Tomographic aspects of xanthogranulomatous pyelonephritis and related complications

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The authors present their experience involving seven patients with histopathologic diagnosis of xanthogranulomatous pyelonephritis who were submitted to preoperative computed tomography (CT). The results are the following: a) stones (86 percent of the cases), b) increase in renal volume, c) hydronephrosis, d) density measurements (from 14 to 29 HU), e) enhancement found in all cases, f) extrarenal involvement (all cases). CT has shown to be a reliable method in characterizing xanthogranulomatous pyelonephritis and extrarenal involvement.


INTRODUCTION

Xanthogranulomatous pyelonephritis (XGP) is an atypical variety of a chronic renal infection which is usually unilateral and frequently associated with urinary obstruction and stones. Before ultrasonography (US) and computed tomography (CT) preoperative diagnosis was not certain in 44-64 percent of all cases, due especially to uncharacterized symptoms, and laboratory exams that showed no alterations. Radiographic exams such as excretory urogram (EU) and antegrade pyelography (APG) allow the diagnosis of a renal mass with or without functional elimination; however, they do not indicate the degree of inflammation caused by the disease. Although it is relatively rare (820 cases described up to 1993), the tomographic aspects have already been described and have been said by some authors to be pathognomonic.

Therefore, it is possible not only to diagnose the disease, but also to evaluate its extrarenal extension and to differentiate the focal and the diffuse forms, allowing for precise surgical planning. Treatment consists of total nephrectomy in the diffuse form and a partial nephrectomy when only a limited area is affected.

The purpose of our study is to present seven cases, describing their tomographic aspects and frequency, so as to help differentiate this disease from others.

MATERIAL AND METHODS

We reviewed CT studies of seven patients with an histopathologic diagnosis of XGP made between 1991 and 1993. Six of these seven patients were female and one was male, with ages ranging from 13 to 71 (mean age - 41 years).
Table 1
Patients according to the presence and the site of stones, degree of hydronephrosis, and measurements of the less dense area of renal lesion

<table>
<thead>
<tr>
<th>Patient</th>
<th>Stones</th>
<th>Coraliform Stones</th>
<th>Site</th>
<th>Degree of Hydronephrosis</th>
<th>Density (UH)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AR</td>
<td>+</td>
<td>-</td>
<td>pyeloureteral junction</td>
<td>IV</td>
<td>21-25</td>
</tr>
<tr>
<td>BFS</td>
<td>+</td>
<td>+</td>
<td>pyelo-calix</td>
<td>IV</td>
<td>17-29</td>
</tr>
<tr>
<td>FLN</td>
<td>+</td>
<td>-</td>
<td>parenchyma/ureteral</td>
<td>I</td>
<td>14</td>
</tr>
<tr>
<td>JCM</td>
<td>+</td>
<td>+</td>
<td>pyelo-calix</td>
<td>IV</td>
<td>14</td>
</tr>
<tr>
<td>LACC</td>
<td>+</td>
<td>-</td>
<td>ureteral</td>
<td>IV</td>
<td>22-24</td>
</tr>
<tr>
<td>MLF</td>
<td>+</td>
<td>+</td>
<td>pyelo-calix</td>
<td>IV</td>
<td>-</td>
</tr>
<tr>
<td>MVS</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>IV</td>
<td>23-25</td>
</tr>
</tbody>
</table>

RESULTS

The exams were carried out using the Somatom DR (Siemens Medical System), with 8 mm wide cuts and an 8 or 16 mm increment, before and after one endovenous injection of 100-150 ml iodine hydrosoluble contrast, measuring densities before and after contrast injection.

The tomographic parameters assessed were:

a) presence of stones and their sites.
b) renal dimensions. We considered the kidney "enlarged" when its longitudinal diameter exceeded 13 cm.
c) degree of hydronephrosis, classification I-IV.
d) density measured in low-density areas.
e) presence of enhancement after endovenous contrast injection.
f) extra-renal damage (perirenal, pararenal and abdominal sites), with presence of heterogeneity or blurring of peritoneal fat due to hyperdense thick layers, mass images with the aspect of soft tissue or circundating liquid collections. Heterogeneity and asymmetric enlargement of paravertebral muscle and or subcutaneous cellular tissue were considered indications of abdominal involvement.

Unilateral renal malfunction was evident in all patients; left malfunction in 5 (71 percent) and right in 2 (29 percent). In six cases, kidney enlargement was observed with grade IV hydronephrosis (86 percent), while in only one case (14 percent) there was a volumetric decrease with hydronephrosis grade I (Fig. 5). In six (86 percent), there were stones in the collecting system; in two (28 percent), there were stones in the renal parenchyma; and only one patient (14 percent) did not present any stones. In three of the six patients with stones in the collecting system, the stone was coraliform (Table 1).

The density of the low-density component of the lesion, measured from 14 to 29 UH, with no fat or gas-type densities being observed. After the intravenous injection of contrast, there was a peripheral enhancement of the affected kidney (Fig. 10), as well as extrarenal damage.

Figure 1 - Increase of left kidney volume showing peripheral enhancement and dilation of the collecting system with central calcification. This patient showed a normal inferior segment of the kidney.

Figure 2 - Left renal site occupied by a large collection of multiloci liquid which extended to the perirenal posterior pararenal and dorsal wall sites. A left enlargement of the psoas muscle can be observed.
Table 2
Patients according to presence and site of extrarenal involvement

<table>
<thead>
<tr>
<th>Patient</th>
<th>perirenal</th>
<th>posterior pararenal</th>
<th>abdominal wall</th>
</tr>
</thead>
<tbody>
<tr>
<td>AR</td>
<td>+</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>BFS</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<tr>
<td>FLN</td>
<td>+</td>
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</tr>
<tr>
<td>JCM</td>
<td>+</td>
<td>+</td>
<td>-</td>
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<tr>
<td>LACC</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>MLF</td>
<td>+</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>MVS</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
</tbody>
</table>

DISCUSSION

XGP was described in 1916 by Schlogenhaufer, who called it staphylomycosis due to its resemblance to actinomycosis and to the presence of staphylococci. The term XGP was used by Oberling, in 1935, due to its yellow color and to its granulomatous characteristic. XGP can be defined as a rare form of a chronic renal infection with diffuse or segmented damage, characterized by the destruction of parenchyma and the accumulation of macrophages containing lipids associated with intrarenal, peripylcal and perirenal fibrosis. There is an association of XGP with a lipomatous degeneration that substitutes necrotic renal tissue, which originates in the renal sinus and spreads to the perirenal site, where it is encapsulated by the renal fascia.

XGP can be classified as diffuse, segmented or focal, all showing similar clinical manifestations. In the diffuse form the kidney is usually enlarged and distorted by the presence of multiple liquid collections which correspond to dilated calix or parenchymal destruction foci filled with purulent material. These alterations are frequently associated with extrarenal extension of the inflammation and perisinus fibrosis. Segmental XGP is formed by two or more inflammatory masses that communicate with a calix or a duplicated unit, which usually is not dilated, although it does contain small obstructive stones close to the papilla. The focal form shows the same alterations as the segmental form, although there is no communication with the collecting system and with a so-called focal xanthogranulomatous inflammation. In our study, we observed six cases of the diffuse form and one case of the segmental form; all were treated with complete nephrectomy.

XGP pathogenesis is not yet fully understood, although the infection and obstruction of the urinary tract (13 are considered to be essential for this to occur). Other factors that seem to contribute to its installation are the alteration of lipid metabolism, immunity and lymphatic drainage, in addition to diabetes and renal papillary...
The incidence of XGP is variable, occurring at any age, with the description of a case at 48 days of age; however, it is more frequent in women in their fifties and sixties.

Clinical manifestations are colic pain in the lower back (84 percent), fever (55 percent), macroscopic hematuria (24 percent), weight loss (10 percent) and a palpable mass in the lower back (39 percent); all these symptoms may be acute or subacute and occur without a previous history of urinary tract infection.

Laboratory exams may be as nonspecific as the clinical manifestations, with a normal uroculture in 40 percent of the patients, especially due to the renal exclusion found in these cases. On the other hand, positive uroculture more frequently indicates the presence of E. coli (49-67 percent), P. mirabilis (26-31 percent), S. aureus (19 percent) and P. aeruginosa (20 percent).

Other laboratory alterations found are increased ESR (100 percent), leukocytosis (70 percent), a decrease in hematocrit (67 percent), and creatinine alterations (46 percent).

Imagery diagnosis such as EU, APG, US and angiography suggest XGP, although there are some limitations which are especially due to the extrarenal extension of the disease.

EU may show renal exclusion (71-96 percent), nephromegaly (100 percent), lithiasis (71-82 percent) and focal dilation of the collecting system (9 percent). APG is used for cases involving renal exclusion, and demonstrates the level of obstruction, which is more frequently located at the ureteropyelic junction.

Ultrasonography has, as its most important limitation, an inadequate retroperitoneal assessment. It is interesting to mention that some authors had difficulties in describing the pyelocalix dilation in XGP patients. On the other hand, it is very easy to diagnose the presence of liquid collections and renal stones, which even allows for a suggestion of the type of liquid.

There are two described ultrasonographic patterns in diffuse XPG. In most cases the kidney is enlarged, with smooth contours, and architecturally disarrayed due to the presence of several liquid masses which have a tendency to join and, in these cases, characterizes to a dilated collecting system with debris, corresponding to purulent material. Stones are frequently identified, except in those cases in which there is peripyletic fibrosis that may mimic the posterior acoustic shadows produced by calcifications.

In another less frequent and non-specific form of presentation, the renal site is noted to have been occupied by an extensive heterogenic formation that is predominantly liquid. The segmented form is very similar to the first form described, although part of the affected kidney is preserved. Difficulty in diagnosis occurs in the focal form, because its presentation in the ultrasonogram is often indistinguishable from a solid or cystic kidney neoplasm or even from an abscess.

Finally, it is important to remember that angiography, when used to distinguish an XGP from a renal carcinoma, only does so in 25-75 percent of the cases, those with a hypovascular (50 percent) or an avascular (25 percent) pattern, different from what is usually found in the hypernephroma. This presentation depends on the quantity of neovascularization of granulation tissue.
Magnetic resonance has not shown any significant advances up to this moment in the diagnosis of XGP, and is not able to provide any information beyond that given by CT (9). However, it is important to remember that the low toxicity of the paramagnetic contrast, as well as the possibility of receiving orthogonal images in three planes, may make MR useful for those patients who are allergic to iodine or need more detailed surgical planning.

In our study, CT showed a constant pattern similar to those described by other authors (10, 11, 12). We observed a diffuse increase in renal volume in most of the patients (6-7), except for one who presented the segmented form of the disease. In this case, besides being enlarged, the kidney kept its usual form, and peripheral enhancement was noted that may correspond to compressed residual parenchyma or to a capsule of inflammatory tissue (Fig. 1).

It is important to observe that the radial distribution of the liquid cavities which were found in the kidney resembles the distribution of the collecting system and has the aspect of a "bunch of grapes" or a "bear claw" as described by some authors (12, 13) (Figs. 2 and 3).

The density measures obtained (14-29 UH) do not differ significantly from those found by Goldman et al. (10-15 UH) (11), if we remember that all these rates indicate a thick liquid and that different calibrations of the equipment used may lead to small differences. On the other hand, we emphasize that in no case did we find densities similar to fat, as was suggested by Acunas et al., who considered this infrequent (1).

It was not surprising to find a frequent extrarenal extension of XGP that was drained in all studied cases and easily identified in several retroperitoneal sites and dorsal wall (Fig. 4). The importance in defining extrarenal damage resides in adequate surgical planning, thus avoiding any undesirable fistulae (12, 13), which has even suggested a classification of the XGP through CT in: State I when the disease is restricted to renal parenchyma, in State II when there is perirenal involvement, and in State III when there is peri and pararenal involvement. Therefore, it may be interesting to define a State IV when there is damage to the abdominal wall.

Despite the characteristic aspect of the XGP in the CT, a differential diagnosis with hypernephroma, renal tuberculosis (Tb), and pyohydronephrosis must be made. Hypernephroma may be similar to XGP in its focal form when studied by the US and CT, and, even more, 50 when in its cystic form, although this is relatively rare in adults. In these cases, a more crude type calcification with a coraliform aspect may help in the diagnosis of XGP (13, 16). When this is not possible and there are no retroperitoneal ganglia in the tomographic exam (rare in XGP) or other malignity signs (e.g. hepatic metastases), selective arteriography may be useful if it demonstrates an avascular pattern therefore ruling out the blastomatose origin of the process. On the other hand, a hypo or hypervascular pattern may be found in XGP and hypernephroma (15).

Renal Tb usually evolves with a decrease in renal volume and with calcifications somewhat different from XGP which are more pointed in shape. When XGP leads to renal reduction, differential diagnosis with Tb may be extremely difficult as in one of the studied cases (Fig. 5).

Finally, pyohydronephrosis is considered by many authors to be an initial stage or a precursor of XGP, and has a very similar pattern.

Other differential diagnoses, which are more rare that should be remembered, are lipoma and liposarcoma, in which XGP coexists with an intense gradual replacement of granulation tissue for adipose tissue of an unknown origin (1).

CONCLUSION

Although some authors suggest that preoperative diagnosis through imagery exams should not be done in any case, but cannot justify this (3), we believe that, according to the results found in our study and literary review, that computed tomography is a method which allows identification of very characteristic signs indicating a preoperative diagnosis of XGP. These signs are: a) increase in renal volume; b) hydronephrosis; c) renal/ureteral lithiasis; c) collections of thick fluid; d) peripheral enhancement after contrast injection; f) extrarenal involvement of perirenal, posterior pararenal and abdominal wall sites.
RESUMO

Os autores apresentam a sua experiência em 7 pacientes com diagnóstico anátomopatológico de Pielonefrite Xantogranulomatosa (PXG) submetidos a tomografia computadorizada (TC) pré-operatória. Os parâmetros estudados e seus resultados foram: a) presença de cálculos em 86% dos casos, b) volume renal frequentemente aumentado, c) hidronefrose, d) medidas de densidade variando entre 14 e 29 HU, e) presença de realce em todos os casos, f) comprometimento extra-renal, também presente em todos os casos. A TC demonstrou ser um método bastante fidedigno na caracterização da PXG e sua extensão extra-renal.

REFERENCES