

ANALYTE:
COL4A3

NAME:	collagen type IV alpha 3 chain
SYMBOL:	COL4A3
VERSION OF ORPHANET:	2023-06-22 14:14:43
SYNONYMS:	tumstatin
XREF(S):	Orphanet Ensembl Genatlas HGNC OMIM Reactome SwissProt
CREATED:	13 May 2019 - 01:01
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RELATED CONTENT

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- [Alport autosomal recessive and X-linked and hematuria \(3 genes\)](#)
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- [End-stage renal disease, ESRD \(gene panel\)](#)
- [Hearing loss \(deafness\), \(gene panel\)](#)
- [Inherited Kidney Diseases \(Gene Panel\)](#)
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- [Nephropathies, hereditary \(gene panel\)](#)
- [Nephrotic syndrome, Focal Segmental Glomerulosclerosis \(FSGS\) , Alport syndrome and podocytopathy \(gene panel\)](#)

Related Diseases

- [Autosomal dominant Alport syndrome](#)
- [Autosomal recessive Alport syndrome](#)
- [Familial idiopathic steroid-resistant nephrotic syndrome with focal segmental hyalinosis](#)
- [Genetic steroid-resistant nephrotic syndrome](#)
- [NON RARE IN EUROPE: Benign familial hematuria](#)

Related Gene Panels

- [Alport \(X-linked and recessive\) \(3 genes\) - IPG](#)
- [Alport \(X-linked and recessive\) \(3 genes\) - UZA](#)
- [Ciliopathy, polycystic kidney and liver diseases, ADTKD, nephronophtisis, Bardet-Biedl syndromes and kidney cancers \(146 genes\) - IPG](#)
- [End-stage renal disease \(106 genes\) - IPG](#)

- [Hearing loss \(deafness\) \(genepanel\) - UZA](#)
- [Hearing loss \(deafness\) syndromic \(59 genes\) - UZA](#)
- [Intellectual disability \(gene panel\)](#)
- [Nephropathies, hereditary \(219 genes\) - KUL](#)
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- [Nephrotic syndrome, FSGS, Alport syndrome \(76 genes\) - IPG](#)
- [Panel Nephro-ULG-V1](#)

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