A novel loss-of-function mutation of PBK associated with human kidney stone disease

Choochai Nettuwakul1, Nunghathai Sawasdee1, Oranud Praditsap2, Nanyawan Rungroj2, Arnat Pasena1, Thanyaporn Dechtawewat1, Nipaporn Deejai1, Suchai Sritippayawan3, Santi Rojsatapong4, Wipada Chaowagul4 & Pa-thai Yenchitsomanus1✉

Kidney stone disease (KSD) is a prevalent disorder that causes human morbidity worldwide. The etiology of KSD is heterogeneous, ranging from monogenic defect to complex interaction between genetic and environmental factors. Since mutations of genes responsible for KSD in a majority of families are still unknown, our group is identifying mutations of these genes by means of genomic and genetic analyses. In this study, we identified a novel loss-of-function mutation of PBK, encoding the PDZ binding kinase, that was found to be associated with KSD in an affected Thai family. Glycine (Gly) substituted by arginine (Arg) at position 43 (p.Gly43Arg) in PBK cosegregated with the disease in affected members of this family, but was absent in 180 normal control subjects from the same local population. Gly43 is highly evolutionarily conserved in vertebrates, and its substitution affects protein structure by alterations in H-bond forming patterns. This p.Gly43Arg substitution results in instability of the variant PBK protein as examined in HEK293T cells. The variant PBK protein (p.Gly43Arg) demonstrated decreased kinase activity to phosphorylate p38 MAPK as analyzed by immunoblotting and antibody microarray techniques. Taken together, these findings suggest a possible new mechanism of KSD associated with pathogenic PBK variation.

Human kidney stone disease (KSD) is a group of common disorders that cause morbidity in human populations and a global public health burden1. The causes of KSD are heterogeneous, ranging from single-gene defect to complex combination of genetic and environmental factors2. The majority of kidney stones (~80%) are composed of calcium oxalate (CaOX) and calcium phosphate (CaP)3,4. Intrarenal and urinary ion imbalance is the known risk factor for KSD5. Growth of calcium oxalate stone over Randall’s plaque site appears to be the predominance of stone formation for the patients with hypercalciuria6,7. Exposure of renal tubular cells to oxalate, calcium oxalate monohydrate (COM), and CaP causes increases in reactive oxygen species (ROS) production and oxidative stress leading to cell injury and inflammation. These accelerate crystal formation, growth, and aggregation, which ultimately leads to stone formation8,9.

Previous studies using family-based or case-control analyses identified genetic variations of several genes in KSD, including variations in calcitonin receptor (CTR)10, vitamin D receptor (VDR)11,12, calcium-sensing receptor (CaSR)13 and osteopontin (OPN)14,15. Genetic variations in claudin 14 (CLDN14), which encodes for the tight junction protein, was also discovered by a high-throughput genome-wide association study (GWAS)16. Our group previously reported KSD in Northeastern Thailand where the prevalence of this disease is high. The relative risk of KSD among the members of the affected families was greater than that of the general population (λR = 3.18), which suggests the genetic contribution to kidney stone in the Northeastern Thai population17. We also found genetic variations to be associated with KSD risk in Thai patients in case-control studies18–21. However, these common variants were likely a modifying factor, not a disease-causing factor. Genome-wide linkage analysis and exome sequencing were later employed to identify a disease-causing gene in families affected by KSD. These analyses showed loss-of-function alterations of SCN10A, encoding Naν1.8 α subunit of voltage-gated sodium channel, to be associated with KSD in the families studied. Naν1.8 was initially reported to be preferentially

1Division of Molecular Medicine, Research Department, Faculty of Medicine Siriraj Hospital, Mahidol University, Bangkok, Thailand. 2Division of Medical Genetics Research and Laboratory, Research Department, Faculty of Medicine Siriraj Hospital, Mahidol University, Bangkok, Thailand. 3Division of Nephrology, Department of Medicine, Faculty of Medicine Siriraj Hospital, Mahidol University, Bangkok, Thailand. 4Sappasithiprasong Hospital, Ubon Ratchathani, Thailand. ✉e-mail: pathai.yen@mahidol.edu
expressed in peripheral sensory neurons and heart tissue. Gain-of-function alterations of \( \text{SCN10A} \) cause painful peripheral neuropathy\(^{22}\), while its loss-of-function alterations contribute to prolonged cardiac conduction disease and Brugada syndrome\(^{23,24}\). In our study, we found that Na\(^{+}\)\text{V}1.8 \( \alpha \) subunit protein expressed in proximal tubules and collecting ducts of nephron in human kidney. The variant protein (p.N909K and p.K1809R in the same allele) expressed in cultured cells was unstable, resulting in reduced current density as analyzed by whole-cell patch clamp technique. We propose that Na\(^{+}\) channel may be activated by the depolarization of Na\(^{+}\) and help to increase Na\(^{+}\) reabsorption\(^{25}\).

In the present work, we continued our search for disease-causing genes in KSD via exome sequencing and genetic analysis in a different family that is affected by KSD. We discovered that a novel loss-of-function alteration (p.Gly43Arg, or p.G43R) in \( \text{PBK} \), encoding the PDZ binding kinase, cosegregated with KSD in affected members of this family. The p.Gly43Arg substitution results in instability of PBK, which is a member of the mitogen-activated protein kinase kinase (MAPKK) family\(^{26–28}\), and the activated form of PBK phosphorylates p38 MAPK\(^{26,29}\). These findings suggest a not previously reported potentially new mechanism of KSD associated with PBK alteration.

**Results**

**Subjects and clinical study.** We recruited patients and members of 180 families affected with KSD for genetic study. The disease in these patients and the members of their families was not secondary to all known causes, including renal tubular acidosis, primary hyperparathyroidism, inflammatory bowel disease, Cushing disease, hyperthyroidism, or drug-induced KSD, as diagnosed by clinical history and symptoms, physical and laboratory examinations, acute acid loading test, and serum electrolytes. The normal control subjects included in this study (\( n = 180 \)) were recruited from the same geographical area (Ubon Ratchathani Province in Northeastern Thailand) that the KSD patients and their families were recruited from. Normal controls and study group subjects were similarly examined, and controls were investigated by radiography of kidney-ureter-bladder (KUB) to confirm the absence of kidney stone. Gender and age data of study subjects and normal controls are shown in Supplementary Table S1.

To identify disease-causing genes in KSD, we selected a large family with many members (UBRS033 family) for exome sequencing and genetic study. The pedigree showed 8 affected, including a twin pair (III:3 and III:4) and 20 unaffected family members and that the KSD phenotype that was inherited as autosomal dominant model (Fig. 1). Five of affected members had opaque stones while another three had no stone detected by both KUB and ultrasound because the stone had been removed by surgery, extracorporeal shock wave lithotripsy (ESWL), or spontaneous passing. Therefore, stones from these patients were not available for the analysis of their compositions. The results of investigation for KSD by KUB radiography, clinical history, and physical examination in 16 family members are shown in Table 1. We collected blood and urine samples from proband (II:5) and his sister (II:1) (Supplementary Table S2). The serum creatinine concentrations of proband (II:5) and his sister (II:1) were 3.7 and 5.4 mg/dL, respectively, which indicated the presence of kidney failure. There were no hypercalciuria, hyperoxaluria or hyperphosphaturia presented in the proband (Supplementary Table S2).
DNA samples from five affected members (II:1, II:3, II:4, III:3, and III:4) and three unaffected members (I:2, II:6 and II:7) from the UBRS033 family were selected for exome sequencing. The nucleotide reads were mapped against UCSC hg19 by Burrows-Wheeler Aligner (BWA), and variations (SNPs and Indels) were detected by Sequence Alignment/Map (SAMTOOLS). Genetic variations acquired from exome sequencing in these eight family members, which included 224,121 variations with an average read depth of $38 \times$, were analyzed. Initially, the variations outside exonic regions were excluded. Since KSD in this family inherited as the autosomal dominant model, heterozygous variations that were shared among the five affected members, but absent in the unaffected members, were reserved for subsequent analysis. After filtration by exclusion of synonymous variations, but selection for non-synonymous, stop, gain/loss, short insertion or deletion (indel) variations, and rare variants (frequency < 0.01), 17 variations remained for analysis in the next step (Supplementary Fig. S1 and Table 2).

The impact of amino acid changes on the protein structures and functions of the 17 candidate variations was predicted by 6 web-based programs, including Polymorphism Phenotyping v2 (PolyPhen-2) 30, VarioWatch31, MutationTaster32, Sorting Intolerant From Tolerant (SIFT)33, Mutation Assessor34, and Likelihood Ratio Test (LRT)35. Two novel variations in PBK (c.127 G > A, p.Gly43Arg) and SORL1 (c.4501 C > T, p.Arg1501Trp), and one reported variation in ANKRD36 (rs571299992, c.202 G > A, p.Ala68Thr) were predicted to be disease-causing or damaging by 5, 3, and 5 of 6 programs, respectively (Table 2).

Exome sequencing and analysis of genetic variations. DNA samples from five affected members (II:1, II:3, II:4, III:3, and III:4) and three unaffected members (I:2, II:6 and II:7) from the UBRS033 family were selected for exome sequencing. The nucleotide reads were mapped against UCSC hg19 by Burrows-Wheeler Aligner (BWA), and variations (SNPs and Indels) were detected by Sequence Alignment/Map (SAMTOOLS). Genetic variations acquired from exome sequencing in these eight family members, which included 224,121 variations with an average read depth of $38 \times$, were analyzed. Initially, the variations outside exonic regions were excluded. Since KSD in this family inherited as the autosomal dominant model, heterozygous variations that were shared among the five affected members, but absent in the unaffected members, were reserved for subsequent analysis. After filtration by exclusion of synonymous variations, but selection for non-synonymous, stop, gain/loss, short insertion or deletion (indel) variations, and rare variants (frequency < 0.01), 17 variations remained for analysis in the next step (Supplementary Fig. S1 and Table 2).

The impact of amino acid changes on the protein structures and functions of the 17 candidate variations was predicted by 6 web-based programs, including Polymorphism Phenotyping v2 (PolyPhen-2) 30, VarioWatch31, MutationTaster32, Sorting Intolerant From Tolerant (SIFT)33, Mutation Assessor34, and Likelihood Ratio Test (LRT)35. Two novel variations in PBK (c.127 G > A, p.Gly43Arg) and SORL1 (c.4501 C > T, p.Arg1501Trp), and one reported variation in ANKRD36 (rs571299992, c.202 G > A, p.Ala68Thr) were predicted to be disease-causing or damaging by 5, 3, and 5 of 6 programs, respectively (Table 2).

Genetic analyses revealed a novel substitution in PBK as disease-associated variant. Three variations predicted to be pathogenic or damaging were genotyped in all affected and unaffected members of the index family. Of those, only p.Gly43Arg variation in PBK gene was cosegregated with KSD in the family (LOD scores = 2.36) (Table 3 and Fig. 1). One family member (III:6; aged 22 years) carried the variation without having KSD, which is possible since the disease usually develops at an age > 25 years. The PBK p.Gly43Arg variation was

| Sample no. | Gender | Age | Age of onset* | KUB result | Site/side of stone | No. of stones/size** (cm.) | Treatment | Urine pH | Other Symptoms |
|------------|--------|-----|--------------|------------|-------------------|------------------------|-----------|---------|----------------|
| II:11      | Male   | 63  | 45           | Positive/opaque stones | Renal/both        | 2/0.5                  | No        | 5.3     | Yes            | No | No | Renal stone (KUB) |
| II:1       | Female | 59  | 40           | Positive/opaque stones | Renal/both        | 1/2.0                  | Surgery/ESWL | 5.3     | No | No | Renal stone (KUB) |
| II:3       | Male   | 54  | 43           | Positive/opaque stones | Renal, ureter/both | 2/2.7                  | No        | 5.8     | Yes | Yes | Renal and ureteric stone (KUB) |
| II:4       | Female | 52  | 40           | Positive/opaque stones | Renal/left        | 1/1.0                  | Surgery/ESWL | 5.4     | Yes | Yes | Renal stone (KUB) |
| II:12      | Female | 48  | 28           | Negative            | No                | 6.4                    | Yes        | No      | Yes | No | Renal stone (strong history) |
| II:5       | Male   | 48  | 23           | Positive/opaque stones | Renal/right       | 2/4.0                  | Surgery/ESWL | 5.8     | No | No | Renal stone (KUB) |
| III:3      | Female | 34  | 5            | Negative            | No                | 5.3                    | Yes        | Yes     | Yes | No | Renal stone (strong history) |
| III:4      | Female | 34  | 19           | Negative            | Unknown/both      | NA                    | Surgery/ESWL | 5.7     | No | No | Renal stone (surgical scar) |
| I:2        | Female | 78  | Negative     | No                  | No                | 5.6                    | Yes        | No      | Yes | No | No stone |
| I:6        | Male   | 47  | Negative     | No                  | No                | 5.9                    | No         | No      | No | No | No stone |
| I:7        | Male   | 44  | Negative     | No                  | No                | 5.6                    | No         | No      | No | No | No stone |
| I:13       | Female | 42  | Negative     | No                  | No                | 5.4                    | No         | No      | No | No | No stone |
| III:1      | Female | 38  | Negative     | No                  | No                | 5.3                    | No         | No      | No | No | No stone |
| III:5      | Female | 29  | Negative     | No                  | No                | 5.7                    | No         | No      | No | No | No stone |
| III:6      | Male   | 22  | Negative     | No                  | No                | 6.5                    | No         | No      | No | No | No stone |
| III:7      | Female | 20  | Negative     | No                  | No                | 6.3                    | No         | No      | No | No | No stone |

Table 1. Some clinical and laboratory data of the members of the UBRS033 family. ESWL = Extracorporeal shock wave lithotripsy for kidney stones. *The patient had positive results of kidney–ureter–bladder (KUB) radiography. The patient had strong clinical history as justified from the presence of several symptoms associated with kidney stone, especially hematuria and stone passage. The patient had surgical scar for removal of stone. *Age of onset of stone, **Size of the biggest stone.
"Probably damaging", "Possibly damaging", "Disease causing", or "Deleterious". predict the impact of amino acid changes on the protein structures and functions as "Pathogenic", "Damaging", "Probably damaging", "Possibly damaging", "Disease causing", or "Deleterious".

their impact on protein structure and function using the 6 previously mentioned web-based programs, and the identified (Supplementary Fig. S4 and Supplementary Table S3). All exonic variations in

PBK DNA sequencing. A total of 13 variations in (including one sample from the index family as a positive control sample) by PCR-HRM method, followed by PBK in the DNA samples from 180 patients with KSD other variations of PBK in the DNA samples of 180 normal control subjects as screened by polymerase chain reaction and not present in the DNA samples of 180 normal control subjects as screened by polymerase chain reaction and high resolution melting (PCR-HRM) analysis (Supplementary Fig. S3). Haplotype analysis suggested that the allele of PBK p.Gly43Arg variation was inherited from the father (I:1) who is deceased (Fig. 1 and Supplementary

Gene scanning and analysis of genetic variations. To examine whether PBK p.Gly43Arg variation or other variations of PBK were present in additional patients with KSD, we genotyped these variations and screened all 8 exons (including their exon-intron boundaries) of PBK in the DNA samples from 180 patients with KSD (including one sample from the index family as a positive control sample) by PCR-HRM method, followed by DNA sequencing. A total of 13 variations in PBK (6 novel and 7 reported variations) including p.Gly43Arg were identified (Supplementary Fig. S4 and Supplementary Table S3). All exonic variations in PBK were analyzed for their impact on protein structure and function using the 6 previously mentioned web-based programs, and the impact of exon-intron boundaries on mRNA splicing process was evaluated using ESEfinder 2.0.

The six web-based programs revealed p.Gly43Arg to be the only variation that was predicted to affect function of the protein (Supplementary Table S3). A non-synonymous variation (c.320 A>G [rs3779620]) was observed, but it is also present in high frequency in the 1,000 Genomes Project database, so it was excluded as a disease-associated variant. We found 4 variations (c.−99C>T [rs3775744], c.14G<C, c.295+4C>A [rs3728713] and c.*683_685delIAAG) that were predicted to affect exonic splicing enhancer and cryptic splice site; thus, these variations were analyzed in the 180 normal control subjects by PCR-HRM. The variations c.−99C>T, c.295+4C>A, and c.*683_685delIAAG were present in 29, 88, and 90 normal control subjects, respectively (Supplementary Fig. S5).

| CHR. | POS. | Gene name | dbSNP   | 1000G Frequency | Effect | Nucleotide changes | Amino acid changes | Prediction* (of 6 programs) |
|------|------|-----------|---------|----------------|--------|--------------------|--------------------|--------------------------|
| 2    | 97783805 | ANKR3D6   | rs571299992 | 0.00199681 | Missense variant | c.202 G>A | p.Ala68Thr | 5 |
| 8    | 2768586 | PBK       | —       | —            | Missense variant | c.127 G>A | p.Gly43Arg | 5 |
| 1    | 85028969 | CT8S      | rs3768249 | 0.0091853 | Missense variant | c.928 G>T | p.Asp310Tyr | 3 |
| 12   | 7742793 | EIF7      | rs139349075 | 0.0019744 | Missense variant | c.1253 A>G | p.Glu418Gly | 3 |
| 5    | 156769914 | FNDC9    | rs20701167 | 0.000399361 | Missense variant | c.631 G>T | p.Gly211Trp | 3 |
| 12   | 10585530 | KLRQ2     | rs575545535 | —            | Missense variant | c.56 G>C | p.Arg19Pro | 3 |
| 11   | 110944398 | RAD9B    | rs552692137 | 0.000599402 | Missense variant | c.288 C>G | p.Ile96Met | 3 |
| 11   | 121466463 | SORL1    | —       | —            | Missense variant | c.4501 C>T | p.Arg1501Trp | 3 |
| 8    | 30701641 | TEX1S    | rs142485241 | 0.00638978 | Missense variant | c.4893 G>C | p.Gln1631His | 3 |
| 2    | 98525148 | TMEM131   | rs535701294 | 0.00197712 | Missense variant | c.263 T>C | p.Leu88Pro | 3 |
| 22   | 20761028 | ZNF74     | rs190749586 | 0.00098403 | Missense variant | c.1705 T>C | p.Ser569Pro | 3 |
| 9    | 34725069 | FAM205A   | rs114933270 | —            | Missense variant | c.2168 A>T | p.Glu723Val | 2 |
| 12   | 15035203 | MGP       | rs374434209 | —            | Missense variant | c.257 G>T | p.Arg86His | 2 |
| 17   | 77895740 | ADNP2     | rs113879497 | 0.00359425 | Missense variant | c.244 A>G | p.Asn815Ser | 1 |
| 5    | 140029086 | PCDHA9   | rs251354 | —            | Missense variant | c.1006 C>G | p.Leu336Val | 1 |
| 2    | 103149147 | SLC9A4    | rs79918239 | 0.00179712 | Stop retained variant | c.2397 G>A | p.Ter799= | 1 |
| 12   | 123810152 | SORL1    | rs537314152 | 0.000199681 | Splice region variant & intron variant | c.1876-6C>T | — | 0 |

Table 2. Prediction of the impact of amino acid changes on the protein structures and functions of 17 candidate variations by 6 web-based programs. CHR. = chromosome, POS. = position, *Number of programs that predict the impact of amino acid changes on the protein structures and functions as “Pathogenic”, “Damaging”, “Probably damaging”, “Possibly damaging”, “Disease causing”, or “Deleterious”.

| Chromosome | Gene | Variation | Segregation in UBR5033 family | LOD | Possible causative variants |
|------------|------|-----------|-------------------------------|-----|---------------------------|
| 2          | ANKR3D6 | p.Ala68Thr | No                           | 0.29 | No                        |
| 8          | PBK   | p.Gly43Arg | Yes                          | 2.36 | Yes*                      |
| 11         | SORL1 | p.Arg1501Trp | No                           | 0.29 | No                        |

Table 3. Genotyping and segregation analyses of 3 selected variations in the UBR5033 family. *This variation was genotyped by PCR-HRM and was not present in the DNA samples of 179 subjects with KSD and 180 normal control subjects.
Even though the variation c.14 G < C was not present in normal control subjects and could not be tested for cosegregation with KSD in the families because only one affected individual was available for testing, the nucleotide and amino-acid sequence alignment revealed them not to be conserved in the evolution of vertebrates.

To investigate PBK expression in human kidney tissues, we examined its mRNA by reverse transcriptase – polymerase chain reaction (RT-PCR), and its protein by immunoblotting analysis. Figure 3A,B shows the expression of PBK mRNA and protein in human kidney tissues and cell lines, respectively.

**mRNA and protein expression, immunohistochemistry, and immunofluorescence staining in human kidney tissues.** To investigate PBK expression in human kidney tissues, we examined its mRNA by reverse transcriptase – polymerase chain reaction (RT-PCR), and its protein by immunoblotting analysis. Figure 3A,B shows the expression of PBK mRNA and protein in human kidney tissues and cell lines, respectively.
The expected size of PBK protein is 36 kDa, but there was an upper band (at about 43 kDa) visible in HEK293 and HEK293T cell lines. The larger size might result from post-translational modifications of PBK protein in HEK293 and HEK293T cell lines, which may not occur in human kidney tissues. In addition, we stained PBK protein in human kidney tissue sections by immunohistochemistry (IHC) and immunofluorescence (IFA) technique (Fig. 3C,D). The PBK protein is coexpressed with Na⁺/K⁺-ATPase, which is a tubular marker, indicating that PBK is expressed in tubules of nephron. Furthermore, we stained protein in human kidney tissue sections with segment specific marker proteins. PBK was not observed with AQP1 and AQP2 that are expressed in proximal tubule and collecting duct, respectively, suggesting that PBK is expressed in the distal tubule of nephrons (Fig. 3C and Supplementary Fig. S6).

Evaluation of wild-type and p.Gly43Arg PBK proteins expressed in HEK293T cells. To examine the stability of the wild-type (WT) and variant PBK (p.Gly43Arg) proteins, transient transfection of FLAG-tagged WT and p.Gly43Arg PBK were performed in HEK293T cells, and the newly synthesized proteins were inhibited with cycloheximide – a protein synthesis inhibitor. The level of WT PBK was initially constant, and then it was slightly reduced after cycloheximide treatment for more than 12 hours (Fig. 4A). In contrast, the level of variant p.Gly43Arg PBK was more rapidly reduced than that of WT PBK. The time required to achieve a 50% reduction of variant p.Gly43Arg PBK was more than 6 hours, and it seemed to demonstrate time-dependent reduction during 0–36 hours (Fig. 4A). Moreover, we investigated the effect of variant p.Gly43Arg PBK on cell proliferation and cell apoptosis. There was no difference in cell viability and cell apoptosis of HEK293T transfected with plasmid construct containing either WT-PBK cDNA or mutant-p.G43R-PBK cDNA (Supplementary Fig. S7A–C). It is possible that endogenous PBK in HEK293T might contribute to the indifferent cell viability. However, in the conditions with the presence of 500μM H₂O₂, the cell viability of HEK293T transfected with plasmid construct containing mutant-p.G43R-PBK cDNA trended to be slightly lower than that transfected with plasmid construct containing WT-PBK cDNA (Supplementary Fig. S7D).

Downstream signaling analysis. Activation of p38 MAPK was evaluated by immunoblotting analysis of HEK293T transfected with plasmid construct containing either WT-PBK cDNA or mutant-p.G43R-PBK cDNA. The results showed that phosphorylation of p38 MAPK in the HEK293T expressing variant p.Gly43Arg PBK was 40% lower than that in the HEK293 expressing WT PBK (Fig. 4B).

The antibody microarray analysis surveyed over 613 signaling components from the major established signaling pathways, and it probed both phosphorylated and total protein levels. Using HEK293T transfected with either WT-PBK cDNA or mutant-p.G43R-PBK cDNA constructs, signaling hits between cells expressing WT PBK and variant p.Gly43Arg PBK (control and treatment) for 24 hours were compared. We used the changes in globally normalized data between control and treatment samples as the percent change from control (%CFC) to filter the data points with %CFC ≥ 0, sum of % error ranges <0.85 × %CFC value and at least one globally normalized intensity value ≥ 1,000. That screening generated several hits that were different between the PBK expressions (Table 4). We noticed that several of these hits were signaling proteins that are implicated in cell cycle pathways (MDM2, p53, CHK1, CDK1, CDK2, CDK7, CCNE1, and ABL1). We explored the expression of the p38 MAPK family (p38-alpha, p38-beta, p38-delta) in HEK293T expressing variant p.Gly43Arg PBK compared to WT PBK. The results showed that phosphorylation of p38 MAPK in the HEK293T-expressing variant p.Gly43Arg PBK was 39–46% lower than that of p38 MAPK in the HEK293T-expressing WT PBK (Fig. 4C), and this result is consistent with that of the immunoblotting analysis.

Discussion

Genetic variations have been reported to cause KSD,[13,16,25–40] however, the cause of KSD in many families is still unknown. Our group is attempting to identify the disease-causing genes of KSD in Thai families affected with this disease. In this study, using exome sequencing and genetic analysis, we discovered a novel loss-of-function variation in PBK (p.Gly43Arg) cosegregating with KSD in the affected family with a calculated log of odds ratio (LOD) of 2.36 (Fig. 1 and Table 3). This PBK (p.Gly43Arg) variation was observed in only one family, and there was no evidence of its presence in 180 normal control individuals (Supplementary Fig. S3) who live in the same geographical region. This or other pathogenic variations in all 8 exons of PBK was/were not detected in DNA samples from a group of 179 patients with KSD as examined by PCR-HRM and sequencing methods (Supplementary Table S3). Although the variations (c.–99C > A, c.14G < C, c.295 + 4C > A and c.683 *685delAAG) were predicted to affect exonic splicing enhancer and cryptic splice site, they were present in normal control subjects (Supplementary Fig. S5), or the nucleotide and amino-acid sequence alignment demonstrated that they are not conserved in the evolution of vertebrates. Thus, it is unlikely that these variations would be pathogenic.

The p.Gly43Arg substitution in PBK identified in the affected family with KSD is most likely a disease-associated variation because it cosegregated with the disease in 6 affected members, but it was absent in 6 unaffected members. However, one member of this family (II:6) who was 22-years-old carried the p.Gly43Arg variation without having KSD (Fig. 1). This is possible because KSD usually develops in individuals older than 25 years. This family member is being followed-up to observe whether KSD occurs in this case or not. Additionally, other family members in generation III should be regularly screened for asymptomatic stones. This family is being followed-up to observe whether KSD occurs in this case or not. Additionally, other family members in generation III should be regularly screened for asymptomatic stones and informed for the risk of having the disease. The finding of genetic factor contributing to the family may be useful to identify the target for prevention and treatment. Since the two family members, III:3 and III:4, were rather young at first kidney stone manifestation, we searched for additional variations that might modify the disease from their exome sequencing data in comparison with the data from other affected members. A set of 175 candidate genes reported to be involved in KSD, gathered from public databases and candidate gene association study databases, were evaluated for their variations associated with KSD. We found 26 variations in 19 genes that match with this set of genes (Supplementary Table S4). Only p.Ala99Ser variation in CASR was reported
Figure 3. Expression of PBK mRNA and PBK protein in kidney cell lines and human kidney tissue. (A) PBK mRNA expression in HEK293 cells, HEK293T cells, human kidney tissue, and kidney cDNA library detected by RT-PCR method. Two regions of PBK mRNA covering exons 5–6 and 7–8 were analyzed, and mRNA of the housekeeping gene ACTB was used as an internal control. (B) PBK protein expression (expected size: 36 kDa) in HEK293 cells, HEK293T cells, and human kidney tissue detected by immunoblot analysis. GAPDH was used as loading control. (C) Staining of PBK in human kidney tubules by immunohistochemistry (IHC) method using PBK-specific antibody compared to that using isotype control antibody. Na\(^+\)/K\(^+\)-ATPase was stained as a marker to localize tubules (Upper panel). AQP1 was used as protein marker of proximal tubule, V-ATPase as protein marker of distal tubule and collecting duct, and AQP2 as protein marker of collecting duct (Two lower panels). (D) Staining of PBK (green) and Na\(^+\)/K\(^+\)-ATPase (red) in human kidney tubules by immunofluorescence (IFA) method showing their colocalization at the basolateral membrane in tubular cells. Blue indicates DAPI staining of nuclei. Signals were visualized by confocal microscopy (scale bars: top 100 \(\mu\)m, and bottom 50 \(\mu\)m). Images of the gels and blots cropped from different parts of the same gel/blot, or from different gels/blots were separated by white space. The full-length gels and blots are presented in Supplementary Fig. S9.
to be associated with serum calcium level and KSD⁴¹. However, the levels of serum calcium in these two family members were not measured; at this stage it is not known whether p.Ala996Ser variation in \( \text{CASR} \) would be a modifying variation in these two members or not. Although most of the variations are not reported to be involved in KSD, these variants and environmental factors may be involved in the pathogenesis of early onset of KSD. The affected II:11 and II:12 members, who married into the family, did not carry the p.Gly43Arg variation of \( \text{PBK} \). It is possible that they had different genetic variations or some different type of KSD. However, in the calculation of

**Figure 4.** Evaluation of wild-type and p.Gly43Arg PBK expressed in HEK293T cells. (A) The wild-type and p.Gly43Arg PBK proteins in transfected HEK293T cells after treatment with 100 \( \mu \)g/ml of cycloheximide (CHX) for 0, 6, 12, 24, and 36 hours as detected by immunoblot method (left). The wild-type and p.Gly43Arg PBK proteins were quantified by ImageJ 1.50i and plotted as relative intensities. The data were shown as the mean ± SEM of three independent experiments (right). (B) Level of p38MAPK (total p38) and phosphorylation of p38MAPK (p-p38) in HEK293T transfected with wild-type, p.Gly43Arg, or empty vector as detected by immunoblot method (left). GAPDH was used as loading control. The bar graph shows staining densities of the proteins normalized by that of GAPDH. Results are shown as the mean ± SEM of the relative signal intensity from three independent experiments (right). (C) The antibody microarray analysis showed the spot intensities of the p38 MAPK family (p38-alpha, p38-beta, p38-delta) in HEK293T expressing p.Gly43Arg PBK compared to wild-type PBK (left). The bar graphs represent relative globally normalized intensity of the p38 MAPKs in HEK293T expressing p.Gly43Arg PBK compared to wild-type PBK. The results are shown as means ± % error ranges of the globally normalized intensity values in duplicate (right). Images of the blots cropped from different parts of the same blot, or from different blots were separated by white space. The full-length blots, multiple exposures, and full scanned-images of antibody microarray are presented in Supplementary Fig. S10–12.
LOD score and the analysis to identify the disease-causing gene, we did not take these two members (II:11 and II:12) into the calculation and analysis as the founding parents. Thus, they would not affect the overall genetic analysis to identify the disease-causing gene in our study.

PBK or T-LAK cell-originated protein kinase (TOPK) is a 322 amino-acid MAPKK-like serine/threonine protein kinase that was identified as an interleukin-2-induced gene in T-lymphokine-activated killer cells, and as an interaction partner with the human tumor suppressor hDlg27. Multiple studies have found that PBK is over-expressed in tumor and cell lines27,29,42–44. PBK expression is most abundant in the placenta, also present in heart muscle and the pancreas, and present at low levels in skeletal muscle, kidney, liver, and lung27, but there was no report on the PBK protein expression in the adult human kidney and the human protein atlas showed PBK as not expressed in the tubules. While, in our study, we found low level of PBK protein expression in distal tubules of nephron in the human kidney tissue as examined by immunohistochemistry and immunofluorescence staining techniques (Fig. 3C,D). One simple explanation is that the level of PBK protein expression in distal tubules is low.

Table 4. Downstream signaling hits between cells expressing wild-type PBK and cells expressing p.Gly43Arg PBK (control and treatment) for 24 hours as analyzed by KAM-900P Antibody Microarray. \%CFC = Percent change from control, this value was calculated from the changes in globally normalized data between control and treatment.
and there might be variable in sensitivities of PBK antibodies used to detect this protein in human kidney tissues in different laboratories. PBK function may be important for cell cycle regulation since it is phosphorylated in a cell cycle-dependent manner at mitosis. Moreover, phosphorylation of p38 MAPK by PBK was also reported to mediate cell survival.

In this study, phosphorylation of p38 MAPK was evaluated by immunoblot analysis and antibody microarray. The results showed that phosphorylation of p38 MAPK in the HEK293T-expressing p.Gly43Arg PBK is 40% lower than that in the HEK293T-expressing wild-type (WT) PBK (Fig. 4B). A number of previous studies have indicated that p38 MAPK plays an important role as a mediator of apoptosis in response to stress stimuli, such as oxidative stress. Conversely, p38 MAPK has a prosurvival function by upregulating antioxidant gene expression and preventing a high accumulation of reactive oxygen species (ROS) upon exposure to low or moderate doses of H2O2. Renal tubular epithelial cells with overproduction of ROS is attributable to oxidative stress under conditions of high oxalate stimulation, which triggers epithelial cell injury, inflammation, and cell apoptosis. We propose that p38 MAPK activation in renal tubular epithelial cells might upregulate antioxidant genes to prevent oxidative stress, cell injury, inflammation, and cell apoptosis, which are initial components of the mechanism of stone formation.

We found that in the inhibition of newly synthesized proteins by cycloheximide, the variant p.Gly43Arg PBK was more rapidly decreased than that of the WT PBK (Fig. 4A), which suggests instability of the variant p.Gly43Arg PBK that results from amino acid changes. The variant p.Gly43Arg PBK might also impair cell viability from oxidative stress (Supplementary Fig. S7D). Previously, gene knockdown of PBK led to decreased viability and increased apoptosis. The instability of the variant p.Gly43Arg PBK observed in this study might decrease renal tubular epithelial cell viability and increase apoptosis under the oxidative stress, causing an initial risk of crystal deposition and kidney stone formation. Moreover, PBK could play a role as an AR (androgen receptor)-regulated protein. There is an evidence that AR could directly upregulate hepatic glycolate oxidase and kidney epithelial NADPH oxidase subunit p22-PHOX expression. This regulation might then increase oxalate biosynthesis and oxidative stress that results in CaOx crystal formation and retention. However, the roles of WT PBK and variant p.Gly43Arg PBK related to AR regulation on oxalate biosynthesis and oxidative stress through the controls of hepatic glycolate oxidase and NADPH oxidase subunit p22-PHOX expression are still unknown.

Although no kidney stones from this family were available for chemical analysis, they were opaque stones as detected by radiography, and the opaque stones from patients in this region of Thailand were generally found to consist primarily of calcium oxalate. The risk factor for calcium oxalate stone is oxidative stress and inflammation in the condition of high oxalate stimulation. Exposure of renal epithelial cells to high oxalate levels and CaOx crystals leads to the production of ROS, development of oxidative stress, and cellular injury. As found in this study, the instability of the variant p.Gly43Arg PBK may not allow cell survival from oxidative stress or may affect AR signaling, resulting in renal epithelial injury. The apoptosis of renal epithelial cells probably plays an important role in kidney stone formation via apoptotic cell surface, which has clusters of phosphatidylserine that can attract calcium and act as sites for the attachment of CaOx crystals. The mechanism involving retention of these crystals inside the kidney or renal collecting system and further aggregation and/or secondary nucleation ultimately results in the formation of a kidney stone.

The p.Gly43Arg variation of PBK was novel and rare. This alteration and other pathogenic variations of PBK were not found by our genotyping in the 179 unrelated kidney stone patients. However, genotyping in a larger number of patients from different populations may help to validate this result. Furthermore, while the mechanism of tubular epithelial cell injury responded to ROS is important, the roles of CaOx-induced ROS and inflammatory responses are of interest to be studied. Also, urinary biomarkers such as 8-hydroxydeoxyguanosine levels combined with metabolic profile may be used for identifying people at risk of kidney stone development.

In conclusion, we discovered a novel loss-of-function variation (p.Gly43Arg) in PBK associated with KSD in a Thai family with several affected and unaffected members. The unstable variant p.Gly43Arg PBK reduces phosphorylation of p38 MAPK that regulates the downstream signaling pathway, including cell viability and apoptosis. The instability of the variant p.Gly43Arg PBK may affect cell survival from oxidative stress, resulting in renal epithelial injury and kidney stone formation. Further studies on the roles of wild-type and variant p.Gly43Arg or knock-out PBK in animal models to investigate the impact of tubular epithelial cell injury upon high oxidative stress and exposure to high oxalate and crystals of CaOx may elucidate the molecular mechanism of KSD associated with the PBK variation.

Methods

Ethics approval. This study was approved by the Human Research Ethics Committee of the Siriraj Institutional Review Board (SIRB), Faculty of Medicine Siriraj Hospital, Mahidol University, Bangkok, Thailand (COA no. Si 392/2012, and COA no. Si 133/2015). Written informed consent was obtained from all subjects before conducting the study. All methods were performed in accordance with the relevant guidelines and regulations.

Subjects and clinical study. This part was similarly conducted in our previous projects. Two hundred and fifty-six patients with kidney stone disease (KSD) and their family members were recruited at Sappasitthiprasong Hospital in Ubon Ratchathani Province, which is located in Northeastern Thailand, during 2004–2006. Patient and family demographic and clinical data were collected. Blood and urine samples were also collected. All patients and relatives were investigated for kidney stones by radiography of kidneys, ureters, and bladder (plain KUB). In suspicious cases, ultrasonography was also performed. Diagnosis of kidney stone was made based on radiography of KUB, ultrasonography, surgical scar with medical record of kidney stone operation, and/or clinical history of kidney stone and associated symptoms (i.e., back and abdominal pain, hematuria, and stone passage).
The unrelated normal control subjects for this study (n = 180) were recruited from local villages in the same areas where the KSD patients live. All control subjects were radiographically examined similar to the protocol used in KSD patients and family to ensure no presence of KSD. Gender and age data of the normal control subjects are shown in Supplementary Table S1. Genomic DNA samples from patients and control subjects were extracted from peripheral blood cells using standard phenol-chloroform method.

**Human kidney tissues and cell lines.** The use of human kidney tissues in this study was approved by the Human Research Ethics Committee of the Siriraj Institutional Review Board, Faculty of Medicine Siriraj Hospital, Mahidol University. Human fresh frozen tissues from patients without KSD were obtained from remaining specimens from routine pathological examinations. Human kidney tissues, HEK293 cell line, and HEK293T cell line were used for the studies of PBK mRNA expression by reverse transcription and polymerase chain reaction (RT-PCR) method. The encoded PBK protein expression was determined by immunoblot analysis, double immunofluorescence staining, and immunohistochemistry staining.

**Genetic analysis.** Exome sequencing and data analysis was performed in a selected family (UBRS033) with a maximal ELOD of 2.66 that includes 28 members (8 affected and 20 unaffected) and an affected twin. DNA samples from five affected members (II:1, II:3, II:4, III:3 and III:4) and three unaffected members (I:2, II:6 and II:7) were sent to Macrogen (Seoul, South Korea) for exome sequencing analysis. Detailed methods are given in the Supplementary Methods section.

Nucleotide sequences of genes of interest were acquired from the GenBank database to facilitate the design of polymerase chain reaction (PCR) primers (Supplementary Table S5). Exons of the genes of interest were amplified from DNA samples from KSD patients and normal control subjects using specific primer-pairs (Supplementary Table S6), and then they were genotyped and screened for genetic variations (more details are provided in the Supplementary Methods section).

**Protein structure modeling.** The crystal structure of PDZ-binding kinase (5J0A) was used to serve as the wild-type PBK structure, and to generate the substitution protein structure (p.Gly43Arg) using the Swiss-model homology modeling server. Alterations in the H-bond-forming pattern caused by amino acid variation were examined using PyMOL 1.7.5.0 (DeLano Scientific LLC, Palo Alto, California, USA).

**Gene expression in human kidney, and transient transfection in HEK293T cells.** Total RNA was extracted from human fresh frozen kidney tissues, HEK293 cells, and HEK293T cells as previously described by Nettuwakul et al. PBK cDNA was examined by RT-PCR. The primer sequences are shown in Supplementary Table S7. Immunoblot analysis of proteins extracted from human kidney tissues, and immunohistochemistry, double immunofluorescence staining, and the stability of wild-type and variant p.Gly43Arg PBK proteins expressed in HEK293T cells were performed as described in the Supplementary Methods section.

**Examination of protein phosphorylation by antibody microarray.** Protein phosphorylation in HEK293T cells transfected with plasmid construct containing either WT-PBK cDNA or mutant-p.G43R-PBK cDNA was examined by KAM-900P Antibody Microarray Service (Kinexus Bioinformatics Corporation, Vancouver, Canada). The KAM-900P Antibody Microarray features 613 phosphosite-specific antibodies (for phosphorylation) and 265 pan-specific antibodies (for expression levels of these phosphoproteins) in duplicate using two samples on the same microarray slide. The transfected cells were harvested with chemical cleavage buffer according to standard Kinexus recommendations and then shipped to Kinexus Bioinformatics Corporation. The globally normalized data from Kinexus was further filtered to remove data points with error ranges, and the changes in spot intensity between control and treatment samples represent the percent change from control (%CFC).

**Statistics.** Data sets were collected from at least three independent experiments, and values are expressed as mean ± SEM (standard error of the mean). One sample t-test was used to identify significant differences between the means of control and test. Differences with a p-value <0.05 were considered statistically significant.

**Data availability** The datasets generated during and/or analyzed during the current study are available from the corresponding authors on reasonable request.

Received: 6 February 2020; Accepted: 12 May 2020;
Published online: 24 June 2020

**References**
1. Romero, V., Akpinar, H. & Assimos, D. G. Kidney stones: a global picture of prevalence, incidence, and associated risk factors. *Rev Urol. 12*, e86–96 (2010).
2. Gambaro, G. et al. Genetics of hypercalciuria and calcium nephrolithiasis: from the rare monogenic to the common polygenic forms. *Am J Kidney Dis. 44*, 963–986 (2004).
3. Asplin, J. R. Hyperoxaluric calcium nephrolithiasis. *Endocrinal Metab Clin North Am. 31*, 927–949 (2002).
4. Cee, F. L., Evan, A. & Worcester, E. Kidney stone disease. *Clin Invest. 115*, 2598–2608, https://doi.org/10.1172/JCI26662 (2005).
5. Ratkalkar, V. N. & Kleinman, J. G. Mechanisms of Stone Formation. *Clin Rev Bone Miner Metab. 9*, 187–197, https://doi.org/10.1007/s12028-011-9104-8 (2011).
6. Evan, A. P., Cee, F. L., Lingeman, J. E. & Worcester, E. Insights on the pathology of kidney stone formation. *Urol Res. 33*, 383–389, https://doi.org/10.1007/s00240-005-0488-0 (2005).
7. Evan, A. P., Worcester, E. M., Cee, F. L., Williams, J. Jr. & Lingeman, J. E. Mechanisms of human kidney stone formation. *Urolithiasis 43*(Suppl 1), 19–32, https://doi.org/10.1007/s00240-014-0701-0 (2015).
44. Zhu, F.  
42. Hu, F.  
20. Rungroj, N.  
39. Webster, K. E., Ferree, P. M., Holmes, R. P. & Cramer, S. D. Identification of missense, nonsense, and deletion mutations in the GRHPR gene in patients with primary hyperoxaluria type II (PH2).  
23. Chambers, J. C.  
19. Rungroj, N.  
21. Rungroj, N.  
18. Rungroj, N., Nettuwakul, C., Sawasdee, N., Sritippayawan, S. & Yenchitsomanus, P. T. Correlation between genotypes of F2 rs5896 mutation and urinary prothrombin fragment 1.  
35. Chun, S. & Fay, J. C. Identification of deleterious mutations within three human genomes.  
36. Cartegni, L., Wang, J., Zhu, Z., Zhang, M. Q. & Krainer, A. R. ESEfinder: A web resource to identify exonic splicing enhancers.  
37. Belostotsky, R.  
13. Vezzoli, G.  
38. Cregeen, D. P., Williams, E. L., Hulton, S. & Rumsby, G. Molecular analysis of the glyoxylate reductase (GRHPR) gene and its restriction fragment length polymorphism in patients from the Eastern Part of India.  PLoS One 10, e0130790, https://doi.org/10.1371/journal.pone.0130790 (2015).  
40. Williams, E. L., et al. A reciprocal feedback between the PDZ binding kinase and androgen receptor drives prostate cancer.  Oncogene 39, e118, https://doi.org/10.1038/ncomms1407 (2011).  
34. Chun, S. & Fay, J. C. Identification of deleterious mutations within three human genomes.  Genome Res. 19, 1553–1561, https://doi.org/10.1101/gr.092619.109 (2009).  
36. Cartegni, L., Wang, J., Zhu, Z., Zhang, M. Q. & Krainer, A. R. ESEfinder: A web resource to identify exonic splicing enhancers.  Nucleic Acids Res. 31, 3588–3571 (2003).  
37. Belostotsky, R. et al. Mutations in DHDPSL are responsible for primary hyperoxaluria type III.  Am J Hum Genet. 87, 392–399, https://doi.org/10.1016/j.ajhg.2010.07.023 (2010).  
38. Cregan, D. P., Williams, E. L., Hulton, S. & Rumsby, G. Molecular analysis of the glyoxylate reductase (GRHPR) gene and description of mutations underlying primary hyperoxaluria type 2.  Hum Mutat. 22, 497, https://doi.org/10.1002/humu.9200 (2003).  
39. Webster, K. E., Ferree, P. M., Holmes, R. P. & Cramer, S. D. Identification of missense, nonsense, and deletion mutations in the GRHPR gene in patients with primary hyperoxaluria type II (PH2).  Hum Mutat. 170, 176–185, https://doi.org/10.1007/s00494-003-0351-0 (2000).  
40. Williams, E. L. et al. Primary hyperoxaluria type 1: update and additional mutation analysis of the AGXT gene.  Hum Mutat. 30, 910–917, https://doi.org/10.1002/humu.21021 (2009).  
41. Guha, M. et al. Polymorphisms in CaSR and CLDN14 Genes Associated with Increased Risk of Kidney Stone Disease in Patients from the Eastern Part of India.  PLoS One 10, e0130790, https://doi.org/10.1371/journal.pone.0130790 (2015).  
42. Hu, F. et al. c-Myc and E2F1 drive PBK/TOPK expression in high-grade malignant lymphomas.  Leuk Res. 37, 447–454, https://doi.org/10.1016/j.leukres.2012.11.010 (2013).  
43. Warren, A. Y. et al. A reciprocal feedback between the PDZ binding kinase and androgen receptor drives prostate cancer.  Oncogene 38, 1136–1130, https://doi.org/10.1038/onc.2017.438 (2019).  
44. Zhu, Z. et al. Bidirectional signals transduced by TOPK-ERK interaction increase tumorigenesis of HCT116 colorectal cancer cells.  Gastroenterology 133, 219–231, https://doi.org/10.1053/j.gastro.2007.04.048 (2007).  
45. Abe, Y. et al. A mitotic kinase TOPK enhances Cdk1/cyclin B1-dependent phosphorylation of PRC1 and promotes cytokinesis.  J Mol Biol. 370, 231–245, https://doi.org/10.1016/j.jmb.2007.04.067 (2007).
46. Bragado, P., Armesilla, A., Silva, A. & Porras, A. Apoptosis by cisplatin requires p53 mediated p38alpha MAPK activation through ROS generation. *Apoptosis* **12**, 1733–1742, https://doi.org/10.1007/s10495-007-0082-8 (2007).

47. Porras, A. *et al.* P38 alpha mitogen-activated protein kinase sensitizes cells to apoptosis induced by different stimuli. *Mol Biol Cell.* **15**, 922–933, https://doi.org/10.1091/mbc.e03-08-0592 (2004).

48. Gutierrez-Uzquiza, A., Areccherrera, M., Bragado, P., Aguirre-Ghiso, J. A. & Porras, A. p38alpha mediates cell survival in response to oxidative stress via induction of antioxidant genes: effect on the p70S6K pathway. *J Biol Chem.* **287**, 2632–2642, https://doi.org/10.1074/jbc.M111.323709 (2012).

49. Chaiyarat, S. & Thongboonked, V. Changes in mitochondrial proteome of renal tubular cells induced by calcium oxalate monohydrate crystal adhesion and internalization are related to mitochondrial dysfunction. *J Proteome Res.* **11**, 3269–3280, https://doi.org/10.1021/pr300018c (2012).

50. Habbabdeagah-Tari, P., Byer, K. G. & Khan, S. R. Reactive oxygen species mediated calcium oxalate crystal-induced expression of MCP-1 in HK-2 cells. *Urol Res.* **34**, 26–36, https://doi.org/10.1007/s00240-005-0007-3 (2006).

51. Joel, M. *et al.* Targeting PKB/Topk decreases growth and survival of glioma initiating cells *in vitro* and attenuates tumor growth *in vivo*. *Mol Cancer.* **14**, 121, https://doi.org/10.1186/s12943-015-0398-x (2015).

52. Liang, L. *et al.* Androgen receptor enhances kidney stone-CaOx crystal formation via modulation of oxalate biosynthesis & oxidative stress. *Mol Endocrinol.* **28**, 1291–1303, https://doi.org/10.1210/me.2014-1047 (2014).

53. Wiesnser, I. H., Hasegawa, A. T., Hung, L. Y., Mandel, G. S. & Mandell, N. S. Mechanisms of calcium oxalate crystal attachment to injured renal collecting duct cells. *Kidney Int.* **59**, 637–644, https://doi.org/10.1046/j.1523-1755.2001.059002637.x (2001).

54. Boonla, C., Wunsuwan, R., Tungsanga, K. & Tosukhowong, P. Urinary 8-hydroxydeoxyguanosine is elevated in patients with nephrolithiasis. *Urol Res.* **35**, 185–191, https://doi.org/10.1007/s00240-007-0098-0 (2007).

55. Biasini, M. *et al.* SWISS-MODEL: modelling protein tertiary and quaternary structure using evolutionary information. *Nucleic Acids Res.* **42**, W252–258, https://doi.org/10.1093/nar/gka340 (2014).

56. Schneider, C. A., Rasband, W. S. & Eliceiri, K. W. NIH Image to ImageJ: 25 years of image analysis. *Nat Methods.* **9**, 671–675 (2012).

Acknowledgements

We thank the patients, family members, and normal control individuals for their participation in this project; the staff members of the Department of Medicine, Sappasithiprasong Hospital for their assistance with logistics and field studies; the staff members of the Division of Nephrology, Department of Medicine Siriraj Hospital, Mahidol University for their assistance with laboratory data and investigations; and, Miss Pucharan Songprakon for technical assistance in confocal laser-scanning microscopy. We are also grateful to Professor Sookkasem Khositseth of Thammasat University for providing laboratory materials for experimental studies. This work was financially supported by a Mahidol University Grant (R015910004), Faculty of Medicine Siriraj Hospital Grant (R016034007), and a grant from the Thailand Research Fund (TRF IRG5980006). NR and PY were supported by Chalermprakiat Grants from the Faculty of Medicine Siriraj Hospital, Mahidol University. We also thank Assistant Professor Kevin P. Jones for critical reading and editing the manuscript.

Author contributions

P.Y. and N.R. coordinated the project, and conceived and designed the experiments. C.N., O.P., N.S., A.P., T.D., N.D. and N.R. performed the experiments. P.Y., C.N., O.P., N.R., N.S., A.P., T.D., N.D., W.C., S.R. and S.S. contributed reagents/materials/analysis tools. C.N., O.P., N.S., A.P., N.R., and P.Y. prepared the manuscript. C.N. and P.Y. edited the manuscript and responded to the editors and reviewers. All authors have read and approved the final version of the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

Supplementary information is available for this paper at https://doi.org/10.1038/s41598-020-66936-4.

Correspondence and requests for materials should be addressed to P.Y.

Reprints and permissions information is available at www.nature.com/reprints.

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2020