When to worry about incidental renal and adrenal masses

Greater use of imaging has led to a corresponding rise in the detection of renal and adrenal incidentalomas—and left many primary care physicians unsure of what to do about the masses they’ve found.

**CASE** Jane C, a 76-year-old patient, reports lower abdominal discomfort and increased bowel movements. Her left lower quadrant is tender to palpation, without signs of a surgical abdomen, and vital signs are normal. Laboratory studies are also normal, except for mild anemia and a positive fecal occult blood test. Abdominal and pelvic computed tomography (CT), with and without contrast, are negative for acute pathology, but a 1.7-cm lesion is found in the upper pole of the left kidney. What is your next step?

Renal or adrenal masses may be discovered during imaging studies for complaints unrelated to the kidneys or adrenals. Detection of incidentalomas has increased dramatically, keeping pace with the growing use of ultrasonography, CT, and magnetic resonance imaging (MRI) for abdominal, chest, and back complaints.¹

Family physicians can evaluate most of these masses and determine the need for referral by using clinical judgment, appropriate imaging studies, and screening laboratory tests. In the pages that follow, we present a systematic approach for evaluating these incidentalomas and determining when consultation or referral is needed.

**Incidental renal masses are common**

Lesions are commonly found in normal kidneys, and the incidence increases with age. Approximately one-third of individuals age 50 and older will have at least one renal cyst on CT.²

Most incidental renal masses are benign cysts requiring no further evaluation. Other possibilities include indeterminate or malignant cysts or solid masses, which may be malignant or benign. Inflammatory renal lesions from infection,
Classification of renal cysts—
not based on size
Cysts are the most common adult renal masses. Typically they are unilocular and located in the renal cortex, frequently extending to the renal surface. Renal function is usually preserved, regardless of the cyst’s location or size. Careful examination of adjacent tissue is essential, as secondary cysts may form when solid tumors obstruct tubules of normal parenchyma. Cystic lesions containing enhancing soft tissue unattached to the wall or septa likely are malignant.

The Bosniak classification system, with 5 classes based on CT characteristics (TABLE 1), is a useful guide for managing renal cystic lesions. Size is not an important feature in the Bosniak system; small cysts may be malignant and larger ones benign. Small cysts may grow into larger benign lesions, occasionally causing flank or abdominal pain, palpable masses, or hematuria.

Simple cysts. Renal cysts that meet Bosniak class I criteria can be confidently labeled benign and need no further evaluation (FIGURE 1). Simple renal cysts on CT have homogenous low-attenuating fluid and thin nonenhancing walls without septa.

On ultrasound, simple renal cysts show spherical or ovoid shape without internal echoes, a thin smooth wall separate from the surrounding parenchyma, and posterior wall enhancement caused by increased transmission through the water-filled cyst. The likelihood of malignancy is extremely low in a renal cyst that meets these criteria, which have a reported accuracy of 98% to 100%.

Thus, no further evaluation is required if an obviously benign simple cyst is first noted on an adequate ultrasound. Inadequate ultrasound visualization or evidence of calcifications, septa, or multiple chambers calls for prompt renal CT.

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**TABLE 1**

In the Bosniak system, CT results guide incidental renal cyst management

<table>
<thead>
<tr>
<th>Category</th>
<th>Imaging features</th>
<th>Clinical management</th>
<th>Likelihood of malignancy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Class I</td>
<td>Simple cyst, benign</td>
<td>Low-attenuating (0-20 HU) fluid; hairline-thin smooth wall without septa, calcifications, or enhancing soft tissue</td>
<td>Renal CT if symptoms occur</td>
</tr>
<tr>
<td>Class II</td>
<td>Minimally complex cyst, probably benign</td>
<td>Single or few thin (&lt;1 mm) septa; may be minimally calcified; perceived but no measurable enhancement; includes high-attenuating (&gt;20 HU) cysts &lt;3 cm that are well marginated, homogeneous, and nonenhancing</td>
<td>Renal CT if symptoms occur</td>
</tr>
<tr>
<td>Class IIIF</td>
<td>Moderately complicated cyst, probably benign but requires follow-up</td>
<td>Thin walls may contain multiple thin septa; may have smooth, minimally thickened wall or septa, with irregular or nodular calcifications or perceived but no measurable enhancement or soft tissue enhancement; includes hyperdense completely intrarenal cysts &gt;3 cm with no enhancement</td>
<td>Renal CT at 6 and 12 months, then annually for 5 years</td>
</tr>
<tr>
<td>Class III</td>
<td>Indeterminate renal lesion</td>
<td>Thickened irregular or smooth walls or septa with measurable enhancement</td>
<td>Renal CT/MRI and referral for surgical evaluation</td>
</tr>
<tr>
<td>Class IV</td>
<td>Presumed malignant cystic mass</td>
<td>Contains enhancing soft tissue components adjacent/separate from wall or septa</td>
<td>Renal CT/MRI and referral for surgical evaluation</td>
</tr>
</tbody>
</table>

CT, computed tomography; HU, Hounsfield units, a density measurement; MRI, magnetic resonance imaging.
The mass on Ms. C’s left kidney is hypoattenuating and nonenhancing on CT. It meets Bosniak criteria for a benign simple cyst (class I) and requires no further evaluation or follow-up. Colonoscopy detects multiple colonic polyps that are removed, and the patient does well.

**Mildly complicated cysts.** Less diagnostic certainty characterizes cysts with mild abnormalities that keep them from being labeled as simple. Bosniak classes II and IIF describe mildly abnormal renal cysts. Class II cysts can be dismissed, whereas class IIF cysts require follow-up.

Class II cysts may contain a few hairline septa, fine calcium deposits in walls or septa, or an unmeasurable enhancement of the walls. A hyperattenuating but nonenhancing fluid also is described as category II. Small homogeneous cysts <3 cm, without enhancement but hyperattenuated, are reliably considered benign and need not be evaluated.

Class IIF cysts may have multiple hairline-thin septa with unmeasurable enhancement or minimal smooth thickening or irregular/nodular calcifications of wall or septa without enhancing soft tissue components. Hyperattenuating cystic lesions >3 cm and intrarenal “noncortical” cysts are included in this category. Class IIF cysts require follow-up at 6 months with CT or MRI, then annually for at least 5 years.

**Obviously complicated cysts.** Bosniak class III is indeterminate—neither benign nor clearly malignant. Class III cysts may have thickened borders or septa with measurable enhancement, or they may be multilocular, hemorrhagic, or infected. In 5 case series, 29 of 57 class III lesions proved to be malignant. MRI may characterize these lesions more definitively than CT prior to urologic referral.

**Malignant cysts.** Bosniak class IV renal lesions are clearly malignant, with large heterogeneous cysts or necrotic components, shaggy thickened walls, or enhancing soft tissue components separate from the wall or septa. Their unequivocal appearance results from solid tumor necrosis and liquefaction. Diagnosis is straightforward, and excision is indicated.

### TABLE 2

<table>
<thead>
<tr>
<th>Incidental adrenal masses: Symptoms that suggest pathology&lt;sup&gt;29&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Consider this pathology. . .</strong></td>
</tr>
<tr>
<td>Hypercortisolism</td>
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<tr>
<td>Hyperaldosteronism</td>
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<tr>
<td>Pheochromocytoma</td>
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<tr>
<td>Hyperandrogenism</td>
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<tr>
<td>Carcinoma or metastatic disease</td>
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</table>

In a retrospective review of inapparent adrenal masses, ≤13% of pheochromocytomas were clinically silent. Therefore, laboratory testing is necessary for an incidental adrenal mass.

**A closer look at solid renal masses**

Solid renal masses usually consist of enhancing tissue with little or no fluid. The goal of evaluation is to exclude malignancies, such as renal cell cancer, lymphomas, sarcomas, or metastasis. Benign solid masses include renal adenomas, angiomylipomas, and oncocytomas, among others.

Several lesions can be diagnosed by appearance or symptoms:

**Angiomyolipomas** are recognized by their fat content within a noncalcified mass. Unenhanced CT usually is sufficient for diagnosis, unless the mass is very small or has atypical features.

**Vascular lesions** can be identified because they enhance to the same degree as the vasculature. With the exception of inflammatory or vascular abnormalities, all enhancing lesions that do not contain fat should be presumed to be malignant.
Bosniak class IV renal lesions are clearly malignant, with large heterogeneous cysts or necrotic components, shaggy thickened walls, or enhancing soft tissue components separate from the wall or septa.

In patients with a known extrarenal primary malignancy, 50% to 85% of incidental solid renal masses will represent metastatic disease. Percutaneous biopsy may be warranted to differentiate metastatic lesions from a secondary, primary (ie, renal cell carcinoma), or benign process.

A study of 2770 solid renal mass excisions revealed that 12.8% were benign, with a direct relationship between malignancy and size. Masses <1 cm were benign 44% of the time. Early identification of small renal carcinomas may improve survival rates. Although renal cell carcinomas <3 cm in diameter have low metastatic potential, a solid, nonfat-containing mass should be evaluated for aggressive nephron-sparing surgery.

**Incidental adrenal masses occur infrequently**

Adrenal incidentalomas are defined as radiographically identified masses >1 cm in diameter. They are much less common than their renal counterparts, with a reported prevalence of 0.35% to 5% on CT. Because the adrenal glands are hormonally active and receive substantial blood flow, metastatic, hormonally active, and nonfunctional causes for adrenal masses need to be considered.

**Adrenal pathology**

Adrenal masses may be characterized by increased or normal adrenal function. Hyperfunctioning syndromes include hypercortisolism, hyperaldosteronism, adrenogenital hypersecretion of adrenocortical origin, and pheochromocytomas of the medulla. Symptom evaluation of these syndromes is important, but not sufficient to rule out a hyperfunctioning syndrome.

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- **Nonfunctional lesions** include adenomas, metastases, cysts, myelolipomas, hemorrhage, and adrenal carcinomas. These masses require evaluation for the possibility of cancer, the most common of which is metastasis. In patients with an extra-adrenal malignancy, the likelihood of malignancy in an incidental adrenal mass is at least 50%. An adrenal mass representing metastasis of a previously unrecognized cancer is exceedingly rare.

- **Primary adrenal carcinoma** is also rare, with an estimated incidence of 2 cases per one million in the general population. For patients with adrenal masses, the prevalence of carcinoma increases with lesion size (2% for tumors <4 cm, 6% for tumors 4-6 cm, and 25% for tumors >6 cm in diameter). For this reason, tumors >4 cm in diameter are usually surgically resected in patients with no previous cancer history, unless radiologic criteria demonstrate clearly benign characteristics.

Although adrenal carcinomas are considered nonfunctioning, some evidence suggests they produce low levels of cortisol that may be associated with clinical features of metabolic syndrome.

**CT is first choice for adrenal mass evaluation**

Dedicated adrenal CT with both unenhanced and delayed contrast-enhanced images is the most reliable study to evaluate an adren...
Before any adrenal biopsy, measure plasma-free metanephrines to exclude undiagnosed pheochromocytoma, which could precipitate a hypertensive crisis if untreated.

Other imaging options. MRI is an alternative to CT for patients with contraindications for contrast or radiation exposure. MRI provides less spatial resolution than CT, but chemical shift imaging can measure cytoplasmic lipid content similar to unenhanced CT. A small study found chemical shift MRI more reliable than unenhanced CT, but less reliable than CT with delayed contrast enhancement.

Positron emission tomography (PET) is useful to noninvasively evaluate biochemical and physiologic processes. PET-CT incorporates unenhanced CT density measurements to improve PET accuracy. In a patient with a history of cancer, PET-CT has a sensitivity of 93% to 100% and a specificity of 95% in differentiating benign from malignant adrenal tumors.

When to order a biopsy
The need for biopsy has decreased as imaging has improved, but biopsy is required when diagnostic imaging fails to differentiate a lesion as benign or malignant. CT-guided biopsy provides diagnostic accuracy of 85% to 95%. Complications such as pneumothorax, hemorrhage, and bacteremia occur in 3% to 9% of biopsies. Before any adrenal biopsy, measure plasma-free metanephrines to exclude undiagnosed pheochromocytoma, which could precipitate a hypertensive crisis if untreated.

These 3 laboratory screening tests are critical
Family physicians can perform the initial biochemical evaluation of an adrenal incidentaloma. Guidance is available from the National Institutes of Health (NIH) and the American Academy of Clinical Endocrinologists (AACE) (FIGURE 2).

Regardless of signs or symptoms, perform screening laboratory tests for 3 types of adrenal hyperfunction: hypercortisolism, hyperaldosteronism, and hypersecretion of catecholamines (pheochromocytoma). Screening tests are not recommended for androgen hypersecretion, which is extremely rare and causes recognizable symptoms such as hirsutism (TABLE 2).

Hypercortisolism occurs in approximately 5% of adrenal incidentalomas. An overnight dexamethasone suppression test (DST) is most reliable for screening, with sensitivity >95% for Cushing syndrome. The patient takes a 1-mg dose of oral dexamethasone at 11 pm, and a fasting plasma cortisol sample is drawn the next day at 8 am.

Dexamethasone binds to glucocorticoid receptors in the pituitary gland, suppressing adrenocorticotropic hormone secretion. Cortisol will be depressed the next morning unless the adrenal mass produces cortisol autonomously. Patients with a DST >5 mcg/dL—highly suggestive of Cushing syndrome—require further evaluation, and we suggest referral to an endocrinologist.

Hyperaldosteronism is seen in 1% to 2% of adrenal incidentalomas. The aldosterone-to-renin ratio (ARR) is recommended as a screening test for hyperaldosteronism, with an ARR >20 requiring further testing. Medications that may affect the ARR include beta-blockers, spironolactone, clonidine, diuretics, angiotensin-converting enzyme inhibitors, and angiotensin receptor blockers.

Refer a patient with evidence of hyperaldosteronism to an endocrinologist and a surgeon with experience in managing these lesions. If the ARR test result suggests an aldosterone excess, a salt-loading test is used to verify failure of aldosterone suppression. Adrenal venous sampling is often performed prior to surgical removal to confirm that an incidentaloma is the source of hyperaldosteronism.

Pheochromocytoma. Approximately 5% of incidental adrenal lesions are pheochromocytomas. Many patients with these epineph-
rine/norepinephrine-secreting tumors do not show the classic symptom triad of headache, palpitations, and diaphoresis, and approximately half have normal blood pressure.34

Identifying a pheochromocytoma is important in any patient requiring surgery or biopsy, as surgical manipulation can cause a potentially fatal intraoperative catecholamine surge. Presurgical medical management can mitigate this reaction.

A plasma-free metanephrines test, which has 95% sensitivity, is the most reliable test for pheochromocytoma.35 Medications, including tricyclic antidepressants, decongestants, amphetamines, reserpine, and phenoxybenzamine, can cause false-positive results.36 Confirm a positive plasma-free metanephrines test with a 24-hour fractionated urine metanephrines test, and refer the patient to an endocrinologist.

continued

FIGURE 2
How to manage incidental adrenal masses: A systematic approach

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continued

FIGURE 2
How to manage incidental adrenal masses: A systematic approach

Does report identify mass as definite cyst or myelolipoma?

Yes  No workup necessary

No

Any cancer history?

Yes  Refer for oncologic evaluation

No

Incidental adrenal mass workup
1. CT adrenal with delayed contrast enhancement
2. Biochemical workup
  • Dexamethasone suppression test (DST)
  • Plasma-free metanephrines test
  • Aldosterone-renin ratio

Mass >4 cm or radiographic findings of concern?

Yes  Refer for surgical evaluation

No

Any definitive lab abnormality?

Yes  Refer for endocrine evaluation

No

Repeat evaluation for negative workup
1. Radiologic: Repeat CT or MRI evaluation at 3 to 6 months, then annually for 2 years.
2. Biochemical workup: Repeat screening evaluation annually for 3 years.
Managing adrenal incidentalomas

Refer all patients with adrenal masses >4 cm for surgical evaluation because of the risk of malignancy; all patients who have a history of malignancy and an adrenal mass of any size require a referral to an oncologist. Perform the AACE-recommended 3-element biochemical workup for all masses, with the exception of definitively diagnosed cysts or myelolipomas.

Refer to an endocrinologist all patients with abnormal screening laboratory results, regardless of adrenal mass size, as well as patients with concerning clinical findings. Initiate cardiovascular, diabetes, and bone density evaluation and management for metabolic syndrome.20

Monitoring after a negative workup

Little evidence exists to guide monitoring of small adrenal incidentalomas (<4 cm) with a negative workup. The 2002 NIH report recommended annual radiologic follow-up for 5 years, whereas the 2009 AACE guidelines recommend radiographic follow-up at 3 to 6 months, then at one and 2 years.29 Evidence indicates that 14% of lesions will enlarge in 2 years, although the clinical significance of enlargement is unknown. Some authors argue against CT monitoring because the risk of adrenal mass progression is similar to the malignancy risk posed by 3 years of radiation exposure with CT.20

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Some guidelines recommend repeat biochemical screening every 3 to 4 years.26,29 AACE guidelines quote a 47% rate of progression over 3 years, but most adrenal masses progress to subclinical Cushing syndrome—a condition of uncertain significance. Subclinical Cushing’s has not been reported to progress to the overt syndrome, and new catecholamine or aldosterone secretion is rare.

Many endocrinologists reduce the frequency of follow-up, depending on the type of adrenal mass (cyst or solid) and its size. AACE suggests CT for adenomas one to 4 cm at 12 months. AACE and NIH recommend hormonal evaluation annually for 4 years. Adrenal cysts or myelolipoma in patients without cancer need no follow-up.29

References

INCIDENTAL RENAL AND ADRENAL MASSES


