the YgfZ-dependent metabolites, was found in the culture supernatant of the wild type E. coli but not in the $\Delta ygfZ$ mutant. Our results showed that the presence of ygfZ is not only critical for the *E coli* resistance to plumbagin but also facilitates the plumbagin degradation.

Background

5-Hydroxy-2-methyl-1,4-naphthoquinone (5-hydroxyl-2-methyl-naphthalene-1,4-dione, IUPAC), known as plumbagin, is found in many herbal plants. It has been found to have antibacterial [1], antifungal [2], anticancer [3], and antimutagenic activities [4]. Similar to redox-cycling chemicals such as paraquat and menadione (vitamin K3), plumbagin generates superoxide or reactive oxygen species that trigger the oxidative stress response [5]. The genes controlled by oxyR and mar/sox are known as the major regulons responsive to the oxidative stress in bacteria. In subtle differences, oxyR is robustly activated in response to oxidative stress [6] while mar/sox are activated by inhibition of the MarR repressor [7] and by oxidization of SoxR [8, 9]. Currently, several lines of evidence suggest that the toxicity of plumbagin is not simply due to production of reactive oxygen species. Plumbagin modifies the lactose carrier, which results in a loss of galactoside-binding ability [10]. Furthermore, high concentration of plumbagin (greater than 100 µM) disrupts bacterial respiratory activity through inactivation of NADH dehydrogenase [11].

In a previous proteomic analysis, plumbagin has been shown to up-regulate the expressions of many proteins belonging to the *oxyR* and *mar/sox* regulons in *E. coli*, such as AhpC, MdaB, NfnB, Nfo, SodA, YggX and YgfZ [12]. The function of AhpC, alkyl hydroperoxidase C, is to detoxify endogenous and exogenous peroxides [13]. MdaB (modulator of drug activity B) and NfnB (a predicted oxygen insensitive NAD(P)H nitroreductase) are members of the mar regulon [14, 15]. The gene nfo encodes endonuclease IV, which participates in the repair of H₂O₂-induced DNA lesions [16]. SodA, a manganese-containing superoxide dismutase, scavenges and coverts O₂⁻ to H₂O₂ [17]. YggX, an iron-binding protein that is involved in intracellular Fe(II) trafficking, is induced by oxidative stress in order to protect DNA from damage [18, 19]. Genes nfo, sodA, yggX and ygfZ are regulated by marbox sequences that are evidently driven by SoxS [12, 20, 21]. Genetic deletion of ygfZ in E. coli has been reported to affect the bacterial tRNA modification and initiation of chromosomal replication [22]. Analysis of the crystallized structure of YgfZ has suggested that the protein may participate in one-carbon metabolism that involves folate or folate derivatives [23]. While ygfZ is regulated by SoxS [12], the role of YgfZ in bacteria facing the challenge of plumbagin remains unresolved.

Theoretically, the above types of responses are triggered in order to resolve an immediate threat of the stress. In such circumstances, plumbagin-responsive genes are likely to be involved in either eliminating the toxicity of the chemical or repairing the damage caused by the drug. It is not known whether any of these plumbagin-responsive genes are directly involved in the detoxification of plumbagin. In this study, we identified the genes that are required for *E coli* to resist plumbagin by analyzing the growth of various *E. coli* mutants in the presence of plumbagin.

We demonstrated that, among these plumbagin-responsive genes, *ygfZ* and *sodA* are the ones required for counteracting plumbagin toxicity. Furthermore, we provided evidence that YgfZ is needed for the degradation of plumbagin. A methylated and less toxic compound found in the media may represent one of the degradation products. Molecularly, Cys228 in the conserved region of *E. coli* YgfZ is essential for this anti-plumbagin activity.

Methods

Bacterial strains, chemicals, and culture conditions

Mutants of *E. coli* K12 with single gene disruption at *ahpC*, *marA*, *mdaB*, *nfnB*, *nfo*, *sodA*, *soxS*, *soxR*, *ygfZ*, *yggX*, *and lpp*, respectively, were gifted from Dr. Hirotada Mori at Nara Institute of Science (Japan), and the parental strain BW25113 was used as the wild-type strain in all comparison experiments. The genotype of BW25113 is $lacI^{q}$ $rrnB_{T14} \Delta lacZ_{WJ16}$ *hsdR514* $\Delta araBAD_{AH33} \Delta rhaBAD_{LD78}$. *E. coli* K-12 JM109 was used as the cloning host. Bacteria were cultured in the Luria-Bertani (LB) broth (Difco) at 37 °C with vigorous rotating (150 rpm, Firstek Scientific S306R). Plumbagin (Sigma) was dissolved in dimethyl sulfoxide as a 10 mg/ml stock.

Primers and expression plasmids

Primers used in this study are listed in Table 1. Plasmid pMH-ygfZ has been described

previously [12]. To induce the expression of SodA by IPTG, pQE-sodA was constructed by amplifying the *sodA* fragment from the *E. coli* genomic DNA with primers PsodAF and PsodAR; the amplified fragment was then digested with *Ba*mHI and ligated into pQE60 (Qiagen) previously digested with the same enzyme. Similarly, pQE-ygfZ was constructed by PCR amplification of the *ygfZ* fragment using primers PygfZF and PygfZR (Table 1), which was followed by insertion of the fragment into *NcoI/BgI*II-digested pQE60. In this way, two plasmids were created to express the SodA and YgfZ proteins, respectively, both with hexahistidine (His_{x6}) tagged at the C-termini. pQE-Kp_ygfZ, and pQE-Mtb_Rv0811c were generated by a similar strategy, except that the genomic DNAs used for amplification were extracted from *Klebsiella pneumoniae* and *Mycobacterium tuberculosis*, respectively, and the primer pairs separately used were PkpygfZF/PkpygfZR and PRv0811cF/PRv0811cR (Table 1).

Site-directed mutagenesis and deletion

Mutagenesis was carried out by PCR. Construction of a variant of *E. coli* YgfZ (K226A) with Lys at residue 226 replaced with Ala was given as an example. In brief, *ygfZ* in pQE-ygfZ was first PCR amplified separately with two primer pairs, PQEF/PygfZK226AR and PygfZK226AF/PQER (Table 1). Due to the design of the sequences of PygfZK226AR and PygfZK226AF, the two so-amplified PCR products have overlapping termini where the mutated codon is embedded. After mixing and melting the two PCR products, the overlapping regions

were annealed to each other. After this, primers PQEF and PQER were added and PCR amplification was carried out to give a fragment containing the full-length *yg/Z* with the designated K226A mutation. The amplicon was then digested with *NcoI* and *BglII*, and ligated into a similarly restricted pQE60 vector to give pQE-ygfZK226A. All the other substitution-mutation plasmids that encode the mutated YgfZ variants were constructed in a similar way by selecting appropriate primer pairs (Table 1).

Immunoblotting

Total protein lysates were prepared as described previously [12]. Electrophoretically separated proteins blotted on nitrocellulose membrane were analyzed by Western blotting using specific antibodies. Anti-YgfZ antibody was generated by immunizing mice with nickel-column purified His_{x6}-YgfZ. Rabbit anti-His_{x6} antibody (Bethyl) was used for detecting His_{x6}-tagged proteins. Mouse monoclonal anti-DnaK has been described previously [24]. Horseradish peroxidase-conjugated secondary antibodies (Sigma) were used to detect the primary antibodies bound on the membrane. The antibody-bound blots were finally developed using chemiluminescence reagent (Perkin-Elmer) and the signals were obtained by exposing the membrane to X-ray film (Fuji).

Inhibition zone analysis

Overnight cultures of the various bacterial strains in LB broth were diluted 100-fold into fresh

LB broth and grown with aeration at 37 °C for 2 h. The turbidity of the cultured bacteria was adjusted to OD_{600} at 0.4 and the resulting bacteria were spread on Mueller-Hinton (MH) agar (Difco) plates using sterile cotton buds. Filter paper discs (8 mm in diameter) containing various chemicals at appropriate amounts were applied to the top of the agar. The diameters of inhibition zones around the filter discs on the plates were measured after overnight incubation at 37 °C.

Minimal inhibitory concentration (MIC) assay

The method described by the Clinical Laboratory Standards Institute (formerly the National Committee for Clinical Laboratory Standards) was followed. In brief, overnight-cultured bacteria in LB broth were diluted 100-fold into MH broth and grown at 37 °C for 2 h. The density of refreshed bacteria was adjusted with MH medium to OD_{600} at 0.05. One ml of the diluted bacterial culture was added to 1 ml of MH broth in a glass tube containing an appropriate concentration of plumbagin and then cultured at 37 °C with agitation for 20 h. Bacterial turbidity was measured at 600 nm by spectrophotometry.

Superoxide detection

A previous method [25] was modified to monitor the changes of superoxide level in *E. coli*. In brief, *E. coli* (*lpp*-deleted) was used for transformation with pQE-sodA or pQE-ygfZ. Then, bacteria at early log phase ($OD_{600} = 0.4$) were loaded with 10 µg/ml of dihydroethidium for 15 min before addition of superoxide inducing agents. Thereafter, the fluorescence of the cultures was followed by monitoring with a fluorescence spectrometer (TECAN) at excitation wavelength 488 nm and emission wavelength 575 nm.

Isolation of the organic soluble plumbagin metabolite

Overnight culture of the wild-type *E. coli* strain in LB broth was refreshed with aeration at 37 °C for 2 h. After adjusting the turbidity to OD_{600} at 0.5, plumbagin was added to the culture to a final concentration at 25 µg/ml. The bacteria were then further agitated at 37 °C for 20 h. After removing the bacteria by centrifugation, the spent media (50 ml) were extracted with chloroform (17.5 ml) three times. The combined chloroform extract was dried over anhydrous Na₂SO₄ and vacuum-concentrated. The resulted residue was dissolved in minimal chloroform and subjected to high performance liquid chromatography (HPLC) using E. Merck Lobar RP-C18 column (40-63 µm).

Identification of the structure of plumbagin metabolite

Infrared spectra were obtained with a Nicolet Avatar 320 FTIR spectrophotometer. UV spectra were measured with a Hitachi U-3310 spectrophotometer. Nuclear magnetic resonance spectra were recorded on a Varian VNMRS-600 spectrometer. The electron impact mass spectra were measured with the direct insertion probe on a Finnigan DSQ II mass spectrometer at 70 eV.

Statistics

All data were taken from at least three independent experiments. Differences between groups

were determined using the two-tail Student *t*-test and were considered statistically significant if p was <0.05.

Results

ygfZ critical for counteracting plumbagin toxicity

To examine the importance of the up-regulated genes previously found [12] in counteracting the plumbagin toxicity, we examined the relative sensitivity of mutant strains with each gene (*ahpC*, mdaB, nfnB, nfo, sodA, ygfZ, and yggX) disrupted individually. Also included in these experiments were three strains with similar disruptions at the upstream regulators soxR, soxS, and marA. The effects on growth inhibition zones surrounding plumbagin-containing discs on the MH agar plates are listed in Table 2. Compared to that of the parental strain, a remarkable increase in plumbagin sensitivity was observed with the $\Delta ygfZ$ and $\Delta sodA$ mutants and to a lesser extent with the $\Delta soxR$, $\Delta soxS$, and $\Delta ahpC$ strains whereas no effect was seen with the other strains. The MICs of the bacteria toward plumbagin were then determined. The MIC of the parental strain was expectedly much higher than those of the $\Delta ygfZ$ and $\Delta sodA$ mutants (Table 3). To ensure that the plumbagin-sensitivity of the $\Delta yg Z$ and $\Delta sodA$ mutants were readily due to the specific gene disruption, complementation assays were carried out. Fig. 1A shows a representative result. Upon transformation with pMH-ygfZ, the $\Delta ygfZ$ mutant showed a diminished inhibition zone, which is similar to that of the parental strain. This reversion of plumbagin-resistance was observed in the presence of different concentrations of plumbagin ranging from 20 to 100 µg per disc (Fig. 1B). Similarly, the increased inhibition zone of the $\Delta sodA$ mutant in an agar diffusion plate could be reduced to that of the wild type by expressing SodA from pQE-sodA (Fig. 2, right panel). Therefore, these results confirm that *ygfZ* and *sodA* are involved in the resistance to plumbagin in *E. coli*.

ygfZ required for the plumbagin breakdown

To test whether degradation of plumbagin occurs by the bacteria, the amounts of plumbagin remained in the culture media of $\Delta ygfZ$ and the parental strains were compared by using chloroform extraction and HPLC analysis. After 20-h aerobic cultivation, the concentration of plumbagin remained in the media with the $\Delta ygfZ$ mutant (5.78 µg/ml) was at least 10 fold higher than that derived from the parental strain (0.49 µg/ml), a fact suggesting a role of ygfZ involved in the degradation of plumbagin.

YgfZ and SodA independently required for resolving plumbagin toxicity

Since both ygfZ and *sodA* were found critical for *E. coli* to resolve the plumbagin toxicity, we examined whether they acted independently. Gene *sodA* encodes a manganese superoxide dismutase that converts superoxide anions to molecular oxygen and hydrogen peroxide [26]. As the action of plumbagin has been attributed to superoxide generation [5], SodA is likely to

combat plumbagin toxicity by detoxifying the superoxide. On the other hand, in view of the fact that plumbagin is degraded by E. coli, it is then reasonable to hypothesize that YgfZ and SodA may counteract plumbagin toxicity in two distinct ways. To test this hypothesis, we addressed whether expressing extra SodA could compensate the absence of YgfZ when E. coli is challenged with plumbagin. As shown in Fig. 2, when SodA was ectopically expressed from pQE-sodA in the $\Delta ygfZ$ strain, the inhibition zone remained large and did not differ significantly from that seen with the control plasmid-transformed $\Delta ygfZ$ strain (Fig. 2, left panel). These observations suggest that increasing expression of SodA in bacteria is not sufficient to overcome the plumbagin stress once YgfZ is absent. Reciprocally, increasingly expressed YgfZ in the $\Delta sodA$ mutant did not reduce the inhibition zone originally seen with the $\Delta sodA$ strain (Fig. 2, right panel). This result indicated that E. coli, in the absence of SodA but with ectopically expressed YgfZ, remained incapable of resisting plumbagin toxicity. A doubly mutated strain at both ygfZ and sodA was then created and MICs toward plumbagin were compared (Table 3). Apparently, the double mutant $(\Delta yg/Z/\Delta sodA)$ was the most sensitive strain and its MIC was smaller than either one of the singly disrupted strains. It is then concluded that ygfZ and sodA both contribute to the resistance of *E. coli* toward plumbagin toxicity but act independently.

To substantiate the notion that different roles are played by YgfZ and SodA in facing the plumbagin challenge, the superoxide levels in the bacteria after receiving chemicals were followed by monitoring the fluorescence change of dihydroethidium. Figure 3A shows that plumbagin tended to increase the superoxide level in bacteria as the known superoxide generator paraquat did. On the other hand, when the bacteria ectopically produced SodA, the original stimulation of superoxide production by either paraquat or plumbagin diminished (compare Fig. 3A with 3B). However, this was not the case when *E. coli* was transformed to produce extra YgfZ (Fig. 3C); the trend of increasing superoxide production after paraqaut/plumbagin treatment remained the same (compare Fig. 3A and 3C). Therefore, these results consolidated the conception that YgfZ behaves in a mechanism different from that of SodA as to resolving the threat of plumbagin. One of the likely roles of YgfZ involved is possibly to accelerate the breakdown of plumbagin.

Determining the ygfZ-dependent metabolites of plumbagin

To confirm the plumbagin degradation happened in E coli, an effort was made to identify any degraded product of plumbagin. In the HPLC profile of an organic extract prepared from the plumbagin-containing culture media of the parental E. coli strain, two extra peaks (peaks II and III in Fig. 4A) were found. These peak fractions were collected and subjected to analysis with electron impact mass spectroscopy. A molecule with a molecular weight of 14 Daltons more than that of plumbagin was found from peak II (see Additional file 1 – Chemical identification data). Further analysis nuclear magnetic identified this molecule with resonance as 2,3-dimethyl-5-hydroxy-1,4-naphthoquinone (2,3-dimethyl-5-hydroxyl-naphthalene-1,4-dione, IUPAC), whose structure is shown in Fig. 4D. This compound is referred as methylated plumbagin hereafter. This compound was then prepared by organic synthesis and compared with that extracted from the spent medium using HPLC (Fig. 4A and 4D), infrared, UV and nuclear magnetic resonance analyses. All data obtained supported that the compound from the culture media and that from synthesis were identical. Identification of the compound in peak III was not successful due to a low yield after purification. Furthermore, this methylated plambagin was not seen in the HPLC profile (Fig. 4B) generated from the $\Delta ygfZ$ strain culture and neither found in the repeated experiment.

To examine whether there is any anti-bacterial activity left with methylated plumbagin, MIC was measured, and no apparent activity was found with concentrations up to 200 µg/ml when *E. coli* of the $\Delta sodA$ and the $\Delta ygfZ$ strains and the parental strain were tested (Table 3). Therefore, adding a methyl group to the 3-position of naphthoquinone ring apparently diminishes the plumbagin toxicity against *E. coli*.

Homologues of YgfZ

To analyze the critical region(s) of ygfZ, we searched for the conserved residues among the homologues of YgfZ. Alignment of the sequences from *E. coli*, *K. pneumoniae*, and *M. tuberculosis* is shown in Fig. 5A. The identity between the two YgfZ homologues from *E. coli*

and K. pneumoniae is 81.9%, whereas it is only 20.1% between Rv0811c of M. tuberculosis and YgfZ of E. coli (insert in Fig. 5A). In the agar diffusion assay (Fig. 5B), Kp_YgfZ from the K. pneumoniae ygfZ was able to restore fully the plumbagin resistance in the E. coli $\Delta ygfZ$ strain. When Mtb_Rv0811c, which is an open reading frame annotated as an aminomethyltransferase-related gene [27], was used in a similar complementation assay, the plumbagin resistance in the $\Delta y g f Z$ strain was regained partially (Fig. 5B). Since there is only a low degree of identity between Rv0811c and YgfZ, it is not clear whether the former is a real counterpart of the latter. Therefore, additional genes annotated as aminomethyltransferases, namely the gcvT gene from E. coli and Rv2211c from M. tuberculosis, were cloned and used in similar assays. No function was observed with either of the two constructs. Therefore, it is believed that Rv0811c is the homologue of YgfZ in M. tuberculosis and the commonly conserved regions among all sequences must play an essential role.

Cys 228 in YgfZ critical for plumbagin resistance

Additional experiments were performed to dissect the critical residue(s) in the highly conserved region from K226 to R237, which contains a stretch (²²⁶K-G-C-Y-T-G-Q-E²³³) of the *E. coli* YgfZ molecule, a region described as fingerprint previously [22, 23]. To address the importance of this highly conserved region, amino acid residues 226-237 were deleted and the so-truncated YgfZ was then used in the complementation assay (Fig. 5B). The truncated YgfZ totally lost the

ability to rescue plumbagin resistance in the $\Delta ygfZ$ strain. This result is consistent with the expectation that this region is crucial for the YgfZ function.

To further narrow down to which residue is critical, single alanine-substitution mutants of YgfZ were created in the fingerprint region. These YgfZ variants were then assessed for the ability to restore plumbagin resistance in the $\Delta ygfZ$ strain. As shown in Fig. 6A, most of these mutated YgfZ constructs (gray bars) readily reduced the inhibition zones and behaved as active as the authentic YgfZ molecule (black bar) in this agar diffusion assay. Two exceptions were mutation at Cys228 and Tyr229 (hatched bars). The C228A mutant performed poorest among these single-point variants. The authentic YgfZ reduced the plumbagin inhibition zone from 40 mm to 10 mm (in diameter), whereas the inhibition zone remained large at 17 mm with C228A and at 12 mm with Y229A (Fig. 6A). Not shown in Fig. 6A, C228A/Y229A (with double substitutions at residues 228 and 229) lost the complementation activity one step further and resulted in a 28-mm inhibition zone. These results together suggest that C228 is the most critical residue in the fingerprint region of YgfZ followed by Y229 that contributes to the protein's functional integrity but to a lesser extent.

The critical role of C228 in YgfZ was previously predicted to form disulfide bridge [23]. There are two cysteine residues in the *E. coli* YgfZ molecule and the second one is located at residue 63. To test whether C228 is critical for the formation of an intra-molecular disulfide in YgfZ, a single-point mutation at C63 was constructed. The YgfZ variant C63G was found to retain the full authentic YgfZ function in the $\Delta ygfZ$ complementation assay (data not shown), suggesting that the critical role of C228 in YgfZ does not rely on forming an intra-molecular disulfide bond with C63. Further efforts were made to explore mechanisms of C228 function in YgfZ by replacing C228 with either Ser or Met. The resulting variants C228S and C228M were then side-by-side compared with C228A in the $\Delta ygfZ$ complementation assay. Fig. 6B shows that C228S was able to complement to the same degree as the authentic YgfZ and their plumbagin resistances were indistinguishable at three increasing amounts of plumbagin (from 20 µg up to 100 µg per disc). C228M, similar to C228A, was indistinguishable from the authentic construct when assayed at 20 µg or 50 µg of plumbagin, but it gave less resistance when plumbagin was applied at 100 µg. Therefore, residues with thiol and hydroxyl groups play equivalent role at position 228 of YgfZ in term of plumbagin resistance and this biological role could only be partially replaced by residues with a methyl group.

Discussion

Among the *E. coli* genes whose products are up-regulated by plumbagin [12], ygfZ and sodA readily contribute to resisting the plumbagin's toxicity. When tested with plumbagin at 100 µg

per disc, the inhibition zone of the $\Delta ygfZ$ strain was apparently greater than that of the $\Delta sodA$ strain (Table 2). On the other hand, when paraquat was applied at 1.28 μ g per disc, the $\Delta ygfZ$ strain showed the same resistance as the parental strain whereas the inhibition zone of the $\Delta sodA$ strain increased substantially (data not shown). It is known that the expression of *sodA* is elevated when E. coli is treated with plumbagin and paraquat separately [12, 28]. Up-regulation of ygfZ expression also occurs when E. coli is treated with plumbagin, but not seen with the paraquat treatment [12, 29]. Consistently, we have seen that the superoxide induction resulted from encountering plumbagin were severely repressed by an additional expression of SodA (Fig. 3B), but not by YgfZ (Fig. 3C). It is then conceivable that in the response to the challenge of plumbagin, E. coli could not handle the toxicity simply by increasing the amount of SodA. An additional amour with more YgfZ is apparently needed. The mutual irreplaceable roles of SodA and YgfZ for bacteria to resolve the plumbagin challenge (Fig. 2) support the notion that the function of YgfZ is acting independently from SodA.

YgfZ homologues are found among many Gram (-) bacteria and in the mitochondria of eukaryotes but are not found in Archaea [22, 23]. No counterpart has been found in Gram (+) bacteria except for those in the chromosomes of high-GC Actinobacteria such as *Streptomyces* spp. and *Mycobacteria* spp. The levels of identity among the YgfZ sequences of the enterobacteria are around 80% or higher whereas that between *E. coli* and *M. tuberculosis* is as

low as 20%. Interestingly, the anti-plumbagin activity of these YgfZ homologues seems to be well preserved although to different degrees. A stretch (from K226 to R237 in E. coli YgfZ) comprising the previously described fingerprint (K-G-C-Y/F-X-G-Q-E) [23] is conserved across these protein sequences. Within this fingerprint region, we have identified C228 as the most imperative residue for plumbagin detoxification (Fig. 6A). However, the effects of single residue site-directed mutants were not as profound as that seen with YgfZ Δ 226-237, which completely lost its anti-plumbagin ability in $\Delta ygfZ$ mutant (Fig 5B). Although other possibilities could not be excluded, a worst explanation for these observations is that the structure of YgfZ could be completely distorted when the segment of residues 226-237 was deleted. Nevertheless, when residues at 228 and 229 of YgfZ were simultaneously mutated to Ala in the construct of C228A/Y229A, the effect on YgfZ was further amplified; the inhibition zone was close to 28 mm, a size similar to that seen with the Mtb Rv0811c complementation (Fig. 5B). This result revealed that these two residues have synergistic effect for anti-plumbagin activity. The possibility of C228 forming a disulfide linkage [23] has been excluded by the substitution experiment of the second Cys at residue 63, which showed no impact on plumbagin resistance. By substituting the thiol group in C228 with a hydroxyl group, we found authentic YgfZ molecule and the C228S variant were functionally comparable (Fig. 6B). In our mass spectroscopy analysis of a vinyl-palmitic acid-reacted sample, the C228 of E. coli YgfZ was

found to be labeled with palmitate (data not shown). Therefore, C228 is concluded to possess a free thiol side-chain. Since we have observed that this cysteine residue could be functionally replaced by Ser but only to a partial extent by Met or Ala (Fig. 6B), the role of Cys at residue 228 is likely to provide a lone pair of electrons during the spatial molecular interactions.

The resistance of bacteria to antimicrobial agents is mediated by a variety of mechanisms [30]. By protein fractionation, we found that YgfZ is located in the cytoplasmic fraction (see Additional file 2 – Localization of the ygfZ gene product to the cytoplasm), a fact suggesting that YgfZ is unlikely to be a part of an efflux/influx system. Furthermore, by comparing HPLC profiles of organic extracts prepared from the culture media of the parental bacteria and the $\Delta ygfZ$ strain, discovered possible metabolite of plumbagin, we a 2,3-dimethyl-5-hydroxy-1,4-naphthoquinone. This methylated plambagin in peak II simply constituted a small portion of the plumbagin metabolites after cultivation for 20 h (compare Fig. 4A and Fig. 4C), an observation suggesting that there may be more breakdown products not recovered or detected by these The identified processes. 2,3-dimethyl-5-hydroxy-1,4-naphthoquinone appears to be non-toxic to bacteria, up to a concentration of 200 μ g/ml (Table 3). In a preliminary experiment, we have found that this compound prepared from our synthesis disappeared gradually when added to the bacterial culture, a fact corroborating the notion that this methylated product is not the final breakdown of

plumbagin in E. coli.

Conclusion

We found that YgfZ plays a critical role in plumbagin resistance in *E coli*. Based on our current findings, we suggest that the mechanisms of plumbagin resistance in *E. coli* may involve at least two independent gene products. SodA is induced to resolve the plumbagin-induced oxidation stress whereas YgfZ is induced to facilitate the plumbagin breakdown. The latter mechanism involves at least the methylation of plumbagin that yields non-toxic 2,3-dimethyl-5-hydroxy-1,4-naphthoquinone.

Competing interests

The authors declare that they have no competing interests.

Authors' contributions

CNL designed and performed the majority of works in this research. WJS was a research supervisor and coordinator. WWS performed the site-directed mutagenesis assays. JWC generated some expression plamsids and initiated the early works in this research. THC carried

out the plumbagin metabolite analysis. CNL and SHW wrote the manuscript. MJD and SHW were research group leaders who contributed to data interpretation. All authors were involved in reviewing and updating the text associated with the manuscript. All authors have read and approved the final manuscript.

Acknowledgements

The technical support from Yang-Ming Proteomic Center is acknowledged. We also thank Dr. CS Chen from Ohio State University (USA) for the useful discussion and Dr. R Kirby for critical reading of this manuscript. This work was supported in part by a grant from Ministry of Education, Aim for the Top University Plan (http://english.moe.gov.tw/). WJS was supported by grants 97-2627-M-010-003 and 97-2320-B-010-005-MY3 from the National Science Council. SHW was supported by 98-2320-B-415-004-MY3 from the National Science Council, Taiwan. Hereby, we claim that this is an independent study and has no connection to the recent report by Waller et al., (*Proc Natl Acad Sci USA* 2010).

References

1. de Paiva SR, Figueiredo MR, Aragao TV, Kaplan MA: Antimicrobial activity in vitro of plumbagin isolated from *Plumbago* species. *Memorias do Instituto Oswaldo Cruz* 2003,

98:959-961.

- Curreli N, Sollai F, Massa L, Comandini O, Rufo A, Sanjust E, Rinaldi A, Rinaldi AC: Effects of plant-derived naphthoquinones on the growth of *Pleurotus sajor-caju* and degradation of the compounds by fungal cultures. *Journal of basic microbiology* 2001, 41:253-259.
- Krishnaswamy M, Purushothaman KK: Plumbagin: A study of its anticancer, antibacterial & antifungal properties. Indian journal of experimental biology 1980, 18:876-877.
- Edenharder R, Tang X: Inhibition of the mutagenicity of 2-nitrofluorene,
 3-nitrofluoranthene and 1-nitropyrene by flavonoids, coumarins, quinones and other phenolic compounds. *Food Chem Toxicol* 1997, 35:357-372.
- Hassan HM, Fridovich I: Intracellular production of superoxide radical and of hydrogen peroxide by redox active compounds. Archives of biochemistry and biophysics 1979, 196:385-395.
- 6. Ieva R, Roncarati D, Metruccio MM, Seib KL, Scarlato V, Delany I: OxyR tightly regulates catalase expression in *Neisseria meningitidis* through both repression and activation mechanisms. *Molecular microbiology* 2008, **70**:1152-1165.
- 7. Alekshun MN, Levy SB: Alteration of the repressor activity of MarR, the negative

regulator of the *Escherichia coli marRAB* locus, by multiple chemicals *in vitro*. *Journal* of bacteriology 1999, **181:**4669-4672.

- Gaudu P, Moon N, Weiss B: Regulation of the soxRS oxidative stress regulon. Reversible oxidation of the Fe-S centers of SoxR in vivo. The Journal of biological chemistry 1997, 272:5082-5086.
- 9. Koh YS, Roe JH: Isolation of a novel paraquat-inducible (*pqi*) gene regulated by the *soxRS* locus in *Escherichia coli*. *Journal of bacteriology* 1995, **177:**2673-2678.
- Neuhaus JM, Wright JK: Chemical modification of the lactose carrier of *Escherichia coli* by plumbagin, phenylarsinoxide or diethylpyrocarbonate affects the binding of galactoside. *European journal of biochemistry / FEBS* 1983, 137:615-621.
- Imlay J, Fridovich I: Exogenous quinones directly inhibit the respiratory NADH dehydrogenase in *Escherichia coli*. Archives of biochemistry and biophysics 1992, 296:337-346.
- Chen JW, Sun CM, Sheng WL, Wang YC, Syu WJ: Expression analysis of up-regulated genes responding to plumbagin in *Escherichia* coli. *Journal of bacteriology* 2006, 188:456-463.
- Dubbs JM, Mongkolsuk S: Peroxiredoxins in bacterial antioxidant defense. Sub-cellular biochemistry 2007, 44:143-193.

- 14. Barbosa TM, Levy SB: Activation of the Escherichia coli nfnB gene by MarA through a highly divergent marbox in a class II promoter. Molecular microbiology 2002, 45:191-202.
- 15. Martin RG, Rosner JL: Genomics of the marA/soxS/rob regulon of Escherichia coli: identification of directly activated promoters by application of molecular genetics and informatics to microarray data. Molecular microbiology 2002, 44:1611-1624.
- 16. Galhardo RS, Almeida CE, Leitao AC, Cabral-Neto JB: Repair of DNA lesions induced by hydrogen peroxide in the presence of iron chelators in *Escherichia coli*: participation of endonuclease IV and Fpg. *Journal of bacteriology* 2000, 182:1964-1968.
- 17. Iuchi S, Weiner L: Cellular and molecular physiology of *Escherichia coli* in the adaptation to aerobic environments. *Journal of biochemistry* 1996, **120**:1055-1063.
- Osborne MJ, Siddiqui N, Landgraf D, Pomposiello PJ, Gehring K: The solution structure of the oxidative stress-related protein YggX from *Escherichia coli*. Protein Sci 2005, 14:1673-1678.
- Gralnick JA, Downs DM: The YggX protein of Salmonella enterica is involved in Fe(II) trafficking and minimizes the DNA damage caused by hydroxyl radicals: residue CYS-7 is essential for YggX function. The Journal of biological chemistry 2003, 278:20708-20715.

- Pomposiello PJ, Koutsolioutsou A, Carrasco D, Demple B: SoxRS-regulated expression and genetic analysis of the yggX gene of Escherichia coli. Journal of bacteriology 2003, 185:6624-6632.
- 21. Martin RG, Bartlett ES, Rosner JL, Wall ME: Activation of the Escherichia coli marA/soxS/rob regulon in response to transcriptional activator concentration. Journal of molecular biology 2008, 380:278-284.
- 22. Ote T, Hashimoto M, Ikeuchi Y, Su'etsugu M, Suzuki T, Katayama T, Kato J: Involvement of the *Escherichia coli* folate-binding protein YgfZ in RNA modification and regulation of chromosomal replication initiation. *Molecular microbiology* 2006, **59**:265-275.
- 23. Teplyakov A, Obmolova G, Sarikaya E, Pullalarevu S, Krajewski W, Galkin A, Howard AJ, Herzberg O, Gilliland GL: Crystal structure of the YgfZ protein from *Escherichia coli* suggests a folate-dependent regulatory role in one-carbon metabolism. *Journal of bacteriology* 2004, 186:7134-7140.
- 24. Ku CP, Lio JC, Wang SH, Lin CN, Syu WJ: Identification of a third EspA-binding protein that forms part of the type III secretion system of enterohemorrhagic *Escherichia coli*. *The Journal of biological chemistry* 2009, 284:1686-1693.
- 25. Herrera G, Martinez A, O'Cornor JE, Blanco M: Functional assays of oxidative stress using genetically engineered *Escherichia coli* strains. *Current Protocols in Cytometry*

2003, Chapter 11:Unit 11 16.

- 26. Fridovich I: The biology of oxygen radicals. Science 1978, 201:875-880.
- 27. Camus JC, Pryor MJ, Medigue C, Cole ST: Re-annotation of the genome sequence of *Mycobacterium tuberculosis* H37Rv. *Microbiology (Reading, England)* 2002, 148:2967-2973.
- 28. Naik SM, Hassan HM: Use of site-directed mutagenesis to identify an upstream regulatory sequence of sodA gene of Escherichia coli K-12. Proceedings of the National Academy of Sciences of the United States of America 1990, 87:2618-2622.
- 29. Pomposiello PJ, Bennik MH, Demple B: Genome-wide transcriptional profiling of the *Escherichia coli* responses to superoxide stress and sodium salicylate. *Journal of bacteriology* 2001, 183:3890-3902.
- 30. Tenover FC: Mechanisms of antimicrobial resistance in bacteria. *The American journal of medicine* 2006, **119:S**3-10; discussion S62-70.

 Table 1. Primers used and their sequences

Name	Sequence (5' to 3')	Used in construction	
PygfZF	CCATGGCTTTTACACCTTTTCCTCCCCG	pQE-ygfZ	
PygfZR	AGATCTCTCTCGAGCGAATACGGCAGC		
PsodAF	GGACTTATGAGCTATACCCTGCCATC	POE and A	
PsodAR	GGATCCTTTTTTCGCCGCAAAACGTA	pQE-sodA	
PkpygfZF	CCATGGGTATGGCTTTTACACCTTTTCC	nOE Kn. vaf7	
PkpygfZR	AGATCTATTTTCTTCCAGCGAATACGGC	pQE-kp_ygiz	
PRv0811cF	CCATGGCCGCAGTCCCTGCCCCAGACCC	$\mathbf{p} \mathbf{O} \mathbf{E} \mathbf{D}_{\mathbf{V}} \mathbf{O} \mathbf{V} 1 1_{\mathbf{C}}$	
PRv0811cR	AGATCTCCGAATACCGCCGCGCAGCCGC	pQE-Kv_08110	
PygfZK226AF	CAGCTTTAAGGCCGGCTGTTATACCG	pOE y of 7K226A	
PygfZk226AR	CGGTATAACAGCCGGCCTTAAAGCTG	pQE-ygiZK220A	
PygfZG227AF	CTTTAAGAAAGCCTGTTATACCGGAC	nOF-vafZG227A	
PygfZG227AR	GTCCGGTATAACAGGCTTTCTTAAAG	pQL-ygiZO22/A	
PygfZC228AF	CTTTAAGAAAGGGGCTTATACCGGACAAG	nOF-vaf7C228A	
PygfZC228AR	CTTGTCCGGTATAAGCCCCTTTCTTAAAG	pQL-ygize220A	
PygfZC228SF	CTTTAAGAAAGGCTCGTATACCGGAC	nOF-vof7C228S	
PygfZC228SR	GTCCGGTATACGAGCCTTTCTTAAAG	pQL-ygize2200	
PygfZC228MF	CTTTAAGAAAGGCATGTATACCGGAC	nOF-vgf7C228M	
PygfZC228MR	GTCCGGTATACATGCCTTTCTTAAAG	pQL-ygiZC220W	
PygfZY229AF	TAAGAAAGGCTGTGCTACCGGACAAG	nOF-vafZV2294	
PygfZY229AR	CTTGTCCGGTAGCACAGCCTTTCTTA		
PygfZT230AF	AAGGCTGTTATGCCGGACAAGAGATG	nOF-vgf7T230A	
PygfZT230AR	CATCTCTTGTCCGGCATAACAGCCTT	pQL-ygiL1250A	
PygfZG231AF	GCTGTTATACCGCGCAAGAGATGGTG	nOF-vofZG231A	
PygfZG231AR	CACCATCTCTTGCGCGGTATAACAGC	pqL-ygiZ02511	
PygfZQ232AF	CTGTTATACCGGAGCAGAGATGGTGG	nOF-vofZO232A	
PygfZQ232AR	CCACCATCTCTGCTCCGGTATAACAG		
PygfZE233AF	GTTATACCGGACAGGCCATGGTGGCGCGA	nOF-vgfZF233A	
PygfZE233AR	TCGCGCCACCATGGCCTGTCCGGTATAAC	PQL ygill2551	
PygfZ∆226-237F	GGGCGGTATCAGCTTTAAGGCCAAATTCC	pQE-ygfZΔ226-237	
PygfZ∆226-237R	GGAATTTGGCCTTAAAGCTGATACCGCCC		
PQEF	GGCGTATCACGAGGCCCTTTTCG	Fragment amplification	
PQER	CATTACTGGATCTATCAACAGG	Fragment amplification	

Strain tested	Relative sensitivity to plumbagin at different amounts [*]			
	20 µg	50 µg	100 µg	
WT, ΔmdaB, ΔnfnB Δnfo, ΔyggX or ΔmarA	, _	-	-	
ΔsoxR, ΔsoxS, or ΔahpC	-	-	+	
$\Delta sodA$	+	++	++	
$\Delta ygfZ$	+	++	+++	

Table 2. Growth inhibitory effect of plumbagin against different *E. coli* mutants

* Bacteria were plated on MH agar plates with plumbagin absorbed on an 8-mm filter paper disc.

-: inhibition zone < 15 mm; +: 15 mm < inhibition zone < 25 mm; ++: 25 mm < inhibition zone

< 35 mm; +++: inhibition zone > 35 mm

Strains	nlasmid	MIC (µg/ml)		
		plumbagin	methylated plumbagin	
WT	-	50	>200	
$\Delta sodA$	-	16	>200	
$\Delta ygfZ$	-	8	>200	
$\Delta ygfZ/\Delta sodA$	-	4	Not tested	
WT	pМH	50	Not tested	
$\Delta ygfZ$	pMH-ygfZ	50	Not tested	
WT	pQE60	40	Not tested	
$\Delta ygfZ$	pQE-ygfZ	40	Not tested	
$\Delta ygfZ$	pQE-ygfZK226A	40	Not tested	
$\Delta ygfZ$	pQE-ygfZG227A	40	Not tested	
$\Delta ygfZ$	pQE-ygfZC228A	30	Not tested	
$\Delta ygfZ$	pQE-ygfZC228S	40	Not tested	
$\Delta ygfZ$	pQE-ygfZC228M	30	Not tested	
$\Delta ygfZ$	pQE-ygfZY229A	30	Not tested	
$\Delta ygfZ$	pQE-ygfZT230A	40	Not tested	
$\Delta ygfZ$	pQE-ygfZG231A	40	Not tested	
$\Delta ygfZ$	pQE-ygfZQ232A	40	Not tested	
$\Delta ygfZ$	pQE-ygfZE233A	40	Not tested	
$\Delta ygfZ$	pQE-ygfZ∆226-237	8	Not tested	
$\Delta ygfZ$	pQE-Kp_ygfZ	40	Not tested	
$\Delta ygfZ$	pQE-Rv_0811c	10	Not tested	
$\Delta ygfZ$	pQE-sodA	8	Not tested	
$\Delta sodA$	pQE-sodA	40	Not tested	
$\Delta sodA$	pQE-ygfZ	16	Not tested	

Table 3. MICs for different *E. coli* mutants

Figure legends

Figure 1. YgfZ is critical for resolving plumbagin toxicity.

(A) Growth inhibition assay on the agar diffusion plates. Bacteria harboring the indicated plasmids were plated overnight at 37 °C on MH plates in the presence of plumbagin-containing filter discs (8 mm in diameter). (B) Diameters of the inhibition zones seen in (A) at different plumbagin concentrations. Note: strain BW25113 (WT) is the parental strain of the $\Delta ygfZ$ mutant whereas pMH-ygfZ differs from the promoterless pMH vector by carrying ygfZ as well its upstream promoter region. NS: no significance; * p < 0.05

Figure 2. Different roles played by YgfZ and SodA in counteracting plumbagin.

The $\Delta ygfZ$ and $\Delta sodA$ strains were transformed with pQE-sodA and pQE-ygfZ to express SodA and YgfZ, respectively, and the agar diffusion assay was performed similar to that described in legend to Fig. 1. Note: pQE60 was the vector used for expression construction. Inset: the plasmid-encoded His_{x6}-tagged proteins were well expressed in the transformants as revealed by Western blotting; antibody-detected DnaK served as a protein-loading control. NS: no significance

Figure 3. Superoxide level in E. coli.

E. coli (*lpp*-deleted) was transformed with pQE-sodA and pQE-ygfZ to express recombinant SodA and YgfZ, respectively, and the superoxide levels in bacteria were determined by monitoring the fluorescence changes after loading with dihydroethidium [25]. Data were taken after 120-min treatments with chemicals. (A) Both paraquat (50 μ M) and plumbagin (50 μ M) stimulated the levels of superoxide detected. (B) The superoxide stimulation seen in (A) was suppressed by SodA expression. (C) The same experiments in (B) were repeated with bacteria expressing YgfZ. Note: pQE60 was the vector control.

Figure 4. HPLC analysis of the metabolized plumbagin.

Samples were subjected to RP-C18 column chromatography that was run with a mixture of methanol/H₂O (7:3, v/v). Compounds eluted were detected with UV absorbance at λ_{254} . Samples were chloroform extract of: (**A**) the plumbagin-containing cultivation media of the wild-type *E*. *coli*; (**B**) the same preparation as (A) but with the $\Delta ygfZ$ strain; (C) the same preparation as (A) but without bacteria; (D) synthesized 2,3-dimethyl-5-hydroxy-1,4-naphthoquinone extracted from media as described for (C). Compounds identification: I, plumbagin; II, 2,3-dimethyl-5-hydroxy-1,4-naphthoquinone; III, unidentified.

Figure 5. Complementation to assay the resistance of the $\triangle ygfZ$ strain toward plumbagin after expressing homologous constructs.

(A) Amino-acid-sequence alignment of E. coli YgfZ (ref|NP_417374), K. pneumoniae YgfZ (Kp_YgfZ; reflBAH65109), and *M. tuberculosis* Rv0811c (reflNP_215326). Residues conserved in all three sequences are marked in black whereas those semi-conserved are boxed in gray; labeled above the alignment are residue numbers of the longest Rv0811c sequence and exceptions are those italicized for which represent the YgfZ residues in E. coli and K. pneumoniae. The cysteine residue in the conserved fingerprint region [23] is asterisked. Inset: amino acid identity between pairs of the three proteins as calculated by Vector NTI (InforMax). (B) Comparison of the activities of different YgfZ constructs to support the growth of the $\Delta ygfZ$ E. coli strain in the presence of plumbagin. Plasmids were separately transformed into the $\Delta ygfZ$ strain and assayed for the diameters of the growth inhibition zone as in Fig. 1B. Inset: the plasmid-encoded proteins expressed in the transformants were detected by Western blotting using anti-His_{x6} antibody; Dank was detected in parallel, to assure a comparable protein loading. Note: pQE60 served as a negative control. NS: no significance; * p < 0.05

Figure 6. Analysis of critical residues in the fingerprint region of YgfZ.

(A) Inhibition zone assay for the plumbagin-countering activity of amino acid-substituted YgfZ.

The $\Delta ygfZ$ mutant was transformed with pQE-ygfZ-derived plasmids to express variants of E. coli YgfZ. K226A, G227A, C228A, Y229A, T230A, G231A, Q232A, and E233A are constructs with single-amino acid substitution at the indicated residue. Hatched bars mark the substitution mutants with the properties obviously different from the authentic control (black bar). (B) Analysis of the substitutability of C228 with structurally similar amino acids. Complementation transformation of the $\Delta ygfZ$ mutant was done as in (A) except that plumbagin was applied at three different levels. Note: the construct with the Cys to Ser mutation (C228S) behaved indistinguishable from the authentic YgfZ at all different plumbagin amounts applied while C228M and C228A mutants apparently deviated from the authentic when plumbagin was applied at 100 µg per disc. Insets: exogenous His_{x6}-tagged YgfZ constructs were expressed in the transformed $\Delta ygfZ$ strain comparably as revealed by Western blotting; DnaK served as a protein-loading control. Note: pQE60 served as negative control. To compare the significance of the data, results from the authentic YgfZ were used as a reference. NS: no significance; * p <0.05

Additional files

Additional file 1 – Chemical identification data

File format: PDF format

Description: The general chemical properties, IR and UV absorption spectra and NMR analysis

of 2,3-dimethyl-5-hydroxy-1,4-naphthoquinone.

Additional file 2 – Localization of the *ygfZ* gene product to the cytoplasm

File format: PDF files

Description: Western bolt analysis showed the cytoplasmic distribution of YgfZ in E. coli.

WT + pMH



∆*ygfZ* + pMH-ygfZ



 $\Delta ygfZ + pMH$





(B)





Figure 2



Figure 3



Time (min)

Absorption (mAU)

1	А.	N	
1	Δ	1	
٩.	6 B	- J.	
×.		1	

1 YgtZ (1) MAPTEREPROPTA KP_YgtZ (1) MAPTERPROPSS	LTUMTLDDWAL LTUMTLDDWAL DAVEVDRSHRAV	AMITCADSEKYM AMISCPUSEKYL LULDCGLRQTWL	QGQVTADVSQMAEDQ QGQITADVSHLTDAQ ISISTQHVSDLPEGA	8 HLLAAHCDAKCKMWSN HLLAAHCDAKCKMWSN STQNLSLDGQCRVEDH	8 LR LR WI
89 YgIZ (75) LFRDGDCFAWIERRSVR-EPOTTERKYAVFS KP_YgIZ (75) VFRREGCFAWIERRSLR-DAOTTERKYAVFS Mtb_Rv0811c (89) QTELG-CTTYLDTEPWRGEPLLAYLRRMVFWS	KVDIAPDDERV KVDIAANDDLV MVDPRAADMAV	L LGVAGFOAR L LGVAGFOAR LSLLCPRLAEER	AALANLESEMPSKEK AALAPLEAAMPDAAT /LDALGLDVEPAEWL	17 OVVKECATTLLWFEHP PVVSECATSLLWFEHP AVPLACGGIVRRMPDG	16 AE GE
177 YgfZ (160) RFLIVTDEATANMLTDKLEGEAELNN KP_YgfZ (160) RFLLVTDVDTANRVTDALEGEAQFNN Mtb_Rv0811c (176) GQIELDVVVKRGDRADWQERLTQAGVRPAGIW	SQQWLALNI SQQWLALNI MYEAHRVAHRV	EAGFFVIDAANS(EAGLFVIDSANS(PARRERLGVDTD)	GOFIFQATNLQA GOFIFQATNLQA ERT <mark>IP</mark> HEVGWIGGPG	226 * 20 LGGISFRKGCYTGQEM LGGISFRKGCYTGQEM AGAVHLMKGCYRGQET	54 VA VA VA
237 265 YgiZ (237) FAKFRCANKFALWILAGSASRLPEAGELLETK KP_YgiZ (237) FAKFRCANKFALWILSGTASRVPESGELLETK Mtb_Rv0811c (264) FVHNLCRPPFMLVLLHLDESVQRPSTGTAVLA	MGENWRRIGTV MGDNWRRIGTV GGRTVGRLGTV	LAAVKUEDGOVVI LAAVOUDDGSLM VEHVEVGPVALAI	VQVVMNNDMEPDSIF VQVVMNNDMEPDSVF LLKRGLPGDTALVTG	3: RVRDDANTLHIEPUPY RVRDDAGSLSIKPUPY PEAEVAAVIDVDSUPP	52 SL SL AD
353 369 YgtZ (325) EE KP_YgtZ (325) EE		automater of	Identity (%)		
Mtb_Rv0811c (353) DVGAGRRAVERLRGGIR		YgfZ	Kp_YgfZ	Mtb_Rv0811c	

YgfZ

100

81.9

20.1





Additional files provided with this submission:

Additional file 1: Additional_file_1-Chemical_identification_data.pdf, 36K <u>http://www.jbiomedsci.com/imedia/1849308646406918/supp1.pdf</u> Additional file 2: Additional_file_2-Localization_of_the_ygfZ_gene_product_to_the_c, 53K <u>http://www.jbiomedsci.com/imedia/4995266504069181/supp2.pdf</u>