Management of incidental adrenal tumours

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What you need to know

• An adrenal incidentaloma is an adrenal lesion found incidentally in asymptomatic patients undergoing imaging scans not performed for suspected adrenal disease. Adrenal incidentalomas are becoming more common due to increased scanning, as well as an older population.

• Most incidentalomas are benign and non-functioning (~85%) and require only minimal evaluation (but are frequently overinvestigated). The remaining 15% (functioning and/or malignant lesions) are at risk of under investigation, delay, or even being missed.

• Patients with adrenal incidentalomas can experience anxiety once an adrenal incidentaloma is detected and while awaiting investigations or appointments.

What is an adrenal incidentaloma?

Adrenal incidentalomas are defined as adrenal masses incidentally discovered during imaging that was not performed for suspected adrenal disease. Hence, an adrenal lesion discovered while investigating raised catecholamines would not be classified as an adrenal incidentaloma. Most definitions have restricted the diagnosis to lesions ≥10 mm3 4 unless there are clinical stigmas of adrenal hormone over-production identified after the detection of the lesion that warrant further investigation.

In view of variation in diagnostic criteria, referral patterns, and surgical rates, it is challenging to precisely classify the underlying adrenal incidentaloma pathology from published literature.23 provides a simplified description of the adrenal gland components, their role, and the consequences of excess hormone production.

Who gets it?

Prevalence is estimated between ~2% (range 1.0%-8.7%) from autopsy studies3 4 and ~4% in radiological studies, rising to 10% in elderly patients.6 8 Prevalence is higher in patients with diabetes, hypertension, or obesity.6 Adrenal incidentalomas are rare in children (<0.04% of tumours).5

Most adrenal incidentalomas (~85%) are benign, with no hormonal over-production (1).1 4 The remaining ~15%, however, are either malignant or associated with hormone over-production (excess cortisol, aldosterone, or catecholamine) and higher morbidity-mortality, thereby requiring rigorous investigation.

The commonest functional abnormality for adrenal incidentalomas is cortisol hyper-secretion, with pheochromocytoma and hyper-aldosteronism being much less common. A 50 year autopsy review of more than 40 000 cases revealed pheochromocytomas in 0.13%, 76% of which were only identified after death.11 Regardless of whether...
pheochromocytoma had been diagnosed or not, a common cause of death was the development of hypertensive/hypotensive crisis, often precipitated by unrelated surgery.

How are adrenal incidentalomas managed?

The most recent recommendations from the European Society of Endocrinology (2016) on managing adrenal incidentalomas are summarised in [1]. Comparisons with guidance from the American Association of Clinical Endocrinologists (2009),[2] Italian Association of Clinical Endocrinologists (2011),[3] and Canadian Urological Association (2011),[4] is available in supplementary table 1. These recommendations are mostly derived from retrospective studies, case series, and consensus opinion, and they largely agree that

- Screening is necessary for cortisol, aldosterone, and catecholamines hyper-secretion; sex hormones are not routinely measured unless clinically indicated
- Lipid-rich lesions are benign and require no further action
- Indeterminate or lipid-poor lesions require more detailed attention.

Decision making approach

Is the lesion high risk?

When assessing patients with newly identified adrenal incidentalomas to determine treatment options and prioritisation, it is important to consider the risk of malignancy, and whether the lesion is hyper-functioning (ie, secreting excessive amounts of adrenal hormones). The risk of malignancy is small in patients not previously known to have cancer. Two large retrospective cohort studies have shown the risk of primary adrenocortical carcinoma to be 4.7% and 5%, and that of metastasis 0.7% and 2.5%. In a pooled analysis from 26 international studies, the prevalence of adrenocortical carcinomas was ≤5%. There is a good correlation between risk and tumour size: 2% in lesions ≤4 cm, 6% in lesions 4.1–6 cm and 25% in lesions >6 cm.

What is the risk of malignant transformation?

Size of lesion and its growth over time are key surrogates for malignancy. Of those, size is the most reliable, with >40 mm having the highest sensitivity (93%). It is unclear what would qualify as a worrying rate of growth, especially as benign adenomas also grow over time. Retrospective data showed that, regardless of hormone over-secretion, benign adrenal incidentalomas grew by 10–20 mm over three years.

If it is not making excess hormones now, could that change in the future?

Given the limited prospective data, it is not possible to conclusively document the progression towards hyper-secretion. A literature review of longer term follow-up studies (>1 year), showed that, of 1147 patients with originally non-functional adrenal incidentalomas, 20 (1.7%) developed hyper-function. The most common evolution is cortisol over-production leading to subclinical hyper-cortisolism. In a longitudinal follow-up study of 284 consecutive patients with non-secreting adrenal incidentalomas, the cumulative risk of developing subclinical hyper-function was 6.6% after five years. Those >30 mm at diagnosis were associated with a higher risk of hyper-function. Relative to excess cortisol production, future development of excess aldosterone or catecholamines is unlikely (<0.1%).

How might patients with malignant tumours present?

Most adrenocortical carcinomas are discovered on incidental imaging, and these patients are asymptomatic from an adrenal viewpoint. Malignant adrenal tumours can be functional, and patients might therefore display previously unrecognized features of the corresponding hormone excess, mostly hypercortisolism and/or virilisation. Patients rarely present with signs of feminisation. The sudden onset of virilising clinical features, gynaecomastia, or Cushing’s syndrome, might, respectively, point towards an androgen, oestrogen, or cortisol producing adrenocortical carcinoma. Some patients with adrenocortical carcinomas might present with abdominal or flank pain, abdominal fullness, and occasionally with constitutional symptoms (eg, lethargy, fever, weight loss) secondary to haemorrhage within the tumour.

Investigations to rule out malignancy

Guidelines agree that no further imaging is necessary where initial scans indicate lipid-rich lesions of less than 10 Hounsfield units (HU) (the measure of tissue density) in patients presenting with no known malignancy; these are classified as benign adenomas. Unfortunately, about one third of lesions will be lipid-poor (>10 HU) and therefore it is difficult to exclude malignancy. In these cases, a contrast enhanced triphasic adrenal computed tomography scan should be undertaken, with the calculation of the absolute and relative washout percentages. An absolute washout >60% or relative washout >40% suggest a benign lesion with excellent sensitivity (88%-96%) and specificity (96%-100%). shows computed tomography examples of (a) lipid-rich and (b) lipid-poor lesions.

MRI is similarly effective, with the advantage of no radiation exposure and avoiding the use of iodine based contrast agents. MRI is a safer option for younger patients and if repeated imaging is required (eg, adrenal incidentalomas confirmed to be pheochromocytomas with underlying genetic mutations; these constitute 15%-20% of all catecholamine-secreting tumours).

2-Deoxy-2-[fluorine-18] fluoro-D-glucose positron emission tomography/computed tomography (18F-FDG PET/CT), a technique that combines nuclear medicine with cross sectional imaging, is particularly useful in detecting metastatic disease in patients with primary adrenal cancer or secondaries from other extra-adrenal malignancies. The newest imaging option is 68Ga-dotatate PET, which has a higher specificity for pheochromocytoma compared with FDG PET/CT.

2174 patients concluded that adrenal biopsy is useful in the diagnosis of adrenal metastasis in patients with known extra-adrenal malignancy. The authors concluded that adrenal biopsy should only be performed if the findings would alter patient management and after biochemical exclusion of a pheochromocytoma.

There are no well established sensitive biochemical markers for the diagnosis of adrenocortical carcinoma, or indeed adrenal metastasis. A study including 45 individuals with adrenocortical carcinomas and 102 patients with adrenocortical adenomas showed that detailed serum and 24 hour urinary steroid profiling had a sensitivity and specificity of 90% in differentiating benign adrenal adenomas from adrenocortical carcinomas. Further work is required to clarify whether such profiling could be useful for monitoring.

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Hormone over-production in adrenal incidentaloma

Previously unrecognised clinical features of Cushing’s syndrome, pheochromocytoma, or hyper-aldosteronism are suggestive of functional lesions. In reality, however, most adrenal incidentalomas are identified in patients who are asymptomatic from an adrenal viewpoint.

All guidelines agree that hormonal evaluation by specialists is crucial in all adrenal incidentalomas to exclude the three major conditions associated with functional adrenal incidentaloma:

1. Glucocorticoid excess—There is consensus that the 1 mg overnight dexamethasone suppression test should be the initial screening test. Values <50 nmol/L (1.8 µg/dL) are normal, while those >138 nmol/L (5 µg/dL) indicate autonomous cortisol production (defined as subclinical Cushing’s in the absence of clinical features). However, there is lack of agreement on the interpretation of values between 50 nmol/L and 138 nmol/L in asymptomatic patients.

European guidelines define this as “possible autonomous cortisol secretion,” and US, Italian, and Canadian guidelines acknowledge its significance, despite the absence of unequivocal data on the long term consequences. Another potentially useful test for autonomous cortisol secretion is serum dehydroepiandrosterone sulphate, which is reduced in cortisol hyper-secretion because of adrenocorticotropic hormone suppression. The dehydroepiandrosterone sulphate ratio (measured as dehydroepiandrosterone sulphate divided by the lower reference limit) has been shown to have good sensitivity and specificity in identifying adenomas with autonomous cortisol secretion.

2. Mineralocorticoid excess—The guidelines agree about screening using plasma aldosterone (ng/dL)/renin (ng/mL/h) ratio only in the presence of hypertension or unexplained hypokalaemia. Values >20 (US guidelines) and >30–50 (Italian guidelines) require further confirmatory tests. The sample should be taken in the morning, two hours after getting out of bed, and after 5-15 minutes’ rest. A careful assessment of the patient’s antihypertensive medication should be undertaken (to evaluate potential impact on the ratio) and some might need to be stopped before the test.

Poor adherence to sampling criteria often leads to challenges in interpretation.

3. Catecholamine excess—All guidelines highlight the importance of screening for pheochromocytoma with plasma or urinary metanephrines, whether or not the patient is hypertensive, as some patients with pheochromocytomas are asymptomatic. Excluding a functioning pheochromocytoma is crucial, as surgery without adequate α then β blockade might be seriously detrimental to the patient. A value greater than threefold above the upper reference range limit confirms the diagnosis. Lower values might indicate a false positive test, but pheochromocytoma should be considered in patients with borderline values and indeterminate computed tomography imaging features.

In addition to the above investigations, patients with clinical features of virilisation or gynaecomastia also need assessment of androgen profile and oestrogen levels, respectively.

Some tumours secrete more than one hormone. Patients with lesions causing both primary hyper-aldosteronism and autonomous cortisol production have been reported.

Follow-up

Follow-up of adrenal incidentalomas is dependent on the characteristics of the original lesion. Current guidelines agree that patients with benign lesions that are non-functional require no further follow-up. Supplementary table 2 outlines variations on this. However, prospective research studies should be undertaken, as these will strengthen (or challenge) these recommendations.

Present challenges and future direction

Despite current guidance, there remain two major challenges:

1. Most patients with adrenal incidentalomas are not referred to relevant specialists (endocrinologists/diabetologists), so there is potential to miss functional or malignant lesions. That said, investigating every lesion could overload any health system. Therefore, an evidence based, streamlined approach is required to ensure clinically effective and prompt evaluation, while avoiding unnecessary investigations and clinic appointments.

We are developing an electronic adrenal incidentaloma management system linked to the latest guidance. This will provide more streamlined and timely management by bringing all the key information together electronically and guiding the management process, shortening the hands-on time for healthcare professionals. This increased efficiency will facilitate the management of all adrenal incidentaloma cases, including those currently missed thereby enhancing patient safety.

2. It is important to recognise that detection of adrenal incidentaloma is associated with substantial patient anxiety (unexpected nature, lack of standardised management pathway, often poor communication from non-specialists); delay in decision making should be avoided (see Tips for the non-specialist).

Future developments in imaging and biochemical marker (eg, steroid metabolites) assays might help differentiate benign from malignant lesions and avoid unnecessary surgery.

How patients were involved in the creation of this article

A patient with an adrenal incidentaloma provided a patient’s perspective on their experience with diagnosis and treatment. Our innovation project work (funded by the Health Foundation) included focus group discussion with patients, which highlighted patient experience (anxiety given the unexpected nature of the finding, the need for clear information on what adrenal incidentaloma is, and the optimum timelines for interaction with healthcare professionals) and helped shape the article, in particular the section on what to tell the patient (see box Tips for the non-specialist).
A patient’s perspective

Diagnosis

“Before diagnosis, I suffered sudden fluctuations in body temperature and palpitations. My doctor sent me for blood tests, which were all normal, and an ultrasound scan. I was informed that the scan showed they had found a lump and I was being referred to a consultant.

“Once under hospital care, I felt totally swept along with different scans and tests. I was informed that I had an uncommon adrenal lump that can be serious. The experience was overwhelming; I took in very little information. I did my own research, but found the internet confusing.”

Removal and beyond

“Blood pressure tablets were introduced in the weeks before the surgery. While encouraging talks kept us going; with hindsight, I realise that the operation was risky. I was lucky to have a good outcome, going home one week after surgery. I feel 100% better and no longer get palpitations or temperature fluctuations.

“Family members were tested and none was found to be carrying the broken gene: a relief to everyone. “My care during and after diagnosis was excellent. My advice to other patients is to trust the healthcare professionals, feel free to ask questions, and ensure internet information is from credible websites.”

Tips for the non-specialist

What to say to patients

• The scan that we arranged for you showed a swelling on the adrenal gland. The adrenal gland is a small but important gland that sits on top of the kidney. It secretes a group of hormones that are important for our wellbeing.

• We are increasingly detecting these swellings because our scanners are becoming better.

• We know that these swellings could have been present for many years. They are likely to stay largely stable with no harm to you. We are reassured by small swellings (<4 cm).

• We will therefore refer you to an endocrinologist/diabetologist (hormone specialist) to do some tests. They will discuss the results and scans in a meeting with other specialists, such as radiologists, surgeons, and clinical biochemists.

When to refer patients

Patients will be anxious while awaiting the conclusion of the investigations required by the hormone specialist. Hence, any investigations need to be timely. Lipid-rich swelling less than 4 cm in diameter with no evidence of hormone over-production will require no further intervention, and patients should be reassured as soon as the biochemical results are available.

When to refer to the endocrinology team

• Refer all detected adrenal incidentalomas.

• Where there is evidence of excess aldosterone or cortisol, refer within 6-8 weeks.

• Where there are large (or increasing in size) lipid-poor lesions (suspected malignancy) or evidence of excess catecholamines, refer within two weeks.

Monitoring and prognosis

Lipid-rich swelling less than 4 cm in diameter with no evidence of hormone over-production requires no further action. Other lesions require specialist input.

Education into practice

• Can you measure the time from adrenal incidentaloma detection to decision?

• How would you measure the effectiveness and efficiency of your current process? How could that be improved?

• Are there any cases that are currently missed/inadequately followed up?

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The authors wish to express their gratitude to the patient who participated in the creation of this article for agreeing to share their experience, having been identified with an adrenal incidentaloma. We also thank Cherian George for kindly identifying and providing the scan images included (for which patient consent was obtained).

Competing interests

All authors have completed the Unified Competing Interest form (available on request from the corresponding author) and declare no support from any organisation for the submitted work; no financial relationships with any organisations that might have an interest in the submitted work in the previous three years, and no other relationships or activities that could appear to have influenced the submitted work.

Contributions and guarantor

FWFH conceived the original idea and wrote the first draft of the manuscript, particularly the clinical and natural history sections of the manuscript. BGI provided a critical appraisal of alternative guidelines. JS and BK provided an independent critical review of the methodological and biochemical aspects of the manuscript, respectively.

AAF wrote the biochemical components and critically reviewed the manuscript as a whole. All authors reviewed and approved the final version of the manuscript. FWFH is the guarantor of the work.

Provenance and peer review: commissioned; externally peer reviewed.

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<table>
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<th>Type</th>
<th>Subtype</th>
<th>Percentage of adrenal incidentalomas</th>
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<tr>
<td>Non-functional</td>
<td>Apparent non-functioning adenoma</td>
<td>71.2–82.4</td>
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<tr>
<td>Functional (hormonally active)</td>
<td>Subclinical (autonomous cortisol secretion) Cushing's syndrome</td>
<td>5.3–7.9</td>
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<td></td>
<td>Pheochromocytoma</td>
<td>5.1–5.6</td>
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<td></td>
<td>Aldosterone producing adenoma</td>
<td>1.0–1.2</td>
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<td>Malignant</td>
<td>Adrenocortical cancer</td>
<td>4.4–4.7</td>
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<td></td>
<td>Metastatic cancer</td>
<td>2.1–2.5</td>
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<tr>
<td>Table 2</td>
<td>Summary of key recommendations from recent European guidance on managing patients with a finding of an incidental adrenal tumour</td>
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<tr>
<td><strong>Criterion</strong></td>
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<td><strong>General recommendations</strong></td>
<td>Recommendations for discussion of adrenal incidentalomas in MDT (any of the following):</td>
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<td></td>
<td>Imaging not consistent with a benign lesion</td>
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<td></td>
<td>Evidence of hormone excess (including autonomous cortisol secretion)</td>
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<td></td>
<td>Evidence of substantial tumour growth during follow-up imaging (more than 20%, or at least a 5 mm increase in maximum diameter during the following 6-12 months)</td>
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<td></td>
<td>Adrenal surgery could be a consideration.</td>
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<td></td>
<td>Non-contrast computed tomography with determination of mass density to assess for malignancy.</td>
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<td></td>
<td>One mg ODST and plasma/urinary metanephrines in all patients. ARR in all patients with concomitant hypertension and/or unexplained hypokalaemia. Sex hormones and steroid precursors in patients with clinical or imaging features of ACC</td>
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<tr>
<td><strong>Surgical recommendation for subclinical Cushing syndrome</strong></td>
<td>Clinical and biochemical assessment for cortisol production and co-morbidities. If worsening then consider surgery</td>
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<tr>
<td><strong>Indications for consideration for surgical intervention</strong></td>
<td>Unilateral adrenal tumour with clinically significant hormone excess</td>
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<td>Suspicion of malignancy</td>
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<td><strong>Specialist screening test for autonomous cortisol production/subclinical Cushing's syndrome</strong></td>
<td>1 mg ODST</td>
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<td><strong>Diagnostic cut-off values for subclinical Cushing's syndrome</strong></td>
<td>Following 1 mg ODST:</td>
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<td></td>
<td>If cortisol between 51 and 138 nmol/L: “possible autonomous cortisol secretion.”</td>
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<td></td>
<td>If cortisol &gt;138 nmol/L (and without clinical features of Cushing's): “autonomous cortisol secretion”</td>
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<tr>
<td><strong>Specialist screening test for pheochromocytoma</strong></td>
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<td><strong>Indications for computed tomography guided needle biopsy</strong></td>
<td>In the setting of a newly diagnosed adrenal mass and a history of extra-adrenal malignancy:</td>
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<tr>
<td></td>
<td>Lesion hormonally inactive</td>
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<td></td>
<td>Lesion has not been conclusively characterised as benign by imaging</td>
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<td>Management would be altered by the knowledge of histology</td>
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<tr>
<td><strong>Imaging</strong></td>
<td>Non-contrast computed tomography scan to determine whether the mass is homogenous and lipid rich (&lt;10 HU).</td>
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<td>If indeterminate on non-contrast computed tomography, imaging with another modality (contrast computed tomography, MRI, or FDG-PET)</td>
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<td><strong>Follow-up if not resected</strong></td>
<td>No further imaging if lesion &lt;4 cm and clear benign features on imaging studies. Indeterminate mass repeat non-contrast computed tomography or MRI after 6-12 months. Resect if growth by &gt;20% in addition to at least a 5 mm increase in max diameter. If growth below this threshold additional imaging after 6-12 months. No further hormonal investigation if initial evaluation normal unless new clinical signs of endocrine activity or there is worsening of comorbidities — eg, hypertension or diabetes</td>
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MDT: multidisciplinary team, ODST: overnight dexamethasone suppression test, ARR: plasma aldosterone (ng/dL)/renin (ng/mL/h) ratio, ACC: adrenocortical carcinoma, SCS: subclinical Cushing's syndrome, HU: Hounsfield units, MRI: magnetic resonance imaging, FDG-PET: 2-deoxy-2-[fluorine-18]fluoro-D-glucose positron emission tomography/computed tomography
Figures

The components of the adrenal gland and consequences of excess hormone production

<table>
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<th>Hormones</th>
<th>Function</th>
<th>Consequences of excess production</th>
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<td>Aldosterone</td>
<td>Regulates mineral balance</td>
<td>Hyperaldosteronism (Conn's syndrome)</td>
</tr>
<tr>
<td>Cortisol</td>
<td>Regulates metabolism, facilitates response to stress</td>
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</tr>
<tr>
<td>Dehydroepiandrosterone (DHEAS)</td>
<td>Facilitates early pubic/axillary hair growth, contributes to libido in females</td>
<td>Virilisation in females (usually associated with adrenal cancer)</td>
</tr>
<tr>
<td>Catecholamines</td>
<td>Fear, fight, and flight reaction</td>
<td>Pheochromocytoma</td>
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</table>
Computed tomography showing lipid-rich and lipid-poor lesions. (a) A 2 cm left adrenal lesion showing a computed tomography density of less than 10 HU, which is suggestive of a lipid-rich adenoma. (b) A 7.7 cm hypervascular left adrenal mass that is suspicious for pheochromocytoma. Inset shows a metaiodobenzylguanidine scan demonstrating intense uptake in the lesion.