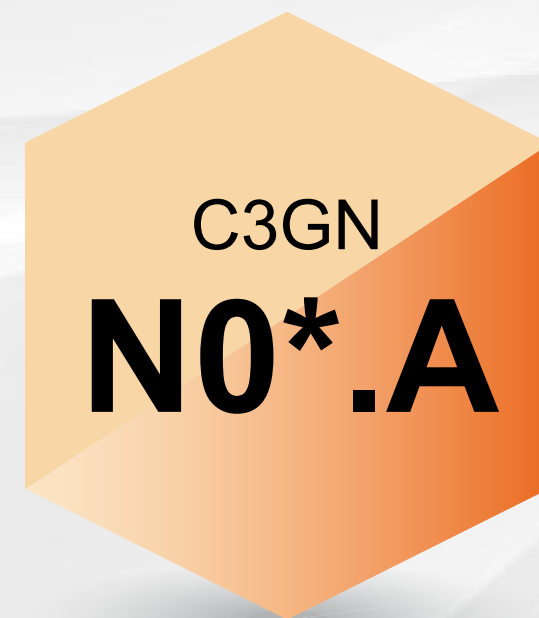


DID YOU KNOW THERE ARE NEW ICD-10 CODES FOR CMKDs?



The release of new ICD-10 codes aims to help address gaps in diagnosis, surveillance, and research of complement-mediated kidney diseases (CMKDs)¹⁻³



Between 2020 and 2023, new ICD-10 codes have been launched for C3G (including C3GN and DDD) and aHUS, allowing for greater specificity in coding the unique clinical features that can present with these CMKDs⁴

aHUS, atypical hemolytic uremic syndrome; C3G, complement 3 glomerulopathy; C3GN, complement 3 glomerulonephritis; CMKD, complement-mediated kidney disease; DDD, dense deposit disease; HUS, hemolytic uremic syndrome; ICD-10, International Statistical Classification of Diseases and Related Health Problems 10th Revision.

*This asterisk denotes a variable character; a complete list of codes can be found in the panel on the right.

1. Sun AZ et al. *Perm J.* 2020;24(2):19.126. doi:10.7812/TPP/19.126; 2. Dendooven A et al. *BMC Nephrol.* 2021;22(1):193. doi:10.1186/s12882-021-02365-3; 3. Kwon CS et al. *J Health Econ Outcomes Res.* 2021;8(2):36-45. doi:10.36469/001c.26129; 4. Centers for Medicare & Medicaid Services. Accessed August 30, 2023. <https://www.cms.gov/medicare/coding/icd10>

NEW ICD-10 CODES FOR CMKDs⁴

C3GN

- N00.A:** acute nephritic syndrome with C3GN
- N01.A:** rapidly progressive nephritic syndrome with C3GN
- N02.A:** recurrent and persistent hematuria with C3GN
- N03.A:** chronic nephritic syndrome with C3GN
- N04.A:** nephrotic syndrome with C3GN
- N05.A:** unspecified nephritic syndrome with C3GN
- N06.A:** isolated proteinuria with C3GN
- N07.A:** hereditary nephropathy, not elsewhere classified with C3GN

DDD

- N00.6:** acute nephritic syndrome with DDD
- N01.6:** rapidly progressive nephritic syndrome with DDD
- N02.6:** recurrent and persistent hematuria with DDD
- N03.6:** chronic nephritic syndrome with DDD
- N04.6:** nephrotic syndrome with DDD
- N05.6:** unspecified nephritic syndrome with DDD
- N06.6:** isolated proteinuria with DDD
- N07.6:** hereditary nephropathy, not elsewhere classified with DDD

aHUS

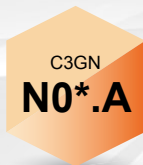
- D59.30:** HUS unspecified
- D59.31:** infection-associated HUS
- D59.32:** aHUS with an identified genetic cause
- D59.39:** aHUS (nongenetic)



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Learn more
about CMKDs on
The Renal Halls
of Science



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