

# Dynamic Proteome Response of *Pseudomonas aeruginosa* to Tobramycin Antibiotic Treatment\*

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Genetically susceptible bacteria become antibiotic tolerant during chronic infections, and the mechanisms responsible are poorly understood. One factor that may contribute to differential sensitivity *in vitro* and *in vivo* is differences in the time-dependent tobramycin concentration profile experienced by the bacteria. Here, we examine the proteome response induced by subinhibitory concentrations of tobramycin in *Pseudomonas aeruginosa* cells grown under planktonic conditions. These efforts revealed increased levels of heat shock proteins and proteases were present at higher dosage treatments (0.5 and 1  $\mu\text{g/ml}$ ), while less dramatic at 0.1  $\mu\text{g/ml}$  dosage. In contrast, many metabolic enzymes were significantly induced by lower dosages (0.1 and 0.5  $\mu\text{g/ml}$ ) but not at 1  $\mu\text{g/ml}$  dosage. Time course proteome analysis further revealed that the increase of heat shock proteins and proteases was most rapid from 15 min to 60 min, and the increased levels sustained till 6 h (last time point tested). Heat shock protein IbpA exhibited the greatest induction by tobramycin, up to 90-fold. Nevertheless, deletion of *ibpA* did not enhance sensitivity to tobramycin. It seemed possible that the absence of sensitization could be due to redundant functioning of IbpA with other proteins that protect cells from tobramycin. Indeed, inactivation of two heat shock chaperones/proteases in addition to *ibpA* in double mutants (*ibpA/clpB*, *ibpA/PA0779* and *ibpA/hsIV*) did increase tobramycin sensitivity. Collectively, these results demonstrate the time- and concentration-dependent nature of the *P. aeruginosa* proteome response to tobramycin and that proteome modulation and protein redundancy are protective mechanisms to help bacteria resist antibiotic

treatments. *Molecular & Cellular Proteomics* 14: 10.1074/mcp.M115.050161, 2126–2137, 2015.

The opportunistic pathogen *Pseudomonas aeruginosa* is ubiquitous in the natural environment and causes human infections (1). *P. aeruginosa* can metabolize various carbon and nitrogen compounds and persists under nutrient-poor and hostile growth environments (2, 3). One example is *P. aeruginosa* pulmonary infection of cystic fibrosis (CF) patients. Despite stress induced by host defenses and high concentrations of antibiotics, *P. aeruginosa* cells are able to persistently colonize CF airways (4).

The aminoglycoside tobramycin is a front-line drug currently used in the treatment of *P. aeruginosa* in CF and other diseases. It is supplied in the forms of inhaled solution (TOBI) and intravenous injection. The tobramycin concentrations in airways after 300-mg dosage TOBI inhalation can reach 1,000  $\mu\text{g}$  per g of sputum (5, 6). This concentration is in the range of 10 to 1,000 times of the minimal inhibitory concentration (MIC) for *P. aeruginosa* clinical isolates tested *ex vivo* (6). However, even with such high tobramycin concentrations, chronic *P. aeruginosa* infections are rarely eradicated (6). This is true even when the infecting bacteria are antibiotic sensitive, as is the case early in disease (7).

One possible reason for *P. aeruginosa* persistence *in vivo* could relate to the time dependence of local concentrations of tobramycin experienced by *P. aeruginosa* in CF patient airways. Many factors, including inflammatory responses, blood and lymphatic circulations, and air flow distribution (for inhaled antibiotics), can alter the local antibiotic concentrations. In addition, *P. aeruginosa* cells can form biofilms in CF lungs and other infection sites (8), and biofilm exopolysaccharide layers may slow the diffusion of tobramycin (9, 10). *P. aeruginosa* cells in the inner layers of biofilms may experience lower concentrations and more gradual increase of tobramycin levels than those in outer layers (10, 11). Furthermore, even if final tobramycin concentration levels inside the biofilm eventually grow to match the highest levels experienced elsewhere, bacteria in these inner regions have experienced a

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Received March 27, 2015, and in revised form, March 27, 2015

Published, MCP Papers in Press, May 27, 2015, DOI 10.1074/mcp.M115.050161

Author contributions: X.W., P.K.S., C.M., and J.E.B. designed the research; X.W., K.H., B.J.S., and J.D.C. performed the research; C.R.W. contributed new reagents or analytic tools; X.W., K.H., C.Z., J.K.E., and J.E.B. analyzed the data; and X.W., P.K.S., C.M., and J.E.B. wrote the paper.

slower increase, during which time proteome levels could be altered to promote the “adapted resistant state” (12). Adaptive resistance can also be induced in planktonic (free-living) *P. aeruginosa* (13, 14), and conventional MIC assays are not designed to measure this.

Once induced, the adaptive resistance confers bacteria higher resistance to antibiotic treatments (13, 14) and is associated with decreased clinical antibiotic treatment efficacy (15). Interestingly, the adaptive resistance is time dependent and reversible. Typical adaptive resistance was observed starting 1 h after antibiotic exposure, and the drug susceptibility was regained after 36 h intervals (14, 15). Thus, adaptive resistance mechanisms may contribute in part to the disparity of *in vivo* persistence and *ex vivo* susceptibility to antibiotics in MIC tests.

As an initial step toward defining adaptive resistance mechanisms, we investigated the time- and concentration-dependence of *P. aeruginosa* proteome response to tobramycin in planktonic conditions. Since the most effective protective responses may operate before killing begins and the rate of change of drug levels is likely to depend on ambient conditions, we studied bacteria exposed to low, subinhibitory levels of tobramycin (0.1, 0.5, and 1.0  $\mu\text{g/ml}$ ) at a range of time points (15, 60, 120, and 360 min) after exposure. The candidate proteome marker of *P. aeruginosa* for tobramycin response, heat shock protein *lbpA*, was further investigated with genetic mutagenesis and MIC assays.

#### EXPERIMENTAL PROCEDURES

*P. aeruginosa* Strains—*P. aeruginosa* strain MPAO1 (16) was used for proteome analysis. Single transposon insertion mutants were obtained from the two-allele mutant library (17). The specific mutants examined were as follows: PA0779 (PW2411, PW2413, PW2414), *clpB* (PW8651, PW8652), *hslV* (PW9485, PW9486, PW9487), and the transposon-containing control PA3303 (18).

Two *ibpA* loss-of-function mutant alleles were used in this study. The first *ibpA::luxCDABE* was constructed in two recombination steps by replacing wild-type *ibpA* with an *ibpA-lux* gene fusion carried on plasmid pEX19Tc (19). The *lux* insertion site was at the N-terminal fifth amino acid of *ibpA*. The structure of the constructed allele was confirmed by PCR, and the absence of *lbpA* expression was confirmed by the selected reaction monitoring (SRM)<sup>1</sup>. The *lux* control strain *PKH181* was constructed by insertion of pUC18-mini-Tn7T-Gm-*lux* vector (without a cloned promoter) into MPAO1, according to methods of Choi and Schweizer (20). The second *ibpA* mutant ( $\Delta ibpA$ ) corresponded to an in-frame deletion that joined sequences coding for the N-terminal fifth amino acid to the fourth amino acid from the C terminus. The deletion was generated by recombination in MPAO1 with pEX19Tc plasmid carrying the deletion allele. The in-frame  $\Delta ftsH$  deletion mutant has been previously described (21).

To generate the double mutants (*ibpA/PA0779*, *ibpA/clpB*, and *ibpA/hslV*), chromosomal DNA isolated from the transposon-containing strains was transformed into *ibpA-lux* insertion inactivation mutant

via lambda red recombination. Positive transformants were selected by tetracycline resistance as previously described (22).

*Tobramycin Treatment for Proteomics Samples*—MPAO1 cells were grown in salt-free LB containing 10 g tryptone, 5 g yeast extract per liter at 37 °C to optical density  $\text{OD}_{600} = 1.0$ , and aliquotted to 5 ml volumes. For the dosage-dependent treatments, tobramycin (Sigma-Aldrich, St. Louis, MO) at 0.1, 0.5, or 1.0  $\mu\text{g/ml}$  (final concentration) was added to the cell cultures with shaking for 60 min at 37 °C. For the time course treatments, cells were treated with 1.0  $\mu\text{g/ml}$  tobramycin (final concentration) were exposed for 15, 60, 120, and 360 min at 37 °C. Cells harvested at time zero (at the time that tobramycin treatment started) were used as controls. Three biological replicates for dosage treatments and two biological replicates for the time course were analyzed.

*Tobramycin MIC Assays*—Minimum inhibitory concentration (MIC) assays were performed as described by Lee *et al.* (18). Single colonies were inoculated into 96-well plates containing 250  $\mu\text{l}$  salt-free LB per well and grown for 18 h at 37 °C in a humidity chamber. 200  $\mu\text{l}$  of these cultures were removed, added to a new plate, and 10-fold dilutions were made with salt-free LB. The salt was not added in LB for initial liquid culture because  $\Delta ftsH$  mutant is salt sensitive (21), and salt-free conditions resulted in relatively uniform growth for strains. Diluted cells were allowed to grow for 90 min at 37 °C in a humidity chamber, at which point cells were spotted to tobramycin LB agar (including 137 mM NaCl) to deposit approximate  $10^5$  cells per spot. Plates were incubated at 37 °C for 18 h and photographed. MIC was defined as the lowest antibiotic concentration preventing the lawn growth of the spotted cells.

*Protein Extraction of P. aeruginosa Cells*—MPAO1 cells were harvested by centrifugation at 3,500 rpm for 10 min. Cell pellets were resuspended with lysis buffer (50 mM ammonium bicarbonate, pH 8.0; 8 M urea; and 10 mM DTT). Cells were lysed by incubation in lysis buffer on ice for 30 min, with vortexing for 30 s every 5 min. Cells in lysis buffer underwent one  $-80$  °C freeze-thaw cycle to maximize cells lysis. Protein extracts were briefly sonicated to shear DNA. Cell debris was removed by centrifugation at 20,000 g for 20 min. Extracted protein concentration was measured with Bradford assay (Thermo Fisher Scientific, Waltham, MA).

100  $\mu\text{g}$  proteins from each extraction were diluted twofold with digestion buffer (40 mM ammonium bicarbonate, 5% acetonitrile) for alkylation by 10 mM iodoacetamide for 30 min in the dark. The mixture then was further diluted to urea concentration less than 1.5 M with digestion buffer for overnight trypsin digestion (Promega, Madison, WI). Peptides were purified with Sep-Pak cartridges (Waters Corporation, Milford, MA), dried, and resuspended with 0.1% formic acid. 1  $\mu\text{g}$  of tryptic digest was loaded in each LC-MS/MS analysis.

*LC-MS/MS Analysis*—The proteomics samples were analyzed with NanoAcquity UPLC (Waters) coupled to a linear quadrupole ion trap mass spectrometer (LTQ-XL from Thermo Fisher Scientific). The column length of 3 cm trap column packed in house with 200 Å C18 magic beads (Bruker-Michrom Inc, Auburn, CA) and 30 cm analytical column packed with 100 Å C18 magic beads were used in all LC/MS-MS analysis to maintain the uniformity of peptide chromatography profiles. A 120 min linear gradient (5–35% acetonitrile) at a flow rate of 300 nl/min was used, and peptides were ionized by electrospray.

Data-dependent acquisition (DDA) mode was used in the spectral counting analysis. The top five most intense ions in the MS<sup>1</sup> scan were selected for tandem mass spectrometry analysis.

For the selected reaction monitoring (SRM) analysis, scan-type SRM was used. Targeted peptides and transition ions were manually selected from DDA identification spectra. Ion intensity and preferentially larger fragment masses were the main selection criteria. Five to eight fragment ions were included per peptide. SRM instrument set-

<sup>1</sup> The abbreviations used are: SRM, selected reaction monitoring; DDA, data dependent acquisition; emPAI, exponentially modified protein abundance index; CF, cystic fibrosis; Tob, tobramycin; MIC, minimum inhibitory concentrations; BSA, bovine serum albumin.

tings for this study were standardized, which included parent ions isolation width 3 *m/z*, transition ions scan width of 1 *m/z*, collision-induced disassociation (CID) collision energy 35, activation Q 0.25, activation time 30 ms, and automatic gain control (AGC) MS<sup>n</sup> value 1 × 10<sup>4</sup>, maximum ion time 500 ms.

**Data Analysis**—The DDA raw files were converted to mgf peak list files using ReAdW (Ver. 4.3.1) (23). The data were searched in Mascot (version 2.3.02) against the *Pseudomonas aeruginosa* (PAO1) database (consisting of 5,680 genes or splicing fragments, annotation version 2012-Nov) (24). The search parameters included 3.0 Da precursor mass tolerance, 0.6 Da fragment mass tolerance, fixed modification cysteine carbamidomethylation, variable modification methionine oxidation, digestion enzyme trypsin, and maximum missed cleavages 2. The peptide expectation value 0.05 was used as the cutoff, and the false discovery rate (FDR) was determined by searching the same mgf file in the randomized decoy database in Mascot. The FDR of the presented dataset was ~1%.

The Exponentially Modified Protein Abundance Index (emPAI) values derived from Mascot search were used for spectral quantitation analysis (25). The data of a total of seven LC-MS/MS analyses for each dosage treatment and five LC-MS/MS analyses for each time course treatment were used for spectral counting statistical analysis.

We required proteins for quantitation analysis to be quantified minimally by three unique peptides from at least two biological replicates. Proteins with significant fold changes ( $P \leq 0.05$ ) as well as proteins with no detection in one state but detected more than four times in another state were considered statistically altered proteins and were further subject to gene ontology analysis.

The SRM data in this experiment were processed in Skyline (version 1.30) (26). Selection of chromatogram peaks for quantitation was manually verified. Peak smoothing method Savitzky–Golay was used. Chromatogram extracted peak areas for transition ions were summed up to yield total peak areas. The ratio of the total peak areas represented the abundance ratio of the peptides. Peptide ratios of each protein were further averaged to estimate the protein fold changes. The quantitation normalization was based upon equivalent total injection amount (1 μg) in each SRM run. A quantitation linear range of four orders of magnitude was shown (down to femtomole) (Fig. S1).

## RESULTS

### *Evaluation of P. aeruginosa* Proteome Quantitation Methods—Spectral Counting and Selected Reaction Monitoring (SRM)

We first evaluated the reliability of our proteome quantitation methods. Spectral counting quantitation allows broad coverage of *P. aeruginosa* proteins and was used to identify potential protein targets. More than 1,000 *P. aeruginosa* proteins were examined in these assays (Fig. 1A). The Exponentially Modified Protein Abundance Index (emPAI) (25) values of 921 proteins commonly observed in all cell samples were used to calculate the fold change for each protein. The linear quantitation range of emPAI for the *P. aeruginosa* proteome was assessed with bovine serum albumin (BSA) spike-in assays, and a linear range over three orders of magnitude (down to 1:10,000 dilution w/w) was observed for BSA proteins (Fig. S1A). The detected coefficient of variation (CV) of emPAI for these proteins appeared protein-dependent as one might expect and scored a median CV of ~40% (Fig. S2). The majority of protein emPAI ratios localized at the center of the scatter plot ( $\log_2 1 = 0$ ), while the proteins with altered abundance

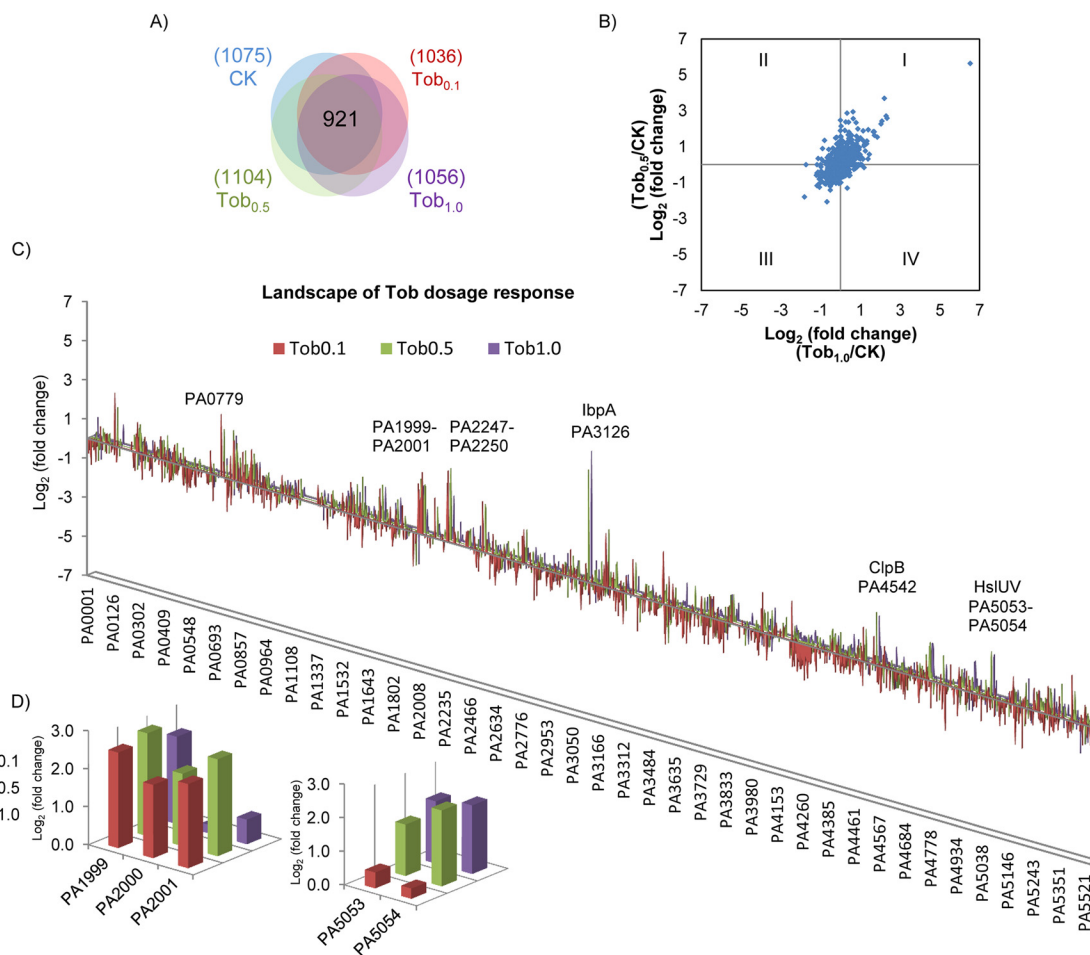
appeared to be consistent between treatments, showing enrichment in Quadrant I and III of the Cartesian plane (Fig. 1B).

Selected reaction monitoring (SRM) achieves high specificity and accurate quantitation and was used for validation of targets identified by spectral counting. The linear quantitation range of SRM was also accessed by BSA spike-in assays, a linear quantitation range of four orders magnitude was achieved for BSA peptides (Fig. S1B), which is consistent with other reports (27, 28).

To determine if the protein-level changes measured by spectral counting or SRM were consistent, quantitation data of a set of 10 *P. aeruginosa* proteins (including 49 SRM peptide assays) at different tobramycin treatment conditions measured with both methods were compared. A general correlation ( $R^2 = 0.61$ ) was observed (Fig. S3). Therefore, emPAI spectral counting and SRM provide consistent measurements for *P. aeruginosa* proteins and allow us to systematically evaluate the proteome response of *P. aeruginosa* to tobramycin antibiotic treatment.

**Concentration- and Time-Dependent Proteome Changes of *P. aeruginosa* in Response to Tobramycin Treatment**—Since we are interested in the adaptive proteome response as a mechanism in protecting cells from antibiotic killing, *P. aeruginosa* cells grown under planktonic conditions were challenged with low and subinhibitory levels of tobramycin (0.1, 0.5, and 1.0 μg/ml) for periods of (15, 60, 120, and 360 min), and the proteome response was assessed. We constructed proteome landscape plots to visualize the global protein fold change as a function of gene locus and treatment dosage (Fig. 1C) or time (Fig. 2A). These plots show groups or clusters of genes that appear similarly increased or decreased with tobramycin treatment concentration or duration. Bar heights in the plot indicate the level of protein abundance change, and grouping of the bars implies the coregulation of protein changes among neighboring genes. The landscape readily distinguished heat shock protein lbpA (PA3126) as the highest fold change protein. It increased 50-fold at 0.5 μg/ml dosage and 90-fold at 1 μg/ml dosage and was observed with increased levels throughout the time course measurement from 60 min to 360 min. In addition to lbpA, the landscape also captured significant increases in PA0779, PA1999–2001 (DhcA-B, AtoB), PA2247–2250 (BkdA1-A2-B, LpdV), PA4542 (ClpB), PA4759–4762 (DapB, DnaJ-DnaK-GrpE), and PA5053–5054 (HslU-V) (Figs. 1C and 2A). Based on emPAI quantitation, many of these increases were fourfold or higher. Examination of their induction patterns revealed concentration- and time-dependent expression (Figs. 1D and 2B).

To validate the tobramycin dosage effects on the observed proteome induction, we used SRM to analyze the dosage-dependent levels of a number of these proteins (Fig. 3A–3C). The proteins can be categorized into three groups based on their biological functions, heat shock proteins, proteases, and amino acid catabolic enzymes (29). Heat shock proteins (lbpA, ClpB, DnaJ, DnaK, GrpE, and HtpG) were significantly

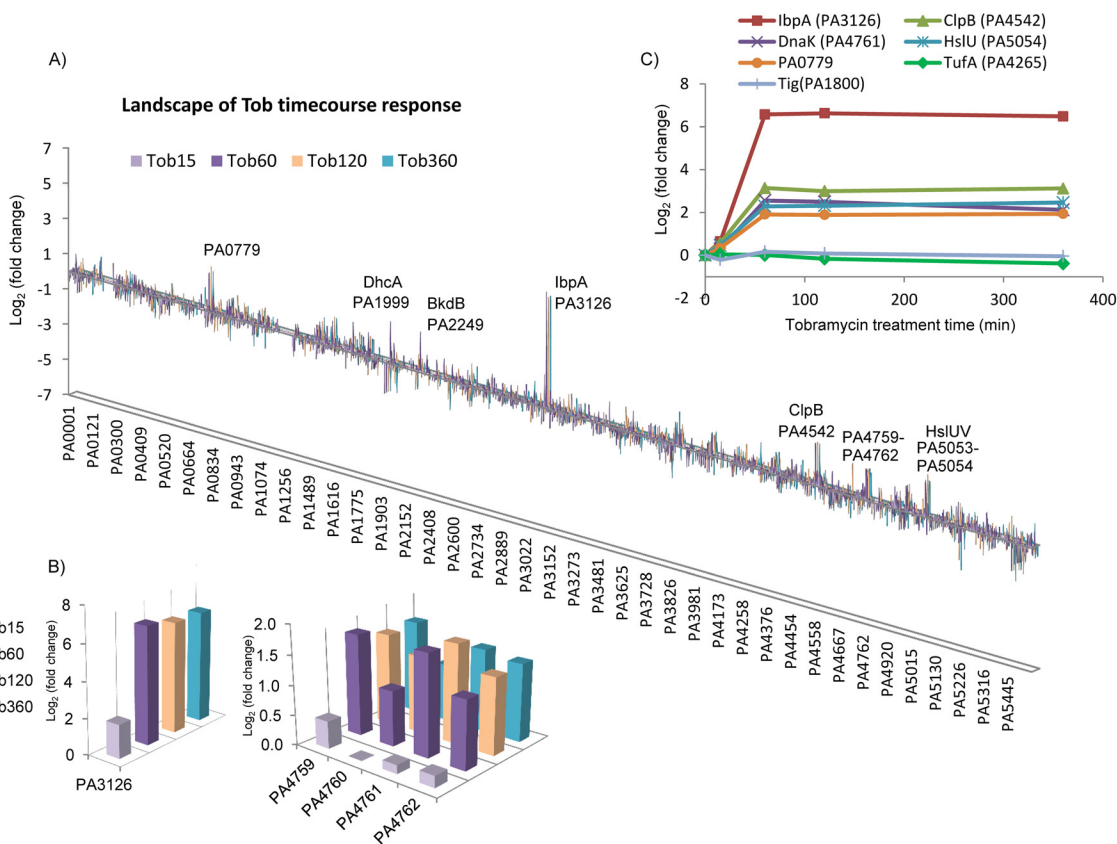


**FIG. 1. Landscape of *P. aeruginosa* proteome detection and quantitation.** (A) Venn diagram showing over 1,000 *P. aeruginosa* proteins were quantified in each cellular state, including 921 common proteins quantified in all four dosages. Tobramycin dosage level is indicated with colors. CK: no tobramycin treatment ( $T_0$  collected cells);  $Tob_{0.1}$ ,  $Tob_{0.5}$ , and  $Tob_{1.0}$ : tobramycin treatment of 0.1, 0.5, and 1.0  $\mu\text{g/ml}$ . (B) The distribution of  $\log_2$  (fold change) of the 921 common proteins. Comparison of  $Tob_{1.0}/CK$  is shown at x axis, and  $Tob_{0.5}/CK$  at y axis. (C) Landscape of protein abundance changes in *P. aeruginosa* during tobramycin dosage treatment. X axis: *P. aeruginosa* gene locus. Y axis:  $\log_2$  protein fold change of  $Tob_{0.1}/CK$ ,  $Tob_{0.5}/CK$  or  $Tob_{1.0}/CK$ . The bar colors illustrate the concentration of Tobramycin in experiment and bar heights indicate the change in each protein level. The genes or operons with the most dramatic increase in protein abundance were IbpA, ClpB, PA0779, HslU-V, PA1999-PA2001, and PA2247-2250. (D) Inset of proteome landscape showing concentration dependent changes of PA1999-PA2001 and PA5053-5054.

increased by tobramycin at all three dosage levels, with the increases at the higher dosage levels (0.5 and 1.0  $\mu\text{g/ml}$ ) most marked. IbpA was again observed with largest fold change, with 84-fold induction at 0.5  $\mu\text{g/ml}$  and 95-fold at 1.0  $\mu\text{g/ml}$  tobramycin treatment, consistent with the quantitation from spectral counting. For the proteases (PA0779, HslU, and HslV), larger increases were also observed at the higher dosage treatment (0.5 and 1.0  $\mu\text{g/ml}$ ), while FtsH and PA1803 (Lon protease) exhibited similar increases at all three dosage levels. For the amino acid catabolic enzymes, only DapB was significantly increased at all three tobramycin concentrations, while DhcA, DhcB, BkdB, and Sdr were significantly increased only at 0.1 and 0.5  $\mu\text{g/ml}$  treatments. For the control proteins (TufA, Tig, and PolA), no significant abundance level change was observed at any dosage treatment (Fig. 3D).

To interrogate the time dependence of inducible proteome changes, we used SRM to measure the tobramycin-induced proteins (IbpA, ClpB, DnaK, PA0779, and HslU) at the time points of 0, 15, 60, 120, and 360 min after 1.0  $\mu\text{g/ml}$  treatment (Fig. 2C). Interestingly, for all five proteins, the rate of increase was most marked from 15 min to 60 min after tobramycin treatment and leveled off at 120 and 360 min. The temporal characteristics of IbpA increase resembles that seen after heat treatment (30). Again, the control proteins TufA and Tig did not significantly change.

We compared the proteome changes with published transcriptome results (18) in order to further comprehend the temporal regulation of proteome response. The transcriptome analysis by Lee *et al.* (18) was performed with similar tobramycin treatment conditions (1  $\mu\text{g/ml}$  tobramycin 15 min treat-



**FIG. 2. Time-dependent proteome response of *P. aeruginosa* to tobramycin.** (A) Landscape of *P. aeruginosa* proteome response showing genes/operons that were most drastically changed in tobramycin time course treatment. X axis: *P. aeruginosa* gene locus. Y axis: log<sub>2</sub> protein fold change of Tob<sub>15min</sub>/CK, Tob<sub>60min</sub>/CK, Tob<sub>120min</sub>/CK, and Tob<sub>360min</sub>/CK. Duration of tobramycin treatment is indicated with bar colors. PA0779, DhcA, BkdB, IbpA, ClpB, PA4759-PA4762, HslU-V were up-regulated, consistent with the results in the dosage treatment shown in Fig. 1C. (B) Inset of the proteome landscape showing time-dependent changes of IbpA and PA4759-PA4762. (C) Selected reaction monitoring (SRM) analysis for proteins that were dramatically increased in the tobramycin time course treatment. The SRM protein abundance ratio is shown, which represents the average of the peptide ratios from the protein. Interestingly, rapid increase of heat shock proteins and proteases occurred from 15 min to 60 min after tobramycin treatment, and the abundance level sustained till 360 min (last time point measured).

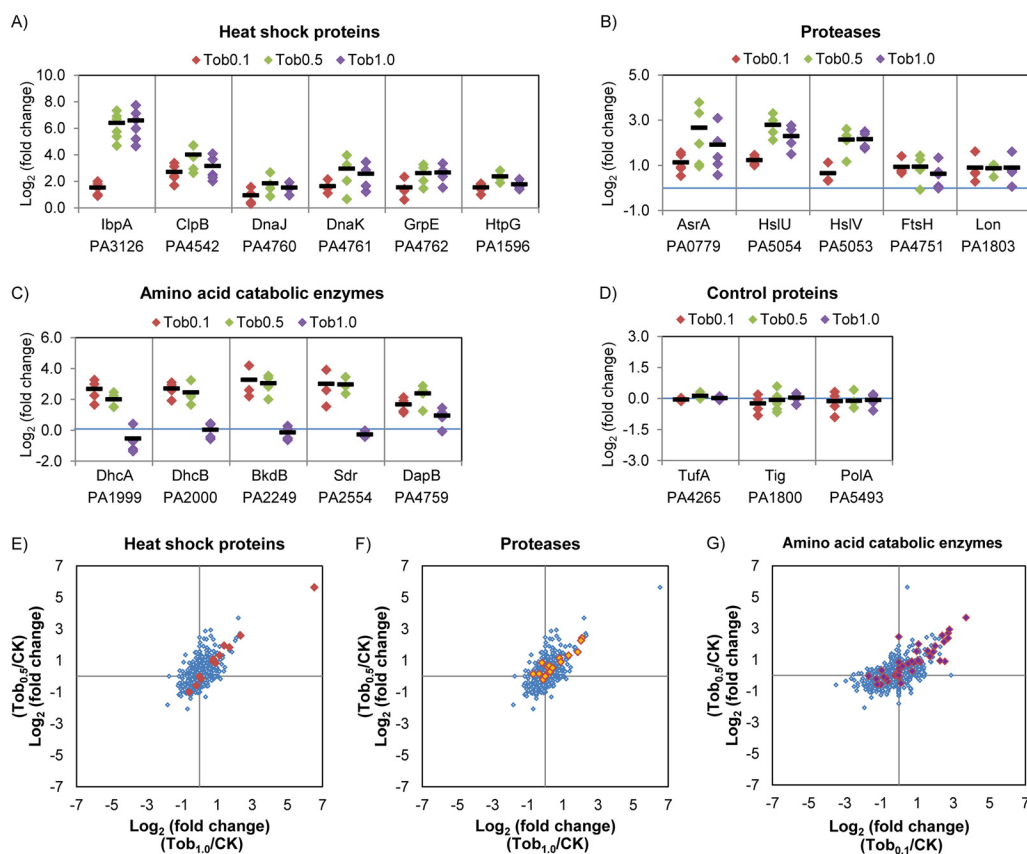
ment, OD<sub>600</sub> = 1.0 MPAO1 cells) as in the present study so provided a useful reference fold change of RNAs. As shown in Fig. S4, 11 genes were found to be tobramycin induced in both transcriptome and proteome analyses, including seven of the genes that corresponded to proteins showing the greatest increase in proteome changes (Figs. 1C and 2A): PA0779, *ibpA*, *clpB*, *dnaJ*, *grpE*, *hslU*, and *hslV*.

Other genes that were observed to increase at the RNA level were not detected at the protein level, which may be due to low protein abundance levels, proteome sample collection, or regulatory changes that dissociate mRNA and protein expression. For genes identified as induced using both approaches, smaller changes were generally observed at the protein level than RNA level at the 15 min time point. This may reflect the longer time required to increase protein than RNA levels after transcriptional induction. However, at 60 min, the levels of protein fold changes became more pronounced. In particular, the increase of five genes (*ibpA*, *clpB*, *hslU-V*, and PA0779) was higher at the protein level than RNA level. The

additional increase suggests the importance of protein-level regulation of tobramycin response in *P. aeruginosa*.

**Functional Enrichment of Tobramycin-Induced Proteins in *P. aeruginosa***—To further define the proteome response induced by tobramycin, we used a one-way unpaired *t* test (also known as a one-way ANOVA test) to distinguish statistically significant changes of *P. aeruginosa* protein levels in different treatment states. In addition to the statistical cutoff (*P* value ≤ 0.05), proteins with significantly altered levels were also required to be repetitively quantified minimally by three unique peptides from at least two of the seven replicates in dosage treatments or two of the five replicates in time course treatments. Proteins that were found by spectral counting with statistically significant level changes but lower than twofold changes were further validated by other quantitation methods (Fig. S5).

Using this approach, as many as 200 proteins appeared with statistically significant altered levels in different tobramycin-



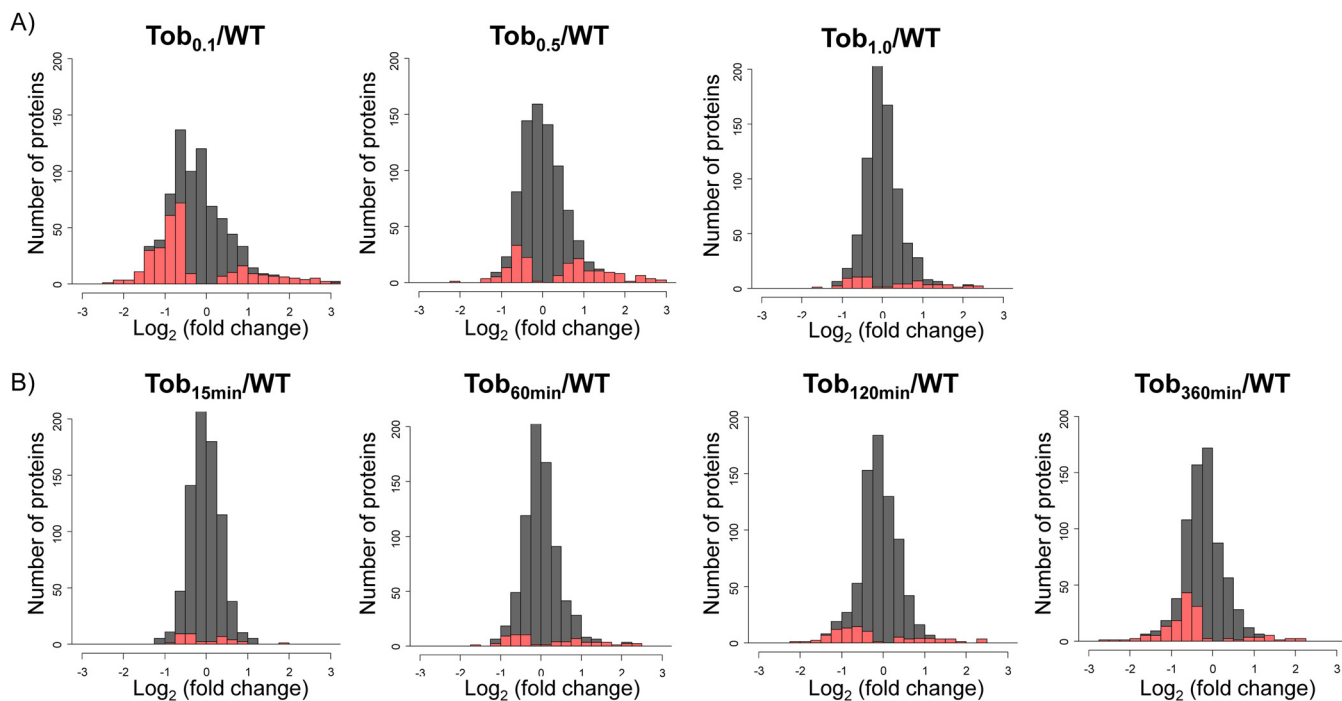
**FIG. 3. Confirmation of the up-regulation of heat shock proteins, proteases and amino acid catabolic enzymes in *P. aeruginosa* during tobramycin treatments with selected reaction monitoring (SRM).** (A–D) SRM analysis confirmed heat shock proteins (IbpA, ClpB, DnaJ–DnaK–GrpE, and HtpG) and proteases (PA0779, HslU–V, FtsH, and PA1803) were significantly increased in tobramycin treatment of all three dosages, and the extent of increase was more dramatic at the higher dosages 0.5 and 1.0  $\mu\text{g/ml}$ . Amino acid catabolic enzymes DhcA, DhcB, BkdB, and Sdr were significantly increased at 0.1 and 0.5  $\mu\text{g/ml}$  dosages, and enzyme DapB was significantly increased in all three dosages. Control proteins TufA, Tig, and PolA did not show significant fold changes in tobramycin treatment. Three to five SRM peptides (each containing 5–8 transition ion quantitation) were used to quantify each protein. The diamond represents the measured fold change for each peptide. The black line indicates the fold change of the protein by averaging the fold change of all the SRM peptides. (E–G) Scatter plot showing the induction for heat shock proteins, proteases, and amino acid catabolic enzymes. The spectral counting data log<sub>2</sub> (fold change) of Tob<sub>1.0</sub>/CK (x axis) and Tob<sub>0.5</sub>/CK (y axis) are shown in (E) and (F), and log<sub>2</sub> (fold change) of Tob<sub>0.1</sub>/CK (x axis) and Tob<sub>0.5</sub>/CK (y axis) are shown in (G). The whole quantified proteome is provided in blue diamonds as background. The heat shock proteins are highlighted in red, proteases in orange, amino acid catabolic enzymes in purple. Proteins in these three pathways contributed most of the largest fold changes in the proteome.

cin treatment states. Consistent with SRM quantitation that showed tobramycin dosage- and time-dependent effects on individual protein level changes (Figs. 2C and 3A, 3B, and 3C), the global proteome induction also showed systematic tobramycin dosage and duration effects (Fig. 4). For instance, a higher number of proteins (and higher percentage of proteome) was observed with significant altered levels with 0.1  $\mu\text{g/ml}$  tobramycin compared with the higher dosage 1.0  $\mu\text{g/ml}$ , while 0.5  $\mu\text{g/ml}$  treatment exhibited intermediate changes (Fig. 4A). Extending 1.0  $\mu\text{g/ml}$  tobramycin treatment to 360 min also gradually increased the number of proteins that were observed with statistically significant decreased levels (Fig. 4B).

Interestingly, examination of the top 30 proteins with greatest fold changes from the three tobramycin dosage treat-

ments indicated that very few proteins were common among the three dosage levels, and the proteins that were common in two dosage levels were more likely to be shared between more similar doses (*i.e.* 0.1 and 0.5  $\mu\text{g/ml}$  or 0.5 and 1.0  $\mu\text{g/ml}$ ) (Fig. 5A). These results suggest that bacterial responses become less similar as the differences in the tobramycin dose increase. Because significant proteome induction was observed at the very low dosage levels (0.1  $\mu\text{g/ml}$ , 214 nM, 1/10 MIC), it also illustrates that *P. aeruginosa* cells can sense a very low amount of tobramycin molecules, supporting the notion that cells first exposed to lower-level tobramycin treatment could yield acclimation phenotypes.

We next characterized the gene ontology of proteins that were observed to increase with tobramycin (Fig. 6A–6C). At higher tobramycin dosages of 0.5 and 1.0  $\mu\text{g/ml}$ , protein



**FIG. 4. Global protein fold change showing the dosage- and time-dependent effects of tobramycin treatments.** The  $\log_2$  protein fold change by spectral counting was shown. The statistically altered proteins ( $P \leq 0.05$ ) are highlighted in red, and the entire quantified proteome is shown in gray. (A) Dosage-dependent effects. The lower dosage tobramycin treatment 0.1 and 0.5  $\mu\text{g/ml}$  for 60 min induced wider proteome changes compared with 1.0  $\mu\text{g/ml}$ . (B) Time-dependent effects. Extending 1.0  $\mu\text{g/ml}$  tobramycin treatment to 360 min gradually increased the number of proteins that were statistically significantly altered.

folding function (heat shock proteins) and proteolysis function (proteases) were significantly overrepresented as compared with other gene ontology categories. These functions were temporally up-regulated at 60, 120, and 360 min with 1.0  $\mu\text{g/ml}$  tobramycin treatments. The up-regulation of protein folding and proteolysis functions highlights the importance of protein quality control after higher tobramycin exposures (21, 31). At lower dosages of 0.1 and 0.5  $\mu\text{g/ml}$ , amino acid metabolic/catabolic (but not biosynthetic) pathways were statistically overrepresented. Thus, different functional activation was observed dependent on tobramycin treatment dosages, and one would anticipate differences in cell phenotypes when specific subsets of proteins were up-regulated (32).

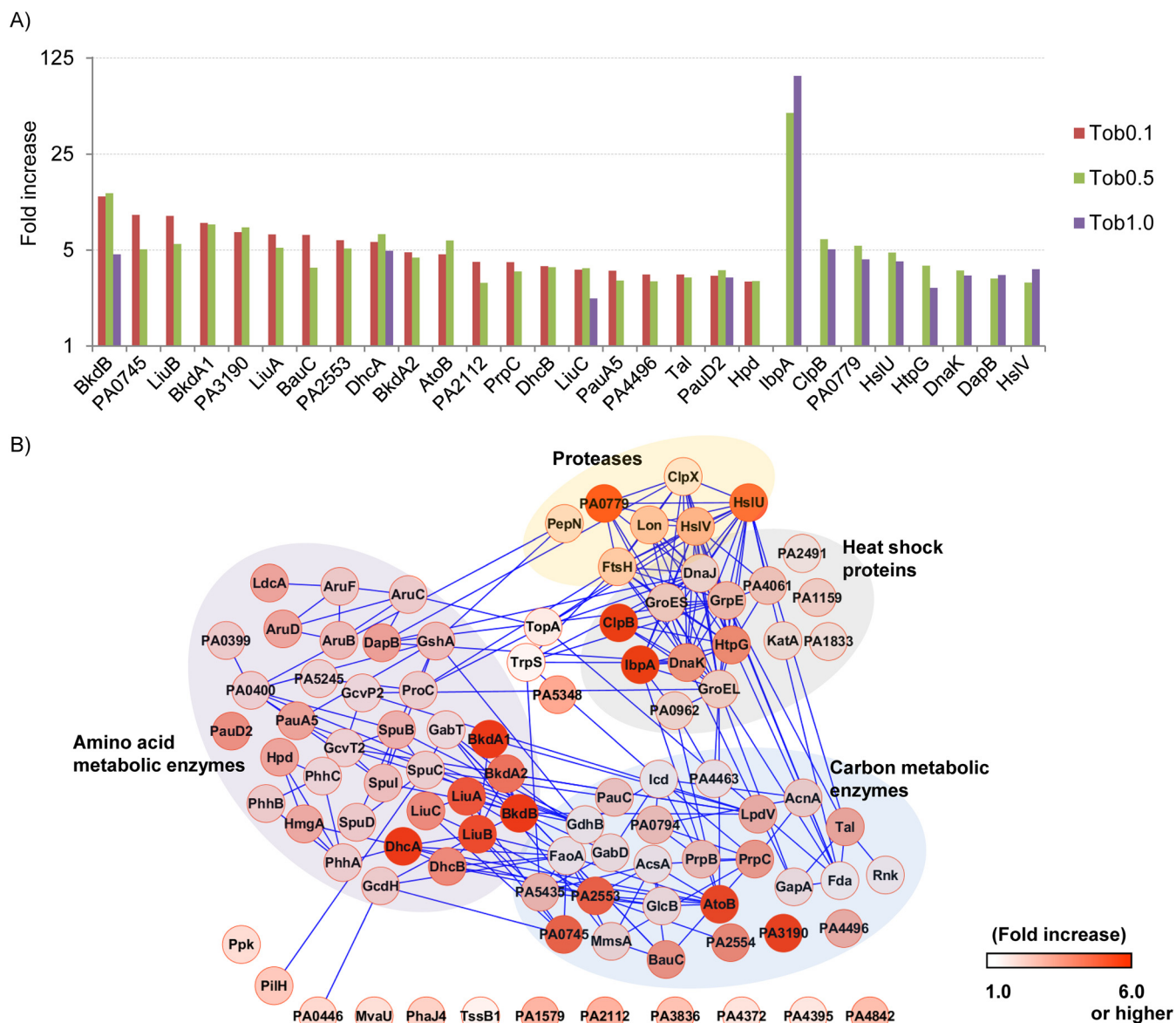
To further interrogate the response of amino acid metabolic enzymes, the metabolic subpathways for different amino acid families were analyzed (Fig. 6B). Interestingly, overrepresentation for the amino acid metabolic pathway was largely due to the catabolic pathways rather than the synthesis pathways. In terms of the amino acid family, the induction was common to enzymes in all five families, but the increases in the aromatic and branched chain amino acid families were more pronounced than the other families and achieved level changes with  $P \leq 0.01$ . Such increases were most apparent with the low level of tobramycin treatment 0.1  $\mu\text{g/ml}$ .

To characterize the functional relationship of tobramycin-induced proteins, we analyzed the gene coregulation referenced from the STRING database (33). The induced pro-

teins observed with 0.5  $\mu\text{g/ml}$  tobramycin dosage are shown in Fig. 5B. Fourteen proteins related to heat shock and proteases functions were represented, and they were tightly connected in the network. Amino acid metabolic enzymes PA1999–2001 (DhcAB, AtoB) in carnitine catabolism, PA2013–2015 (LiuABC) in leucine degradation, and PA2247–2250 (BkdA1A2B, LpdV) in valine degradation (29) were also significantly induced. Although many carbon metabolic enzymes were also increased by tobramycin, the enrichment of the pathway was not statistically significant (Fig. 5B), considering the large base number of carbon metabolic enzymes in *P. aeruginosa* genome.

To verify the widespread increase for heat shock proteins, proteases, and amino acid catabolic enzymes, we examined the protein fold change of all the proteins quantified in the three pathways. As shown in Figs. 3E–3G, proteins in the three pathways contributed most of the highest fold changes in *P. aeruginosa* proteome, indicating that the three pathways are critical to *P. aeruginosa* tobramycin response.

As a complementary analysis, we also analyzed the tobramycin repressed proteome in *P. aeruginosa*. Proteins with levels that decreased during tobramycin treatment also showed dosage and time dependent effects (Fig. 4). Those proteins were functionally enriched in protein synthesis, nucleotide metabolism, tricarboxylic acid (TCA) carbon metabolism and energy derivation, and electron transport activities (Fig. S6). Gene coregulation analysis demonstrated that the



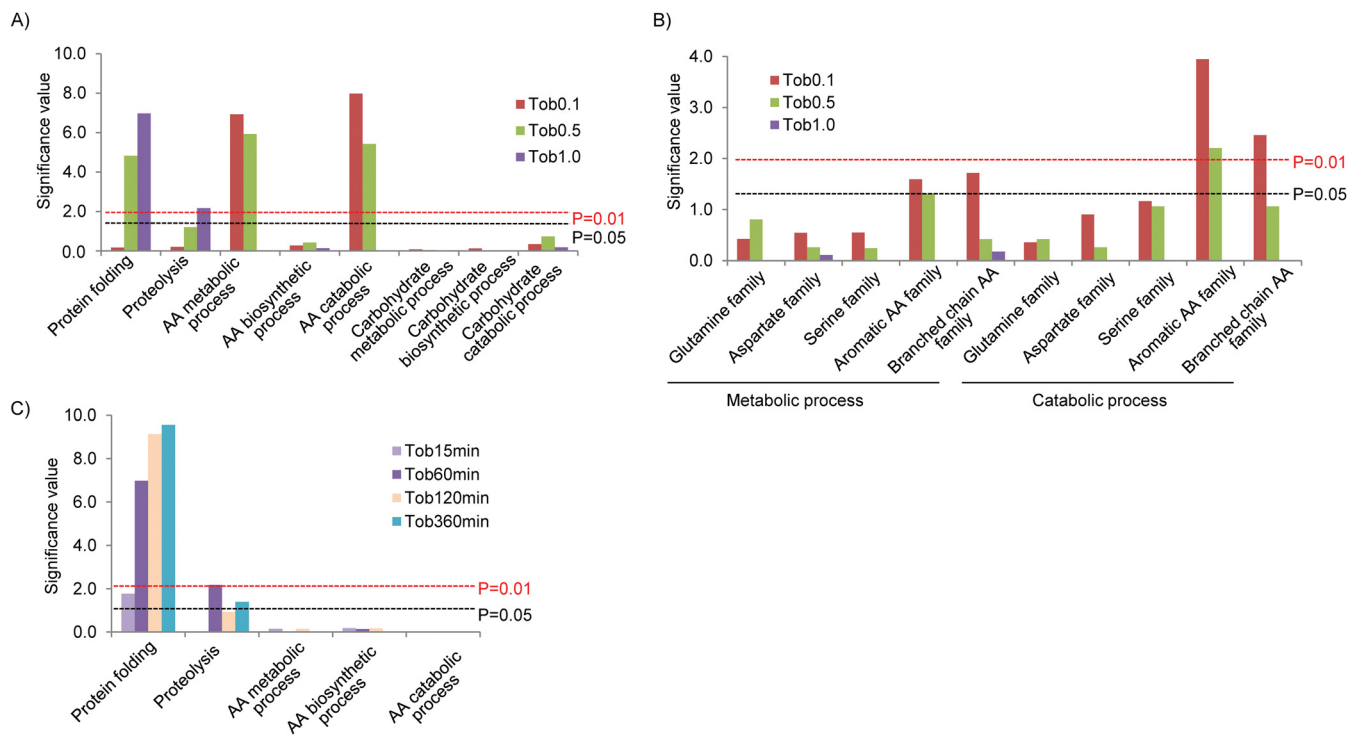
**FIG. 5. Dosage-dependent induction of proteome subsets and quantitative proteome interaction network.** (A) The top 30 proteins with greatest fold changes from the three tobramycin dosage treatments were compared. The emPAI fold changes of those common in at least two dosages were plotted. Interestingly, very few proteins were common to be the top 30 in all three dosages, and the proteins that were common in two dosages were more likely to be shared between 0.1 and 0.5  $\mu\text{g/ml}$  or 0.5 and 1.0  $\mu\text{g/ml}$ . (B) Quantitative proteome interaction network for proteins significantly increased ( $P \leq 0.05$ ) in 0.5  $\mu\text{g/ml}$  tobramycin treatment. The network was visualized with Cytoscape (version 2.8) (47). Each node indicates the protein identity, and the extent of increase is shown in red. The protein fold change from spectral counting was used. The edges indicate the protein co-regulation evidence of the connected nodes referenced from STRING database (33). Four functional clusters are observed in the network.

down-regulated proteins were functionally connected (Fig. S7).

**Tobramycin Sensitivity Phenotypes of *ibpA* Single and Double Mutants**—Because *IbpA* levels were increased by the greatest amount in the observed proteome response to tobramycin, we constructed *ibpA* gene-inactivation mutants to determine if the absence of *IbpA* proteins may result in changes of *P. aeruginosa* tobramycin sensitivity. Surprisingly, neither *ibpA-lux* inactivation mutant nor  $\Delta ibpA$  in-frame dele-

tion mutant exhibited significant changes in tobramycin sensitivity compared with MPAO1 WT (Table I and Fig. S8). This observation suggested that either *IbpA* is not directly involved in tobramycin resistance or the function of *IbpA* is compensated by other heat shock chaperones (34), leading to no obvious changes in tobramycin sensitivity. To examine these possibilities, we further constructed the double mutants of *ibpA* with *clpB*, (another heat shock protein) or PA0779 (*lon* protease) or *hslV* (ATP-dependent protease subunit), all of





**FIG. 6. Gene ontology analysis of the significantly up-regulated proteins in *P. aeruginosa* by tobramycin.** (A) Up-regulated proteins are enriched in the pathways of protein folding, proteolysis, and amino acid metabolic (catabolic) process. Significance test was performed at BiNGO (version 2.44) (48) with protein annotations from Gene Ontology (49). Significance level was determined by hypergeometric test with Benjamini–Hochberg FDR correction. The resultant values were converted to y axis significance value by  $-\log_{10}$ , and the corresponding 0.01 and 0.05 significance levels are indicated in the figure. (B) The amino acid catabolic pathways rather than biosynthetic pathways were significantly up-regulated by tobramycin. The catabolism of all amino acid families was up-regulated, but the increase in aromatic and branch chain amino acid family was more significant. (C) Protein folding and proteolysis pathways were also up-regulated in tobramycin time course treatment.

which showed significantly increased protein levels with tobramycin treatments (Figs. 3A and 3B) and exposure time (Figs. 2A and 2C).

Detectable increases in tobramycin sensitivity were observed for most of the *ibpA* double mutants. The double mutants *ibpA/hsIV* exhibited tobramycin MIC of 0.25  $\mu\text{g/ml}$ , which was twofold more sensitive as compared with single mutant *ibpA* and *MPAO1 WT* (Table I and Fig. S8). Two of the *ibpA/hsIV* strains also showed twofold enhancement in tobramycin sensitivity compared with *hsIV* single mutant. For double mutants *ibpA/clpB*, *ibpA/PA0079*, tobramycin MIC of 0.5  $\mu\text{g/ml}$  was identified. Although the extent of tobramycin sensitivity change was relatively small, close examination of colony lawn density showed that *ibpA/clpB* and *ibpA/PA0079* strains were almost absent at tobramycin MIC of 0.5  $\mu\text{g/ml}$ , whereas a clear thin lawn was still observed for single mutants *ibpA*, *clpB*, and *PA0079* and *MPAO1 WT* at the same plates (Table I and Fig. S8). These sensitivity differences were reproducible for different transformants of double mutants tested. Thus, *ibpA/clpB* and *ibpA/PA0079* double mutants were also more tobramycin sensitive compared with the single mutants. The MIC results support the hypothesis that *IbpA*, as well as *ClpB*, *HslV*, and *PA0079* proteins, indeed play a role in tobra-

mycin resistance in *P. aeruginosa* cells; however, some of these chaperones and proteases may share overlapped function, which preclude the observation of strong tobramycin sensitivity changes for single-gene inactivation mutants.

#### DISCUSSION

In this study, we investigated the dynamic concentration- and time-dependent proteome response of *P. aeruginosa* to the aminoglycoside antibiotic tobramycin. Marked proteome changes were observed after 60 min of tobramycin exposure and in response to the dosage as low as 0.1  $\mu\text{g/ml}$  (1/10<sup>th</sup> MIC). The concerted proteome changes led to the functional induction of heat shock proteins and proteases at higher dosages (1.0 and 0.5  $\mu\text{g/ml}$ ) and amino acid catabolic enzymes at lower dosages (0.1 and 0.5  $\mu\text{g/ml}$ ). We further showed that inactivation of proteome markers *ibpA/clpB*, *ibpA/PA0079*, or *ibpA/hsIV* was observed with increased tobramycin sensitivity changes in *P. aeruginosa*, which supports the notion that proteome response indeed has effects on antibiotic resistance in *P. aeruginosa*.

Our proteomics approach complements previous genetic analyses from the *P. aeruginosa* transposon mutant library (16, 18, 35, 36) and reveals the dynamic changes of many

TABLE I  
Summary of tobramycin MIC assays for *P. aeruginosa* mutants<sup>a</sup>

Genotype and alleles	Strains	Tob MIC (μg/ml)
WT	MPAO1	0.5–1
<i>ftsH</i> gene deletion	Δ <i>ftsH</i>	0.03–0.06
<i>ibpA</i> in-frame deletion	Δ <i>ibpA</i>	0.5–1
<i>ibpA</i> gene inactivation with <i>lux</i> insert	<i>ibpA-lux</i>	0.5–1
Control containing <i>lux</i> insert	<i>PKH181</i>	0.5–1
Control for Tn insertion	<i>PA3303</i>	0.5–1
<i>clpB</i> single mutant, Tn 1076(2565)	<i>clpB</i> (PW8651)	1
<i>clpB</i> single mutant, Tn 1076(2565)	<i>clpB</i> (PW8652)	1
<i>PA0779</i> single mutant, Tn 2245(2400)	<i>PA0779</i> (PW2411)	1
<i>PA0779</i> single mutant, Tn 1351(2400)	<i>PA0779</i> (PW2414)	1
<i>PA0779</i> single mutant, Tn 1345(2400)	<i>PA0779</i> (PW2413)	1
<i>hsIV</i> single mutant, Tn 223(534)	<i>hsIV</i> (PW9487)	0.25
<i>hsIV</i> single mutant, Tn 168(534)	<i>hsIV</i> (PW9485)	0.5
<i>hsIV</i> single mutant, Tn 244(534)	<i>hsIV</i> (PW9486)	0.5–1
double mutant <i>ibpA/clpB</i>	<i>ibpA/PW8651</i> (1,2,3)	0.5, 0.5, 0.5
double mutant <i>ibpA/clpB</i>	<i>ibpA/PW8652</i> (1,2,3)	0.5, 0.5, 0.5
double mutant <i>ibpA/PA0779</i>	<i>ibpA/PW2411</i> (1,2,3)	0.5, 0.5, 0.5
double mutant <i>ibpA/PA0779</i>	<i>ibpA/PW2414</i> (1,2,3)	0.5, 0.5, 0.5
double mutant <i>ibpA/PA0779</i>	<i>ibpA/PW2413</i> (1,2,3)	0.5–1, 0.5, 0.5
double mutant <i>ibpA/hsIV</i>	<i>ibpA/PW9487</i> (1,2,3)	0.25, 0.25, 0.25
double mutant <i>ibpA/hsIV</i>	<i>ibpA/PW9485</i> (1,2,3)	0.25, 0.25, 0.25
double mutant <i>ibpA/hsIV</i>	<i>ibpA/PW9486</i> (1,2,3)	0.25, 0.25, 0.25

<sup>a</sup> The sequence-verified transposon insertion mutants (Tn) are as reported in Held et al. (17), and information of mutant-specific insertion sites (gene lengths), e.g. 1,076(2,565) and library names, e.g. *clpB* (PW8651) are indicated in the table. Two lines of *ibpA/clpB* mutants, three lines of *ibpA/PA0779*, and *ibpA/hsIV* were examined. For each double mutant line, three independent transformants (1,2,3) were assayed. Tobramycin MIC was defined as the lowest antibiotic concentration preventing the lawn growth of the spotted cells. A listing of a range of concentration levels indicates that a very thin lawn was observed at the lower concentration and this lawn was entirely abolished at the higher concentration in the range. See Figure S8 for additional details.

essential candidate genes during tobramycin treatment. For example, protein levels for essential candidate genes DnaK, GrpE, GroEL, and GroES heat shock proteins were significantly increased at dosages 0.5 and 1.0 μg/ml and time points 60, 120, and 360 min. Likewise, essential metabolic enzymes DhcA, DhcB, and DapB were significantly up-regulated by 0.1 and 0.5 μg/ml dosages. The observed protein-level changes support the hypothesis that many of these essential genes are not only indispensable for *P. aeruginosa* normal growth but also play important roles in tobramycin response.

Adaptive resistance of *P. aeruginosa* to aminoglycoside antibiotics has been described both *in vivo* and *in vitro* (13–15). It is associated with enhanced bacterial drug resistance and reduced clinical treatment efficacy (15). Some previously described mechanisms of adaptive response involved membrane impermeability (37) and the up-regulation of efflux pump MexXY (38). We hypothesize that such adaptive resistance may also involve other changes at the proteome level. Results presented in this manuscript demonstrate that the *P. aeruginosa* proteome response is dependent on the tobramycin levels experienced by cells, with induction of different groups of proteins being observed with exposure to various subinhibitory levels (0.1–1 μg/ml) of tobramycin. Thus, *P. aeruginosa* sense and respond differently to tobramycin, de-

pending on the antibiotic concentration, supporting the adaptive resistance concept (12).

The observed time scale of proteome response is likely sufficient to produce functional impact in antibiotic defense. For example, time-lapse confocal microscopy analyses indicate that exopolysaccharide layers of biofilms can delay the biocide penetration to the central cell clusters up to 60 min (and in some cases, even longer) (10, 11, 39). In addition, CF patients that inhaled a second dose tobramycin 1 h after an initial dose also experienced the time-dependent decrease in drug efficacy (15). This effect was hypothesized to be resultant from induced adaptive resistance of *P. aeruginosa* *in vivo*. Thus, this time dependence of bacterial adaptive resistance is consistent with the observed time dependence in proteome changes in the present study. We also show that prolonged exposure of *P. aeruginosa* to tobramycin up to 6 h is associated with sustained proteome acclimation, during which time adaptive resistance also occurs (13–15).

Distinct proteome subsets were induced when *P. aeruginosa* cells were exposed to increasing concentrations of tobramycin. Such concentration-dependent changes underscore the diverse mechanisms in *P. aeruginosa* adaptive response. Widespread induction of heat shock proteins and proteases were observed with 0.5 and 1.0 μg/ml tobramycin treatments, consistent with previous transcriptome analysis

(31, 40). Recent *Escherichia coli* proteome analysis of streptomycin treatments also highlights the roles of heat shock chaperones and proteases (41, 42), suggesting the response may be conserved among bacterial species and aminoglycoside drugs.

In particular, heat shock protein lbpA was the highest fold increased protein in the present study. Inactivation of *ibpA* did not yield significant tobramycin MIC changes. However, inactivation of two heat shock proteins/proteases *ibpA/clpB*, *ibpA/PA0779*, or *ibpA/hsIV* led to increased tobramycin sensitivity changes in *P. aeruginosa*. The lack of strong changes in tobramycin sensitivity for  $\Delta$ *ibpA* may be owing to the overlapped function of lbpA with other heat shock chaperones and proteases in *P. aeruginosa*. In *E. coli*, lbpA was found to perform concerted actions with ClpB and DnaK-DnaJ-GrpE to reverse protein aggregation induced by high temperatures (34, 43).

In addition to the induction of heat shock chaperones and proteases, changes in amino acid metabolic and biosynthesis functions were observed with tobramycin treatments at the lower dosages 0.1 and 0.5  $\mu$ g/ml and longer time points 120 and 360 min. These include the up-regulation of essential candidate genes of *dhcA* and *dhcB*, which are dehydrocarnitine CoA transferase subunits involved in metabolism of the amino acid carnitine, and *dapB*, which is a 4-hydroxy-tetrahydrodipicolinate reductase involved in lysine biosynthesis. Consistent with our findings that show amino acid metabolic enzyme levels were altered during tobramycin exposure, the interaction effects of amino acids and aminoglycoside antibiotics have been reported. For instance, supplementing 0.4% L-arginine in tryptic soy agar plate enhances the tobramycin killing of *P. aeruginosa* biofilm colonies over 10-fold (44). Conversely, the presence of 20 mM cadaverine (a decarboxylation product of lysine amino acid) promotes *P. aeruginosa* resistance to aminoglycoside kanamycin and gentamycin by fourfold (45), while suppressing its resistance to carboxypenicillins (46). Although much still needs to be understood, the results demonstrated here show that bacterial cells exposed to low levels of tobramycin exhibit increased levels of enzymes that metabolize and synthesize amino acids that could alter drug sensitivity. If so, manipulation of amino acid homeostasis pathways may provide new ways to modulate proteome response in *P. aeruginosa* and improve antibiotic treatment effects.

*Acknowledgments*—We thank Dr. Priska D. von Haller for helpful discussions, and Samuel Lee for experimental contributions.

\* This research was supported in part by National Institutes of Health grants 5R01HL110879, 5R01AI101307, 5R01GM086688, and 7S10RR025107, and the University of Washington's Proteomics Resource (UWPR95794).

 This article contains supplemental material Figs. S1-S8.

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