

Autosomal Dominant Polycystic Kidney Disease

Carolynn DeBenedectis September 15, 2005 BIDMC





 47 year old female with a history of hypertension, hepatomegaly, early satiety, and some abdominal discomfort.

 Family history is significant for a father who died of renal failure.

 CT abdominal/pelvis was performed for a chief complaint of abdominal discomfort



CT Abdominal/Pelvis with IV contrast



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DEBILITIES/GANG & 10	[F]	
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W342





Small sliding hiatal hernia



Stomach protruding into the left hemithorax

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Hiatal Hernias

 Definition – proximal portion of the stomach slips thru the esophageal hiatus and into the thorax

- <u>Sliding type</u> the gastroesophageal (GE) junction slides into the mediastinum above the diaphragm
 - Most common type

Paraesophageal type – the GE junction is in the normal location but the gastric fundus "rolls" into the mediastinum

Can become infarcted, which is a surgical emergency



Specific CT findings in the liver Multiple fluid-filled cysts are seen throughout and largely replacing the right lobe of the liver

There is diffuse hypertrophy of the liver parenchyma, with lateral displacement of the stomach wall





CT findings in the liver

The largest liver cyst



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CT findings in the liver

Se:2 Im:60

Liver extending into the pelvis





CT findings in the liver

Note the left lobe ^{Se:2} m:43 of the liver is partially clear of cysts (note the amount of liver parenchyma)



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Specific CT findings in the kidneys

 Multiple large cysts are seen replacing bilateral kidneys.

 Remaining renal parenchyma enhances normally

- There is symmetric excretion of contrast
- No enhancement of any of the visualized cysts
 - Cyst enhancement would indicate cyst rupture, hemorrhage, or infection



Se:2 Im:47

Asymmetric cystic involvement of the kidneys

[A]

[L]

C+ ABD/PEL DEBILITIES/GANG & 10... [P] C56 W342 PACS, BIDMC More cysts, less Fewer cysts, more parenchyma parenchyma

Wide variation in the size of the cysts

[R] [L] C+ ABD/PEL C56 DEBILITIES/GANG & 10... [P] W342 PACS, BIDMC Large cyst Small cyst



Kidney Cysts

 Fluid-filled sacs arising from a dilatation in any part of the nephron or collecting duct.

+Renal cysts are quite common.

The incidence of renal cysts increases with age, most people over the age of 60 have one or more simple renal cyst.



Differential Diagnosis

<u>Nongenetic</u>

- Acquired disorders
 - Simple renal cysts
 - Cysts of the renal sinus
 - Acquired cystic kidney disease (i.e. CRF)
 - Multilocular cysts
 - Hypokalemia-related cysts

Developmental disorders

- Medullary sponge kidney
- Multicystic dysplastic kidney
- Pyelocalyoceal cysts

<u>Genetic</u>

- Autosomal-dominant
 - Autosomal-dominant polycystic kidney disease (ADPKD)
 - Tuberous sclerosis complex
 - 🔹 von Hippel-Lindau disease
 - Medullary cystic disease
 - Glomerulocystic kidney disease
- Autosomal-recessive
 - Autosomal-recessive polycystic kidney disease (ARPKD)
 - Nephronophthisis
- X-linked
 - Orofaciodigital syndrome, type I



Diseases that have renal cysts and liver cysts

ADPKD

- + Adult onset (4th or 5th decade)
- + Large "bumpy" kidneys (due to renal cysts)
- Hepatic cysts
- + Autosomal dominant inheritance

ARPKD

- + Childhood onset (at birth , and die young)
- + Large, smooth kidneys (due to numerous small cysts)
- Hepatic cysts, congenital hepatic fibrosis, and biliary duct ectasia
- + Autosomal recessive inheritance
- + Tuberous sclerosis complex
 - Renal cysts, liver cysts, and liver angiomylipomas
- Nephronophthisis
 - + Liver fibrosis, renal cysts, and hepatic cysts



IMAGING CHARACTERISTICS OF THE MOST COMMON RENAL CYSTIC DISEASES

Kidney Size	Cyst Size	Cyst Location	Liver
Normal	Variable (mm-10 cm)	AL	Normal
Most often small, sometimes large	0.5–2 cm	AL	Normal
Normal or slightly enlarged	mm	Precalyceal	Normal (most often)
Enlarged	Variable (mm-10 cm)	AL	Cysts (most often)
Enlarged	mm increase with age	AL	CHF
Small	mm-2 cm (when present)	Medullary	Normal
	Kiciney Size Normal Most often small, sometimes large Normal or slightly enlarged Enlarged Enlarged Small	Kiciney Size Cyst Size Normal Variable (mm-10 cm) Most often small, sometimes large 0.5-2 cm Normal or slightly enlarged mm Enlarged Variable (mm-10 cm) Enlarged mm increase with age Small mm-2 cm (when present)	Kidney Size Cyst Size Cyst Location Normal Variable (mm-10 cm) All Most often small, sometimes large 0.5-2 cm All Normal or slightly enlarged mm Precatyceal Enlarged Variable (mm-10 cm) All Enlarged mm increase with age All Small mm-2 cm (when present) Medullary

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Imaging of Renal Cysts

Plain abdominal radiography + IVP Radionucleotide imaging + Ultrasound Computed tomography + MRI Renal angiography Retrograde pyelography



Best Modalities for assessing Renal Cysts

 U/S is best for differentiating solid masses from cystic ones

CT scan is best for finding small cysts and assessing for cystic rupture, infection, or hemorrhage (due to the use of contrast to get differential enhancement of cysts)

 MRI is useful for further characterizing renal cysts and detecting renal neoplasms



More Information About the Case

 Further research revealed that this patient's father died of renal failure secondary to Autosomal Dominant Polycystic Kidney Disease

 This patient had been diagnosed with and worked up for ADPKD at an outside hospital previously



ADPKD

- Autosomal dominant hereditary condition characterized by multiple expanding cysts of both kidneys that ultimately destroy the renal parenchyma and cause renal failure
- 100% penetrance and affects 1 of every 400-1000 live births
- The majority of cases are caused by a mutation in the PKD1 gene on chromosome 16, which produces polycystin 1 (involved in cell-to cell interactions)
 The remainder of cases are caused by a mutation in PKD2 on chromosome 4, which produces polycystin 2 (less severe)
- Has variable expression and does not produce symptoms until adult life (4th or 5th decade of life)



PKD1

PKD1 vs. PKD2

Earlier
 presentation
 (average age 35)

 Earlier end-stage renal failure (average age 60)

 Earlier death (average age 70) <u>PKD2</u>
Later presentation (average age 61)

 Later end-stage renal failure (average age 74)

Later death (average age 75)



 Cysts may also occur within the liver, pancreas, and spleen

+ Cysts are more common in the liver and tend to be less common in the pancreas and spleen

Hepatic failure does not tend to occur despite extensive infiltration of the liver with cysts

+ However, cysts in the liver and pancreas can lead to obstructive jaundice

+ Circle of Willis aneurysms are present in up to 41% of patients that undergo cerebral angiography



ADPKD and imaging

- The imaging appearance varies with the severity of the disease
- Full-blown disease shows the presence of bilaterally enlarged kidneys with numerous cysts of varying sizes
- Ultrasound demonstrates cysts in the adolescent and young adult, who are usually asymptomatic
- Urography may be normal
- Ultrasound and CT are most sensitive
- MRI is useful for differentiating between simple cysts, hemorrhagic cysts, and neoplasms when CT and U/S are equivocal



Ultrasonographic Diagnostic Criteria for ADPKD

- The absence of cysts before age 30 does not rule out the diagnosis of ADPKD
- False negative rate is inversely proportional to age
- If U/S diagnosis remains equivocal, CT should be used next (more sensitive for small cysts)



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Example of the cystic involvement of the kidneys

Classic Renal Imaging Findings in ADPKD

 Note the marked asymmetry in the number and size of cysts between the 2 kidneys

 CT is more sensitive at detecting small cysts than U/S



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Small kidney with minimal cystic involvement Large kidney with multiple cysts of varying size



Classic Hepatic Imaging Findings in ADPKD





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cysts

 Note the cystic involvement of the liver

The presence of liver cysts helps to establish the diagnosis of ADPKD, especially if kidney involvement is mild

Massive cystic disease of the liver can lead to early satiety, abdominal /diaphragmatic hernia, supine dyspnea, and rarely jaundice.



Prevalence of asymptomatic ICA in ADPKD is 8% (1.2% in the general population), increases to 16% with a family history of ICA

Intracranial aneurysms (ICAs) and ADPKD

ICA on the anterior communicating artery

 ICA rupture entail a mortality of 30-50% (results in SAH)

 First-line diagnostic procedure for ICA rupture is CT



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Indications for Screening for ICA in ADPKD



 Screening in ADPKD for ICA can be achieved with either MRA or spiral CT angiography

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Treatment

Treatment is largely supportive
The associated hypertension can be managed medically
Renal failure can be managed with dialysis
Kidney transplantation is the only way to "cure" the renal disease



Summary

ADPKD is a disease with onset in adulthood

 ADKPD is associated with enlarged, cystic kidneys, hepatic cysts, pancreatic cysts, splenic cysts, and ICAs.

 U/S is the best imaging method for detecting renal cysts, however, CT may be more sensitive for small renal cysts

 It is important to screen ADPKD patients for ICA with spiral CT angiography or MRA in accordance with the appropriate indications.



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