Introduction

The widespread use of F has contributed to the caries decline, but excessive intake may affect both bone metabolism and enamel development, causing skeletal and dental fluorosis, respectively. There are many sources of F intake, such as drinking water, dental products, dietary supplements and infant formulas [1]. There is evidence that the prevalence of dental fluorosis (DF) is increasing worldwide both in fluoridated and non-fluoridated communities [2]. In the US, 23% of 6- to 39-yr-old subjects present enamel fluorosis, ranging from very low to relatively high in severity [3]. However, the exact mechanisms by which F affects biomineralization are not completely understood [4,5]. It has been proposed that genetic determinants influence the susceptibility to DF in humans [6] and mice [7]. Two strains of mice have been identified with distinct responses to the effects of F in the mineralized tissues. The A/J strain is "susceptible", with a rapid onset and severe development of DF, while the 129P3/J is "resistant", with minimum development of DF [7]. These strains also differ regarding their susceptibilities to the effects of F in bone [8,9].

Renal Proteome in Mice with Different Susceptibilities to Fluorosis

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Abstract

A/J and 129P3/J mouse strains have different susceptibilities to dental fluorosis due to their genetic backgrounds. They also differ with respect to several features of fluoride (F) metabolism and metabolic handling of water. This study was done to determine whether differences in F metabolism could be explained by diversities in the profile of protein expression in kidneys. Weanling, male A/J mice (susceptible to dental fluorosis, n = 18) and 129P3/J mice (resistant, n = 18) were housed in pairs and assigned to three groups given low-F food and drinking water containing 0, 10 or 50 ppm (F) for 7 weeks. Renal proteome profiles were examined using 2D-PAGE and LC-MS/MS. Quantitative intensity analysis detected between A/J and 129P3/J strains 122, 126 and 134 spots differentially expressed in the groups receiving 0, 10 and 50 ppmF, respectively. From these, 25, 30 and 32, respectively, were successfully identified. Most of the proteins were related to metabolic and cellular processes, followed by response to stimuli, development and regulation of cellular processes. In F-treated groups, PDZK-1, a protein involved in the regulation of renal tubular reabsorption capacity was down-modulated in the kidney of 129P3/J mice. A/J and 129P3/J mice exhibited 11 and 3 exclusive proteins, respectively, regardless of F exposure. In conclusion, proteomic analysis was able to identify proteins potentially involved in metabolic handling of F and water that are differentially expressed or even not expressed in the strains evaluated. This can contribute to understanding the molecular mechanisms underlying genetic susceptibility to dental fluorosis, by indicating key-proteins that should be better addressed in future studies.

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To determine whether such differences were due to differences in F metabolism, we conducted a metabolic study in which total F intake and excretion were measured. Our results showed that, compared to A/J mice, 129P3/J mice ingested less water, excreted less urine, had lower urinary F excretion and consequently had higher F retention and plasma and femur F levels [10]. However, these findings were not able to explain the mechanisms underlying the differences in the metabolic handling of F.

Kidneys represent the major route of removal of F from the body [11]. After F enters the renal tubules, a variable amount is reabsorbed, depending on the urinary pH because transmembrane migration occurs by diffusion of HF [12]. Thus, any factor that affects urinary pH will have an impact on the amount of F that is excreted in urine [11]. Urinary F excretion is also influenced by glomerular filtration rate since its reduction, as occurs in chronic renal dysfunction as well as in the last decades of life, results in lower excretion and increased plasma F levels [13]. Considering that kidney is a key organ in the metabolism of F, we then sought to investigate the molecular mechanisms underlying the renal F metabolism in A/J and 129P3/J mice that may account for their...
differential metabolic handling of F. To address this, proteomic analyses were performed on kidneys of A/J and 129P3/J mice receiving both low and high level of F-containing water.

**Materials and Methods**

**Animals and Treatment**

Male mice from the A/J and 129P3/J inbred strains (3-week-old) were randomly distributed into three groups (n = 6/strain) based on the F concentrations in the drinking water. All animals were housed in pairs in metabolic cages with *ad libitum* access to low-F food (AIN76A, PMI Nutrition, Richmond, IN, USA, 0.95 mg/Kg F) and water, to allow analysis of water and food consumption [10]. The temperature and humidity in the climate-controlled room, which had a 12-h light/dark cycle, were 23 ± 1°C and 40%–80%, respectively. All experimental protocols were approved by the Ethics Committee for Animal Experiments of Bauru Dental School, University of Sao Paulo (Protocol # 026/2007). Experimental groups received drinking water containing 10 (low) or 50 (high) ppm F ion (as NaF), for 60 days. Control group received deionized water for the same period. At the end of the study, the mice were anesthetized with ketamine/xylazine and kidneys were collected. The left kidney was washed with cold buffer containing Tris 100 mM, EDTA 1 mM, PMSF 1 mM, pH 7.4, frozen at liquid nitrogen and kept at -80°C until proteomic analysis. The right kidney was collected for F analysis.

**F analysis in kidney**

Kidneys were homogenized in deionized water for 2 min using a homogenizer (Marconi, Model MA 102). Kidney F concentrations were determined in duplicate (100 mg of kidney tissue) after overnight hexamethyldisiloxane (HMDS)-facilitated diffusion [14,15] using the ion-specific electrode (Orion Research, Model 9409) and a miniature calomel electrode (Accumet, #13-620-79) both coupled to a potentiometer (Orion Research, Model EA 940). F standards (0.00475 to 0.19 µgF) were prepared in triplicate and diffused in the same manner as the samples. In addition, nondiffused standards were prepared to have exactly the same F concentrations as the diffused standards. Comparison of the mV readings demonstrated that the F in the diffused standards had been completely trapped and analyzed (recovery >95%). The mV potentials were converted to µg F using a standard curve with a coefficient correlation of r=0.99.

**Sample Preparation for 2DE**

Kidney samples were homogenized using mortar and pestle in liquid nitrogen. Denaturation buffer (7 M urea, 2 M de thiouriea, 4% CHAPS, 1% DTT and 0.5% IPG pH 3–10, GE Healthcare, Uppsala, Sweden) was added. After 1 h vortexing at 4°C, samples were centrifuged at 25000 x g for 30 min for 4°C for supernatants collection. The proteins were precipitated by using the kit PlusOne 2D Cleanup (GE Healthcare, Uppsala, Sweden), as recommended by the manufacturer. The pellets were resuspended in rehydration buffer (8 M urea, 0.5% CHAPS, 10% glycerol, 0.5% IPG buffer pH 3–10, 7 mg/2.5 mL DTT, 0.002% bromophenol blue). Protein concentration was measured in each sample by Bradford assay. After quantification, 1000 µg of kidney proteins from each animal of the same group and strain were combined to constitute a pool [16] that was submitted to proteomic analysis in triplicate, as described below.

**2-DE Separation**

Renal proteins (1000 µg) were taken from each pooled sample and mixed in rehydration buffer to a volume of 400 µL which was then loaded onto 24-cm IPG strips (linear pH 3–10). Rehydration and first-dimensional IEF were performed on IPGphor IEF system at 20°C with the following parameters: 50V for 12 h, 500V for 1 h, 1000V gradient for 1 h, then 10000 V for a total 40,000 V. *Etion DAMTsix* (GE Healthcare) (GE Healthcare, Uppsala, Sweden) with homemade 12.5% acrylamide gels was used for the second dimension separation. Electrophoresis was performed at 15 mA/gel (30V) for 1 h and at 60 mA/gel (500V) until bromophenol blue line had reached the bottom of the gels. The resolved protein spots were stained with Colloidal Coomassie Brilliant Blue G-250 [17]. Gels were scanned with an ImageMaster scanner, and all images were analyzed by ImageMATHER 2D Platinum 7.0 software (GE Healthcare, Uppsala, Sweden). Parameters used for spot detection were minimal area = 5 pixels; smooth factor = 4; and saliency = 100. The gel chosen as the reference had the highest number of spots. The reference gel was then used for matching of corresponding protein spots between gels. Following average mode of background subtraction, individual spot intensity volume was normalized with total intensity volume (summation of the intensity volumes obtained from all spots in the same 2-DE gel). The normalized intensity volume values of individual protein spots were then used to determine differential protein expression between control and experimental groups. 2D spots that exhibited a twofold or more decrease or increase were tested for statistical significance. Analysis of 2D-gel variability among the replicates of each experimental condition was taken by using the relative volume (% vol). The correlation coefficients among the triplicates are shown to vary from 0.9385 to 0.9821 (Figure S4 - Supplementary information).

**LC-MS/MS Analysis**

Protein spots of interest were excised from the gel and destained three times with 25 mM Ambic/Acetonitrile (50:50 v/v) for 30 min. The destained gel pieces were dehydrated twice with acetonitrile (ACN) for 10 min and dried in a vacuum concentrator (Eppendorf, Hamburg, Germany). The dried gel pieces were rehydrated with 20 mM DTT in 50 mM Ambic for 40 min at 56°C. Excess of reagent was removed and 55 mM iodoacetamide (IAA) in Ambic 50 mM was added for 30 min RT at dark. The remaining liquid was removed and washed out with 25 mM Ambic, followed by dehydration with ACN. After its removal, the gels were dried again. For digestion, dried gels were incubated with 10 ng/µL trypsin in 25 mM Ambic for 15 min (Trypsin Gold Mass spectrometry, Promega, Madison, USA). Peptides were sequentially extracted from the gels initially in 50% ACN (v/v) with 5% formic acid for 14 h at 37°C, then in 50% ACN (v/v) with 1% formic acid for 15 min, followed by 60% methanol (v/v) with 1% formic acid for 15 min and twice with 100% ACN at 45°C under sonication (40kHz/30W, Branson, Danbury, USA). Extracts were dried using a vacuum concentrator (Eppendorf, Hamburg, Germany) and kept at -20°C. Prior to MS identification, digested peptides were dissolved in 12 µL 0.1% formic acid. The peptides were identified and quantified by LC-ESI-Q-TOF MS (Liquid Chromatography Electrospray Ionization Quadrupole Time of Flight Mass Spectrometry) (Waters, Milford, USA). MassLynx 4.1 SCN662 software (Waters, Milford, USA) was used to submit the combined MS and MS/MS data to MASCOT database search engine (http://www.matrixscience.com) (version 2007.12.04) based on IPI (International Protein Index) protein database restricted to taxonomies *Mus musculus* (Mouse). The search was limited with a mass tolerance of 100 ppm and only one missed cleavage per peptide was allowed. For modification of peptides, cysteine carbamido-methylation (fixed) and methionine oxidation (variable) were considered. Significant matching protein
required score of >60. Accuracy between the theoretical and experimental obtained mass and pI were also considered. When 2 or more proteins with high scores were identified in the same spot, they are excluded from analysis. Identified proteins were classified into 6 different categories according to their primary function [10].

Statistical Analysis
For kidney F concentration, the software GraphPad InStat version 4.0 for Windows (GraphPad software Inc., La Jolla, USA) was used. Data were analysed by 2-way ANOVA and Bonferroni test for individual comparisons (p<0.05).

For proteomic data, statistical analysis was performed using ANOVA available at ImageMaster 2D Platinum 7.0 software (GE Healthcare, Uppsala, Sweden). Only proteins with significantly altered levels were excised for identification by LC-MS/MS (p<0.05).

Results
Renal F Concentration
Mean kidney F (± S.E.) concentrations for A/J mice for control, 10 ppmF and 50 ppmF groups were: 0.126±0.008, 0.174±0.007 and 0.296±0.026 μg/g. The corresponding values for 129P3/J mice were 0.139±0.015, 0.163±0.010 and 0.198±0.046 μg/g, respectively. Two-way ANOVA revealed a significant difference among the treatments (F = 35.13, p<0.0001), but not between the strains (F = 0.099, p = 0.756) without significant interaction between these criteria (F = 0.124, p = 0.484). For both strains, significantly higher kidney F concentrations were found for the 50 ppmF group, when compared with control and 10 ppmF groups that did not significantly differ from each other.

Identification of Differentially Expressed Proteins
For the statistical analysis, comparisons were performed between the strains as follows: Control groups (A/J vs 129P3/J mice), 10 ppmF groups (A/J vs 129P3/J mice), and 50 ppmF groups (A/J vs 129P3/J mice). Tables 1–3 show the proteins that were differentially expressed (p<0.05) in each comparison. Representative 2D map of each comparison is also shown in the Supplementary Information (Figures S1-S3). Quantitative intensity analysis showed 26 changed spots between control groups (Table 1). Among them, 14 spots were up-modulated, while 12 were down-regulated in control 129P3/J mice, when compared to control A/J mice. In general, the kidney proteome dataset was found to be significantly related with several metabolic and cellular processes pathways. Most of the 14 proteins up-modulated in the kidney of 129P3/J mice are related with metabolism (57.2%), while 28.6% are involved in cell processes and the remainder in information pathways (7.1%) and transport (7.1%). A similar pattern was observed for the proteins that were down-regulated in kidney 129P3/J mice. The respective percentages were 50.0, 25.0, 16.7 and 8.3 (Table 1). From the differentially expressed proteins in control groups, 10 were exclusively expressed in this comparison whereas 2, 6, and 8 proteins were also present in either 10 ppmF or 50 ppmF or both F-treated groups, respectively (Figure 1).

For the comparison between the A/J and 129P3/J mice treated with 10 ppmF, 14 proteins were increased and 17 diminished in kidney of 129P3/J. Among the increased proteins, 64.3% are related with metabolism, while 35.7% are associated with cell processes. Most of the decreased proteins are also related to metabolism (41.1%), followed by information pathways (23.6%), cell processes (17.6%), transport (11.8%) and structure (5.9%) (Table 2). From the differentially expressed proteins in 10 ppmF group, 15 were exclusively expressed in this comparison whereas 2, 6 and 8 proteins were also present in either control or 50 ppmF or in both groups, respectively (Figure 1).

Regarding the comparison between the groups treated with 50 ppmF, 18 proteins were significantly up-regulated and 13 down-modulated in kidney of 129P3/J mice when compared with A/J mice. Fourteen of eighteen enhanced proteins are associated with metabolism (77.8%), followed by processes (11.1%), information pathways (5.6%) and processes pathways (5.6%). Among the down-modulated proteins, most are also related to metabolism (46.2%), followed by cell processes (23.0%), transport (15.4%), information pathways (7.7%) and structure (7.7%) (Table 3). Among the differentially expressed proteins in kidney of animals treated with 50 ppmF, 11 proteins are exclusively expressed in this group while 6, 6 and 8 proteins are also present in either control or 10 ppmF or both groups, respectively (Figure 1).

Among the 8 proteins differentially expressed between the mice strains, regardless of the treatment with F, catalase, medium-chain specific acyl-CoA dehydrogenase and alpha-aminoadipic semialdehyde dehydrogenase were up-regulated, while isovaleryl-CoA dehydrogenase, ornithine aminotransferase, lactoylglutathione lyase, meprin A subunit alpha and albumin were down-regulated in the kidney of 129P3/J mice.

Identification of Unique Proteins
A/J and 129P3/J mice exhibited 11 and 3 exclusive proteins, respectively. From these, 9 (64.3%) are related to metabolism, followed by cell processes (4 or 28.6%) and information pathways (1 or 7.1%). This profile was not altered upon exposure to F (Table 4).

Discussion
In the present study, we identified proteins potentially involved in renal F metabolism that are either exclusively or differentially expressed in A/J and 129P3/J mice. This highlights the molecular mechanisms underlying the differential metabolic handling of F by these two strains of mice. Exclusive proteins expressed in A/J or 129P3/J mice exhibited the same profile, regardless exposure to F. This suggests that the genetic background per se accounts for such differences between these two strains of mice. We have focused on identified proteins that may be associated with metabolic handling of F and water and renal functions. Unique metabolic proteins in kidney from A/J mice are involved in carbohydrate (probable D-lactate dehydrogenase), carbon (transaldolase), aminoacid (isobutyryl-CoA dehydrogenase, hydroxymethylglutaryl-CoA syn-
Table 1. Expression of differentially significant kidney proteins between control A/J vs control 129P3/J mice.

| Spot n°. | Protein                                                                 | \( M_w \) (kDa)/p | Expt. | * Theor. | Number of peptides/Score | Difference (P value) | Uniprot ID | Biological Process |
|---------|-------------------------------------------------------------------------|-------------------|-------|----------|--------------------------|----------------------|------------|-------------------|
| 2       | Aconitate hydratase, mitochondrial                                       | 91.4/7.1          |       | 82.5/7.4 | 6/99                     | \( \uparrow \) 1.29 (0.046) | Q9K010     | Metabolism        |
| 119/280 | ATP synthase subunit beta, mitochondrial                                 | 33/5.155          |       | 51.8/5.0 | 16/1129                  | \( \uparrow \) 1.29 (0.038) | P56480     | Metabolism        |
| 151     | Pyruvate dehydrogenase E1 component subunit beta, mitochondrial         | 36.5/5.1          |       | 35.8/5.4 | 8/434                    | \( \uparrow \) 1.29 (0.011) | Q9D051     | Metabolism        |
| 175     | Hydroxynitric oxide 2                                                  | 38.5/7.94         |       | 38.7/7.6 | 11/529                   | \( \uparrow \) 1.29 (0.012) | Q9NYQ2     | Metabolism        |
| 217/221 | Medium-chain specific acyl-CoA dehydrogenase, mitochondrial             | 42.5/8.055        |       | 43.6/7.69| 15/715                   | \( \uparrow \) 1.29 (0.001) | P45952     | Metabolism        |
| 269     | Homogentisate 1,2-dioxygenase                                           | 50/7.2            |       | 50/6.85  | 6/105                    | \( \uparrow \) 1.29 (0.011) | Q09713     | Metabolism        |
| 384     | Sarcoamine dehydrogenase, mitochondrial                                 | 95/6.14           |       | -        | 18/583                   | \( \uparrow \) 1.29 (0.041) | Q99LB7     | Metabolism        |
| 116     | Beta-lactamase-like protein 2                                           | 32.5/8.885        |       | 32.8/5.9 | 14/198                   | \( \uparrow \) 1.29 (0.043) | Q99KR3     | Metabolism        |
| 7       | Serine/threonine-protein phosphatase PPI-alpha catalytic subunit        | 38.5/6.75         |       | 37.4/5.9 | 9/122                    | \( \uparrow \) 1.29 (0.018) | P62137     | Processes/Information pathways |
| 317/321 | Catalase                                                                | 58/7.37           |       | 59.7/7.7 | 6/103                    | \( \uparrow \) 1.29 (0.032) | P24270     | Processes        |
| 396     | Aldehyde dehydrogenase family 1 member L1                              | 98.5/7.6          |       | 75.5/6.6 | 24/571                   | \( \uparrow \) 1.29 (0.013) | Q8ROY6     | Processes        |
| 285     | Alpha-aminoadipic semialdehyde dehydrogenase                           | 52.6/22           |       | 55.9/6.0 | 13/374                   | \( \uparrow \) 1.29 (0.001) | Q9DBF1     | Processes        |
| 324/583 | 60 kDa heat shock protein, mitochondrial                                | 59.5/4.875        |       | 58/5.35  | 22/992                   | \( \uparrow \) 1.29 (0.022) | P63038     | Information Pathways |
| 413     | Cootder subunit delta                                                  | 63/5.2            |       | 57.2/5.9 | 7/206                    | \( \uparrow \) 1.29 (0.009) | Q3XUY5     | Transport        |
| 3       | ATP synthase subunit delta, mitochondrial                               | 15.5/2.82         |       | 15.7/5.0 | 4/188                    | \( \uparrow \) 1.29 (0.022) | P56480     | Metabolism        |
| 171     | Alcohol dehydrogenase (NADP+)                                          | 38.5/7.51         |       | 36.6/5.6 | 17/775                   | \( \uparrow \) 1.29 (0.049) | Q9H16      | Metabolism        |
| 189     | Sorbitol dehydrogenase                                                 | 40.5/6.885        |       | 38.2/6.6 | 13/853                   | \( \uparrow \) 1.29 (0.041) | Q64442     | Metabolism        |
| 209     | Isovaleryl-CoA dehydrogenase, mitochondrial                            | 42.5/6.6          |       | 43/6.3   | 12/535                   | \( \uparrow \) 1.29 (0.000) | Q9H15      | Metabolism        |
| 233     | Ornithine aminotransferase, mitochondrial                               | 45.5/7.2          |       | 45.8/5.7 | 10/255                   | \( \uparrow \) 1.29 (0.024) | P29758     | Metabolism        |
| 237     | Creatine kinase U-type, mitochondrial                                   | 45.8/5.0          |       | 43.2/7.3 | 9/133                    | \( \uparrow \) 1.29 (0.033) | P30275     | Metabolism        |
| 60      | Lactoylglutathione lyase                                                | 27/4.48           |       | 20.7/5.25| 37/365                   | \( \uparrow \) 1.29 (0.001) | Q9CPU0     | Processes        |
| 190     | Phosphotriesterase-related protein                                      | 41/6.58           |       | 39.2/6.2 | 7/293                    | \( \uparrow \) 1.29 (0.029) | Q60886     | Processes        |
| 205     | 40S ribosomal protein SA                                               | 42.5/3.96         |       | 32.7/4.8 | 7/825                    | \( \uparrow \) 1.29 (0.044) | P14206     | Processes        |
| 99      | Proteosome subunit beta type-7                                          | 30/5.885          |       | 25.3/5.8 | 7/85                     | \( \uparrow \) 1.29 (0.041) | P70195     | Information pathways |
| 385/386/520 | Meprin A subunit alpha                         | 93.5/43          |       | 77.2/5.9 | 11/187                   | \( \uparrow \) 1.29 (0.020) | P28825     | Information pathways |
| 355     | Serum albumin                                                          | 69.5/5.33         |       | 65.9/5.5 | 15/635                   | \( \uparrow \) 1.29 (0.022) | P07724     | Transport        |

*Experimental molecular weight (kDa)/p of protein spot in the gel (Mean of min. and max.) based on the coordinates of landmark proteins. *Theoretical molecular weight (kDa)/p of theoretical protein. *Number of peptides identified and score. Differences in expression in relation to 129P3/J mice (\( \downarrow \) down-modulation; \( \uparrow \) up-modulation); Individual P value after ANOVA. *Identification is based on protein ID from IPI (international protein index) protein database (http://www.uniprot.org/). *Category of protein based on its primary biological function according to Rison (2000) [18].

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| Spot n°. | Protein | \(M_w\) (kDa)/pI | Theor. | Number of peptides/Score | Difference (P value) | Uniprot ID | Biological Process |
|---------|---------|------------------|--------|--------------------------|---------------------|------------|-------------------|
| 3       | ATP synthase subunit delta, mitochondrial | 15.5/2.82 | 51.7/5.0 | 4/188 | 129(0.040) | P54680 | Metabolism |
| 182     | Aldose 1-epimerase | 40.6/6.25 | 37.6/6.3 | 4/146 | 129(0.016) | Q8K157 | Metabolism |
| 210     | Aminocyclase-1 | 42.5/5.94 | 45.8/5.9 | 18/831 | 129(0.013) | Q99W2 | Metabolism |
| 221/17  | Medium-chain specific acyl-CoA dehydrogenase, mitochondrial | 43/6.25 | 43.7/6.7 | 15/541 | 129(0.011) | P45952 | Metabolism |
| 223     | Adenosylhomocysteinase | 42.5/6.6 | 43.5/6.3 | 15/559 | 129(0.047) | P50247 | Metabolism |
| 261     | Fumarate hydratase, mitochondrial | 45/5.72 | 50.7/9 | 8/142 | 129(0.015) | P97807 | Metabolism |
| 462     | Short/branched chain specific acyl-CoA dehydrogenase, mitochondrial | 42.6/1.1 | 44/6.0 | 2/66 | 129(0.047) | Q9DBL1 | Metabolism |
| 467     | 3-hydroxyanthranilate 3,4-dioxygenase | 34/6.45 | 32.8/6.1 | 8/181 | 129(0.039) | Q78JT3 | Metabolism |
| 489     | Citrate synthase, mitochondrial | 43/9.1 | 49/8.2 | 3/107 | 129(0.035) | Q9CZU6 | Metabolism |
| 142/146 | Asparagylase-2 | 35.5/4.915 | 35.5/3.9 | 9/488 | 129(0.003) | Q91XE4 | Processes |
| 7       | Serine/threonine-protein phosphatase PP1-alpha catalytic subunit | 38.5/6.75 | 37.5/4.9 | 9/122 | 129(0.018) | P62137 | Processes/Information pathways |
| 285     | Alpha-aminoacidic semialdehyde dehydrogenase | 52.5/6.34 | 55.9/6.0 | 13/374 | 129(0.005) | Q9DBF1 | Processes |
| 317/321 | Catalase | 58.7/4.45 | 59.7/7.7 | 6/103 | 129(0.032) | P24270 | Processes |
| 289     | Selenium-binding protein 1 | 53.5/6.8 | 52.5/5.9 | 13/374 | 129(0.037) | P17563 | Processes |
| 5       | Ester hydrolase C1orf54 homolog | 37/5.95 | 51.7/5.0 | 4/188 | 129(0.022) | Q91V76 | Metabolism |
| 87      | Enoyl-CoA hydratase, mitochondrial | 28/6.15 | 28.5/7.8 | 13/353 | 129(0.017) | Q88H95 | Metabolism |
| 157     | L-lactate dehydrogenase B chain | 37/5.5 | 36.5/7 | 10/705 | 129(0.002) | P16125 | Metabolism |
| 185     | 3’(2’),5’-bisphosphate nucleotidase 1 | 41/5.55 | 33.1/5.0 | 7/216 | 129(0.016) | Q9O251 | Metabolism |
| 209     | Isovaleryl-CoA dehydrogenase, mitochondrial | 43/6.715 | 43.6/6 | 12/535 | 129(0.021) | Q9HIU | Metabolism |
| 233     | Ornithine aminotransferase, mitochondrial | 45/5.72 | 45.8/5.7 | 10/255 | 129(0.024) | P29758 | Metabolism |
| 518     | Nucleoside diphosphate-linked moiety X motif 19, mitochondrial | 42/6.15 | 39.3/6.0 | 7/245 | 129(0.041) | P11930 | Metabolism |
| 60      | Lactoylglutathione lyase | 27/4.48 | 20.7/5.5 | 37/365 | 129(0.001) | Q9CPU0 | Processes |
| 85      | Rho GDP-dissociation inhibitor 1 | 27/4.55 | 23.3/5.1 | 37/365 | 129(0.005) | Q99PT1 | Processes |
| 512     | Nlrp1/12/11 exchange regulatory cofactor NRE-RF3 | 69.5/4.65 | 56.5/5.3 | 13/403 | 129(0.041) | Q9JL4 | Processes |
| 75      | GTP-PAM phosphotransferase, mitochondrial | 26.5/10.135 | 25.3/8.8 | 7/85 | 129(0.050) | Q9WTP7 | Information pathways |
| 320     | Protein disulfide-isomerase A3 | 59.5/7.3 | 54.3/5.7 | 16/362 | 129(0.043) | P2777 | Information pathways |
| 385/386/520 | Meprin A subunit alpha | 89.5/5.5 | 77.2/5.9 | 11/187 | 129(0.002) | P28125 | Information pathways |
| 400     | Heat shock protein 90, beta (Hsp90), member 1 | 99.5/3.9 | 92.5/4.7 | 24/515 | 129(0.011) | Q9V138 | Information Pathways |
| 267     | Actin-related protein 3 | 49.5/4.45 | 47.2/5.6 | 13/354 | 129(0.016) | Q99J9Y | Structure |
| 311     | Vitamin D-binding protein | 57.5/4.76 | 51.9/5.2 | 9/109 | 129(0.028) | P21614 | Transport |
| 355     | Serum albumin | 69.5/4.2 | 65.9/5.3 | 15/635 | 129(0.020) | P07724 | Transport |

*aExperimental molecular weight (kDa)/pI of protein spot in the gel (Mean of min. and max.) based on the coordinates of landmark proteins.
*bTheoretical molecular weight (kDa)/pI of theoretical protein.
*cNumber of peptides identified and score.
*dDifferences in expression in relation to 129P3/J mice (↓ down-modulation; ↑ up-modulation); individual P value after ANOVA.
*eIdentification is based on protein ID from IPI (International protein index) protein database (http://www.uniprot.org). fCategory of protein based on its primary biological function according to Rison (2000) [18].

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Table 3. Expression of differentially significant kidney proteins between control A/J vs control 129P3/J mice.

| Spot n° | Protein                                                                 | $M_w$ (kDa)/pI | Theor. | Number of peptides/Score | Difference (P value) | Uniprot ID | Biological Process |
|---------|-------------------------------------------------------------------------|----------------|--------|--------------------------|----------------------|------------|-------------------|
| 2       | Aconitate hydratase, mitochondrial                                      | 90/4.77        | 82.5/7.4 | 6/99                     | ↑ 129(0.029)         | Q99K00     | Metabolism         |
| 35      | ATP synthase subunit d, mitochondrial                                    | 24/5.16        | 18.6/5.53 | 8/402                    | ↑ 129(0.036)         | Q9DCX2     | Metabolism         |
| 116     | Beta-lactamase-like protein 2                                            | 32.5/6.04      | 32.7/5.9 | 7/198                    | ↑ 129(0.045)         | Q99KB3     | Metabolism         |
| 133     | Hydroxyacyl-Coenzyme A dehydrogenase, mitochondrial                     | 33.5/9.28      | 32995/8.39 | 8/255                    | ↑ 129(0.035)         | Q61425     | Metabolism         |
| 175     | Hydroxocyc oxide 2                                                       | 37/8.1         | 38/8.7  | 11/529                   | ↑ 129(0.003)         | Q9NYQ2     | Metabolism         |
| 184     | Sorbitol dehydrogenase                                                   | 40.5/6.805     | 38.2/6.56 | 10/586                   | ↑ 129(0.003)         | Q64442     | Metabolism         |
| 210     | Aminopyruvylase                                                         | 42.5/5.92      | 45.8/5.9 | 18/831                   | ↑ 129(0.006)         | Q99JW2     | Metabolism         |
| 217/221 | Medium-chain specific acyl-CoA dehydrogenase, mitochondrial            | 41.8/5.22      | 43/5.79 | 15/715                   | ↑ 129(0.009)         | P45952     | Metabolism         |
| 269     | Homogenetase 1,2-dioxynase                                                | 49/7.41        | 50/7.85 | 6/105                    | ↑ 129(0.002)         | Q09173     | Metabolism         |
| 280/119 | ATP synthase subunit beta, mitochondrial                                 | 51/4.345       | 51.7/5.0 | 16/1129                  | ↑ 129(0.001)         | P56480     | Metabolism         |
| 360     | Propionyl-CoA carboxylase alpha chain, mitochondrial                     | 77/6.07        | 74.4/6.0 | 13/315                   | ↑ 129(0.003)         | Q91ZA3     | Metabolism         |
| 378     | Acylamino-acid-releasing enzyme                                          | 86/5.075       | 81.5/5.3 | 10/256                   | ↑ 129(0.001)         | Q8R146     | Metabolism         |
| 491     | Argininosuccinate synthase                                               | 44/9.41        | 46.6/8.4 | 14/321                   | ↑ 129(0.001)         | P16460     | Metabolism         |
| 142/146 | Aspartatecarboxylase-2                                                   | 36.5/4.91      | 35.3/5.3 | 9/488                    | ↑ 129(0.003)         | Q91X64     | Metabolism         |
| 285     | Alpha-aminoacidic semialdehyde dehydrogenase                            | 53/6.55        | 55.9/6.0 | 13/374                   | ↑ 129(0.024)         | Q9DBF1     | Processes          |
| 321     | Catalase                                                                | 56/8.055       | 59.7/7.7 | 6/103                    | ↑ 129(0.008)         | P24270     | Processes          |
| 298     | ATP synthase subunit alpha, mitochondrial                                | 51/4.345       | 51.7/5.0 | 16/1129                  | ↑ 129(0.001)         | P56480     | Metabolism         |
| 372     | Radixin                                                                 | 84/6.305       | 86.5/5.91 | 9/125                    | ↑ 129(0.015)         | P26043     | Structure          |
| 90      | Phosphoglycerate mutase 1                                                | 28.5/7.235     | 28.7/6.75 | 23/324                   | ↑ 129(0.003)         | Q9DBU1     | Metabolism         |
| 157     | L-lactate dehydrogenase B chain                                          | 37.5/5.49      | 36.4/5.7 | 10/705                   | ↑ 129(0.009)         | P16125     | Metabolism         |
| 209     | Isovaleryl-CoA dehydrogenase, mitochondrial                              | 43/6.76        | 43/6.3  | 12/535                   | ↑ 129(0.000)         | Q9JH5      | Metabolism         |
| 233     | Orotic acid aminotransferase, mitochondrial                              | 46/5.855       | 45.8/5.7 | 10/255                   | ↑ 129(0.006)         | P29758     | Metabolism         |
| 244     | Glycine aminotransferase, mitochondrial                                   | 45/7.07        | 42/6.54 | 15/234                   | ↑ 129(0.003)         | Q9D965     | Metabolism         |
| 518     | Nucleoside diphosphate-linked moiety X motif 19, mitochondrial           | 45/5.99        | 39.3/6.0 | 7/245                    | ↑ 129(0.018)         | P11930     | Metabolism         |
| 60      | Lactoylglutathione lyase                                                 | 27/4.252       | 20.7/5.25 | 10/365                   | ↑ 129(0.003)         | Q9CPU0     | Processes          |
| 190     | Phosphotriesterase-related protein                                       | 40.5/6.74      | 39.2/6.2 | 7/293                    | ↑ 129(0.004)         | Q60866     | Processes          |
| 512     | Na(+)/H(+) exchange regulatory cator NHE-RF3                            | 70/4.665       | 56.5/5.5 | 13/403                   | ↑ 129(0.009)         | Q9JL4      | Processes          |
| 267     | Actin-related protein 3                                                  | 49.5/4.44      | 47.2/5.6 | 13/554                   | ↑ 129(0.015)         | Q99J9      | Structure          |
| 355     | Serum albumin                                                           | 70/5.42        | 65.9/5.53 | 15/635                   | ↑ 129(0.026)         | P07724     | Transport          |
| 376     | Serotransferrin                                                          | 82/7.264       | 74.9/6.8 | 12/326                   | ↑ 129(0.019)         | Q92111     | Transport          |
| 385/386/520 | Meprin A subunit alpha                                               | 96.5/5.945 | 77.2/5.9 | 11/187                   | ↑ 129(0.014)         | P28825     | Information pathways |

*Experimental molecular weight (kDa)/pI of protein spot in the gel (Mean of min. and max.) based on the coordinates of landmark proteins. Theoretical molecular weight (kDa)/pI of theoretical protein. Number of peptides identified and score. "Differences in expression in relation to 129P3/J mice (↓ down-modulation; ↑ up-modulation); individual P value after ANOVA. Identification is based on protein ID from IPI (international protein index) protein database (http://www.uniprot.org/). Category of protein based on its primary biological function according to Rison (2000) [18].

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Thus, the higher the urinary pH, the higher the concentration of general, including the walls of the renal tubules in the form of HF. The pH-dependency found for urinary F excretion observed for this strain, when compared with the presence of CSE in A/J mice might contribute to increase the presence of HMGCS2 in kidney of A/J mice might turn these animals more prone to nephropathy, which could impair F reduce renal damage caused by different hostile conditions, such as diabetes. In our data, the presence of HMGCS2 was increased fourfold in diabetic kidneys, which leads to increased biotransformation and elimination in proximal tubule epithelial cells. This enzyme breaks down cystathione into cysteine and acetyl-CoA dehydrogenase, mitochondrial. CSE is an enzyme that catalyses elimination of L-homoserine, L-cystine and L-cysteine of theoretical protein. The presence of HMGCS2 in kidney of A/J mice might turn these animals more prone to nephropathy, which could impair F reabsorption in kidneys [10].

Besides presenting unique proteins involved in metabolism, A/J mice also expresses exclusive proteins involved in cell processes (phenazine biosynthesis-like domain containing protein 2 (PBLD), biliverdin reductase A (BVR) and sorting nexin-5) and information pathways [serum amyloid P-component (SAP)]. Among these, SAP constitutes amyloid deposits characterized by the ordered aggregation of normal globular proteins and peptides into insoluble fibrils, which disrupt tissue architecture and are associated with cell death [22]. The presence of SAP only in A/J mice might increase the probability of kidney damage that could account for their diminished capacity to reabsorb various solutes including F, helping to explain the higher urinary F excretion seen in this strain previously [10].

From those proteins found exclusively in kidneys of 129P3/J mice, the peroxisomal acyl-coenzyme A oxidase 1 (AOX), a fatty acid metabolic protein, is shown to be expressed in proximal tubules and enhancement of its activity is associated with the preservation of kidney function during ischemia [23]. Another exclusive protein called arsenite-methyltransferase, presented only in 129P3/J mice, is a detoxifying protein involved in the arsenic biotransformation and elimination in proximal tubule epithelial cells [24]. The presence of these proteins in 129P3/J but not in A/J mice suggest that the former might have a higher capacity to reduce renal damage caused by different hostile conditions, such as exposure to F. Thus, the 129P3/J mice would be able to maintain F reabsorption in kidneys even under exposure to high F doses [10].

As mentioned above, F exposure did not alter the profile of unique proteins in either strain of mice. However, among the proteins differentially expressed in the comparisons between the two strains, only 8 were present in the control, 10 and 50 ppmF groups (catalase, medium-chain specific acyl-CoA dehydrogenase...
and alpha-aminoacidic semialdehyde dehydrogenase (\(\alpha\)-AASA), isovaleryl-CoA dehydrogenase, ornithine aminotransferase, lactoylglutathione lyase, meprin A subunit alpha and albumin). Some of these significantly altered proteins with potential roles to contribute for the intrinsic differences in F and water handling by A/J and 129P3/J mice are highlighted below. Meprin A, an information pathways related protein, is an enzyme that hydrolyzes protein and peptide substrates including components of the extracellular matrix [25]. It is highly expressed at the brush border membrane of proximal tubule cells of the kidney. Inbred strains of mice subjected to ischemia reperfusion that express low levels of meprin A in kidney have markedly less kidney damage [26]. Our data show that meprin A is consistently reduced in 129P3/J kidney in all experimental conditions. This suggests that this protein could act in concert with SAP to decrease renal damage caused by F in 129P3/J mice. Among the proteins related to cellular processes, it is important to highlight \(\alpha\)-AASA dehydrogenase and catalase. \(\alpha\)-AASA dehydrogenase metabolizes irreversibly betaine aldehyde to betaine, which is the most effective osmoprotectant accumulated by eukariotic organisms to cope with osmotic stress [27]. This enzyme was increased in the 129P3/J kidney, regardless F exposure. This can explain the lower volume of water consistently ingested by the 129P3/J mice throughout the study, which led us to adjust water F concentrations throughout the experiment in order that both strains had the same amount of F intake from the water [10]. The increased expression of the antioxidant enzyme catalase might indicate a higher capacity of the 129P3/J mice to deal with oxidative stress [28].

Two and 6 proteins with differential expression between the two strains in the control group were also identified upon exposure to 10 and 50 ppmF, respectively. Low F level increased the expression of serine/threonine-protein phosphatase PP1 and ATP synthase subunit delta. High F level kidney up-expressed aconitate hydratase, ATP synthase subunit beta, hydroxyacid oxidase 2, homogentisate 1,2-dioxygenase and beta-lactamase-like protein 2 and down-expressed phosphotriesterase-related protein. Besides, 6 proteins presented altered expression only in F-treated groups. Aminoacylase-1 and aspartoacylase-2 were increased, whereas L-lactate dehydrogenase B chain, nucleoside diphosphate-linked moiety X motif 19, Na\(^+\)/H\(^+\) exchange regulatory cofactor NHE-RF3 (PDZK1) and actin-related protein 3 were diminished in 129P3/J kidney. These proteins may act as molecular targets for the differential F metabolism between these strains induced by the treatment. Protein phosphatase 1 (PP1) is a serine/threonine protein phosphatase involved in diverse cellular processes, such as transcription, replication, pre-mRNA splicing, protein synthesis, carbohydrate metabolism, neuronal signaling, cell survival, and cell cycle progression [29,30]. Phosphatases typically function antagonistically with kinases to achieve fine control over the phosphorylation state of proteins. Phosphatases are widely expressed enzymes that mediate the functional regulation of many proteins, including some renal channels and transporters such as the inwardly rectifying K\(^+\) channel, Na\(^+\)-K\(^+\)-Cl\(^-\) cotransporter (NKCC1), CFTR, epithelial Na\(^+\) channel (ENaC), aquaporin-2 (AQP2) and Na\(^+\)/H\(^+\) exchanger 3 (NHE3) [30,31,32,33,34,35,36]. In general, these ions and water channels are responsible to maintain the urine normal volume and 

Supporting Information

**Figure S1 2D gel analysis of renal proteome.** Representative 2D maps of control kidneys. Selected spots in green represent those with differential expression in the comparison between control A/J (A) vs control 129P3/J mice (B). In Figure B, spot identification numbers in boundaries or not represents increases or decreases in protein expression when compared to A/J, respectively (Figure A). Dashed lines represent unique spots in the A/J group (A) and 129P3/J group (B), regardless exposure to F. (TIF)

**Figure S2 2D gel analysis of renal proteome.** Representative 2D maps of 10 ppmF treated-groups. Selected spots in green represent those with differential expression in the comparison between 10 ppmF treated- A/J (A) vs 10 ppmF treated- 129P3/J mice (B). In Figure B, spot identification numbers in boundaries or not represents increases or decreases in protein expression when compared to A/J, respectively (Figure A). (TIF)

**Figure S3 2D gel analysis of renal proteome.** Representative 2D maps of 50 ppmF treated-groups. Selected spots in green represent those with differential expression in the comparison between 50 ppmF treated- A/J (A) vs 50 ppmF treated- 129P3/J mice (B). In Figure B, spot identification numbers in boundaries or not represents increases or decreases in protein expression when compared to A/J, respectively (Figure A). (TIF)
Figure S4 2D gel variability analysis. Scatter plot of binary comparisons among the ratios of relative spot volumes detected in the representative gels (replicate 1), and the respective replicates (replicates 2 and 3). (A) Control A/J mice. (B) 10 ppmF treated-A/J mice. (C) 50 ppmF treated-A/J, (D) Control 129P3/J mice. (E) 10 ppmF treated-129P3/J mice. (F) 50 ppmF treated-129P3/J mice. (TIP).

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Author Contributions
Conceived and designed the experiments: MARB JGC ALL ETE GMW. Performed the experiments: JGC ALL FS GMW. Analyzed the data: CPB CAL. Contributed reagents/materials/analysis tools: MARB ETE GMW. Wrote the paper: CPB MARB.

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