

University of Parma Research Repository

Idiopathic calcium nephrolithiasis with pure calcium oxalate composition: clinical correlates of the calcium oxalate dihydrate/monohydrate (COD/COM) stone ratio

This is the peer reviewd version of the followng article:

Original

Idiopathic calcium nephrolithiasis with pure calcium oxalate composition: clinical correlates of the calcium oxalate dihydrate/monohydrate (COD/COM) stone ratio / Guerra, A.; Ticinesi, A.; Allegri, F.; Pinelli, S.; Aloe, R.; Meschi, T.. - In: UROLITHIASIS. - ISSN 2194-7228. - 48:3(2020), pp. 271-279. [10.1007/s00240-019-01156-8]

Availability: This version is available at: 11381/2884207 since: 2022-01-07T17:14:15Z

Publisher: Springer

Published DOI:10.1007/s00240-019-01156-8

Terms of use: openAccess

Anyone can freely access the full text of works made available as "Open Access". Works made available

Publisher copyright

(Article begins on next page)

Urolithiasis

Idiopathic calcium nephrolithiasis with pure calcium oxalate composition: clinical correlates of the calcium oxalate dihydrate/monohydrate (COD/COM) stone ratio --Manuscript Draft--

correlates of the calcium oxalate dihydrate/monohydrate (COD/COM) stone ratio Article Type: Original Paper Keywords: urolithiasis; Hypercalciuria; Hyperoxaluria; kidney stones; calcium oxalate Corresponding Author: Duniversita degli Studi di Parma Corresponding Author's Institution: Universita degli Studi di Parma Corresponding Author's Institution: Universita degli Studi di Parma Corresponding Author's Institution: Angela Guerra Andrea Ticinesi, M.D. France Allegri Silvana Pinelli Corresponding Author's Secondary Information: Per calcium oxalate is the most frequent type of idiopathic kidney stone composition. Fourier transform infrared spectroscopy (FT-IF) allows to detect the ratio of calcium oxalate dihydrate (COO) and on ange of 456 (322), agget 42 ±14) patients uffering from idiopathic kidney stone discoveration at study was to verify the association of clinical and laboratop parameters of kidney stone consestion or clinical and laboratop parameter scilicon toxalate dihydrate (COO) arado in a govor 456 (322),	Manuscript Number:	URES-D-19-00122R1
Keywords: urolithiasis; Hypercalciuria; Hyperoxaluria; kidney stones; calcium oxalate Corresponding Author: Andrea Ticinesi, M,D. Universita degli Studi di Parma Parma, ITALY Corresponding Author's Secondary Information: Universita degli Studi di Parma Corresponding Author's Institution: Universita degli Studi di Parma Corresponding Author's Secondary Information: Universita degli Studi di Parma Corresponding Author's Secondary Institution: Angela Guerra Andrea Ticinesi, M.D. First Author: Angela Guerra Andrea Ticinesi, M.D. First Authors: Angela Guerra Andrea Ticinesi, M.D. Franca Allegri Silvana Pinelli Rosalia Aloe Tiziana Meschi Tiziana Meschi Order of Authors Secondary Information: Pure calcium oxalate is the most frequent type of idiopathic kidney stone composition. Fourier transform infared spectroscopy (FT-IR) allows to detect the ratio of calcium disparate diftydrate divodrate (COD) and monoydrate (COD) ratio stoody ratio stoods, such and haboratory parameters of kidney stone disease with COD/COD ratio in agroup of 485 (322 M. age 46 ±14) patients suffering from idiopathic calcium nephrolitasis with pure calcium oxalate stones 2000. OS10. 75. 0.76-11, a significante of the parameter remains uncertain. The objective of kidney stone disease with COD/COD ratio in agroup of 485 (322 M. age 46 ±14) patients suffering from idiopathic calcium mep	Full Title:	
Corresponding Author: Andrea Ticinesi, M.D. Universita degli Studi di Parma Parma, ITALY Corresponding Author Secondary Information: Universita degli Studi di Parma Corresponding Author's Institution: Universita degli Studi di Parma Corresponding Author's Secondary Information: Universita degli Studi di Parma Corresponding Author's Secondary Information: Angela Guerra First Author: Angela Guerra First Author: Angela Guerra Andrea Ticinesi, M.D. Franca Allegri Silvana Pinelli Rosalia Aloe Tiziana Meschi Rosalia Aloe Tuziana Meschi Correspond regulation or calcium coxalate is the most frequent type of idiopathic kidney stone composition. Funding Information: Pure calcium oxalate is the most frequent type of idiopathic kidney stone composition. Funding Linformation: Pure calcium coxalate is the most frequent type of idiopathic kidney stone colcium coxalate dihydrain (COD) and monhydrate (COM) crystals in stones, but the clinical significance of this parameter remains uncertain. The objective of this observational study was to verify the association was observed or uncalcum coxales (SPR%). Each participatin turderwent a complete clinical examination, serum chemistry, Albour unive calcium noxalate supersaturation and a stones analyzed by FT-IR. Most (62%) of the stone shas a COD/COM ratio acode of uninary metabolic abnormatities. With increasing COD/COM	Article Type:	Original Paper
Universite degli Studi di Parma Parma, ITALY Corresponding Author's Secondary Information: Universite degli Studi di Parma Corresponding Author's Secondary Information: Universite degli Studi di Parma Corresponding Author's Secondary Institution: Angela Guerra First Author: Angela Guerra First Author Secondary Information: Angela Guerra Order of Authors: Angela Guerra Andrea Ticinesi, M.D. Franca Allegri Silvana Pinelli Rosalia Aloe Tiziana Meschi Tiziana Meschi Order of Authors Secondary Information: Pure calcium exalete is the most frequent type of idiopathic kidney stone composition. Fourier transform dimoratory parameter remains or diversion of calcium cosalate dihydraie (COD) card monohydrate (COM) crystals in stones, but the clinical significance of this parameter remains of clinical examination, serum chemistry, 24-hour urine disease with COD/COM ratio in a group of 465 (322 M. age 46 ± 14) patients suffering edicease with COD/COM ratio in a group of 465 (32 LM. age 46 ± 14) patients suffering roteicoticn for the determination of the profile of lithogenic risk, and had stones analyzed by FT-IK, Most (62%) of the stones has a COD/COM ratio intervals of 25, and the urine chereistry of the corresponding patients shores do 25, and the urine chereistry of the corresponding patients shore of urinary matabolic abnormalities, with increasing COD/COM ratio intervals (0.25, 0.36-0.50, 0.51-0.75, 0.76-1), a significant association was observed for the number of urinary realabolic abnormalities, with incr	Keywords:	urolithiasis; Hypercalciuria; Hyperoxaluria; kidney stones; calcium oxalate
Information: Information: Information: Information: Universita degli Studi di Parma Corresponding Author's Secondary Corresponding Author's Secondary First Author: Angela Guerra First Author: Angela Guerra First Author Secondary Information: Corder of Authors: Angela Guerra Andrea Ticinesi, M.D. Franca Allegri Silvana Pinelli Rosalia Aloe Tiziana Meschi Corder of Authors Secondary Information: Punding Information: Punding Information: Punding Author Secondary Information: Abstract: Punding Information: Punding Information: Punding Information: Punding Information: Abstract: Punding Information: Punding Information Punding Informatinform Punding Information Punding Information Punding In	Corresponding Author:	Universita degli Studi di Parma
Corresponding Author's Secondary Institution: Angela Guerra First Author: Angela Guerra First Author Secondary Information: Angela Guerra Order of Authors: Angela Guerra Andrea Ticinesi, M.D. Franca Allegri Silvana Pinelli Rosalia Alce Tiziana Meschi Tiziana Meschi Order of Authors Secondary Information: Pure calcium oxalate is the most frequent type of idiopathic kidney stone composition. Fourier transform infrared spectroscopy (FT-IR) allows to detect the ratio of calcium oxalate dihydrate (COD) and monohydrate (COM) crystals in stones, but the clinical significance of this parameter remains uncertain. The objective of this observational significance of the stones has a COD/COM ratio alles atomes of kidney stone composition. Formicipant underwent a complete clinical and laboratory parameters of kidney stone disease with COD/COM ratio in a group of 465 (322 M, age 46 ± 14) patients suffering from idopathic calcium mephoribithiasis with pure calcium colaste stones (597%). Each participant underwent a complete clinical avamination, serum chemistry, 24-hour urine collection for the determination of the profile of lithogenic risk, and had stones analyzed by FT-IR. Nots (62%) of the stones has a COD/COM ratio intervals (0-25, 0.26-0.30, 0.51-0.75, 0.75-0.75, 0.75-0.17, a significant association was observed for the number of urological procedures, serum calcium, 24-h urinary calcium excretion, prevalence of hyperaclicuita and relative exceted with COD/COM ratio intervals (0-0.001) and urine pH (standardized p=0.103, p=0.013). In pure calcium	Corresponding Author Secondary Information:	
Institution: Angela Guerra First Author: Angela Guerra First Author Secondary Information: Order of Authors: Angela Guerra Andrea Ticinesi, M.D. Franca Allegri Silvana Pinelli Rosalia Aloe Tiziana Meschi Order of Authors Secondary Information: Funding Information: Funding Information: Funding Information: Abstract: Vertical Context of the secondary of the second	Corresponding Author's Institution:	Universita degli Studi di Parma
First Author Secondary Information: Angela Guerra Andrea Ticinesi, M.D. Franca Allegri Silvana Pinelli Rosalia Aloe Tiziana Meschi Tiziana Meschi Order of Authors Secondary Information: Pure calcium oxalate is the most frequent type of idiopathic kidney stone composition. Funding Information: Pure calcium oxalate is the most frequent type of idiopathic kidney stone composition. Fourier transform infrared spectroscopy (FT-IR) allows to detect the ratio of calcium oxalate dihydrate (COD) and monohydrate (COM) crystals in stones, but the clinical significance of this parameter remains uncertain. The objective of this observational study was to verify the association of clinical and laboratory parameters of kidney stone collection for the determination of the profile of lithogenic risk, and had stones analyzed by FT-IR. Most (62%) of the stones has a COD/COM ratio sol.25, and the urine collection for the determination of the profile of lithogenic risk, and had stones analyzed by FT-IR. Most (62%) of the stones has a COD/COM ratio sol.25, 0.26-0.50, 0.21-0.75, 0.76-1), a significant association was observed for the number of urological procedures, serum calcium oxalate supersaturation, and an egative trend was detected for the age of the first stone episode (all p-values <0.05). A linear regression model showed that the only parameters significantly sascciated with COD/COM ratio was observed for the number of urological procedures, serum calcium oxalate supersaturation, and a negative trend was detected for the age of the first stone episode (all p-values <0.05). A linear regression wordel showed that the only parameters significanthy sascciated with COD/COM ratio were 24-h urinary calcium excreti	Corresponding Author's Secondary Institution:	
Order of Authors: Angela Guerra Andrea Ticinesi, M.D. Franca Allegri Silvana Pinelli Rosalia Aloe Tiziana Meschi Order of Authors Secondary Information: Funding Information: Abstract: Pure calcium oxalate is the most frequent type of idiopathic kidney stone composition. Fourier transform infrared spectroscopy (FT-IR) allows to detect the ratio of calcium oxalate dihydrate (COD) and monohydrate (COM) crystals in stones, but the clinical significance of this parameter remains uncertain. The objective of this observational study was to verify the association of clinical and laboratory parameters of kidney stone disease with COD/COM ratio in a group of 465 (322 M, age 46 ± 14) patients suffering from idiopathic calcium nephrolithiasis with pure calcium oxalate stones (297%). Each participant underwent a complete clinical examination, serum chemistry. 24-hour urine collection for the determination of the profile of ilitogenic risk, and had stones analyzed by FT-IR. Most (62%) of the stones has a COD/COM ratio 50.25, and the urine chemistry of the corresponding patients showed a low prevalence of trypercalciuria and relative calcium oxalate supersaturation, and a negative tred was detected for the age of the first stone episode (all p-values <0.05). A linear regression model showed that the only parameters significantly associated with COD/COM ratio were 24-h urinary calcium excretion (standardized β=0.484, p<0.001) and urine pH (standardized β=0.103, p=0.013). In pure calcium oxalate idiopathic ctores, COD/COM ratio may reflect the presence of urinary metabolic risk factors, and represent a guide for the prescription of urinary analyses	First Author:	Angela Guerra
Andrea Ticinesi, M.D. Franca Allegri Silvana Pinelli Rosalia Aloe Tiziana Meschi Order of Authors Secondary Information: Funding Information: Abstract: Pure calcium oxalate is the most frequent type of idiopathic kidney stone composition. Fourier transform infrared spectroscopy (FT-IR) allows to detect the ratio of calcium oxalate dihydrate (COD) and monchydrate (COM) crystals in stones, but the clinical significance of this parameter remains uncertain. The objective of this observational study was to verify the association of clinical and laboratory parameters of kidney stone disease with COD/COM ratio in a group of 465 (322 M, age 46 ± 14) patients suffering from idiopathic calcium nephrolithiasis with pure calcium oxalate stones (297%). Each participant underwent a complete clinical examination, serum chemistry, 24-hour urine collection for the determination of the profile of lithogenic risk, and had stones analyzed by FT-IR. Most (62%) of the stones has a COD/COM ratio is 0.0.25, 0.26-0.50, 0.51-0.75, 0.76-1), a significant association was obverved for the number of urological procedures, serum calcium, 24-huirary calcium extention, mervelance of urinary metabolic abnormalities. With increasing COD/COM ratio intervals (0-0.25, 0.26-0.50, 0.51-0.75, 0.76-1), a significant association was observed for the number of urological procedures, serum calcium, 24-huirary calcium extention, and a negative trend was detected for the age of the first stone episode (all p-values <0.05). A linear regression model showed that the only parameters significantly associated with COD/COM ratio were 24-h urinary calcium excretion (standardized β=0.464, p<0.001) and urine pH (standardized β=0.103, p=0.013). In pure calcium oxalate idiopathic stones, COD/COM ratio may reflect the presence of urinary metabolic risk factors, and represent a guide for the prescription of urinary analyses.	First Author Secondary Information:	
Franca Allegri Silvana Pinelli Rosalia Aloe Tiziana Meschi Order of Authors Secondary Information: Funding Information: Abstract: Pure calcium oxalate is the most frequent type of idiopathic kidney stone composition. Fourier transform infrared spectroscopy (FT-IR) allows to detect the ratio of calcium oxalate dihydrate (COD) and monohydrate (COM) crystals in stones, but the clinical significance of this parameter remains uncertain. The objective of this observational study was to verify the association of clinical and laboratory parameters of kidney stone disease with COD/COM ratio in a group of 465 (322 M, age 46 ± 14) patients suffering from idiopathic calcium nephrolithiasis with pure calcium oxalate stones (±97%). Each participant underwent a complete clinical examination, serum chemistry, 24-hour urine collection for the determination of the profile of lithogenic risk, and had stones analyzed by FT-IR. Most (62%) of the stones has a COD/COM ratio s0.25, and the urine collection for the determination was observed for the number of urological procedures, serum calcium, 24-h urinary calcium excretion, prevalence of hypercalciuria and relative calcium oxalate supersaturation, and a negative trend was detected for the age of the first stone episode (all p-values 0.05). A linear regression model showed that the only parameters significantly associated with COD/COM ratio were 24-h urinary calcium excretion (standardized §=0.464, p<0.001) and urine pH (standardized §=0.103, p=0.013). In pure calcium coxalate idopathic stones, COD/COM ratio may reflect the presence of urinary metabolic risk factors, and represent a guide for the prescription of urinary analyses.	Order of Authors:	Angela Guerra
Silvana Pinelli Rosalia Aloe Tiziana Meschi Order of Authors Secondary Information: Funding Information: Abstract: Pure calcium oxalate is the most frequent type of idiopathic kidney stone composition. Fourier transform infrared spectroscopy (FT-IR) allows to detect the ratio of calcium oxalate dihydrate (COD) and monohydrate (COM) crystals in stones, but the clinical significance of this parameter remains uncertain. The objective of this observational study was to verify the association of clinical and laboratory parameters of kidney stone disease with COD/COM ratio in a group of 465 (322 M, age 46 ± 14) patients suffering from idiopathic calcium nephrolithiasis with pure calcium oxalate stones (297%). Each participant underwent a complete clinical examination, serum chemistry, 24-hour urine collection for the determination of the profile of lithogenic risk, and had stones analyzed by FT-IR. Most (62%) of the stones has a COD/COM ratio 50.25, and the urine chemistry of the corresponding patients showed a low prevalence of urinary metabolic abnormalities. With increasing COD/COM ratio intervals (0-0.50, 0.51-0.75, 0.76-1), a significant association was observed for the number of urological procedures, serum calcium, 24-h urinary calcium excretion, prevalence of hypercalciuria and relative calcium oxalate supersaturation, and a negative trend was detected for the age of the first stone episod (all p-0.464, p-0.01), and urine pH (standardized \$=0.103, p=0.013). In pure calcium exate idiopathic stones, COD/COM ratio may reflect the presence of urinary metabolic risk factors, and represent a guide for the prescription of urinary analyses.		Andrea Ticinesi, M.D.
Rosalia Aloe Tiziana Meschi Order of Authors Secondary Information: Funding Information: Abstract: Pure calcium oxalate is the most frequent type of idiopathic kidney stone composition. Fourier transform infrared spectroscopy (FT-IR) allows to detect the ratio of calcium oxalate dihydrate (COD) and monohydrate (COM) crystals in stones, but the clinical significance of this parameter remains uncertain. The objective of this observational study was to verify the association of clinical and laboratory parameters of kidney stone disease with COD/COM ratio in a group of 465 (322 M, age 46 ± 14) patients suffering from idiopathic calcium nephrolithiasis with pure calcium oxalate stones (≥97%). Each participant underwent a complete clinical examination, serum chemistry, 24-hour urine collection for the determination of the profile of lithogenic risk, and had stones analyzed by FT-IR. Most (62%) of the stones has a COD/COM ratio 50.25, 0.26-0.50, 0.51-0.75, 0.76-1), a significant association was observed for the number of urological procedures, serum calcium, 24-h urinary calcium excretion, pervalence of hypercalciuria and relative calcium oxalate supersaturation, and a negative trend was detected for the age of the first stone episode (all p-values <0.05). A linear regression model showed that the only parameters significantly associated with COD/COM ratio were 24-h urinary calcium excretion (standardized β=0.464, p<0.001) and urine pH (standardized β=0.103, p=0.013). In pure calcium oxalate idiopathic stones, COD/COM ratio may reflect the presence of urinary metabolic risk factors, and represent a guide for the prescription of urinary analyses.		Franca Allegri
Tiziana Meschi Tiziana Meschi Funding Information: Funding Information: Abstract: Pure calcium oxalate is the most frequent type of idiopathic kidney stone composition. Fourier transform infrared spectroscopy (FT-IR) allows to detect the ratio of calcium oxalate dihydrate (COD) and monohydrate (COM) crystals in stones, but the clinical significance of this parameter remains uncertain. The objective of this observational study was to verify the association of clinical and laboratory parameters of kidney stone disease with COD/COM ratio in a group of 465 (322 M, age 46 ± 14) patients suffering from idiopathic calcium nephrolithiasis with pure calcium oxalate stones (≥97%). Each participant underwent a complete clinical examination, serum chemistry, 24-hour urine collection for the determination of the profile of lithogenic risk, and had stones analyzed by FT-IR. Most (62%) of the stones has a COD/COM ratio 50.25, and the urine chemistry of the corresponding patients showed a low prevalence of urinary metabolic abnormalities. With increasing COD/COM ratio intervals (0-0.25, 0.26-0.50, 0.51-0.75, 0.76-1), a significant association was observed for the number of urological procedures, serum calcium, 24-h urinary calcium excretion, prevalence of hypercalciuria and relative calcium oxalate supersaturation, and a negative trend was detected for the age of the first stone episode (all p-values <0.05). A linear regression model showed that the only parameters significantly associated with COD/COM ratio were 24-h urinary calcium excretion (standardized β=0.464, p<0.001) and urine pH (standardized β=0.103, p=0.013). In pure calcium oxalate idiopathic stones, COD/COM ratio may reflect the presence of urinary metabolic risk factors, and represent a guide for the prescription of urinary analyses.		Silvana Pinelli
Order of Authors Secondary Information: Funding Information: Abstract: Pure calcium oxalate is the most frequent type of idiopathic kidney stone composition. Fourier transform infrared spectroscopy (FT-IR) allows to detect the ratio of calcium oxalate dihydrate (COD) and monohydrate (COM) crystals in stones, but the clinical significance of this parameter remains uncertain. The objective of this observational study was to verify the association of clinical and laboratory parameters of kidney stone disease with COD/COM ratio in a group of 465 (322 M, age 46 ± 14) patients suffering from idiopathic calcium nephrolithasis with pure calcium oxalate stones (≥97%). Each participant underwent a complete clinical examination, serum chemistry, 24-hour urine collection for the determination of the profile of lithogenic risk, and had stones analyzed by FT-IR. Most (62%) of the stones has a COD/COM ratio ≤0.25, and the urine chemistry of the corresponding patients showed a low prevalence of urinary metabolic abnormalities. With increasing COD/COM ratio intervals (0-0.25, 0.26-0.50, 0.51-0.75, 0.76-1), a significant association was observed for the number of urological procedures, serum calcium, 24-h urinary calcium excretion, prevalence of hypercalciuria and relative calcium oxalate supersaturation, and a negative trend was detected for the age of the first stone episode (all p-values <0.05). A linear regression model showed that the only parameters significantly associated with COD/COM ratio were 24-h urinary calcium excretion (standardized β=0.464, p<0.001) and urine pH (standardized β=0.013), p=0.013). In pure calcium oxalate idiopathic stones, COD/COM ratio may reflect the presence of urinary metabolic risk factors, and represent a guide for the prescription of urinary analyses.		Rosalia Aloe
Funding Information:Abstract:Pure calcium oxalate is the most frequent type of idiopathic kidney stone composition. Fourier transform infrared spectroscopy (FT-IR) allows to detect the ratio of calcium oxalate dihydrate (COD) and monohydrate (COM) crystals in stones, but the clinical significance of this parameter remains uncertain. The objective of this observational study was to verify the association of clinical and laboratory parameters of kidney stone disease with COD/COM ratio in a group of 465 (322 M, age 46 \pm 14) patients suffering from idiopathic calcium nephrolithiasis with pure calcium oxalate stones (≥97%). Each participant underwent a complete clinical examination, serum chemistry, 24-hour urine collection for the determination of the profile of lithogenic risk, and had stones analyzed by FT-IR. Most (62%) of the stones has a COD/COM ratio ≤ 0.25 , 0.26-0.50, 0.51-0.75, 0.76-1), a significant association was observed for the number of urological procedures, serum calcium, 24-h urinary calcium excretion, prevalence of hypercalciuria and relative calcium oxalate supersaturation, and a negative trend was detected for the age of the first stone episode (all p-values <0.05). A linear regression model showed that the only parameters significantly associated with COD/COM ratio were 24-h urinary calcium excretion (standardized $\beta = 0.464$, p<0.001) and urine pH (standardized $\beta = 0.103$, p=0.013). In pure calcium oxalate idiopathic stones, COD/COM ratio may reflect the presence of urinary metabolic risk factors, and represent a guide for the prescription of urinary analyses.		Tiziana Meschi
Abstract: Pure calcium oxalate is the most frequent type of idiopathic kidney stone composition. Fourier transform infrared spectroscopy (FT-IR) allows to detect the ratio of calcium oxalate dihydrate (COD) and monohydrate (COM) crystals in stones, but the clinical significance of this parameter remains uncertain. The objective of this observational study was to verify the association of clinical and laboratory parameters of kidney stone disease with COD/COM ratio in a group of 465 (322 M, age 46 ± 14) patients suffering from idiopathic calcium nephrolithiasis with pure calcium oxalate stones (\geq 97%). Each participant underwent a complete clinical examination, serum chemistry, 24-hour urine collection for the determination of the profile of lithogenic risk, and had stones analyzed by FT-IR. Most (62%) of the stones has a COD/COM ratio \leq 0.25, and the urine chemistry of the corresponding patients showed a low prevalence of urinary metabolic abnormalities. With increasing COD/COM ratio intervals (0-0.25, 0.26-0.50, 0.51-0.75, 0.76-1), a significant association was observed for the number of urological procedures, serum calcium, 24-h urinary calcium excretion, prevalence of hypercalciuria and relative calcium oxalate supersaturation, and a negative trend was detected for the age of the first stone episode (all p-values <0.05). A linear regression model showed that the only parameters significantly associated with COD/COM ratio were 24-h urinary calcium excretion (standardized β =0.464, p<0.001) and urine pH (standardized β =0.103, p=0.013). In pure calcium oxalate idiopathic stones, COD/COM ratio may reflect the presence of urinary metabolic risk factors, and represent a guide for the prescription of urinary analyses.	Order of Authors Secondary Information:	
Fourier transform infrared spectroscopy (FT-IR) allows to detect the ratio of calcium oxalate dihydrate (COD) and monohydrate (COM) crystals in stones, but the clinical significance of this parameter remains uncertain. The objective of this observational study was to verify the association of clinical and laboratory parameters of kidney stone disease with COD/COM ratio in a group of 465 (322 M, age 46 ± 14) patients suffering from idiopathic calcium nephrolithiasis with pure calcium oxalate stones (≥97%). Each participant underwent a complete clinical examination, serum chemistry, 24-hour urine collection for the determination of the profile of lithogenic risk, and had stones analyzed by FT-IR. Most (62%) of the stones has a COD/COM ratio ≤0.25, and the urine chemistry of the corresponding patients showed a low prevalence of urinary metabolic abnormalities. With increasing COD/COM ratio intervals (0-0.25, 0.26-0.50, 0.51-0.75, 0.76-1), a significant association was observed for the number of urological procedures, serum calcium, 24-h urinary calcium excretion, prevalence of hypercalciuria and relative calcium oxalate supersaturation, and a negative trend was detected for the age of the first stone episode (all p-values <0.05). A linear regression model showed that the only parameters significantly associated with COD/COM ratio were 24-h urinary calcium excretion (standardized β =0.464, p<0.001) and urine pH (standardized β =0.103, p=0.013). In pure calcium oxalate idiopathic stones, COD/COM ratio may reflect the presence of urinary metabolic risk factors, and represent a guide for the prescription of urinary analyses.	Funding Information:	
	Abstract:	Fourier transform infrared spectroscopy (FT-IR) allows to detect the ratio of calcium oxalate dihydrate (COD) and monohydrate (COM) crystals in stones, but the clinical significance of this parameter remains uncertain. The objective of this observational study was to verify the association of clinical and laboratory parameters of kidney stone disease with COD/COM ratio in a group of 465 (322 M, age 46 ± 14) patients suffering from idiopathic calcium nephrolithiasis with pure calcium oxalate stones (≥97%). Each participant underwent a complete clinical examination, serum chemistry, 24-hour urine collection for the determination of the profile of lithogenic risk, and had stones analyzed by FT-IR. Most (62%) of the stones has a COD/COM ratio 0.25 , $0.26-0.50$, $0.51-0.75$, $0.76-1$), a significant association was observed for the number of urological procedures, serum calcium, 24-h urinary calcium excretion, prevalence of hypercalciuria and relative calcium oxalate supersaturation, and a negative trend was detected for the age of the first stone episode (all p-values <0.05). A linear regression model showed that the only parameters significantly associated with COD/COM ratio were 24-h urinary calcium excretion (standardized β =0.464, p<0.001) and urine pH (standardized β =0.103, p=0.013). In pure calcium oxalate idiopathic stones, COD/COM ratio may reflect the presence of urinary metabolic risk factors, and represent a guide
	Response to Reviewers:	RESPONSES TO REVIEWER #1

Powered by Editorial Manager® and ProduXion Manager® from Aries Systems Corporation

"The authors of this submission have performed an investigation in order to detect a possible association between calcium oxalate stone (COM & COD) and clinical characteristics, including 24-hour urinary parameters of lithogenic risk, of a large group of patients with idiopathic calcium nephrolithiasis. Chemical composition of kidney stones and thus the COD/COM ratios were determined by FTIR spectroscopy. The data were well analyzed and thus the conclusion is clear. Here are some suggested revisions:

- In the introduction : it is worth to underline that the morphology of crystallites indicates quite precisely the pathology. For example, five morphological subtypes Ia, Ib, Ic, Id and Ie exists for COM kidney stones (Daudon, M., Bader, C. A. & Jungers, P. (1993). Scan. Microsc. 7, 1081-1106 as well as M. Daudon, D. Bazin, G. André, P. Jungers, A. Cousson, P. Chevallier, E. Véron, G. Matzen, Examination of whewellite kidney stones by scanning electron microscopy and powder neutron diffraction techniques, J. Appl. Cryst. (2009). 42, 109-115)."

R: We thank the reviewer for the positive comment and for the important suggestions. In the novel version of the manuscript, we have included a better focus on the relationship between calcium oxalate crystal morphology and stone etiopathogenesis, highlighting the mentioned classification of COM kidney stones (see page 2 lines 7-12). The two mentioned papers are now referenced in the manuscript (number 3 and 9).

"Also there is a study performed on urine which has to be included in the references : M. Daudon, E. Letavernier, V. Frochot, J.-Ph. Haymann, D. Bazin, P. Jungers, Respective influence of calcium and oxalate urine concentration on the formation of calcium oxalate monohydrate or dihydrate crystals, C. R. Chimie 19 (2016) 1504-1513."

R: We apologize for having missed this important study on the clinical correlates of COD and COM stone composition. In the novel version of the manuscript, this study is referenced (number 24) and discussed (see page 8 lines 12-15 and page 9 lines 8-11).

RESPONSES TO REVIEWER #2

"The paper of Guerra et al is an interesting work analysing clinical characteristics of patients forming calcium oxalate monohydrate stones or calcium oxalate dihydrate stones. Although relevant, I seem that the paper needs to be reconsidered in some methodological points."

R: We thank the reviewer for the positive comments and constructive criticism. All the methodological concerns have been carefully considered and the manuscript has been modified accordingly.

"1. Control group characteristics and criteria for their enrolment could be more extensively detailed at page 3, line 29-33."

R: In the novel version of the manuscript, we have introduced more information on the characteristics of controls and criteria for their inclusion in the study (see page 3 lines 15-22). The mandatory criteria for control selection were the absence of a personal history of kidney stones/renal colic and the absence of retained stones at abdominal ultrasound performed immediately before urine collection.

"2. Please specify conditions for 24-hour urine storage in order to measure their pH."

R: In the novel version of the manuscript, more details on the methods of urine collection, preservation and analysis have been added (see page 4 lines 4-11). This methodology is well standardized at our Stone Clinic, and has remained substantially the same over the last 30 years.

"3. References 9-10, reported at page 4, to support limits of hypercalciuria and hyperoxaluria do not appear appropriate to me."

R: The references have been deleted from the novel version of the manuscript. Unfortunately, there are no widely accepted definitions of hypercalciuria and hyperoxaluria, and "convenience" definitions were adopted in this investigations. Hypercalciuria is in fact generally defined as urinary calcium excretion >200 mg/24 h (see Pak CY et al. Kidney Int 2011), but some investigations demonstrated that the risk of stone formation increases continuously with increasing urinary calcium excretion values, so that no threshold for hypercalciuria can be certainly identified (Curhan GC et al. Kidney Int 2001). Moreover, different "convenience" definitions of hypercalciuria were used in nephrolithiasis studies (Escribano J et al. Cochrane Database Syst Rev 2014), and the definition establishing 4 mg/kg/day of urinary calcium excretion as threshold for hypercalciuria is commonplace in clinical practice (see Leslie SW et al. StatPearls 2019). We underline that a rigorous definition of hypercalciura and hyperoxaluria was beyond the aims of the present investigation.

"4. The Authors declared to analyse characteristics of patients grouped in quartiles of COD/COM ratio: according to usual statistical criteria, each quartile has to include 25% of patients (n=116-117 in the present study); on the contrary the present study divided patients in four groups defined according to the value of the COD/COM ratio (0.0.25, 0.26-0.5, 0.51-0.75, 0.76-1), a method which does not identify quartiles. Description of these analyses has to be revised in the abstract, discussion, statistical method and result sections and tables."

R: We agree with the reviewer. In the previous version of the manuscript, the definition of COD/COM ratio quartiles was not appropriated. Each COD/COM ratio quartile should in fact have included 25% of the study population, and thus interquartile limits should have been different. However, we believe that a categorization of COD/COM ratio quartile (0-0.25; 0.26-0.50; 0.51-0.75; 0.76-1) is much more practical for clinical interpretation of the parameter. For this reason, we preferred to maintain this method of categorization, changing the incorrect definition of "quartiles". The manuscript has been revised in accordance with this choice. Each category has been labeled as "COD/COM interval".

"5. In the statistical analysis section it is reported that data were expressed as mean ± standard deviation or mean and 95% confidence intervals or median and interquartile range. However, mean±SD was used to express the large part of the variables in the manuscript, whereas median and IQR was used to describe COD/COM ratio (at page 6) and duration of the disease (in table 1) as non-parametric variables. Mean and 95% confidence intervals was used only at page 9 in the manuscript discussion and not in result section. Therefore, methods to express variables need to be revised at page 5."

R: We agree with the reviewer and apologize for the lack of clarity of the previous version of the "statistical analysis" paragraph. In the novel version of the manuscript, the section on how data were expressed and handled has been modified as follows: "Continuous variables were expressed as mean ± standard deviation or, for non-normally distributed variables, median and interquartile range (IQR). Dichotomous variables were expressed as percentages" (page 5 lines 10-11).

"6. In table 1, ESWL number and stone rate could be considered as non-parametric variables and reported as median and interquartile range."

R: We agree with the reviewer. Table 1 has been modified accordingly. The comparison of ESWL number and stone rate across different groups of stone formers, stratified by COD/COM ratio intervals, was then performed using Kruskal-Wallis test. For this reason, p-values changed.

"7. Significance of findings was reported with 3 p values in tables 2 and 3 but 2 p values in tables 1 and 4. I seem that the Authors considered unadjusted p values raising from ANOVA (the so-called p for trend), p values adjusted for confounding variables (p for trend from ANCOVA) and p values obtained with multiple comparisons between groups (Bonferroni test). It is not immediately clear why 3 p values were not reported in tables 1. The use of these p values could be more extensively explained in statistical methods.

8. In table 4 the Authors compared values of variables estimated using ANCOVA. If this is correct, this method has to be detailed in statistical analysis section at page 5 and specified in the text at page 6 and table 4."

R: The whole statistical analysis paragraph has been revised in order to improve comprehension by readers (see page 5 lines 14-26). In Tables 1, 2 and 3, normally

distributed clinical and laboratory parameters were compared among groups of patients, stratified by intervals of COD/COM ratio, using one-way analysis of variance (ANOVA) for crude comparisons, and analysis of covariance (ANCOVA) for comparisons adjusted for covariates (age, sex, duration of disease, BMI). The Bonferroni test for multiple comparisons was applied if adjusted p values were <0.05. Non-normally distributed variables were compared among groups of patients by Kruskal-Wallis test.

In Table 4, a comparison between stone formers, stratified by COD/COM ratio intervals, and stone-free controls was made. Unfortunately, stone formers and controls showed mild, but statistically significant, differences in age, sex distribution, body mass index, and urinary volume. For this reason, in Table 4 urinary parameters were handled as mean ± standard deviation adjusted for age, sex, BMI, and, for urinary supersaturation indexes, also for urinary volume. Comparisons were made using ANCOVA. Bonferroni test was again applied if adjusted p values were <0.05. Tables 1, 2, 3 and 4 have been carefully revised to improve comprehension on what kind of statistical analyses were applied in each case. In Table 1, crude p values obtained with ANOVA (dichotomous variables or continuous variables with normal distribution) or Kruskal-Wallis test (continuous variables with non-normal distribution) were reported for all lines. Adjusted p values, obtained with ANCOVA, were reported only for those variables where adjustment for covariates is meaningful in clinical terms. Finally, p for trend values obtained with ANOVA (linear trends) were reported only if <0.05 (statistically significant).

RESPONSE TO EDITOR-IN-CHIEF'S ADDITIONAL COMMENT

"Please note that this journal's policy is only to allow a maximum of five to six authors unless the article is the product of a multi-centre research study. Your article currently has 7 authors, which is not permitted. If you decide to submit a revised version of your manuscript, please reduce the number of authors to six or fewer. Also, if you wish to submit any articles to Urolithiasis in future, I would be grateful if you could please adhere to this policy (see Instructions for Authors on journal website)."

R: We apologize that our original submission was not adherent to this editorial policy. After discussion among all authors, we decided to remove from the author team Dr Antonio Nouvenne, who gave advice for study conception and assistance in manuscript drafting, but was not directly involved in data collection, analysis and interpretation. The manuscript has now six authors. Dr Nouvenne has been mentioned in the acknowledgement section.

Idiopathic calcium nephrolithiasis with pure calcium oxalate composition: clinical correlates of the calcium oxalate dihydrate/monohydrate (COD/COM) stone ratio

3

Angela Guerra^{1,2}, Andrea Ticinesi^{1,2*}, Franca Allegri^{1,2}, Silvana Pinelli², Rosalia Aloe³, Tiziana Meschi^{1,2}

¹Geriatric Rehabilitation Department, University-Hospital of Parma, Via Antonio Gramsci 14, 43126 Parma,

Italy

²Department of Medicine and Surgery, University of Parma, Via Antonio Gramsci 14, 43126 Parma, Italy

³Diagnostic Department, U.O. Diagnostica Ematochimica e S.S.D. Biochimica ad Elevata Automazione,

University-Hospital of Parma, Via Antonio Gramsci 14, 43126 Parma, Italy

Corresponding author:

*Andrea Ticinesi, MD, PhD, Geriatric-Rehabilitation Department, Parma University-Hospital (Azienda Ospedaliero-Universitaria di Parma), via Antonio Gramsci 14, 43126 Parma, Italy.

Work Telephone number: 0039-0521703871; Mobile: 00393471845191; Fax: 0039-0521702383 E-mail address: <u>aticinesi@ao.pr.it</u>

Word count: 3110

Abstract: 248; Tables: 5; Figures: 1; References: 40 Running Title: Mono- vs dihydrate calcium oxalate stones

Abstract

Pure calcium oxalate is the most frequent type of idiopathic kidney stone composition. Fourier transform infrared spectroscopy (FT-IR) allows to detect the ratio of calcium oxalate dihydrate (COD) and monohydrate (COM) crystals in stones, but the clinical significance of this parameter remains uncertain. The objective of this observational study was to verify the association of clinical and laboratory parameters of kidney stone disease with COD/COM ratio in a group of 465 (322 M, age 46 ± 14) patients suffering from idiopathic calcium nephrolithiasis with pure calcium oxalate stones ($\geq 97\%$). Each participant underwent a complete clinical examination, serum chemistry, 24-hour urine collection for the determination of the profile of lithogenic risk, and had stones analyzed by FT-IR. Most (62%) of the stones has a COD/COM ratio <0.25, and the urine chemistry of the corresponding patients showed a low prevalence of urinary metabolic abnormalities. With increasing COD/COM ratio intervals (0-0.25, 0.26-0.50, 0.51-0.75, 0.76-1), a significant association was observed for the number of urological procedures, serum calcium, 24-h urinary calcium excretion, prevalence of hypercalciuria and relative calcium oxalate supersaturation, and a negative trend was detected for the age of the first stone episode (all p-values < 0.05). A linear regression model showed that the only parameters significantly associated with COD/COM ratio were 24-h urinary calcium excretion (standardized β =0.464, p<0.001) and urine pH (standardized $\beta=0.103$, p=0.013). In pure calcium oxalate idiopathic stones, COD/COM ratio may reflect the presence of urinary metabolic risk factors, and represent a guide for the prescription of urinary analyses.

Key words: urolithiasis; hypercalciuria; hyperoxaluria; kidney stones; calcium oxalate.

Introduction

Under infrared spectroscopy, calcium oxalate crystals of calcium stones may appear in two distinct molecular forms: whewellite, that is, calcium oxalate monohydrate (COM), and weddellite, that is, calcium oxalate dihydrate (COD). These forms are associated with different etiology of stones [1, 2] and are also associated with different surface morphology of calculi [3].

COM depends on urinary excretion of oxalate and is typical of conditions of hyperoxaluria, such as primary hyperoxaluria, intestinal diseases or dietary regimens with a high oxalate load [4-8]. According to the surface morphology and crystallite appearance at environmental scanning electron microscopy, five different types of COM stones can be identified [3, 9]. Each of them corresponds to different pathophysiological mechanisms: low diuresis or slight intermittent hyperoxaluria (type Ia), low diuresis and slight intermittent hyperoxaluria (type Ia), hyperoxaluria with anatomical alterations (type Id) and enteric hyperoxaluria (type Ie) [9].

COD is instead typically found in stones of patients who have a high urinary calcium excretion, with or without hyperoxaluria, due to primary hyperparathyroidism, Paget bone disease, prolonged immobilization, sarcoidosis, myeloma, bone metastasis, acromegaly, hyperthyroidism, renal or enteric hypercalciuria [3, 10]. In this context, the detection of prevalent COM or COD composition in stones passed by patients with calcium lithiasis may serve as a guide for detecting stone etiology and prescribing appropriated second-level diagnostic tests [11, 12].

However, the most common etiology of calcium stones is idiopathic, accounting for around 80-85% of patients visited in stone clinics [1, 11, 12]. In these patients, the presence of COM, COD, or a combination of the two in passed stones examined by infrared spectroscopy has uncertain significance.

The objective of this observational study was to detect the possible associations between calcium oxalate stone composition, in terms of COM, COD and their ratio, and the clinical characteristics, including 24-hour urinary parameters of lithogenic risk, of a large group of patients with idiopathic calcium nephrolithiasis (ICN).

Materials and methods

Study participants

All subjects over 18 who completed a medical and urinary metabolic evaluation at our Stone Clinic from 2009 to 2017 were eligible for study enrolment. Inclusion criteria were the presence of ICN, infrared spectroscopy analysis of stones completed at our laboratory within three months from urinary metabolic evaluation, and pure calcium oxalate stone composition (defined as calcium oxalate crystals \geq 97%). Subjects with known calcium stone etiology, such as primary hyperoxaluria, enteric hyperoxaluria, primary hyperparathyroidism or other bone diseases associated with hypercalciuria, were excluded from the study. Subjects with chronic kidney disease (creatinine clearance <60 ml/min), renal tubular acidosis, recurrent urinary tract infections, congenital or acquired anomalies of the kidney and the urinary tract, spina bifida, or cystic fibrosis were excluded as well. Subjects with missing clinical or laboratory data were not considered for the final analysis.

Thus, the main study population was composed exclusively of calcium stone formers with documented pure calcium oxalate stone composition and no known etiology. From an epidemiological perspective, this circumstance represents the majority of cases with calcium nephrolithiasis [12, 13].

A database of urinary profiles of lithogenic risk from a group of non-stone forming controls who underwent urinary metabolic evaluation at our Stone Clinic was also considered, to compare the urine composition of patients with different COD/COM ratio in their stone composition with normal standards. These controls (mean age 42 ± 12 years old, male:female ratio 1:2, body mass index [BMI] 24 ± 4 kg/m²) were selected according to the absence of episodes of renal colic in their personal history and absence of retained stones at abdominal ultrasound at the moment of urine collection. Subjects with congenital or acquires anomalies of the urinary tract, recurrent urinary tract infections, creatinine clearance <60 ml/min and suspected diseases of calcium metabolism were not considered.

Clinical and urinary metabolic evaluation

According to the clinical protocol adopted in our stone clinic [13], a comprehensive medical history, with particular focus on the stone disease course and risk factors, was collected from all participants. Family history and age of onset of the first stone episode were carefully collected [14]. The coexistence of kidney stones with hypertension, that represents an important risk factor for urinary metabolic abnormalities [15], was also particularly assessed.

Height, weight, and arterial pressure were measured. Abdominal ultrasound or X-ray were performed to detect retained stones and their radio-opacity. Blood tests, including serum creatinine, calcium, phosphorus, uric acid, parathormone (PTH), and 25-hydroxyvitamin D (25-OH-D) were performed.

Each participant also collected a 24-hour urine sample for the urinary metabolic profile of lithogenic risk [13]. During the collection, urine was equally distributed in two containers: one containing 2 ml of chlorhexidine gluconate 20% and the other 15 ml of 18% hydrochloric acid. The panel of urinary analyses, performed on the same day the collection was concluded, included pH, sodium, potassium, chloride, creatinine, ammonium, urea, uric acid, citrate (all measured from the chlorhexidine container), calcium, magnesium, oxalate, sulfate and phosphate (measured from the hydrochloric acid container). Urine volume was also assessed considering the content of both urine containers. Urinary relative supersaturations for lithogenic salts, representing an index of the risk of stone recurrence [16], were calculated by using the Equil2 software [17].

Hypercalciuria was defined as a 24-hour urinary calcium excretion $\geq 4 \text{ mg/kg/day}$, while hyperoxaluria was defined as a 24-hour urinary oxalate excretion >45 mg/day.

Stone analyses

Stones passed by participants or extracted during urologic procedures were examined at our stone clinic laboratory by Fourier transform infrared spectroscopy (FT-IR). This technique allows the detection and quantification of COM and COD crystals in stones.

Mixtures with different percentages of COD and COM, selected from patients' kidney stones, were prepared and used for the calibration. The pure COM and COD infrared spectra used for calibration, corresponding to the reference ones [18], are shown in Supplementary Material (Figure S1). COM has a band with absorption peak at 1315 cm⁻¹ and COD at 1325 cm⁻¹, respectively. Among kidney stones with spectra corresponding to COM [18], the one with the lowest value of the 1325/1315 cm⁻¹ ratio was chosen as the reference for pure COM. In fact, due to the additivity of the Lambert-Beer law, the presence of minimal traces of COD in the sample increases the absorbance to a greater extent at 1325 cm⁻¹ than at 1315 cm⁻¹, increasing COD/COM ratio. Conversely, pure COD was selected from samples with spectra equal to COD [18] and with the highest value at 1325/1315 cm⁻¹.

For the FT-IR analyses of stones, pellets were prepared mixing pulverized stone (1%) with potassium bromide (99%). Absorbance spectrum was recorded using a Shimadzu FTIR – 8400S spectrophotometer (Shimadzu

Corporation, Kyoto, Japan), with a measurement range between 400 and 4000 cm⁻¹, resolution 4 cm⁻¹, number of scans 45. The absorbance intensity of recorded spectra ranged between 0.2 and 0.8, to avoid deviations from the Lambert-Beer law.

The relationship between the percentage of COD in the mixture (x) and the absorbance ratio at 1325/1315 cm⁻¹ (y) was described by a quadratic equation $(0.078x^2 + 0.352x + 0.734, R^2 = 0.9995)$, shown in the Supplementary Material (Figure S2). This equation was used to determine the COD/COM ratio in the calcium oxalate kidney stones of patients enrolled in this study.

Statistical analyses

Continuous variables were expressed as mean ± standard deviation or, for non-normally distributed variables, median and interquartile range (IQR). Dichotomous variables were expressed as percentages. Stone formers were stratified by COD/COM ratio intervals of their stone composition, as following: 0-0.25 (first interval), 0.26-0.50 (second interval), 0.51-0.75 (third interval), 0.76-1 (fourth interval). Normally distributed continuous and dichotomous clinical and laboratory parameters were compared among groups of patients, stratified by intervals of COD/COM ratio, using one-way analysis of variance (ANOVA) for crude comparisons, and analysis of covariance (ANCOVA) for comparisons adjusted for covariates (age, sex, duration of disease, BMI). The Bonferroni test for multiple comparisons was applied if adjusted p values were <0.05. Non-normally distributed continuous variables were compared among groups of patients by Kruskal-Wallis test.

For comparisons between patients, stratified by intervals of COD/COM ratio, and controls, urinary parameters were handled as mean \pm standard deviation adjusted for age, sex and BMI, since these variables were different between patients and controls. Urinary supersaturations were also adjusted for urinary volume. Comparisons were then made using ANCOVA. Bonferroni test was again applied if adjusted p values were <0.05.

The relationship between urinary parameters and COD/COM ratio in stone composition was also assessed by linear regression models.

All p-values were considered significant for p<0.05. Analyses were performed with the SPSS software v.24 (SPSS Inc., Chicago, IL, USA).

Results

From 2009 to 2017, 947 stone samples (from 677 males and 270 females) with pure calcium oxalate composition were analyzed in our Stone Clinic. However, 482 subjects were excluded from the analysis for missing clinical or laboratory data or not meeting inclusion criteria. Thus, the study was conducted on a group of 465 stone formers (322 males, 143 females, age 46 ± 14) and their stones.

In stone formers, the COD/COM ratio median was 0.20 (IQR 0.10-0.40), so that most participants (62%) fell within the first interval of COD/COM ratio (0-0.25). The clinical characteristics of patients, stratified by COD/COM ratio intervals, are reported in Table 1. Patients with the highest COD/COM ratio had a higher number of extra-corporeal shock-wave lithotripsy procedures (p = 0.017 with Kruskal-Wallis test) and a lower age of onset of kidney stone disease (p for trend = 0.001 with ANOVA) (Table 1). However, the trend for an earlier onset of the disease was confirmed only in those without a family history of stones (p for trend = 0.003 with ANOVA), and not in those self-reporting a family history of stones (Table 1, Figure 1A).

There was also a trend for an increase in serum calcium with increasing COD/COM ratio (p adjusted with ANCOVA for age, sex, duration of disease and BMI = 0.014), while other serum parameters were not different across COD/COM ratio intervals (Table 2).

The analysis of 24-hour urinary parameters of lithogenic risk across COD/COM intervals is depicted in Table 3. With increasing COD/COM ratio, significantly higher levels of 24-h calcium excretion, calcium oxalate and calcium phosphate supersaturation (all p values adjusted with ANCOVA for age, sex, duration of disease and BMI <0.001) could be demonstrated. Moreover, urinary pH was higher in the fourth vs the third interval of COD/COM ratio, while 24-hour urinary oxalate excretion was not related with the COD/COM ratio (Table 3). The comparison of the 24-hour urine parameters of patients belonging to different intervals of COD/COM ratio with non-stone forming controls revealed that subjects in the first interval of COD/COM ratio had a very similar urine composition than controls, exhibiting only a higher volume and excretion of phosphorus and oxalate (Table 4). Conversely, those with a COD/COM ratio >0.25 exhibited a wider range of urinary abnormalities compared to controls, including a higher calcium excretion and a higher calcium oxalate relative supersaturation index (Table 4). The 24-hour urinary calcium excretion was also unaffected by the presence of a family history of stones (Figure 1B).

A linear regression model, exploring the possible clinical and urinary parameters associated with the COD/COM ratio of stone composition, is shown in Table 5. Only 24-hour urinary calcium ($\beta = 0.124, 95\%$ CI

0.102-0.145, standardized β = 0.464, p<0.001) and urine pH (β = 6.402, 95% CI 1.347-11.457, standardized β

= 0.103, p=0.013) were significantly associated with COD/COM ratio.

Discussion

In a group of patients with idiopathic calcium nephrolithiasis and pure calcium oxalate composition, the COD/COM ratio of stone composition, determined by FT-IR, was significantly associated with an earlier onset of the disease, higher number of urologic procedures, higher serum calcium, higher urinary excretion of calcium and pH. Among these parameters, 24-hour urinary calcium excretion exhibited the strongest correlation with COD/COM ratio. Moreover, patients with a COD/COM ratio ≤ 0.25 , representing the majority of subjects with idiopathic calcium nephrolithiasis, showed no clinically relevant metabolic abnormalities in urine chemistry.

This is one of the first studies exploring the clinical correlates of COD/COM ratio of stone composition in idiopathic calcium nephrolithiasis with stones of pure calcium oxalate composition. Previous investigations were in fact focused on patients with known metabolic abnormalities or secondary forms of calcium nephrolithiasis, and showed an association between hypercalciuria and high COD/COM ratio [19-23]. The only study conducted on an unselected population of calcium oxalate stone formers showed the presence of a significant correlation between urinary calcium/oxalate ratio and stone COD/COM ratio, with oxalate dependence of COM crystal formation and calcium dependence of COD crystal formation [24].

In our group of pure calcium oxalate stone formers, the highest values of 24-hour urinary calcium excretion and the highest prevalence of hypercalciuria were found in those with a COD/COM ratio >0.50. Those in the fourth interval of COD/COM ratio (>0.75) also exhibited a higher urinary pH, suggesting a role of pH in determining the COD content of calcium oxalate stones in hypercalciuric patients [25].

Previous studies also suggested a significant association between hyperoxaluria and prevalent COM composition of stones [4-8, 25], that was not confirmed in our group of pure calcium oxalate stone formers with idiopathic calcium nephrolithiasis. This association is probably typical of gastrointestinal diseases with increased oxalate absorption and dietary regimens with high oxalate load [4-8]. Patients with these conditions were not included in our study, since they do not fit with the criteria for diagnosing idiopathic calcium nephrolithiasis. However, participants with a low COD/COM ratio did exhibit a significantly higher 24-hour urinary oxalate excretion than controls, although the difference was mild.

Another point of interest is the circumstance that the relative supersaturation indexes for calcium oxalate were similar between subjects with pure calcium oxalate stones and COD/COM ratio ≤ 0.25 and healthy controls. The supersaturation indexes are well-known predictors of recurrence of kidney stones [16, 26], and depend on

urine volume and urinary metabolic abnormalities. In clinical practice, the finding of a low COD/COM ratio at stone analysis in patients with idiopathic calcium nephrolithiasis may imply that these patients have no urinary metabolic abnormalities and a low risk of stone recurrence. This assumption is also supported by the findings of two studies performed in large groups of stone formers from the United States [27, 28]. In these studies, a large prevalence of COM composition in kidney stones from first-time stone formers was found, and this composition was associated with the lowest risk of recurrence, compared with patients with COD or other stone compositions [27, 28].

In COM stone formers, if the clinical evaluation allows to exclude the presence of a secondary cause of calcium lithiasis, such as primary hyperparathyroidism or gastrointestinal diseases, hypercalciuria and hyperoxaluria are rarely present. The urinary calcium/oxalate ratio may be involved in the pathogenesis of stones in such situations, as suggested by Daudon and colleagues [24]. However, other factors may be implied. Poor hydration may represent the most important one [29]. In fact, this is a very common risk factor for urolithiasis, although not easy to detect since patients correctly tend to increase the fluid intake after an episode of stones even before medical evaluation [29].

Nutritional imbalances, such as excessive salt intake or reduced fruit and vegetable intake, may have not a relevant role in idiopathic COM stone formers, because they are generally associated with recognizable urinary abnormalities [30, 31], that were not detected in our study. Interestingly, nutritional investigations comparing the dietary habits of idiopathic calcium stone formers with controls showed only minor differences [32, 33], supporting the assumption that nutrition plays a central role in the pathogenesis of kidney stones only in selected cases.

Family history may instead be involved. It is well known that a family history of stones is associated with an earlier onset of stone disease irrespective of urinary metabolic abnormalities [14, 34, 35] and even with stone composition [34]. In the present study, the age of onset of patients with and without family history of stones was significantly different in those with COD/COM ratio ≤ 0.25 , who had few metabolic abnormalities and low urinary supersaturations (mean age of onset 35, 95% CI 33-37, vs 42, 95% CI 40-44, respectively, p<0.001).

This effect may depend on the urinary levels of macromolecules involved in the lithogenic process but not detected in routine 24-hour urine chemistry. These molecules may promote aggregation of calcium oxalate crystals or urinary viscosity even in the absence of high urinary calcium excretion [36-38]. Their action could

also explain the absence of a trend on the age of onset in patients with family history of stones with increasing urinary supersaturations (Figure 1A); conversely, in patients without family history of stones, with the increase in urinary supersaturations the age of onset is lowered to values close to those with family history. Higher proportions of COD in kidney stones are associated with an earlier onset of the disease, irrespective of the presence of family history.

From a clinical perspective, our findings may have relevance for defining the best management strategy for patients with idiopathic calcium nephrolithiasis. In those who have a COD/COM ratio >0.25, metabolic evaluation, i.e., 24-hour urinary collection for determination of the profile of lithogenic risk, is mandatory because the risk of metabolic abnormalities is elevated [11, 12]. Follow-ups should be scheduled every 3-6 months [39], due to the elevated risk of stone recurrence [28, 40]. Conversely, our findings suggest that, in patients with a COD/COM ratio \leq 0.25, the prescription of the urinary profile of lithogenic risk should be made only in selected cases, based on a personal history suggesting the presence of risk factors for recurrence. If these risk factors are not present and the patient is a first-time stone former, metabolic evaluation could be avoided, due to the low risk of detecting metabolic abnormalities that can modify the strategy of secondary prevention [40].

The clinical relevance of the COD/COM ratio in pure idiopathic calcium oxalate stone formers should be further investigated in the future. Although our study suggests a potential usefulness of this parameter in guiding the prevention management of kidney stone formers, some limitations should be considered. The most obvious one is the observational design of the study and the absence of a follow-up, not allowing to ascertain whether the COD/COM ratio is able to predict the clinical course of stone disease. Moreover, the sample size was relatively limited, compared with other previous studies [27, 28], although focused on the most common clinical form of urolithiasis.

Conclusions

In a group of idiopathic pure calcium oxalate stone formers, the COD/COM ratio of stone composition, examined by FT-IR, was positively associated with 24-hour urinary calcium excretion and urinary pH. A COD/COM ratio ≤ 0.25 was associated with little urinary metabolic abnormalities, suggesting different management strategies for patients with these characteristics of stone composition. The clinical significance of COD/COM ratio in idiopathic calcium nephrolithiasis deserves further investigation in the future.

Conflict of interest

The authors have nothing to disclose.

Ethical standards

The study protocol was approved by the local Ethics Committee as part of a larger project on the clinical and nutritional correlates of urinary parameters in nephrolithiasis. The study was carried out according to the principles of the Declaration of Helsinki. Informed consent was obtained according to Italian law for retrospective studies.

Acknowledgements

The authors wish to thank Antonio Nouvenne, for important assistance in study design and manuscript drafting, Maurizio Rossi, for the precious statistical consult, and Michele Zenna, for assistance in database management and support in manuscript drafting.

References

- 1. Romero V, Akpinar H, Assimos DG (2010) Kidney stones: a global picture of prevalence, incidence, and associated risk factors. Rev Urol 12(2-3):e86-e96.
- Daudon M, Réveillaud RJ (1984) Whewellite and weddellite: toward a different etiopathogenesis. Nephrologie 5(5):195-201.
- Daudon M, Bader CA, Jungers P (1993) Urinary calculi: review of classification methods and correlations with etiology. Scan Microsc 7(3):1081-1106.
- 4. Jiang D, Geng H (2017) Primary hyperoxaluria. N Engl J Med 376(15):e33.
- Daudon M, Jungers P, Bazin D (2008) Peculiar morphology of stones in primary hyperoxaluria. N Engl J Med 359(1):100-102.
- Sutton RA, Walker VR (1994) Enteric and mild hyperoxaluria. Miner Electrolyte Metab 20(6):352-360.
- Massey LK, Liebman M, Kynast-Gales SA (2005) Ascorbate increases human oxaluria and kidney stone risk. J Nutr 135(7):1673-1677.
- Albert A, Tiwari V, Paul E, Ponnusamy S, Ganesan D, Prabhakaran R et al (2018) Oral administration of oxalate-enriched spinach extract as an improved methodology for the induction of dietary hyperoxaluric nephrocalcinosis in experimental rats. Toxicol Mech Methods 28(3):195-204.
- Daudon M, Bazin D, André G, Jungers P, Cousson A, Chevallier P et al (2009) Examination of whewellite kidney stones by scanning electron microscopy and powder neutron diffraction techniques. J Appl Cryst 42:109-115.
- Park S, Pearle MS (2007) Pathophysiology and management of calcium stones. Urol Clin N Am 34(3):323-334.
- Prezioso D, Strazzullo D, Lotti T, Bianchi G, Borghi L, Caione P et al (2015) Dietary treatment of urinary risk factors for renal stone formation. A review of CLU Working Group. Arch Ital Urol Androl 87(2):105-120.
- Gambaro G, Croppi E, Coe F, Lingeman J, Moe O, Worcester E et al (2016) Metabolic diagnosis and medical prevention of calcium nephrolithiasis and its systemic manifestations: a consensus statement. J Nephrol 29(6):715-734.

- 13. Nouvenne A, Ticinesi A, Allegri F, Guerra A, Guida L, Morelli I et al (2014) Twenty-five years of idiopathic calcium nephrolithiasis: has anything changed? Clin Chem Lab Med 52(3):337-344.
- 14. Guerra A, Folesani G, Nouvenne A, Ticinesi A, Allegri F, Pinelli S et al (2016) Family history influences clinical course of idiopathic calcium nephrolithiasis: case-control study of a large cohort of Italian patients. J Nephrol 29(5):645-651.
- 15. Ticinesi A, Guerra A, Allegri F, Nouvenne A, Cervellin G, Maggio M et al (2018) Determinants of calcium and oxalate excretion in subjects with calcium nephrolithiasis: the role of metabolic syndrome traits. J Nephrol 31(3):395-403.
- 16. Ferraro PM, Ticinesi A, Meschi T, Rodgers A, Di Maio F, Fulignati P et al (2018) Short-term changes in urinary supersaturation predict recurrence of kidney stones: a tool to guide preventive measures in urolithiasis. J Urol 200(5):1082-1087.
- 17. Werness P, Brown CM, Smith LH, Finlayson B (1985) Equil2: a basic computer program for the calculation of urinary saturation. J Urol 134:1242-1244.
- Maurice-Estepa L, Levillain P, Lacour B, Daudon M (2000) Advantage of zero-crossing-point firstderivative spectrophotometry for the quantification of calcium oxalate crystalline phases by infrared spectrophotometry. Clin Chim Acta 298(1-2):1-11.
- Castiglione V, Jouret F, Bruyère O, Dubois B, Thomas A, Waltregny D et al (2015) Epidemiology of urolithiasis in Belgium on the basis of a morpho-constitutional classification. Nephrol Ther 11(1):42-49.
- 20. Daudon M, Traxer O, Lechevallier E, Saussine C (2008) Epidemiology of urolithiasis. Prog Urol 18(12):802-814.
- 21. Pierratos AE, Khalaff H, Cheng PT, Psihramis K, Jewett MA (1994) Clinical and biochemical differences in patients with pure calcium oxalate monohydrate and calcium oxalate dehydrate kidney stones. J Urol 151(3): 571-574.
- 22. Parent X, Boess G, Brignon P (1999) Calcium oxalate lithiasis. Relationship between biochemical risk factors and crystalline phase of the stone. Prog Urol 9(6):1051-1056.
- 23. Asplin JR, Lingeman J, Kahnoski R, Mardis H, Parks JH, Coe FL (1998) Metabolic urinary correlates of calcium oxalate dehydrate in renal stones. J Urol 159(3):664-668.

- 24. Daudon M, Letavernier E, Frochot V, Haymann JP, Bazin D, Jungers P (2016) Respective influence of calcium and oxalate urine concentration on the formation of calcium oxalate kidney monohydrate or dehydrate crystals. C R Chimie 19:1504-1513.
- 25. Manissorn J, Fong-Ngern K, Peerapen P, Thongboonkerd V (2017) Systematic evaluation for effects of urine pH on calcium oxalate crystallization, crystal-cell adhesion and internalization into renal tubular cells. Sci Rep 7(1):1798.
- 26. Parks JH, Coward M, Coe FL (1997) Correspondence between stone composition and urine supersaturation in nephrolithiasis. Kidney Int 51(3):894-900.
- 27. Singh P, Enders FT, Vaughan LE, Bergstralh EJ, Knoedler JJ, Krambeck AE et al (2015) Stone composition among first-time symptomatic kidney stone formers in the community. Mayo Clin Proc 90(10):1356-1365.
- 28. Vaughan LE, Enders FT, Lieske JC, Pais VM, Rivera ME, Mehta RA et al (2019) Predictors of symptomatic kidney stone recurrence after the first and subsequent episodes. Mayo Clin Proc 94(2):202-210.
- 29. Ticinesi A, Nouvenne A, Borghi L, Meschi T (2017) Water and other fluids in nephrolithiasis: state of the art and future challenges. Crit Rev Food Sci Nutr 57(5):963-974.
- Ticinesi A, Nouvenne A, Maalouf NM, Borghi L, Meschi T (2016) Salt and nephrolithiasis. Nephrol Dial Transplant 31(1):39-45.
- 31. Guerra A, Ticinesi A, Allegri F, Nouvenne A, Prati B, Pinelli S et al (2019) Insights about urinary hippuric and citric acid as biomarkers of fruit and vegetable intake in patients with kidney stones: the role of age and gender. Nutrition 59:83-89.
- 32. Meschi T, Nouvenne A, Ticinesi A, Prati B, Guerra A, Allegri F et al (2012) Dietary habits in women with recurrent idiopathic calcium nephrolithiasis. J Transl Med 10:63.
- 33. Ticinesi A, Milani C, Guerra A, Allegri F, Lauretani F, Nouvenne A et al (2018) Understanding the gut-kidney axis in nephrolithiasis: an analysis of the gut microbiota composition and functionality of stone formers. Gut 67(12):2097-2106.
- 34. Guerra A, Ticinesi A, Allegri F, Nouvenne A, Pinelli S, Folesani G et al (2016) The influence of maternal and paternal history on stone composition and clinical course of calcium nephrolithiasis in subjects aged between 15 and 25. Urolithiasis 44(6):521-528.

- 35. Guerra A, Ticinesi A, Allegri F, Nouvenne A, Pinelli S, Lauretani F et al (2017) Calcium urolithiasis course in young stone formers is influenced by the strength of family history: results from a retrospective study. Urolithiasis 45(6):525-533.
- 36. Jaggi M, Nakagawa Y, Zipperle L, Hess B (2007) Tamm-Horsfall protein in recurrent calcium kidney stone formers with positive family history: abnormalities in urinary excretion, molecular structure and function. Urol Res 35(2):55-62.
- 37. Yamate T, Tsuji H, Amasaki N, Iguchi M, Kurita T, Kohri K (2000) Analysis of osteopontin DNA in patients with urolithiasis. Urol Res 28(3):159-166.
- 38. Rimel JD, Kolbach-Mandel AM, Ward MD, Wesson JA (2017) The role of macromolecules in the formation of kidney stones. Urolithiasis 45(1):47-54.
- 39. Wollin DA, Kaplan AG, Preminger GM, Ferraro PM, Nouvenne A, Tasca A et al (2018) Defining metabolic activity of nephrolithiasis - Appropriate evaluation and follow-up of stone formers. Asian J Urol 5(4):235-242.
- 40. Daudon M, Jungers P, Bazin D, Williams jr JC (2018) Recurrence rates of urinary calculi according to stone composition and morphology. Urolithiasis 46(5):459-470.

COD/COM ratio	(0-0.25)	(0.26-0.50)	(0.51-0.75)	(0.76-1)	p*	p** adjusted	p*** value for trend
Number (%)	287 (62%)	86 (18%)	40 (9%)	(0.70-1) 52 (11%)		aujusteu	value for trend
Intervals	(1)	(2)	(3)	(4)			
Females,%	31	37	30	17	0.104		0.029
Age, years	47 ± 13	46 ± 14	46 ± 15	42 ± 16	0.085		0.016
Weight, kg	75 ± 16	74 ± 15	75 ± 14	76 ± 12	0.961		
BMI, kg/m^2	25 ± 4	26 ± 4	25 ± 3	25 ± 3	0.468		
Disease duration	5 [1-14]	5 [1-15]	8 [1-21]	3 [1-19]	0.373		
Family history of stones (FHS), %	52	51	60	48	0.716		
Age of onset of kidney stones	38 ± 14	37 ± 14	32 ± 13	32 ± 12	0.007		0.001
Age of onset of kidney stones in patients without FHS	42 ± 14	<i>39</i> ± <i>14</i>	<i>31</i> ± <i>12</i>	35 ± 13	0.003		0.003
Age of onset of kidney stones in patients with FHS	35 ± 13	35 ± 13	<i>33</i> ± <i>14</i>	<i>30</i> ± <i>11</i>	0.315		
Hypertensive, %	21	26	28	15	0.396	0.369	
Recurrents, %	62	68	73	62	0.495	0.634	
Stones retained, %	55	50	59	41	0.304	0.138	
Bilateral stones,%	45	52	55	43	0.473	0.585	
Extra-Corporeal Shock-Wave Lithotripsy (ESWL), number	0 [0-1]	0 [0-1]	0 [0-2]	0 [0-2]	0.017		
Stone rate, years	0.39 [0.16-1.00]	0.62 [0.20-1.00]	0.35 [0.16-0.97]	0.46 [0.15-1.00]	0.361		

Table 1. Clinical characteristics of the patients with idiopathic calcium nephrolithiasis and pure calcium oxalate stone composition included in the study (n=465, 322 M and 143 F), stratified by the intervals of the ratio of calcium oxalate dihydrate (COD) and calcium oxalate monohydrate (COM) crystals in stones.

Data reported as percentage or median and interquartile range or mean \pm standard deviation. Significant p values (p<0.05) are indicated in bold.

*Crude p values obtained with ANOVA (dichotomous variables or continuous variables with normal distribution) or Kruskal-Wallis test (continuous variables with non-normal distribution).

** p values adjusted for sex, age, BMI and duration of disease with ANCOVA (only variables requiring adjustment for clinical reasons).

***p for trend values obtained with ANOVA (linear trends). Values are reported only if significant (p<0.05).

Table 2 . Blood chemistry parameters of the patients with idiopathic calcium nephrolithiasis and pure calcium oxalate stone composition included in the
study (n=465, 322 M and 143 F), stratified by the intervals of the ratio of calcium oxalate dihydrate (COD) and calcium oxalate monohydrate (COM)
crystals in stones.

COD/COM ratio Number Intervals	(0-0.25) N.287 (1)	(0.26-0.50) N.86 (2)	(0.51-0.75) N.40 (3)	(0.76-1) N.52 (4)	p*	p**	p<0.05 Bonferroni test
Creatinine, mg/dl	0.90 ± 0.18	0.86 ± 0.18	0.90 ± 0.18	0.89 ± 0.14	0.285	0.024	
Uric acid, mg/dl	5.38 ± 1.27	5.02 ± 1.09	5.25 ± 0.96	5.41 ± 1.10	0.127	0.101	
Calcium, mg/dl	9.47 ± 0.38	9.46 ± 0.45	9.51 ± 0.45	9.70 ± 0.44	0.003	0.014	(1) and (2) vs (4)
Phosphorus, mg/dl	3.29 ± 0.53	3.29 ± 0.55	3.17 ± 0.60	3.33 ± 0.62	0.600	0.468	
PTH, pg/ml	44 ± 14	43 ± 13	39 ± 12	40 ± 14	0.226	0.282	
25-OH-D, ng/ml	22 ± 13	23 ± 15	25 ± 12	21 ± 13	0.724	0.887	

25-OH-D: 25-hydroxy-vitamin D.

Data reported as mean ± standard deviation. Significant p values (p<0.05) are indicated in bold. *Crude p values obtained with ANOVA. ** p values adjusted for sex, age, BMI and duration of disease with ANCOVA

Table 3. Urinary chemistry parameters of the patients with idiopathic calcium nephrolithiasis and pure calcium oxalate stone composition included	in the
study (n=465, 322 M and 143 F), stratified by the intervals of the ratio of calcium oxalate dihydrate (COD) and calcium oxalate monohydrate (COD)	COM)
crystals in stones.	

COD/COM ratio Number Intervals	(0-0.25) N.287 (1)	(0.26-0.50) N.86 (2)	(0.51-0.75) N.40 (3)	(0.76-1) N.52 (4)	p*	p**	p<0.05 Bonferroni test
Volume, ml/24h	1905 ± 702	1916 ± 774	1716 ± 640	1969 ± 757	0.367	0.254	
Creatinine, mg/24h	1525 ± 426	1522 ± 452	1643 ± 455	1580 ± 404	0.367	0.125	
Sodium, mEq/24h	171 ± 61	168 ± 59	164 ± 53	166 ± 59	0.867	0.548	
Potassium, mEq/24h	55 ± 18	53 ± 20	55 ± 15	54 ± 15	0.965	0.952	
Calcium, mg/24h	194 ± 85	256 ± 84	315 ± 122	313 ± 119	<0.0001	<0.0001	(1) vs (2) vs (3) vs (4 (2) vs (3) vs (4)
Hypercalciuria $(\geq 4 \text{ mg/kg/24h}), \%$	12	28	53	48	<0.0001	<0.0001	(1) vs (2) vs (3) vs (4); (2) vs (3) vs (4)
Magnesium, mg/24h	87 ± 29	91 ± 24	101 ± 34	95 ± 29	0.013	0.023	(1) vs (3)
Chloride, mEq/24h	167 ± 62	165 ± 56	167 ± 52	168 ± 59	0.984	0.891	
Phosphorus, mg/24h	842 ± 271	848 ± 239	921 ± 235	855 ± 286	0.360	0.195	
Uric acid, mg/24h	571 ± 164	597 ± 178	595 ± 166	567 ± 144	0.518	0.186	
Oxalate, mg/24h	31 ± 11	31 ± 9	33 ± 8	30 ± 9	0.645	0.457	
Hyperoxaluria (>45 mg/24h), %	7	7	8	4	0.859	0.886	
Sulphate, mmol/24h	21 ± 7	21 ± 7	22 ± 7	20 ± 6	0.554	0.151	
Ammonium, mmol/24h	36 ± 12	37 ± 12	38 ± 10	38 ± 13	0.351	0.494	
Urea, g/24h	24 ± 7	23 ± 7	25 ± 7	23 ± 7	0.661	0.240	
Citrate, mg/24h	578 ± 257	633 ± 249	600 ± 258	591 ± 278	0.394	0.169	
Urine pH, 24h	5.88 ± 0.45	5.93 ± 0.43	5.76 ± 0.43	6.04 ± 0.46	0.016	0.015	(3) vs (4)
Calcium oxalate supersaturation	5.26 ± 2.94	6.75 ± 3.83	8.54 ± 3.42	7.22 ± 3.58	<0.0001	<0.0001	(1) vs (2) vs (3) vs (4) (2) vs (3)
Calcium phosphate supersaturation	0.66 ± 0.61	0.94 ± 0.75	1.17 ± 1.08	1.27 ± 0.83	<0.0001	<0.0001	(1) vs (2) vs (3) vs (4

Data reported as percentage or mean ± standard deviation. Significant p values (p<0.05) are indicated in bold. *Crude p values obtained with ANOVA. **p values adjusted with ANCOVA for sex, age, BMI and duration of disease.

Table 4. Comparison of urinary chemistry parameters between patients with idiopathic calcium nephrolithiasis and pure calcium oxalate stone
composition (n=465), stratified by the intervals of the ratio of calcium oxalate dihydrate (COD) and calcium oxalate monohydrate (COM) crystals in
stones, and a group of non-stone forming controls (n=486).

			Stone Formers N.465		р	p < 0.05 Bonferroni test	
	Controls N.486 (c)	COD/COM (0-0.25) N. 287 (1)	COD/COM (0.26-0.50) N.86 (2)	COD/COM (0.50-1) N.92 (3)			
Volume, ml/24h	1516 ± 693	1883 ± 685	1895 ± 668	1843 ± 674	<0.0001	(c) vs (1) vs (2) vs (3)	
Creatinine, mg/24h	1404 ± 293	1432 ± 290	1436 ± 283	1454 ± 285	0.351		
Sodium, mEq/24h	161 ± 57	164 ± 56	160 ± 55	155 ± 55	0.589		
Potassium, mEq/24h Calcium, mg/24h Hypercalciuria,%	57 ± 19 198 ± 95 17	53 ± 18 186 ± 94 12	52 ± 18 247 ± 91 29	53 ± 18 304 ± 92 50	0.014 <0.0001 <0.0001	(c) vs (2) vs (3) (c) vs (2) vs (3)	
Phosphorus, mg/24h	727 ± 236	805 ± 233	809 ± 227	829 ± 229	<0.0001	(c) vs (1) vs (2) vs (3)	
Magnesium, mg/24h Uric acid, mg/24h Citrate, mg/24 h Sulphate, mmol/24h Ammonium, mmol/24h	$\begin{array}{c} 84 \pm \ 29 \\ 537 \pm \ 157 \\ 625 \pm \ 260 \\ 20 \pm \ 6 \\ 35 \pm \ 11 \end{array}$	85 ± 29 550 ± 155 578 ± 257 20 ± 6 34 ± 11	89 ± 28 572 ± 151 626 ± 251 20 ± 5 36 ± 11	94 ± 29 548 ± 153 613 ± 253 20 ± 5 35 ± 11	0.025 0.231 0.095 0.351 0.687	(c) vs (3)	
Oxalate, mg/24 h	27 ± 10	30 ± 10	30 ± 9	30 ± 9	<0.0001	(c) vs (1) vs (3)	
Hyperoxaluria,%	5	6	6	4	0.935		
Urine pH, 24h	5.99 ± 0.51	5.93 ± 0.51	5.97 ± 0.49	5.95 ± 0.49	0.472		
Calcium oxalate supersaturation	5.32 ± 3.04	5.67 ± 3.00	7.18 ± 2.91	8.02 ± 2.93	<0.0001*	(c) vs (2) vs (3)	
Calcium phosphate supersaturation	0.84 ± 0.77	0.74 ± 0.76	1.02 ± 0.74	1.23 ± 0.75	<0.0001*	(c) vs (3)	

supersaturation
 58
 bit 2 of 1 2 o

Table 5. Linear regression model testing the relationship between calcium, oxalate and urine pH with COD/COM ratio in 465 stone formers with idiopathic calcium nephrolithiasis and pure calcium oxalate stone composition.

	β	95%CI	β standardized	р
Calcium, mg/24h	0.124	0.102-0.145	0.464	< 0.0001
Oxalate, mg/24 h	-0.145	-0.370- 0.081	- 0.052	0.209
Urine pH, 24h	6.402	1.347-11.457	0.103	0.013

Significant p values (p<0.05) are indicated in bold

Figure 1. Mean and 95% CI of the age of onset of kidney stones (A) and of calciuria (B) in 461 patients stratified by family history of stones (FHS) (221 without FHS, 240 with FHS) and ratio COD/COM (0-0.25, 0.26-0.50, 0.51-1). A significant trend, for the age of onset of kidney stones, is present in patients without FHS (p = 0.003) increasing the COD/COM ratio and urinary supersaturations, but not in patients with FHS (p = 0.374). Calciuria (B) is not different in patients with and without FHS, p = 0.798.Age of onset of kidney stones is adjusted for BMI and sex. Calciuria by age, sex, duration of disease, BMI, sodium, potassium, ammonium and urinary sulfates. Calcium oxalate supersaturation (srcaox) and calcium phosphate supersaturation (srcap) adjusted for BMI, age, volume and sex, no differences for over-saturation between patients with FHS and without FHS

Abstract

Pure calcium oxalate is the most frequent type of idiopathic kidney stone composition. Fourier transform infrared spectroscopy (FT-IR) allows to detect the ratio of calcium oxalate dihydrate (COD) and monohydrate (COM) crystals in stones, but the clinical significance of this parameter remains uncertain. The objective of this observational study was to verify the association of clinical and laboratory parameters of kidney stone disease with COD/COM ratio in a group of 465 (322 M, age 46 ± 14) patients suffering from idiopathic calcium nephrolithiasis with pure calcium oxalate stones ($\geq 97\%$). Each participant underwent a complete clinical examination, serum chemistry, 24-hour urine collection for the determination of the profile of lithogenic risk, and had stones analyzed by FT-IR. Most (62%) of the stones has a COD/COM ratio <0.25, and the urine chemistry of the corresponding patients showed a very low prevalence of urinary metabolic abnormalities. With increasing quartiles of COD/COM ratio intervals (0-0.25, 0.26-0.50, 0.51-0.75, 0.76-1), a significant positive trend association was observed for the number of urological procedures, serum calcium, 24-h urinary calcium excretion, prevalence of hypercalciuria and relative calcium oxalate supersaturation, and a negative trend was detected for the age of the first stone episode (all p-values for trend <0.05). A linear regression model showed that the only parameters significantly associated with COD/COM ratio were 24-h urinary calcium excretion (standardized $\beta=0.464$, p<0.001) and urine pH (standardized $\beta=0.103$, p=0.013). In pure calcium oxalate idiopathic stones, COD/COM ratio may reflect the presence of urinary metabolic risk factors, and serve asrepresent a guide for the prescription of urinary analyses.

Key words: urolithiasis; hypercalciuria; hyperoxaluria; kidney stones; calcium oxalate.

Introduction

Under infrared spectroscopy, calcium oxalate crystals of calcium stones may appear in two distinct molecular forms: whewellite, that is, calcium oxalate monohydrate (COM), and weddellite, that is, calcium oxalate dihydrate (COD). These forms are associated with different etiology of stones [1, 2] and are also associated with different surface morphology of calculi [3].

COM depends on urinary excretion of oxalate and is typical of conditions of hyperoxaluria, such as primary hyperoxaluria, intestinal diseases or dietary regimens with a high oxalate load [34-78]. According to the surface morphology and crystallite appearance at environmental scanning electron microscopy, five different types of COM stones can be identified [3, 9]. Each of them corresponds to different pathophysiological mechanisms: low diuresis or slight intermittent hyperoxaluria (type Ia), low diuresis and slight intermittent hyperoxaluria (type Ia), hyperoxaluria with anatomical alterations (type Id) and enteric hyperoxaluria (type Ie) [9].

COD is instead typically found in stones of patients who have a high urinary calcium excretion, with or without hyperoxaluria, due to primary hyperparathyroidism, Paget bone disease, prolonged immobilization, sarcoidosis, myeloma, bone metastasis, acromegaly, hyperthyroidism, renal or enteric hypercalciuria [3, 810]. In this context, the detection of prevalent COM or COD composition in stones passed by patients with calcium lithiasis may serve as a guide for detecting stone etiology and prescribing appropriated second-level diagnostic tests [911, 1012].

However, the most common etiology of calcium stones is idiopathic, accounting for around 80-85% of patients visited in stone clinics [1, 911, 1012]. In these patients, the presence of COM, COD, or a combination of the two in passed stones examined by infrared spectroscopy has uncertain significance.

The objective of this observational study was to detect the possible associations between calcium oxalate stone composition, in terms of COM, COD and their ratio, and the clinical characteristics, including 24-hour urinary parameters of lithogenic risk, of a large group of patients with idiopathic calcium nephrolithiasis (ICN).

Materials and methods

Study participants

All subjects over 18 who completed a medical and urinary metabolic evaluation at our Stone Clinic from 2009 to 2017 were eligible for study enrolment. Inclusion criteria were the presence of ICN, infrared spectroscopy analysis of stones completed at our laboratory within three months from urinary metabolic evaluation, and pure calcium oxalate stone composition (defined as calcium oxalate crystals \geq 97%). Subjects with known calcium stone etiology, such as primary hyperoxaluria, enteric hyperoxaluria, primary hyperparathyroidism or other bone diseases associated with hypercalciuria, were excluded from the study. Subjects with chronic kidney disease (creatinine clearance <60 ml/min), renal tubular acidosis, recurrent urinary tract infections, congenital or acquired anomalies of the kidney and the urinary tract, spina bifida, or cystic fibrosis were excluded as well. Subjects with missing clinical or laboratory data were not considered for the final analysis.

Thus, the main study population was composed exclusively of calcium stone formers with documented pure calcium oxalate stone composition and no known etiology. From an epidemiological perspective, this circumstance represents the majority of cases with calcium nephrolithiasis [1012, 113].

A database of urinary profiles of lithogenic risk from a group of non-stone forming controls who underwent urinary metabolic evaluation at our Stone Clinic for reasons other than nephrolithiasis was also considered, to compare the urine composition of patients with different COD/COM ratio in their stone composition with normal standards. These controls (mean age 42±12 years old, male:female ratio 1:2, body mass index [BMI] 24±4 kg/m²) were selected according to the absence of episodes of renal colic in their personal history and absence of retained stones at abdominal ultrasound at the moment of urine collection. Subjects with congenital or acquires anomalies of the urinary tract, recurrent urinary tract infections, creatinine clearance <60 ml/min and suspected diseases of calcium metabolism were not considered.

Clinical and urinary metabolic evaluation

According to the clinical protocol adopted in our stone clinic [1113], a comprehensive medical history, with particular focus on the stone disease course and risk factors, was collected from all participants. Family history and age of onset of the first stone episode were carefully collected [1214]. The coexistence of kidney stones with hypertension, that represents an important risk factor for urinary metabolic abnormalities [1315], was also particularly assessed.

Height, weight, and arterial pressure were measured. Abdominal ultrasound or X-ray were performed to detect retained stones and their radio-opacity. Blood tests, including serum creatinine, calcium, phosphorus, uric acid, parathormone (PTH), and 25-hydroxyvitamin D (25-OH-D) were performed.

Each participant also collected a 24-hour urine sample for the urinary metabolic profile of lithogenic risk [4413]. During the collection, urine was equally distributed in two containers: one containing 2 ml of chlorhexidine gluconate 20% and the other 15 ml of 18% hydrochloric acid. This-The panel of urinary analyses, performed on the same day the collection was concluded, included pH, volume and urinary excretion of calcium, chloride, phosphorus, oxalate, citrate, magnesium, potassium, sodium, sulfate, ammonium, urie acid, urea and creatininesodium, potassium, chloride, creatinine, ammonium, urea, uric acid, citrate (all measured from the chlorhexidine container), calcium, magnesium, oxalate, sulfate and phosphate (measured from the hydrochloric acid container). Urine volume was also assessed considering the content of both urine containers. Urinary relative supersaturations for lithogenic salts, representing an index of the risk of stone recurrence [4416], were calculated by using the Equil2 software [4517].

According to recent consensus, <u>hHypercalciuria</u> was defined as a 24-hour urinary calcium excretion ≥ 4 mg/kg/day, while hyperoxaluria was defined as a 24-hour urinary oxalate excretion >45 mg/day [9, 10].

Stone analyses

Stones passed by participants or extracted during urologic procedures were examined at our stone clinic laboratory by Fourier transform infrared spectroscopy (FT-IR). This technique allows the detection and quantification of COM and COD crystals in stones.

Mixtures with different percentages of COD and COM, selected from patients' kidney stones, were prepared and used for the calibration. The pure COM and COD infrared spectra used for calibration, corresponding to the reference ones [4618], are shown in Supplementary Material (Figure S1). COM has a band with absorption peak at 1315 cm⁻¹ and COD at 1325 cm⁻¹, respectively. Among kidney stones with spectra corresponding to COM [4618], the one with the lowest value of the 1325/1315 cm⁻¹ ratio was chosen as the reference for pure COM. In fact, due to the additivity of the Lambert-Beer law, the presence of minimal traces of COD in the sample increases the absorbance to a greater extent at 1325 cm⁻¹ than at 1315 cm⁻¹, increasing COD/COM ratio. Conversely, pure COD was selected from samples with spectra equal to COD [4618] and with the highest value at 1325/1315 cm⁻¹. For the FT-IR analyses of stones, pellets were prepared mixing pulverized stone (1%) with potassium bromide (99%). Absorbance spectrum was recorded using a Shimadzu FTIR – 8400S spectrophotometer (Shimadzu Corporation, Kyoto, Japan), with a measurement range between 400 and 4000 cm⁻¹, resolution 4 cm⁻¹, number of scans 45. The absorbance intensity of recorded spectra ranged between 0.2 and 0.8, to avoid deviations from the Lambert-Beer law.

The relationship between the percentage of COD in the mixture (x) and the absorbance ratio at 1325/1315 cm⁻¹ (y) was described by a quadratic equation $(0.078x^2 + 0.352x + 0.734, R^2 = 0.9995)$, shown in the Supplementary Material (Figure S2). This equation was used to determine the COD/COM ratio in the calcium oxalate kidney stones of patients enrolled in this study.

Statistical analyses

Data-Continuous variables were expressed as mean ± standard deviation or, for non-normally distributed variables, mean and 95% confidence intervals (CI), median and interquartile range (IQR), and Dichotomous variables were expressed as percentages-as appropriate. Stone formers were stratified by COD/COM ratio quartile-intervals of their stone composition, as following: 0-0.25 (first interval), 0.26-0.50 (second interval), 0.51-0.75 (third interval), 0.76-1 (fourth interval), and clinical parameters were compared among these groups and controls by using Kruskal-Wallis test, Bonferroni test for multiple comparisons, one-way analysis of variance (ANOVA), and covariance analysis (ANCOVA), as appropriate according to the variable type and distribution.Normally distributed continuous and dichotomous clinical and laboratory parameters were compared among groups of patients, stratified by intervals of COD/COM ratio, using one-way analysis of variance (ANOVA) for crude comparisons, and analysis of covariance (ANCOVA) for comparisons adjusted for covariates (age, sex, duration of disease, BMI). The Bonferroni test for multiple comparisons was applied if adjusted p values were <<0.05. Non-normally distributed continuous variables were compared among groups of patients by Kruskal-Wallis test. The relationship between urinary parameters and COD/COM ratio in stone composition was also assessed by linear regression models.

For comparisons between patients, stratified by intervals of COD/COM ratio, and controls, urinary parameters were handled as mean \pm standard deviation adjusted for age, sex and BMI, since these variables were different between patients and controls. Urinary supersaturations were also adjusted for urinary volume. Comparisons were then made using ANCOVA. Bonferroni test was again applied if adjusted p values were <0.05.

 The relationship between urinary parameters and COD/COM ratio in stone composition was also assessed by linear regression models.

All p-values were two tailed and were considered significant for p<0.05. Analyses were performed with the SPSS software v.24 (SPSS Inc., Chicago, IL, USA).

Ethical statement

The study protocol was approved by the local Ethics Committee. All data were obtained and handled in anonymous way. Informed consent was obtained and the study procedures followed the principles contained in the Declaration of Helsinki.

Results

From 2009 to 2017, 947 stone samples (from 677 males and 270 females) with pure calcium oxalate composition were analyzed in our Stone Clinic. However, 482 subjects were excluded from the analysis for missing clinical or laboratory data or not meeting inclusion criteria. Thus, the study was conducted on a group of 465 stone formers (322 males, 143 females, age 46 ± 14) and their stones. A group of 486 non-stone forming controls who performed urine analyses at our center was also considered as a control group.

In stone formers, the COD/COM ratio median was 0.20 (IQR 0.10-0.40), so that most participants (62%) fell within the lowest-quartile<u>first interval</u> of COD/COM ratio (0-0.25). The clinical characteristics of patients, stratified by COD/COM ratio <u>quartileintervals</u>, are reported in Table 1. Patients with the highest COD/COM ratio had a higher number of extra-corporeal shock-wave lithotripsy procedures (p for trend = 0.004017 with Kruskal-Wallis test) and a lower age of onset of kidney stone disease (p for trend = 0.001 with ANOVA) (Table 1). However, the trend for an earlier onset of the disease was confirmed only in those without a family history of stones (p for trend = 0.003 with ANOVA), and not in those self-reporting a family history of stones (Table 1, Figure 1A).

There was also a trend for an increase in serum calcium with increasing COD/COM ratio (p adjusted with <u>ANCOVA</u> for age, sex, duration of disease and <u>body mass index BMI</u> = 0.014), while other serum parameters were not different across COD/COM ratio <u>quartiles-intervals</u> (Table 2).

The analysis of 24-hour urinary parameters of lithogenic risk across COD/COM quartiles-<u>intervals</u> is depicted in Table 3. With increasing COD/COM ratio, significantly higher levels of 24-h calcium excretion, calcium oxalate and calcium phosphate supersaturation (all p values adjusted <u>with ANCOVA</u> for age, sex, duration of disease and <u>body mass indexBMI</u> <0.001) could be demonstrated. Moreover, urinary pH was higher in the <u>highest-fourth</u> vs the third <u>quartile-interval</u> of COD/COM ratio, while 24-hour urinary oxalate excretion was not related with the COD/COM ratio (Table 3).

The comparison of the 24-hour urine parameters of patients belonging to different <u>quartiles-intervals</u> of COD/COM ratio with non-stone forming controls revealed that subjects in the <u>lowest quartilefirst interval</u> of COD/COM ratio had a very similar urine composition than controls, exhibiting only a higher volume and excretion of phosphorus and oxalate (Table 4). Conversely, those with a COD/COM ratio >0.25 exhibited a wider range of urinary abnormalities compared to controls, including a higher calcium excretion and a higher

calcium oxalate relative supersaturation index (Table 4). The 24-hour urinary calcium excretion was also unaffected by the presence of a family history of stones (Figure 1B).

A linear regression model, exploring the possible clinical and urinary parameters associated with the COD/COM ratio of stone composition, is shown in Table 5. Only 24-hour urinary calcium ($\beta = 0.124, 95\%$ CI 0.102-0.145, standardized $\beta = 0.464$, p<0.001) and urine pH ($\beta = 6.402, 95\%$ CI 1.347-11.457, standardized $\beta = 0.103$, p=0.013) were significantly associated with COD/COM ratio.

Discussion

In a group of patients with idiopathic calcium nephrolithiasis and pure calcium oxalate composition, the COD/COM ratio of stone composition, determined by FT-IR, was significantly associated with an earlier onset of the disease, higher number of urologic procedures, higher serum calcium, higher urinary excretion of calcium and pH. Among these parameters, 24-hour urinary calcium excretion exhibited the strongest correlation with COD/COM ratio. Moreover, patients with a COD/COM ratio ≤ 0.25 , representing the majority of subjects with idiopathic calcium nephrolithiasis, showed no clinically relevant metabolic abnormalities in urine chemistry.

This is one of the first studies exploring the clinical correlates of COD/COM ratio of stone composition in idiopathic calcium nephrolithiasis with stones of pure calcium oxalate composition. Previous investigations were in fact focused on patients with known metabolic abnormalities or secondary forms of calcium nephrolithiasis, and showed an association between hypercalciuria and high COD/COM ratio [47<u>19-2423</u>]. The only study conducted on an unselected population of calcium oxalate stone formers showed the presence of a significant correlation between urinary calcium/oxalate ratio and stone COD/COM ratio, with oxalate dependence of COM crystal formation and calcium dependence of COD crystal formation [24].

In our group of pure calcium oxalate stone formers, the highest values of 24-hour urinary calcium excretion and the highest prevalence of hypercalciuria were found in those with a COD/COM ratio >0.50. Those in the highest quartilefourth interval of COD/COM ratio (>0.75) also exhibited a higher urinary pH, suggesting a role of pH in determining the COD content of calcium oxalate stones in hypercalciuric patients [2225].

Previous studies also suggested a significant association between hyperoxaluria and prevalent COM composition of stones [34-78, 2225], that was not confirmed in our group of pure calcium oxalate stone formers with idiopathic calcium nephrolithiasis. This association is probably typical of gastrointestinal diseases with increased oxalate absorption and dietary regimens with high oxalate load [34-78]. Patients with these conditions were not included in our study, since they do not fit with the criteria for diagnosing idiopathic calcium nephrolithiasis. However, participants with a low COD/COM ratio did exhibit a significantly higher 24-hour urinary oxalate excretion than controls, although the difference was mild.

Another point of interest is the circumstance that the relative supersaturation indexes for calcium oxalate were similar between subjects with pure calcium oxalate stones and COD/COM ratio ≤ 0.25 and healthy controls. The supersaturation indexes are well-known predictors of recurrence of kidney stones [1416, 2326], and

depend on urine volume and urinary metabolic abnormalities. In clinical practice, the finding of a low COD/COM ratio at stone analysis in patients with idiopathic calcium nephrolithiasis may imply that these patients have no urinary metabolic abnormalities and a low risk of stone recurrence. This assumption is also supported by the findings of two studies performed in large groups of stone formers from the United States [2427, 2528]. In these studies, a large prevalence of COM composition in kidney stones from first-time stone formers was found, and this composition was associated with the lowest risk of recurrence, compared with patients with COD or other stone compositions [2427, 2528].

In COM stone formers, if the clinical evaluation allows to exclude the presence of a secondary cause of calcium lithiasis, such as primary hyperparathyroidism or gastrointestinal diseases, hypercalciuria and hyperoxaluria are rarely present. The urinary calcium/oxalate ratio may be involved in the pathogenesis of stones in such situations, as suggested by Daudon and colleagues [24]. From a physic pathological perspective, this means thatHowever, other factors may be implied. Poor hydration may represent the most important one [2629]. In fact, this is a very common risk factor for urolithiasis, although not easy to detect since patients correctly tend to increase the fluid intake after an episode of stones even before medical evaluation [2629].

Nutritional imbalances, such as excessive salt intake or reduced fruit and vegetable intake, may have not a relevant role in idiopathic COM stone formers, because they are generally associated with recognizable urinary abnormalities [2730, 2831], that were not detected in our study. Interestingly, nutritional investigations comparing the dietary habits of idiopathic calcium stone formers with controls showed only minor differences [2932, 3033], supporting the assumption that nutrition plays a central role in the pathogenesis of kidney stones only in selected cases.

Family history may instead be involved. It is well known that a family history of stones is associated with an earlier onset of stone disease irrespective of urinary metabolic abnormalities [1214, 3134, 3235] and even with stone composition [3134]. In the present study, the age of onset of patients with and without family history of stones was significantly different in those with COD/COM ratio ≤ 0.25 , who had few metabolic abnormalities and low urinary supersaturations (mean age of onset 35, 95% CI 33-37, vs 42, 95% CI 40-44, respectively, p<0.001).

This effect may depend on the urinary levels of macromolecules involved in the lithogenic process but not detected in routine 24-hour urine chemistry. These molecules may promote aggregation of calcium oxalate crystals or urinary viscosity even in the absence of high urinary calcium excretion [3336-3538]. Their action

could also explain the absence of a trend on the age of onset in patients with family history of stones with increasing urinary supersaturations (Figure 1A); conversely, in patients without family history of stones, with the increase in urinary supersaturations the age of onset is lowered to values close to those with family history. Higher proportions of COD in kidney stones are associated with an earlier onset of the disease, irrespective of the presence of family history.

From a clinical perspective, our findings may have relevance for defining the best management strategy for patients with idiopathic calcium nephrolithiasis. In those who have a COD/COM ratio >0.25, metabolic evaluation, i.e., 24-hour urinary collection for determination of the profile of lithogenic risk, is mandatory because the risk of metabolic abnormalities is elevated [911, 1012]. Follow-ups should be scheduled every 3-6 months [3639], due to the elevated risk of stone recurrence [2528, 3740]. Conversely, our findings suggest that, in patients with a COD/COM ratio \leq 0.25, the prescription of the urinary profile of lithogenic risk should be made only in selected cases, based on a personal history suggesting the presence of risk factors for recurrence. If these risk factors are not present and the patient is a first-time stone former, metabolic evaluation could be avoided, due to the low risk of detecting metabolic abnormalities that can modify the strategy of secondary prevention [3740].

The clinical relevance of the COD/COM ratio in pure idiopathic calcium oxalate stone formers should be further investigated in the future. Although our study suggests a potential usefulness of this parameter in guiding the prevention management of kidney stone formers, some limitations should be considered. The most obvious one is the observational design of the study and the absence of a follow-up, not allowing to ascertain whether the COD/COM ratio is able to predict the clinical course of stone disease. Moreover, the sample size was relatively limited, compared with other previous studies [2427, 2528], although focused on the most common clinical form of urolithiasis.

Conclusions

In a group of idiopathic pure calcium oxalate stone formers, the COD/COM ratio of stone composition, examined by FT-IR, was positively associated with 24-hour urinary calcium excretion and urinary pH. A COD/COM ratio \leq 0.25 was associated with little urinary metabolic abnormalities, suggesting different management strategies for patients with these characteristics of stone composition. The clinical significance of COD/COM ratio in idiopathic calcium nephrolithiasis deserves further investigation in the future.

Conflict of interest

The authors have nothing to disclose.

Ethical standards

The study protocol was approved by the local Ethics Committee as part of a larger project on the clinical and nutritional correlates of urinary parameters in nephrolithiasis. The study was carried out according to the principles of the Declaration of Helsinki. Informed consent was obtained according to Italian law<u>for</u> retrospective studies.

-Acknowledgements

The authors wish to thank <u>Antonio Nouvenne, for important assistance in study design and manuscript drafting.</u> Maurizio Rossi, for the precious statistical consult, and Michele Zenna, for assistance in database management and support in manuscript drafting.

References

- 1. Romero V, Akpinar H, Assimos DG (2010) Kidney stones: a global picture of prevalence, incidence, and associated risk factors. Rev Urol 12(2-3):e86-e96.
- Daudon M, Réveillaud RJ (1984) Whewellite and weddellite: toward a different etiopathogenesis. Nephrologie 5(5):195-201.
- 3. Daudon M, Bader CA, Jungers P (1993) Urinary calculi: review of classification methods and correlations with etiology. Scan Microsc 7(3):1081-1106.
- 3.4. Jiang D, Geng H (2017) Primary hyperoxaluria. N Engl J Med 376(15):e33.
- 4.5. Daudon M, Jungers P, Bazin D (2008) Peculiar morphology of stones in primary hyperoxaluria. N Engl J Med 359(1):100-102.
- 5.6. Sutton RA, Walker VR (1994) Enteric and mild hyperoxaluria. Miner Electrolyte Metab 20(6):352-360.
- 6.7. Massey LK, Liebman M, Kynast-Gales SA (2005) Ascorbate increases human oxaluria and kidney stone risk. J Nutr 135(7):1673-1677.
- 8. Albert A, Tiwari V, Paul E, Ponnusamy S, Ganesan D, Prabhakaran R et al (2018) Oral administration of oxalate-enriched spinach extract as an improved methodology for the induction of dietary hyperoxaluric nephrocalcinosis in experimental rats. Toxicol Mech Methods 28(3):195-204.
- 7.9. Daudon M, Bazin D, André G, Jungers P, Cousson A, Chevallier P et al (2009) Examination of whewellite kidney stones by scanning electron microscopy and powder neutron diffraction techniques. J Appl Cryst 42:109-115.
- 8.10. Park S, Pearle MS (2007) Pathophysiology and management of calcium stones. Urol Clin N Am 34(3):323-334.
- 9.11. Prezioso D, Strazzullo D, Lotti T, Bianchi G, Borghi L, Caione P et al (2015) Dietary treatment of urinary risk factors for renal stone formation. A review of CLU Working Group. Arch Ital Urol Androl 87(2):105-120.
- 10.12. Gambaro G, Croppi E, Coe F, Lingeman J, Moe O, Worcester E et al (2016) Metabolic diagnosis and medical prevention of calcium nephrolithiasis and its systemic manifestations: a consensus statement. J Nephrol 29(6):715-734.

- <u>11.13.</u> Nouvenne A, Ticinesi A, Allegri F, Guerra A, Guida L, Morelli I et al (2014) Twenty-five years of idiopathic calcium nephrolithiasis: has anything changed? Clin Chem Lab Med 52(3):337-344.
- 12.14. Guerra A, Folesani G, Nouvenne A, Ticinesi A, Allegri F, Pinelli S et al (2016) Family history influences clinical course of idiopathic calcium nephrolithiasis: case-control study of a large cohort of Italian patients. J Nephrol 29(5):645-651.
- 13.15. Ticinesi A, Guerra A, Allegri F, Nouvenne A, Cervellin G, Maggio M et al (2018) Determinants of calcium and oxalate excretion in subjects with calcium nephrolithiasis: the role of metabolic syndrome traits. J Nephrol 31(3):395-403.
- 14.16. Ferraro PM, Ticinesi A, Meschi T, Rodgers A, Di Maio F, Fulignati P et al (2018) Short-term changes in urinary supersaturation predict recurrence of kidney stones: a tool to guide preventive measures in urolithiasis. J Urol 200(5):1082-1087.
- 15.17. Werness P, Brown CM, Smith LH, Finlayson B (1985) Equil2: a basic computer program for the calculation of urinary saturation. J Urol 134:1242-1244.
- 16.18. Maurice-Estepa L, Levillain P, Lacour B, Daudon M (2000) Advantage of zero-crossing-point first-derivative spectrophotometry for the quantification of calcium oxalate crystalline phases by infrared spectrophotometry. Clin Chim Acta 298(1-2):1-11.
- <u>17.19.</u> Castiglione V, Jouret F, Bruyère O, Dubois B, Thomas A, Waltregny D et al (2015)
 Epidemiology of urolithiasis in Belgium on the basis of a morpho-constitutional classification.
 Nephrol Ther 11(1):42-49.
- 18.20. Daudon M, Traxer O, Lechevallier E, Saussine C (2008) Epidemiology of urolithiasis. Prog Urol 18(12):802-814.
- 19.21. Pierratos AE, Khalaff H, Cheng PT, Psihramis K, Jewett MA (1994) Clinical and biochemical differences in patients with pure calcium oxalate monohydrate and calcium oxalate dehydrate kidney stones. J Urol 151(3): 571-574.
- 20.22. Parent X, Boess G, Brignon P (1999) Calcium oxalate lithiasis. Relationship between biochemical risk factors and crystalline phase of the stone. Prog Urol 9(6):1051-1056.
- 21.23. Asplin JR, Lingeman J, Kahnoski R, Mardis H, Parks JH, Coe FL (1998) Metabolic urinary correlates of calcium oxalate dehydrate in renal stones. J Urol 159(3):664-668.

- 24. Daudon M, Letavernier E, Frochot V, Haymann JP, Bazin D, Jungers P (2016) Respective influence of calcium and oxalate urine concentration on the formation of calcium oxalate kidney monohydrate or dehydrate crystals. C R Chimie 19:1504-1513.
- 22.25. Manissorn J, Fong-Ngern K, Peerapen P, Thongboonkerd V (2017) Systematic evaluation for effects of urine pH on calcium oxalate crystallization, crystal-cell adhesion and internalization into renal tubular cells. Sci Rep 7(1):1798.
- 23.26. Parks JH, Coward M, Coe FL (1997) Correspondence between stone composition and urine supersaturation in nephrolithiasis. Kidney Int 51(3):894-900.
- 24.27. Singh P, Enders FT, Vaughan LE, Bergstralh EJ, Knoedler JJ, Krambeck AE et al (2015) Stone composition among first-time symptomatic kidney stone formers in the community. Mayo Clin Proc 90(10):1356-1365.
- 25.28. Vaughan LE, Enders FT, Lieske JC, Pais VM, Rivera ME, Mehta RA et al (2019) Predictors of symptomatic kidney stone recurrence after the first and subsequent episodes. Mayo Clin Proc 94(2):202-210.
- 26.29. Ticinesi A, Nouvenne A, Borghi L, Meschi T (2017) Water and other fluids in nephrolithiasis: state of the art and future challenges. Crit Rev Food Sci Nutr 57(5):963-974.
- 27.30. Ticinesi A, Nouvenne A, Maalouf NM, Borghi L, Meschi T (2016) Salt and nephrolithiasis. Nephrol Dial Transplant 31(1):39-45.
- 28.31. Guerra A, Ticinesi A, Allegri F, Nouvenne A, Prati B, Pinelli S et al (2019) Insights about urinary hippuric and citric acid as biomarkers of fruit and vegetable intake in patients with kidney stones: the role of age and gender. Nutrition 59:83-89.
- <u>29.32.</u> Meschi T, Nouvenne A, Ticinesi A, Prati B, Guerra A, Allegri F et al (2012) Dietary habits in women with recurrent idiopathic calcium nephrolithiasis. J Transl Med 10:63.
- 30.33. Ticinesi A, Milani C, Guerra A, Allegri F, Lauretani F, Nouvenne A et al (2018) Understanding the gut-kidney axis in nephrolithiasis: an analysis of the gut microbiota composition and functionality of stone formers. Gut 67(12):2097-2106.
- 31.34. Guerra A, Ticinesi A, Allegri F, Nouvenne A, Pinelli S, Folesani G et al (2016) The influence of maternal and paternal history on stone composition and clinical course of calcium nephrolithiasis in subjects aged between 15 and 25. Urolithiasis 44(6):521-528.

- <u>32.35.</u> Guerra A, Ticinesi A, Allegri F, Nouvenne A, Pinelli S, Lauretani F et al (2017) Calcium urolithiasis course in young stone formers is influenced by the strength of family history: results from a retrospective study. Urolithiasis 45(6):525-533.
- 33.36. Jaggi M, Nakagawa Y, Zipperle L, Hess B (2007) Tamm-Horsfall protein in recurrent calcium kidney stone formers with positive family history: abnormalities in urinary excretion, molecular structure and function. Urol Res 35(2):55-62.
- 34.<u>37.</u> Yamate T, Tsuji H, Amasaki N, Iguchi M, Kurita T, Kohri K (2000) Analysis of osteopontin DNA in patients with urolithiasis. Urol Res 28(3):159-166.
- 35.38. Rimel JD, Kolbach-Mandel AM, Ward MD, Wesson JA (2017) The role of macromolecules in the formation of kidney stones. Urolithiasis 45(1):47-54.
- 36.39. Wollin DA, Kaplan AG, Preminger GM, Ferraro PM, Nouvenne A, Tasca A et al (2018)
 Defining metabolic activity of nephrolithiasis Appropriate evaluation and follow-up of stone formers.
 Asian J Urol 5(4):235-242.
- <u>37.40.</u> Daudon M, Jungers P, Bazin D, Williams jr JC (2018) Recurrence rates of urinary calculi according to stone composition and morphology. Urolithiasis 46(5):459-470.

COD/COM ratio Number (%) Quartiles<u>Intervals</u>	(0-0.25) 287 (62%) (1)	(0.26-0.50) 86 (18%) (2)	(0.51-0.75) 40 (9%) (3)	(0.76-1) 52 (11%) (4)	₽ <u>p*</u>	<u>p**</u> adjusted	Pp*** value for trend (reported only if significant)
Females,%	31	37	30	17	0.104		0.029
Age, years	47 ± 13	46 ± 14	46 ± 15	42 ± 16	0.085		0.016
Weight, kg BMI, kg/m ²	$\begin{array}{c} 75\pm16\\ 25\pm4 \end{array}$	$\begin{array}{c} 74\pm15\\ 26\pm4 \end{array}$	$\begin{array}{c} 75\pm14\\ 25\pm3 \end{array}$	76 ± 12 25 ± 3	0.961 0.468		
Disease duration	5 <mark>([</mark> 1-14)]	5 ([1-15)]	8 ([1-21)]	3 <mark>-[1-19-]</mark>	0.373		
Family history of stones (FHS), %	52	51	60	48	0.716		
Age of onset of kidney stones	38 ± 14	37 ± 14	32 ± 13	32 ± 12	0.007		0.001
Age of onset of kidney stones in patients without FHS	42 ± 14	<i>39</i> ± <i>14</i>	<i>31</i> ± <i>12</i>	35 ± 13	0.003		0.003
Age of onset of kidney stones in patients with FHS	35 ± 13	35 ± 13	<i>33</i> ± <i>14</i>	<i>30</i> ± <i>11</i>	0.315		
Hypertensive, %	21	26	28	15	0.396	<u>0.369</u>	
Recurrents, %	62	68	73	62	0.634*<u>0.</u> 495	<u>0.634</u>	
Stones retained, %	55	50	59	41	0.138* <u>0.</u> 304	<u>0.138</u>	
Bilateral stones,%	45	52	55	43	0.585* 0. 473	<u>0.585</u>	
Extra-Corporeal Shock-Wave Lithotripsy (ESWL), number	0.67 ± 1.28<u>0 [0-1]</u>	0.80 ± 1.710 [0-1]	<u>0.94 ± 1.410 [0-</u> <u>2]</u>	<u>1.42 ± 2.340 [0-</u> 2]	0.022* 0. 017		0.00 4
Stone rate, years	0.79 ± 1.20<u>0.39</u>	0.93 ± 1.40<u>0.62</u>	$0.73 \pm 0.77 0.35$	$0.61 \pm 0.52 \underline{0.46}$	<u>0.948*0.</u>		
-	[0.16-1.00]	[0.20-1.00]	[0.16-0.97]	[0.15-1.00]	361		

Table 1. Clinical characteristics of the patients with idiopathic calcium nephrolithiasis and pure calcium oxalate stone composition included in the study (n=465, 322 M and 143 F), stratified by the <u>quartiles intervals</u> of the ratio of calcium oxalate dihydrate (COD) and calcium oxalate monohydrate (COM) crystals in stones.

Data reported as percentage or median and interquartile range or mean \pm standard deviation. <u>Significant p values (p<0.05) are indicated in bold.</u> *Crude p values obtained with ANOVA (dichotomous variables or continuous variables with normal distribution) or Kruskal-Wallis test (continuous variables with non-normal distribution).

p**-p values adjusted for sex, age, BMI and duration of disease with ANCOVA (only variables requiring adjustment for clinical reasons).

15
16
17
18
19
20
21
21 22
23
24
25
20
26
27 28
28
29
30
31
32
22
33 34 35 36
34
35
36
37
38
39
40
41
42
43 44
45
46
46 47
48
49
50
51
51 52 53 54
52
23
54
55
56
56 57 58
58
59
60

***p for trend values obtained with ANOVA (linear trends). Values are reported only if significant (p<0.05). Significant p values (p<0.05) are indicated in bold.
Significant p values (p<0.05) are indicated in bold.

COD/COM ratio Number QuartileIntervals	(0-0.25) N.287 (1)	(0.26-0.50) N.86 (2)	(0.51-0.75) N.40 (3)	(0.76-1) N.52 (4)	<u>₽p*</u>	p* <u>*</u>	p≛<0.05 Bonferroni test
Creatinine, mg/dl	0.90 ± 0.18	0.86 ± 0.18	0.90 ± 0.18	0.89 ± 0.14	0.285	0.024	
Uric acid, mg/dl	5.38 ± 1.27	5.02 ± 1.09	5.25 ± 0.96	5.41 ± 1.10	0.127	0.101	
Calcium, mg/dl	9.47 ± 0.38	9.46 ± 0.45	9.51 ± 0.45	9.70 ± 0.44	0.003	0.014	(1) and (2) vs (4)
Phosphorus, mg/dl	3.29 ± 0.53	3.29 ± 0.55	3.17 ± 0.60	3.33 ± 0.62	0.600	0.468	
PTH, pg/ml	44 ± 14	43 ± 13	39 ± 12	40 ± 14	0.226	0.282	
25-OH-D, ng/ml	22 ± 13	23 ± 15	25 ± 12	21 ± 13	0.724	0.887	

25-OH-D: 25-hydroxy-vitamin D.

Data reported as mean ± standard deviation. Significant p values (p<0.05) are indicated in bold.

*Crude p values obtained with ANOVA.

** p values adjusted for sex, age, BMI and duration of disease with ANCOVA

Data reported as mean ± standard deviation. p* adjusted for age, duration disease, sex and BMI with ANCOVA. Significant p values (p<0.05) are indicated in bold. 25-OH-D: 25-hydroxy-vitamin D.

Table 3. Urinary chemistry parameters of the patients with idiopathic calcium nephrolithiasis and pure calcium oxalate stone composition included in the study (n=465, 322 M and 143 F), stratified by the quartiles intervals of the ratio of calcium oxalate dihydrate (COD) and calcium oxalate monohydrate (COM) crystals in stones.

COD/COM ratio Number Intervals Quartile	(0-0.25) N.287 (1)	(0.26-0.50) N.86 (2)	(0.51-0.75) N.40 (3)	(0.76-1) N.52 (4)	₽ <u>₽</u> *	p* <u>*</u>	p≛<0.05 Bonferroni test
Volume, ml/24h	1905 ± 702	1916 ± 774	1716 ± 640	1969 ± 757	0.367	0.254	
Creatinine, mg/24h	1525 ± 426	1522 ± 452	1643 ± 455	1580 ± 404	0.367	0.125	
Sodium, mEq/24h	171 ± 61	168 ± 59	164 ± 53	166 ± 59	0.867	0.548	
Potassium, mEq/24h	55 ± 18	53 ± 20	55 ± 15	54 ± 15	0.965	0.952	
Calcium, mg/24h	194 ± 85	256 ± 84	315 ± 122	313 ± 119	<0.0001	<0.0001	(1) vs (2) vs (3) vs (4); (2) vs (3) vs (4)
Hypercalciuria $(\geq 4 \text{ mg/kg/24h}), \%$	12	28	53	48	<0.0001	<0.0001	(1) vs (2) vs (3) vs (4); (2) vs (3) vs (4)
Magnesium, mg/24h	87 ± 29	91 ± 24	101 ± 34	95 ± 29	0.013	0.023	(1) vs (3)
Chloride, mEq/24h	167 ± 62	165 ± 56	167 ± 52	168 ± 59	0.984	0.891	
Phosphorus, mg/24h	842 ± 271	848 ± 239	921 ± 235	855 ± 286	0.360	0.195	
Uric acid, mg/24h	571 ± 164	597 ± 178	595 ± 166	567 ± 144	0.518	0.186	
Oxalate, mg/24h	31 ± 11	31 ± 9	33 ± 8	30 ± 9	0.645	0.457	
Hyperoxaluria (>45 mg/24h), %	7	7	8	4	0.859	0.886	
Sulphate, mmol/24h	21 ± 7	21 ± 7	22 ± 7	20 ± 6	0.554	0.151	
Ammonium, mmol/24h	36 ± 12	37 ± 12	38 ± 10	38 ± 13	0.351	0.494	
Urea, g/24h	24 ± 7	23 ± 7	25 ± 7	23 ± 7	0.661	0.240	
Citrate, mg/24h	578 ± 257	633 ± 249	600 ± 258	591 ± 278	0.394	0.169	
Urine pH, 24h	5.88 ± 0.45	5.93 ± 0.43	5.76 ± 0.43	6.04 ± 0.46	0.016	0.015	(3) vs (4)
Calcium oxalate supersaturation	5.26 ± 2.94	6.75 ± 3.83	8.54 ± 3.42	7.22 ± 3.58	<0.0001	<0.0001	(1) vs (2) vs (3) vs (4) (2) vs (3)
Calcium phosphate supersaturation	0.66 ± 0.61	0.94 ± 0.75	1.17 ± 1.08	1.27 ± 0.83	<0.0001	<0.0001	(1) vs (2) vs (3) vs (4

Data reported as percentage or mean ± standard deviation. Significant p values (p<0.05) are indicated in bold.

*Crude p values obtained with ANOVA.

22

15	
16	
17	
18	
19	
20	
20	<u>**p values adjusted with ANCOVA for sex, age, BMI and duration of disease.</u>
21	<u>**p values adjusted with ANCOVA for sex, age, BMI and duration of disease.</u> p crude, p* adjusted for age, duration disease, sex and BMI with ANCOVA. P<0.05 in bold.
22	r that, r the second seco
20 21 22 23 24	
24	
25	
26	
27	
28	
29	
30	
31	
31	
32	
33	
34	
35	
36	
37	
38	
39	
40	
41	
41	
43	
44	
45	
46	
47	
48	
49	
50	
51	
52	
53	
54	
55	
55	
56	
57	
58	
59	
60	
61	
62	
63	
64	
64 65	
05	

trolsCOD/CO486 $(0-0.25)$ c)N. 287(1) ± 693 1883 ± 68 ± 293 1432 ± 29 ± 57 164 ± 56	$ \begin{array}{c} (0.26-0.50) \\ (0.26-0.50) \\ (0.26) \\ (0.$	$\begin{array}{c} \text{COD/COM} \\ (0.50-1) \\ \text{N.92} \\ \textbf{(3)} \\ \hline 1843 \pm 674 \end{array}$	0.0001	
± 293 1432 ± 29		1843 ± 674	0.0001	
	1426 262		<0.0001	(c) vs (1) vs (2) vs (3)
+ 57 164 + 56	$90 1436 \pm 283$	1454 ± 285	0.351	
± 57 104 ± 50	$5 160 \pm 55$	155 ± 55	0.589	
$\begin{array}{cccc} \pm 19 & 53 \pm 18 \\ \pm 95 & 186 \pm 94 \\ 17 & 12 \end{array}$		53 ± 18 304 ± 92 50	0.014 <0.0001 <0.0001	(c) vs (2) vs (3) (c) vs (2) vs (3)
± 236 805 ± 232	809 ± 227	829 ± 229	<0.0001	(c) vs (1) vs (2) vs (3)
$\begin{array}{c} \pm 29 \\ \pm 157 \\ \pm 260 \\ \pm 16 \\ \pm 11 \\ \pm 10 \\ $	$5 572 \pm 151 \\ 7 626 \pm 251 \\ 20 \pm 5 \\ 36 \pm 11 \\ 30 \pm 9 $	$94\pm 29548\pm 153613\pm 25320\pm 535\pm 1130\pm 9$	0.025 0.231 0.095 0.351 0.687 <0.0001	(c) vs (3) (c) vs (1) vs (3)
5 6	6	4	0.935	
± 0.51 5.93 ± 0.5	$51 5.97 \pm 0.49$	5.95 ± 0.49	0.472	
		8.02 ± 2.93	<0.0001*	(c) vs (2) vs (3) (c) vs (3)
	$\begin{array}{l} \pm 3.04 \\ \pm 0.77 \end{array} \begin{array}{l} 5.67 \pm 3.0 \\ 0.74 \pm 0.77 \end{array}$	$\begin{array}{ll} \pm \ 3.04 & 5.67 \pm 3.00 & 7.18 \pm 2.91 \\ \pm \ 0.77 & 0.74 \pm 0.76 & 1.02 \pm 0.74 \end{array}$	± 3.04 5.67 ± 3.00 7.18 ± 2.91 8.02 ± 2.93 ± 0.77 0.74 ± 0.76 1.02 ± 0.74 1.23 ± 0.75	± 3.04 5.67 ± 3.00 7.18 ± 2.91 8.02 ± 2.93 <0.0001*

Table 4. Comparison of urinary chemistry parameters between patients with idiopathic calcium nephrolithiasis and pure calcium oxalate stone composition (n=465), stratified by the <u>quartiles-intervals</u> of the ratio of calcium oxalate dihydrate (COD) and calcium oxalate monohydrate (COM) crystals in stones, and a group of non-stone forming controls (n=486).

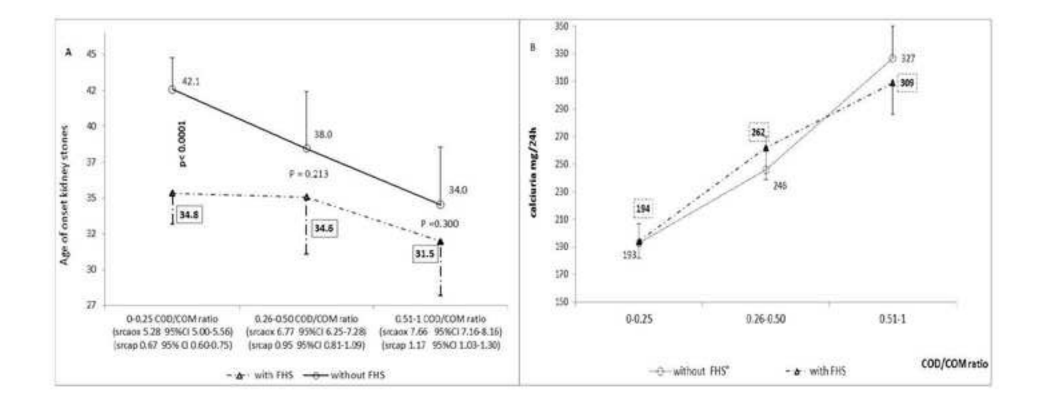
Data reported as percentage or mean \pm standard deviation adjusted for age, sex and BMI with ANCOVA, or mean \pm standard deviation* adjusted for age, sex, volume and BMI with ANCOVA. Significant p values (p<0.05) are indicated in bold.

Table 5. Linear regression model testing the relationship between calcium, oxalate and urine pH with COD/COM ratio in 465 stone formers with idiopathic calcium nephrolithiasis and pure calcium oxalate stone composition.

	β	95%CI	β standardized	р
Calcium, mg/24h	0.124	0.102-0.145	0.464	< 0.0001
Oxalate, mg/24 h	-0.145	-0.370- 0.081	- 0.052	0.209
Urine pH, 24h	6.402	1.347-11.457	0.103	0.013

Significant p values (p<0.05) are indicated in bold

Figure 1. Mean and 95% CI of the age of onset of kidney stones (A) and of calciuria (B) in 461 patients stratified by family history of stones (FHS) (221 without FHS, 240 with FHS) and ratio COD/COM (0-0.25, 0.26-0.50, 0.51-1). A significant trend, for the age of onset of kidney stones, is present in patients without FHS (p = 0.003) increasing the COD/COM ratio and urinary supersaturations, but not in patients with FHS (p = 0.374). Calciuria (B) is not different in patients with and without FHS, p = 0.798.Age of onset of kidney stones is adjusted for BMI and sex. 34 Calciuria by age, sex, duration of disease, body mass indexBMI, sodium, potassium, ammonium and urinary sulfates. Calcium oxalate supersaturation (srcaox) and calcium phosphate supersaturation (srcap) adjusted for BMI, age, volume and sex, no differences for over-saturation between patients with FHS and without FHS



Supplementary Material

Click here to access/download Supplementary Material Supplementary Material.docx Response to reviewers (not for publication)

Click here to access/download Supplementary Material Response to reviewers.doc

Dringer
an A
palgrave macmillan
Bohn Stafleu van Loghum
SS .
Adis
1

Change of authorship request form (pre-acceptance)

Please read the important information on page 4 before you begin

This form should be used by authors to request any change in authorship including changes in corresponding authors. Please fully complete all sections. Use black ink and block capitals and provide each author's full name with the given name first followed by the family name. Please note: In author collaborations where there is formal agreement for representing the collaboration, it is sufficient for the representative or legal guarantor (usually the corresponding author) to complete and sign the Authorship Change Form on behalf of all authors.

	I if avail	(:
Section 1: Please provide the current title of manuscript	(For journals: Please provide the manuscript ID, title and/or DOI if avail	(For books: Please provide the title. ISBN and/or DOI if available.)
ection 1: Please p	For journals: Pleas	For books: Please

lable.)

Manuscript ID no. in case of unpublished manuscript: URDS -D - 49 -	101: (KES-D-19
DOI IN CASE OF PUBLISHED MAINSCRIPT;	
ISBN (for books):	

OF THE MPOJITION; CLINICAL CORRELATES Title: 12 Section 2: Please provide the previous authorship, in the order shown on the manuscript before the changes were introduced. Please indicate the corresponding author by adding (CA) behind the name.

First name(s)	Family name	ORCID or SCOPUS id, if available
1 st author ANI AFLA	6 UE RRA	7102572472 /20001 1D
2 nd author AND REA	TICINESI	Kr143936300 /100PUS 10)
3 rd author FRANCA	ALLEGRI	1559 66595 ADO /SCOPULIAL
4 th author ANTON 10	NOUVENNE	(20 20 307 269 / CORUL 1D)
5 th author 1/1/2 NANA	PINECLI	(21 104 COL 290 / 100 M
6th author Rn (ACIA	ALDE	lai many untur ik ssgr
7 th author 71,21,0 NA	NESCHI	KCO2 MARTAK MANUS INV

SPRINGER NATURE Springer Nature is one of the world's leading global research, educational and professional publishers, created in May 2015 through the combination of Nature Publishing Group, Palgrave Macmillan, Macmillan Education and Springer Science+Business Media.

±

Adis an state	palgrave Springer	Change of authorship request form (pre-acceptance)
Section 3: Please provide a justification authorship? Please refer to the (journal removed on the submitted manuscript.	Section 3: Please provide a justification for change. Please use this section authorship? Please refer to the (journal) policy pages for more information removed on the submitted manuscript.	ı to explain your reasons for changing the authorship of your manuscript, e.g. what necessitated the change in ın about authorship. Please explain why omitted authors were not originally included and/or why authors were
THE CHANGE JOURNAL REG	THE CHANGE WAS NADE ON REQUEST BY JOURNAL REQUIRES THAT A MAXIMUM NUN	Y THE EDITOR IN CHIEF, SINCE THE EDITORIAL POLICY OF THE
REPORTING 1	REPORTING REJULTS FROM JINGLE-CENTER JTUDIES.	TER JTVDIES,
	29 29	
Section 4: Proposed ne adding (CA) behind the	Section 4: Proposed new authorship. Please provide your new authorship adding (CA) behind the name. <u>If the corresponding author has changed, p</u> l	list in the order you would like it to appear on the manuscript. Please indicate the corresponding author by lease indicate the reason under section 3.
First name(s)	ame(s)	Family name (this name will appear in full on the final publication and will be searchable in various abstract and indexing databases)
1 st author ANGELA	ELA	GUERRA
2 nd author ANDREA	REA	TICINECI
3 rd author RRANGA	ICA.	AILEARI
1000	ANTRACO TILVANA	RINELLI
5 th author ROIALIA	L1A	AME
	ANA ANA	MESchi
7 th author		

Please use an additional sheet if there are more than 7 authors.

SPRINGER NATURE Springer Nature is one of the world's leading global research, educational and professional publishers, created in May 2015 through the combination of Nature Publishing Group, Palgrave Macmillan, Macmillan Education and Springer Science+Business Media.

section 5. Author contribution, Actionologiement and Disclosures. Please use this section to provide a new disclosure statement and, if appropriate, actionologies any contribution and new authors and experiments of the point of the control of an operation and the analysis of the approximation of the control of any array with a controllation and new authors and experiments of the control of any array with a controllation and new authors and experiments of the control of any array with a controllation and new authors and experiments of the control of any array with a controllation and new authors and experiments of the control of any array with a controllation any reas with a controllation any reason of the control of a controllation any reason of the control of a controllation any reas with a control of a controllation any reas and reach and reas and	SPRINGER NATURE Springer Nature is one of the world's leading global research, educational and professional publishers, created in May 2015 through the combination of Nature Publishing Group, Palgrave Macmillan, Macmillan Education and Springer Science+Business Media.
--	---

Change of authorship request form (pre-acceptance)

Section 6: Declaration of agreement. All authors, unchanged, new and removed must sign this declaration.

(NB: Please print the form, sign and return a scanned copy. Please note that signatures that have been inserted as an image file are acceptable as long as it is handwritten. Typed names in the signature box are unacceptable.) * Please delete as appropriate. Delete all of the bold if you were on the original authorship list and are remaining as an author.

	First name	Family name		Signature	Affiliated institute		Date
1 st author	ANGELA	GUERRA	I agree to the proposed new authorship shown in section 4 / and the addition/removal*of my name to the authorship list.	Mynne	UNIVERJITY HOIPITAL OF PARMA		6/24/2018
2 nd author	ANDREA	TICINES 1	I agree to the proposed new authorship shown in section 4 / and the addition/removal*of my name to the. authorship list	In Future in	Mutulaine OF PARMA HOJPITAL 6/28/2019	JPI TAL	6/28/2019
3 rd author	FRANCA	ALLEGRI	I agree to the proposed new authorship shown in section 4 / and the _ addition/removal*of my name to the _ authorship list	Ally France	Ally FLOWERD ITY HOSPITAL 6/28/2019	UPITAL	6/28/2019
4 th authors	JILYANA	PINELLI	I agree to the proposed new authorship shown in section 4 / and the addition/removal*of my name to the authorship list-	L'Elec Ruel	Deve River UNIVERJITY HOSPITAL 6/28/2019	PEIdro	6/28/2018
5 th author	Rotalia	A LO E	I agree to the proposed new authorship shown in section 4 / and the addition/removal*of my name to the authorship list.	43-duche	Hoollolle UNIVERITY HOLPITAL 6/28/2019	JPITA2	6/28/2013
6 th author	TIZIANA	MESCHI	I agree to the proposed new authorship BOLELL UNIYERITY HOIPITAL 6/28/2019 addition/removal*of my name to the of PARMA 0. PARMA	backle	UNIVERSITY HO	JUP I EAL	6/28/2019
7 th author	ANTONIO	NOUVENNE	I agree to the proposed new authorship shown in section 4 /and the addition/removal*of my name to the authorship list.	Choun	Oldonu UNIVERSITY HOSPITAL 6/28/2019	JUP ITA2	6/28/2019
Please us	e an additional sheet	Please use an additional sheet if there are more than 7 authors.	authors.				

SPRINGER NATURE Springer Nature is one of the world's leading global research, educational and professional publishers, created in May 2015 through the combination of Nature Publishing Group, Palgrave Macmillan, Macmillan Education and Springer Science+Business Media. 1

Important inform, inform, information. Please read. Preservation inform the form, information or number for orander the information or understate a further investigation, if appropriate, before making a final decision. These requires the content information or numerities a further investigation, if appropriate, before making a final decision. The sing this decisation, and that only authors guarantee that the corder of the authors are information to the work. The sing this decisation, if appropriate, before making a final decision. The sing this decisation, if appropriate, before making a final decision. The sing this decisation is number of the authors guarantee that the corder of the author of the number of the numbe		otance)
 Please return this form, fully completed, to Springer Nature, We will consider the information you have provided to decide whether to approve the proposed change in authorship. We may choose to contact your institution for more information or undertake a further investigation, if appropriate, before making a final decision. By signing this declaration, all authors guarantee that the order of the authors are in accordance with their scientific contribution, if applicable as different conventions apply per decipine, and that only authors have been added who made a meaningful contribution to the work. Bease note, we cannot investigate or mediate any authorship disputes. If you are not able to return a fully completed form within 30 days of the date that it was sent to the author requesting the change, we may have to withdraw your manuscript. We many reducting these who have to with an uscripts were authorship has not been agreed by all authors (including those who you wigh to be removed) ye must refer the matter to your institution for investigation. These inform us if you need to ot bits. If you are not able to return a fully completed form within 30 days of the date that it was sent to the author requesting the change, we may have to withdraw your manuscript. We not publish manuscripts where authorship has not been agreed by all authors (including these who have been removed). Incomplete forms will be rejected. 	Important information. Please read.	
 by signing this declaration, all authors guarantee that the order of the authors are in accordance with their scientific contribution, if applicable as different conventions apply per discipline, and that only authors have been added who made a meaningful contribution to the work. Please note, we cannot investigate or mediate any authorship disputes. If you are unable to obtained agreement from all authors (Including those who you wish to be removed) yo must refer the matter to your institution(s) for investigation. Please inform us fryou need to do this. If you are not able to return a fully completed form within 30 days of the date that it was sent to the author requesting the change, we may have to withdraw your manuscript. We cannot publish manuscripts where authorship has not been agreed by all authors (Including those who have been removed). Incomplete forms will be rejected. Incomplete forms will be rejected. 		change in authorship.
 Please note, we cannot investigate or mediate any authorship disputes. If you are unable to obtained agreement from all authors (Including those who you wish to be removed) you must refer the matter to your institution(s) for investigation. Please inform us if you need to do this. If you are not able to return a fully completed form within 30 days of the date that it was sent to the author requesting the change, we may have to withdraw your manuscript. W cannot publish manuscripts where authorship has not been agreed by all authors (including those who have been removed). Incomplete forms will be rejected. 		nventions apply per
 If you are not able to return a fully completed form within 30 days of the date that it was sent to the author requesting the change, we may have to withdraw your manuscript. We cannot publish manuscripts where authorship has not been agreed by all authors (including those who have been removed). Incomplete forms will be rejected. 		ish to be removed) yo
• Incomplete forms will be rejected.		/ your manuscript. We
FIGURE And TABLE Surger Nature is one of the word's leading global research, educational and professional publishers, created in Nay 2005.	Incomplete forms will be rejected.	
Ender Nature 5 on of the word's leading global research, educational and professional publishers, created in May 2015 Thrush the combination of Nature Science Apartments and Software Stence Abatines Averalian		
SPRINGER NATURE Springer Nature is one of the world's leading global research, educational and professional publishers, created in May 2015 through the combination of Nature Publishine Storus. Plarave Macmillan, Nacmillan, Nacmillan, Ratrost Media.		
Ender Nature Springer Nature is one of the world's leading global research, educational and professional publishers, created in May 2015 Through the combination of Nature Publishing Group, Palerave Macrillian, Macrillian Education and Springer Science+Business Media.		
Ender Nature Springer Nature is one of the world's leading global research, educational and professional publishers, created in May 2015 through the combination of Nature Publishing Group. Palerave Macmillan. Macmillan Education and Science+Business Media.		
SPRINGER NATURE Springer Nature is one of the world's leading global research, educational and professional publishers, created in May 2015 through the combination of Nature Publishing Group, Palerave Macmillan, Macmillan, Education and Socience+Business Media.		
SPRINGER NATURE Springer Nature is one of the world's leading global research, educational and professional publishers, created in May 2015 through the combination of Nature Publishing Group, Palerave Macmillan, Macmillan Education and Socinger Science+Business Media.		
	SPRINGER NATURE Springer Nature is one of the world's leading global research, educational and professional publishers, created in May 2015 through the combination of Nature Publishing Group. Palerave Macmillan. Macmillan Education and Socinger Science+Business Media.	