**A novel variant in *GATM* causes idiopathic renal Fanconi syndrome and predicts progression to end-stage kidney disease**

Eleanor G. Seaby1,2\*, Steven Turner1\*, David J. Bunyan3, Fariba Seyed-Rezai1, Jonathan Essex4, Rodney D. Gilbert1,5^, Sarah Ennis1^

\*,^authors contributed equally

1. Faculty of Medicine, University of Southampton, Southampton, UK
2. Broad Institute of MIT and Harvard, Cambridge, Massachusetts, USA
3. Wessex Regional Genetics Laboratory, Salisbury NHS Foundation Trust, Salisbury, UK
4. School of Chemistry, University of Southampton, Southampton UK
5. Southampton Children’s Hospital, Southampton, UK

**Corresponding author:**

Rodney D Gilbert

Southampton Children’s Hospital

Tremona Road

Southampton SO16 6YD

United Kingdom

Rodney.Gilbert@uhs.nhs.uk

Tel: 02381206254

Orcid: 0000-0001-7426-0188

**Short title**

A novel GATM variant causes Fanconi syndrome

**Funding**

EGS is supported by the Gerald Kerkut Charitable Trust and the University of Southampton’s Presidential Scholarship.

**Author contributions**

EGS processed data, performed data analysis, and wrote the first draft of the manuscript. ST, DJB and JE performed data analysis. FSR assisted with data processing. SE, JE and RDG supervised the project. All authors agreed the final manuscript.

**Ethics statement**

All participants were recruited to the ‘Use of NGS technologies for resolving clinical phenotypes’ study. This study was approved by Leeds East Research Ethics Committee (REC: 17/YH/0069; Protocol Number: RHM NEU0302; IRAS project ID: 212945) on the 17/05/17. Written consent was obtained from the parents of the index patient for genomic testing and from the mother for publication after she had seen the manuscript.

**Conflicts of interest**

The authors have no conflicts to declare.

**Keywords**

Renal Fanconi syndrome, exome sequencing, genetics, end-stage kidney disease, molecular dynamics

**Abstract**

Renal Fanconi syndrome (RFS) is a generalised disorder of the proximal convoluted tubule. Many genes have been associated with RFS including those that cause systemic disorders such as cystinosis, as well as isolated RFS. We discuss the case of a 10-year-old female who presented with leg pain and raised creatinine on a screening blood test. Her mother has RFS and required a kidney transplant in her thirties. Further investigations confirmed RFS in the daughter. Exome sequencing was performed on the affected mother, child, and unaffected father. We identified a novel variant in *GATM*; c.965G>C p.(Arg322Pro) segregating dominantly in the mother and daughter. We validated our finding with molecular dynamics simulations and demonstrated a dynamic signature that differentiates our variant and two previously identified pathogenic variants in *GATM* from wildtype. Genetic testing has uncovered a novel pathogenic variant that predicts progression to end stage kidney failure and has important implications for family planning and cascade testing. We recommend that *GATM* is screened for in children presenting with RFS, in addition to adults, particularly with kidney failure, who may have had previous negative gene testing.

**Introduction**

Renal Fanconi syndrome (RFS) is a disorder of the proximal convoluted tubule, resulting in insufficient reabsorption of solutes in the proximal renal tubule, a process that is highly energy dependent. To date, several genes have been associated with RFS, of which some cause systemic disorders with RFS and others cause isolated RFS with or without kidney failure (**Supplementary Table 1**).

The genetic aetiology of isolated RFS is still being elucidated. In 2001, Lichter-Konecki *et al.*1 used genome-wide linkage analysis to map the locus of autosomal dominant RFS to chromosome 15q15.3; at the time the pathogenic gene was unknown. With advances in genomic sequencing technologies, further genes have been associated with isolated RFS including *SLC34A1,2 EHHADH,3* and *NDUFAF6*.4 In 2018, using linkage analysis and contemporary sequencing technologies, Reichold *et al.5* identified *GATM* as the causal gene in the 15q15.3 locus. *GATM* was previously missed by Lichter-Konecki *et al*. as they had erroneously excluded flanking markers which excluded *GATM*.

**Clinical history**

A 10-year-old white female presented with bilateral knee pain. Screening blood tests showed a raised plasma creatinine with an eGFR of 65 ml/min/1.73m2. Shewas referred to Southampton’s paediatric nephrology service. Her mother presented at a similar age with leg pain, polyuria and polydipsia. Aged 14, she was found to have rickets and was referred for further investigation. These confirmed RFS with a plasma creatinine concentration of 53 µmol/L. Aged 17 her plasma creatinine concentration had risen to 104 µmol/L and her GFR measured by EDTA plasma disappearance was 57 ml/min/1.73m2. She progressed to end-stage kidney disease and had a live, unrelated kidney transplant in 2020 (aged 38). Neither of the maternal grandparents had any features to suggest RFS and neither developed chronic kidney disease. A maternal uncle had completely normal kidney function.

Biochemical investigations of the presenting daughter revealed low plasma concentrations of bicarbonate (13 mmol/L), potassium (3.3 mmol/L), and inorganic phosphate (0.89 mmol/L). Urinary protein/creatinine ratio was elevated (124 mg/mmol) and her lactate was raised at 2.5 mmol/L. Alkaline phosphatase was also raised (540 U/L). Plasma glucose, sodium, magnesium, and parathyroid hormone were within normal ranges. She had heavy generalised aminoaciduria and raised urinary retinol binding protein/creatinine ratio (3850 ug/mmol; normal range 3.9-32 ug/mmol). She had intermittent glucosuria. A lower limb X-ray revealed widening of the distal femoral and proximal tibial growth plates suggestive of rickets. Upper limb X-rays were unremarkable. Abdominal and kidney ultrasound scan revealed bilateral medullary nephrocalcinosis. There were no other features to suggest a mitochondrial disorder. Normal white blood cell cystine (0.05 nmol ½cystine per mg protein) excluded cystinosis. Fructose intolerance was excluded due to absence of jaundice or vomiting. Galactosaemia was ruled out due to absence of cataracts and hepatosplenomegaly. Absence of persistent glycosuria, hypoglycaemia, and hepatomegaly excluded Fanconi-Bickel syndrome. Therefore, the patient was diagnosed with RFS of unknown aetiology. However, with her mother’s progression to end-stage kidney disease and evidence of an impaired eGFR, there was high suspicion for an autosomal dominant genetic cause with a similar disease trajectory.

Trio exome sequencing (affected mother, daughter, and unaffected father) was performed to investigate a molecular cause. Library capture was performed using Agilent SureSelect Human All Exon V6 and DNA were sequenced by Novogene. Data were processed from *fastq* to *vcf* using an automated joint-calling pipeline, aligned to GRCh38. Quality control ensured only high-quality variants remained for downstream analysis. Variants were annotated with Ensembl VEP v103 and the resultant joint-called *vcf* was uploaded to a local installation of seqr (<https://github.com/broadinstitute/seqr>) for data visualisation, analysis, filtering, and reporting.

To filter our data, we applied the Renal Tubulopathies V 2.30 PanelApp gene panel and restricted our analysis to: a dominant inheritance pattern; all exonic variants excluding synonymous; allele frequency <0.001 from gnomAD v2.1.16; and variants that passed variant quality score recalibration.

A total of 96,894, 96,370 and 96,830 variants were called in the daughter, mother, and unaffected father respectively. Average read depth was 53, 49, and 49 respectively. Post-filtering, two variants remained (**Table 1**).

The variant in *KCNJ1* was considered an incidental finding and not causal for RFS. Pathogenic variants in *KCNJ1* are autosomal recessive7 and both affected individuals are heterozygous. The novel missense variant in *GATM*: c.965G>C p.(Arg322Pro), had a genotype quality of 99 in all sequenced individuals and a depth of 74 in the proband, 74 in the mother, and 37 in the father.The variant was confirmed in the proband and mother by Sanger sequencing.

Validation of p.(Arg332Pro)

To validate the pathogenic effect of the p.(Arg322Pro) variant, we performed conventional molecular dynamics simulations on wildtype (WT), p.(Arg322Pro) (our variant), and two pathogenic variants, p.(Thr336Ala) and p.(Pro320Ser), as reported by Reichold *et al*.5 Simulations were tested on homodimers, dimerized at the multimeric B4-B4 interface as per Reichold *et al*.5 (**Figure 1A**) and run for 600 ns per replica, with 3 replicas per *GATM* variant. The classical dimer B2-B2 interaction also required for multimerization was not included in simulations as we expect this interaction to be invariant across both pathogenic and non-pathogenic states due to its existence in the native form, and our prior simulations of B2-B2 dimers did not indicate any notable differences between variants. Further simulation details are provided in the supplementary. Initial principal component analysis showed no discrepancy between *GATM* variants on large scale motions observed in the trajectories (**Supplementary Figure 1**). On visualization of the local environment around the variant sites, p.(Arg322Pro) simulation trajectories identified unique close contacts between residues 320 on each monomer of the homodimer relative to WT (**Figure 1B**). Relative free energy estimations were performed across the residue 320 distance as described in the supplementary for all variants (**Figure 1C & SF1**).

**Discussion**

GATM encodes a proximal tubular nuclear-encoded mitochondrial enzyme, glycine amidinotransferase, which catalyses the transfer of a guanidino group from l-arginine to glycine, resulting in guanidinoacetic acid, an immediate precursor to creatine.8 GATM is most prominently expressed in kidney, liver, pancreas, and brain. In the kidney, *GATM* expression is limited to the highly energy dependent proximal tubular cells, which have abundant mitochondria to support the oxygen-dependent generation of ATP.5 Biallelic loss-of-function variants in *GATM* cause a rare, congenital neurological disorder without kidney dysfunction.9

In 2018, Reichold *et al.5* used genome-wide linkage analysis and sequencing studies on 28 patients from five extended families with childhood-onset autosomal dominant without debilitating rickets. The youngest patient exhibited laboratory features of RFS without glomerular compromise at 18 months old. For all patients, plasma creatinine started to rise during late adolescence or adulthood, with evidence of renal fibrosis and kidney disease. Progression to dialysis or transplant happened in the third to sixth decades. There were no extra-renal features. All affected patients had a single heterozygous missense variant in *GATM* affecting highly conserved residues, segregating in an autosomal dominant pattern. The four causal variants, p.(Pro320Ser); p.(Thr336Ala); p.(Thr336Ile); and p.(Pro341Leu), were fully penetrant and clustered on conserved proline and threonine residues representing <5% of the protein. *In silico* modelling suggested the variants could adversely affect protein folding and cause GATM to form longitudinal multimers. Biopsy samples from affected patients demonstrated fibrosis and extremely large, filament filled mitochondria within proximal tubule cells. These were confirmed to contain GATM using immunogold staining.5 Overexpression of wild-type *GATM* had no impact on mitochondria, but all mutants caused structural deformity to mitochondria consistent with the findings on patient biopsies. LLC-PK1 cells (porcine proximal tubule cell line) transfected with the p.(Thr336Ala) variant had no effect on enzymatic activity and oxidative phosphorylation, but instead caused dimeric GATM to form multimers, resulting in fibrillary aggregation within the mitochondria consistent with the *in silico* modelling.5 These protein aggregates led to dysfunctional elongated mitochondria with increased reactive oxygen species, increased transcription of *NLRP3*, a component of the inflammasome, elevated pro-inflammatory cytokine interleukin-18, increased profibrotic factors and increased cell death. Furthermore, mutant *GATM* showed impaired protein degradation with an increased half-life. Of note, mice haploinsufficient for *GATM* had no significant kidney phenotype, suggesting the four variants cause disease through a dominant negative mechanism. These mice did however display neurological phenotypes consistent with biallelic loss-of-function variants in *GATM*.8 Of interest, rats treated with oral creatine showed 27% reduced kidney Gatm expression and 58% reduced protein levels. Therefore, exogenous creatine may suppress endogenous production of mutant GATM and reduce production of abnormal protein aggregates.5

We identified a novel (absent from population databases) variant, NM\_001482.3: c.965G>C p.(Arg322Pro), in *GATM*, segregating in an affected mother-daughter pair with idiopathic RFS. This variant is two amino acids downstream of a previously described pathogenic variant, p.(Pro320Ser), in *GATM* causing RFS with progression to kidney failure (**Figure 2**).

The p.Arg322 amino acid is extremely conserved across species, and *in silico* predictors suggest high pathogenicity with a CADD score of 32. To validate pathogenicity of our novel variant we performed conventional molecular dynamics simulations. We tested WT, our variant p.(Arg332Pro), and two previously identified pathogenic variants, p.(Thr336Ala) and p.(Pro320Ser). The two previously reported pathogenic variants by Reichold *et al.* and our novel variant behaved differently to wildtype. Disease-associated variants adopted a new “close-contact” global minimum at approximately 3.8 Å, whilst WT uniquely retained a “distant” global minimum at approximately 12.4 Å. Close contacts of residues 320 on both homodimers may be a direct or precursor event to formation of the B4 interaction surface capable of pathogenic *GATM* multimerization as hypothesized by Reichold *et al*.5 This dynamic signature localised directly at the proposed oligomeric interface, clearly stratifies disease-associated mutants from WT in support of the pathogenicity of the p.(Arg322Pro) variant.

The phenotype of our female proband and her affected mother are consistent with the phenotypes reported by Reichold *et al.5* The mother presented in childhood and progressed to end stage kidney disease and received a transplant in her late thirties. The daughter is already showing signs of chronic kidney disease and we expect that she will also develop end stage kidney disease and should be counselled accordingly. This has implications for family planning and the daughter may wish to undergo pre-implantation genetic diagnosis in the future.

Currently, *GATM* (when monoallelic) is included on three gene panels in PanelApp10: Unexplained paediatric onset end-stage renal disease (R257); Tubulointerstitial kidney disease (R202); and Renal tubulopathies (R198). However, it is absent from additional kidney-related panels in PanelApp including the ‘Renal superpanel-broad’, ‘Unexplained kidney failure in young people’ and ‘Proteinuric renal disease’. As GATM is a relatively new disease gene, we expect that there may be patients with chronic kidney disease, or end-stage kidney failure harbouring variants in *GATM*.

**Conclusion**

All patients with known pathogenic variants in *GATM* progress to end stage kidney disease, and therefore we have uncovered a diagnosis that has serious clinical consequences for our patient and will warrant genetic counselling and cascade testing. Early functional work in rats suggests that creatine may be able to significantly reduce expression of the GATM protein and this warrants further investigation.

This case highlights how genetic sequencing and complementary molecular dynamics simulations can identify disease pathogenesis and inform patient care and prognosis. *GATM* variants should be routinely tested for in cases of idiopathic RFS even in the absence of renal failure which develops after initial presentation. We further recommend that adult patients showing signs of RFS and chronic kidney disease should also be screened for *GATM* variants.

**References**

1. Lichter-Konecki U, Broman K, Blau E, Konecki D. Genetic and physical mapping of the locus for autosomal dominant renal Fanconi syndrome, on chromosome 15q15. 3. The American Journal of Human Genetics*.* 2001;68(1):264-268.

2. Magen D, Berger L, Coady MJ, et al. A loss-of-function mutation in NaPi-IIa and renal Fanconi's syndrome. New England Journal of Medicine*.* 2010;362(12):1102-1109.

3. Klootwijk ED, Reichold M, Helip-Wooley A, et al. Mistargeting of peroxisomal EHHADH and inherited renal Fanconi's syndrome. New England Journal of Medicine*.* 2014;370(2):129-138.

4. Hartmannová H, Piherová L, Tauchmannová K, et al. Acadian variant of Fanconi syndrome is caused by mitochondrial respiratory chain complex I deficiency due to a non-coding mutation in complex I assembly factor NDUFAF6. Human molecular genetics*.* 2016;25(18):4062-4079.

5. Reichold M, Klootwijk ED, Reinders J, et al. Glycine Amidinotransferase (GATM), renal Fanconi syndrome, and kidney failure. Journal of the American Society of Nephrology*.* 2018;29(7):1849-1858.

6. Karczewski KJ, Francioli LC, Tiao G, et al. The mutational constraint spectrum quantified from variation in 141,456 humans. Nature*.* 2020;581(7809):434-443.

7. Derst C, Konrad M, Köckerling A, et al. Mutations in the ROMK gene in antenatal Bartter syndrome are associated with impaired K+ channel function. Biochemical and biophysical research communications*.* 1997;230(3):641-645.

8. Forst A-L, Reichold M, Kleta R, Warth R. Distinct Mitochondrial Pathologies Caused by Mutations of the Proximal Tubular Enzymes EHHADH and GATM. Frontiers in Physiology*.* 2021;12 (Review).

9. Clark JF, Cecil KM. Diagnostic methods and recommendations for the cerebral creatine deficiency syndromes. Pediatric research*.* 2015;77(3):398-405.

10. Martin AR, Williams E, Foulger RE, et al. PanelApp crowdsources expert knowledge to establish consensus diagnostic gene panels. Nature genetics*.* 2019;51(11):1560-1565.

**Legends**

**Table 1** | Results of primary analysis

*Abbreviations: chr – chromosome; pos – position; gnomAD(g) – gnomAD genomes frequency; gnomAD – gnomAD v2.1.1 exomes frequency; Het – heterozygous; WT – wild type.*

**Figure 1** | Molecular dynamics results of GATM B4-B4 dimer mutants

*(a) Graphical representation of GATM wild-type (WT) homodimer, each monomer coloured blue and orange, dimerized at the B4-B4 interface (b) Representative cartoon structures of 316-324 loop for WT (blue/orange) and R322P (purple/brown) global minima with respect to residue 320 distances, proline 320 residues are overlaid as a stick representation. (c) Free-energy values for all mutants and WT across residue 320 CB atom distances, averaged across all three repeats. (d/e) Free-energy values for WT and p.(Arg322Pro) for each simulation replica for WT and p.(Arg322Pro) simulations across residue 320 CB atom distances.*

**Figure 2** | Heterozygous variants in *GATM* causing renal Fanconi syndrome with kidney failure

*All known pathogenic variants in GATM, including our variant Arg322Pro (bold and with asterisk). All variants are highly conserved across species. Amino acids discordant with the human reference are shown in red. All variants span a small region of exons 6 and 7.*