

PRIMARY HYPEROXALURIA TYPE 1 (PH1)

Rapid decline in kidney function can occur in PH1¹



HYPOTHETICAL PATIENT PROFILE:

An adult patient with PH1 experiences rapid decline in kidney function without consistent management and monitoring.

This case study is hypothetical and is not representative of all patients with PH1.

BEHIND THE
STONE



PATIENT
TIMELINE :

Hypothetical patient profile: an adult patient with PH1 experiences rapid decline in kidney function without consistent management and monitoring.

AGE
(years)

28

29

30

eGFR

(mL/min/1.73 m²)

62

?

?

CKD STAGE²

2

PATIENT HISTORY :



KIDNEY STONE EVENTS

AGE 24: REFERRAL TO UROLOGY

Patient passed stone fragments, and the episode was managed by a urologist nonsurgically^{3,5}

AGE 26: ED VISIT

Patient passed stone fragments and was again managed by a urologist nonsurgically^{3,5}

AGE 28: ED VISIT

Episode of hematuria accompanied by flank pain associated with stone formation that was passed without complications. Stone fragment identified by a urologist as calcium oxalate monohydrate^{1,6}

The AUA recommends considering genetic testing in any adult without bowel dysfunction with urinary oxalate excretion >75 mg/day (0.83 mmol/1.73 m²/day).⁷

AGXT, alanine glyoxylate aminotransferase;
AUA, American Urological Association;
CKD, chronic kidney disease;
ED, emergency department;
eGFR, estimated glomerular filtration rate.

Normal urinary oxalate level (all ages)⁸:
<0.50 mmol (<45 mg)/24h/1.73 m²
Normal plasma oxalate levels¹²: 1-5 µmol/L

REFERRAL TO NEPHROLOGIST UPON RECURRENT STONE EVENT⁹



PH1 DIAGNOSIS

CKD stage 2²

IMAGING FINDINGS:

Increased medullary echogenicity suggestive of nephrocalcinosis on ultrasound^{8,10}

24-HOUR URINE FINDINGS:

Oxalate¹: 0.97 mmol (85.3 mg)/24h/1.73 m²



GENETIC TEST FINDINGS:

Genetic test identified homozygous AGXT mutation, confirming PH1^{11*}

*It is recommended that siblings be screened.^{4,8}

MEDICAL MANAGEMENT
PLAN ESTABLISHED:

Water (6 L/day), potassium citrate (12 g/day), pyridoxine (B6) (410 mg/day)^{1,12†}

†For an adult patient who is 6 feet tall and weighs 180 pounds.

KIDNEY STATUS UNKNOWN



MANAGEMENT CHALLENGES:

- Patient misses scheduled office visits
- eGFR and urinary oxalate levels are not monitored

PATIENT
TIMELINE:

AGE
(years)

31

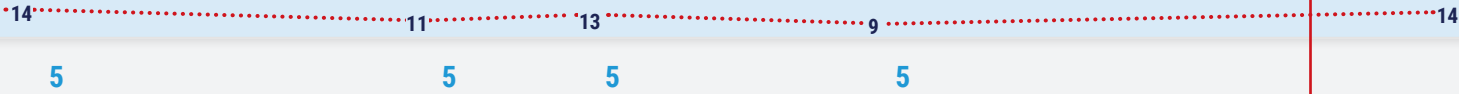
32

33

Hypothetical patient
profile: an adult patient
with PH1 experiences
rapid decline in kidney
function without
consistent management
and monitoring.

eGFR
(mL/min/1.73 m²)

CKD STAGE²



eGFR DECLINES DRAMATICALLY DUE TO DEHYDRATION¹³⁻¹⁵

PLASMA OXALATE MONITORED CLOSELY⁸

PLASMA OXALATE NOT CONTROLLED WITH DIALYSIS



**PATIENT DEVELOPS
COMPLICATED UTI**

Presents to ED with fever and chills¹⁶

DIALYSIS PLANNING:

- Plasma oxalate is measured due to concern of systemic oxalosis¹
- Predialysis plasma oxalate: 130 µmol/L¹⁷
- Goal: <30 µmol/L¹



**PATIENT IS LISTED FOR COMBINED LIVER/
KIDNEY TRANSPLANT**

Since patient is in end-stage kidney disease (ESKD), combined liver/kidney transplant is required¹



INTENSIVE DIALYSIS REGIMEN:

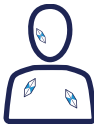
- Patient undergoes hemodialysis 6 times per week¹⁷
- Each session of hemodialysis is 4 hours long¹⁷

MONITORING REGIMEN:

- Routine surveillance for complications of systemic oxalosis and ESKD⁸:
- Long-bone radiographs
 - Echocardiogram
 - Thyroid function
 - Electrocardiogram
 - Hemoglobin

EFFECTS ON QUALITY OF LIFE:

- Work absences
- Emotional stress
- Financial burden
- Intensity of required medical care
- Missed family gatherings
- Inability to plan for future



SIGNS OF SYSTEMIC OXALOSIS

Patient reports painful skin nodules and bone pain¹

PLASMA OXALATE LEVEL:

Minimum plasma oxalate remains around 140 µmol/L despite intensive dialysis¹⁸



**PATIENT IS STILL AWAITING TRANSPLANT
A YEAR LATER¹⁹**

CKD, chronic kidney disease;
ED, emergency department;
eGFR, estimated glomerular filtration rate.

Normal urinary oxalate level (all ages)⁸:
<0.50 mmol (<45 mg)/24h/1.73 m²

Normal plasma oxalate levels¹²: 1-5 µmol/L



TAKEAWAYS:

Patients with PH1 may require continuous management, regardless of their age or symptomatology. Given the progressive nature of the disease, it is important that patients understand the benefit of continued management.^{4,8}

Monitoring for decline in kidney function may be needed. In some instances, kidney function can decline after a single incident of dehydration due to acute illness or intense physical activity. This can occur even in patients with previously stable kidney function.^{4,8,13,15,20}

Active management of PH1 may help slow the progression to ESKD. However, some patients may eventually progress to ESKD when a combined liver/kidney transplant is their only option. This procedure carries significant long-term morbidity and mortality risks.^{1,4,8,11}

PH1 is a very heterogeneous disease. Even siblings with the same genotype experience distinct and variable clinical manifestations.²¹

Consider genetic testing for your patients

when you suspect PH1^{9,11}

**ONE OPTION FOR TESTING IS
THE ALNYLAM ACT® PROGRAM:**
Third-party genetic testing and counseling
programs offered at no charge to patients.



The Alnylam Act® program was created to provide access to genetic testing and counseling to patients as a way to help people make more informed decisions about their health.

- While Alnylam provides financial support for this program, tests and services are performed by independent third parties
- Healthcare professionals must confirm that patients meet certain criteria to use the program
- Alnylam receives de-identified patient data from this program, but at no time does Alnylam receive patient identifiable information. Alnylam may use healthcare professional contact information for research purposes
- Both genetic testing and genetic counseling are available in the US and Canada
- Healthcare professionals or patients who use this program have no obligation to recommend, purchase, order, prescribe, promote, administer, use, or support any Alnylam product
- No patients, healthcare professionals, or payers, including government payers, are billed for this program

**FOR MORE INFORMATION,
VISIT [ALNYLAMACT.COM](https://alnylamact.com) AND [ABOUTPH1.COM](https://aboutph1.com) >**

References: 1. Cochat P, Hulton SA, Acquaviva C, et al. *Nephrol Dial Transplant*. 2012;27(5):1729-1736. doi:10.1093/ndt/gfs078 2. KDIGO 2012 Clinical Practice Guideline for the Evaluation and Management of Chronic Kidney Disease. *Kidney Int Suppl*. 2013;3(1). 3. Chu DI, Tasian GE, Copelovitch L. *Curr Treat Options Pediatr*. 2016;2(2):104-111. doi:10.1007/s40746-016-0046-8 4. Hoppe B, Beck BB, Milliner DS. *Kidney Int*. 2009;75(12):1264-1271. doi:10.1038/ki.2009.32 5. Coll DM, Varanelli MJ, Smith RC. *AJR Am J Roentgenol*. 2002;178(1):101-103. doi:10.2214/ajr.178.1.1780101 6. Frassetto L, Kohlstadt I. *Am Fam Physician*. 2011;84(11):1234-1242. 7. Pearle MS, Goldfarb DS, Assimos DG, et al; American Urological Association. *J Urol*. 2014;192(2):316-324. doi:10.1016/j.juro.2014.05.006 8. Milliner DS, Harris PC, Cogal AG, Lieske JC. In: Adam MP, Ardinger HH, Pagon RA, et al. *GeneReviews*®. University of Washington, Seattle; 1993-2021. 9. Hoppe B. *Nat Rev Nephrol*. 2012;8(8):467-475. doi:10.1038/nrneph.2012.113 10. Ferraro PM, D'Addessi A, Gambaro G. *Nephrol Dial Transplant*. 2013;28(4):811-820. doi:10.1093/ndt/gfs545 11. Cochat P, Rumsby G. *N Engl J Med*. 2013;369(7):649-658. doi:10.1056/NEJMr1301564 12. Bhasin B, Ürekli HM, Atta MG. *World J Nephrol*. 2015;4(2):235-244. doi:10.5527/wjn.v4.i2.235 13. Leumann E, Hoppe B. *J Am Soc Nephrol*. 2001;12(9):1986-1993. doi:10.1681/ASN.V1291986 14. El-Reshaid K, Al-Bader D, Madda JP. *Saudi J Kidney Dis Transpl*. 2016;27(3):606-609. 15. Harambat J, Fargue S, Bacchetta J, Acquaviva C, Cochat P. *Int J Nephrol*. 2011;2011:864580. doi:10.4061/2011/864580 16. Hooton TM, Gupta K. Accessed July 7, 2021. <https://www.uptodate.com/contents/acute-complicated-urinary-tract-infection-including-pyelonephritis-in-adults> 17. Yamauchi T, Quillard M, Takahashi S, Nguyen-Khoa M. *Nephrol Dial Transplant*. 2001;16(12):2407-2411. doi:10.1093/ndt/16.12.2407 18. Kuiper JJ. *West J Med*. 1996;164(1):42-53. 19. Cochat P. *Kidney Int*. 1999;55(6):2533-2547. doi:10.1046/j.1523-1755.1999.00477.x 20. Tintillier M, Pochet J-M, Cosyns J-P, Delgrange E, Donckier J. *Clin Nephrol*. 2004;62(2):155-157. doi:10.5414/cnp62155 21. Hoppe B. *Kidney Int*. 2010;77(5):383-385. doi:10.1038/ki.2009.471