Management of Incidental Liver Lesions on CT: A White Paper of the ACR Incidental Findings Committee

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Abstract

The ACR Committee on Incidental Findings presents recommendations for managing liver lesions that are incidentally detected on CT. These recommendations represent an update from the liver component of the ACR 2010 white paper on managing incidental findings in the pancreas, adrenal glands, kidneys, and liver. The Liver Subcommittee—which included five abdominal radiologists, one hepatologist, and one hepatobiliary surgeon—developed this algorithm. The recommendations draw from published evidence and expert opinion and were finalized by informal iterative consensus. Algorithm branches categorize liver lesions on the basis of patient characteristics and imaging features. They terminate with an assessment of benignity or a specific follow-up recommendation. The algorithm addresses most, but not all, pathologies and clinical scenarios. The goal is to improve the quality of care by providing guidance on how to manage incidentally detected liver lesions.

Key Words: Incidental finding, hepatic cyst, hemangioma, focal nodular hyperplasia, hepatic metastasis, hepatocellular carcinoma

OVERVIEW OF THE ACR INCIDENTAL FINDINGS PROJECT

The core objectives of the Incidental Findings Project are to (1) develop consensus on patient characteristics and imaging features that are required to characterize an incidental finding, (2) provide guidance to manage such findings in ways that balance the risks and benefits to patients, (3) recommend reporting terms that reflect the level of confidence regarding a finding, and (4) focus future research by proposing a generalizable management framework across practice settings. The ACR Committee on Incidental Findings (IFC) generated its first white paper in 2010, addressing methods for managing incidental findings in four organ systems: pancreas, adrenal glands, kidneys, and liver [1].

THE CONSENSUS PROCESS: THE LIVER LESION ALGORITHM

The present report represents the first revision of the IFC’s 2010 recommendations regarding incidental liver
lesions detected on CT. The process of developing this algorithm included naming a subcommittee chair, who appointed four additional abdominal radiologists, one hepatologist, and one hepatobiliary surgeon. The subcommittee then developed and gained consensus on a preliminary version of the algorithm using published evidence as their primary source. Where evidence was not available, they invoked the collective expertise of their team. The preliminary algorithm underwent review by additional members within the IFC, including the Body Commission chair, the IFC chair, and additional IFC subcommittee chairs. The revised algorithm and corresponding white paper draft were submitted to additional ACR stakeholders to gain input and feedback. Consensus was obtained iteratively after successive reviews and revisions. After completion of this process, the algorithm and white paper were finalized. The IFC’s consensus processes meet policy standards of the ACR. However, they do not meet any specific, formal national standards. This algorithm and set of recommendations does not represent policy of the ACR Practice Guidelines or the ACR Appropriateness Criteria. Our consensus may be termed “guidance” and “recommendations” rather than “guidelines,” which has a more formal definition.

ELEMENTS OF THE FLOWCHARTS: COLOR CODING

The proposed algorithm for incidental liver lesions is included in Figure 1. Yellow boxes indicate using or acquiring clinical data (eg, lesion size), green boxes describe recommendations for action (eg, follow-up MRI), and red boxes indicate that no follow-up is needed (eg, the finding is benign). To minimize complexity, the algorithm addresses most, but not all, imaging appearances and clinical scenarios. Radiologists should feel comfortable deviating from the algorithm in

![Algorithm for incidental liver lesions](image-url)
circumstances that are not represented in the algorithm, on the basis of the specific imaging appearance of the finding in question and patient characteristics. The algorithm content must be viewed as recommendations and should not be considered as “standard of care.”

**NATURE AND SCOPE OF THE PROBLEM**

Advances in cross-sectional imaging have led to the discovery of innumerable incidental liver lesions [2-4]. Such lesions will be detected in up to 30% of individuals older than 40 years [5-23]. Although most are benign, in many cases, further workup can be difficult to avoid. Conversely, it is well recognized that overdiagnosis of benign or indolent lesions places patients at risk for potentially dangerous and expensive follow-up care [24-29]. We provide recommendations for managing incidental liver lesions that appropriately balance the benefits and risks of further workup.

**Definition of an Incidentally Detected Liver Lesion**

An incidentally detected liver lesion is one that is identified in a patient imaged for an unrelated reason. As a result, we address patients with a wide spectrum of risk for a malignant liver lesion: from an otherwise healthy patient with right lower quadrant pain to a patient with a history of a primary malignancy or cirrhosis with pain after a motor vehicle collision. Although the latter patient is at increased risk for a malignant hepatic lesion—for metastasis or hepatocellular carcinoma, respectively—benign incidental liver lesions are also common in such patients, which makes management decisions particularly difficult in the absence of guidance [30-36]. Therefore, our recommendations are based on both the imaging appearance of the incidental lesion in question and the patient’s risk for having a malignant lesion (Fig. 1). Importantly, our algorithm was developed to distinguish benign from potentially malignant incidental findings, and not hepatic infections or abscesses, given that the latter are very likely to be associated with clinical signs or symptoms.

**Risk Categories for Patients With Incidental Liver Lesions: “Low” Versus “High”**

Our algorithm requires designation of patients as low risk or high risk for having a malignant hepatic lesion (Table 1). These categories, defined later, stratify the need for, and nature of, further workup. Within each category, “hepatic risk factors” refer to conditions that place patients at risk for primary hepatic malignancy and include hepatitis, alcoholism, nonalcoholic steatohepatitis, sclerosing cholangitis, primary biliary cirrhosis, choledochal cysts, hemochromatosis and other hereditary hepatic conditions, and anabolic steroid use [37,38]. Low-risk patients have no known malignancy, hepatic dysfunction, or hepatic risk factors. Within the low-risk category, older patients (>40 years of age) are at higher risk than younger patients for malignancy [28]. High-risk patients have known malignancies with a propensity to metastasize to the liver, cirrhosis, and/or other hepatic risk factors. Therefore, when evaluating an incidental hepatic lesion, it is critical to know the patient’s clinical history.

**Commonly Encountered Benign Lesions**

Independent of patient-level risk, our recommendations are based on the premise that the absence of a benign signature in most incidental lesions ≥1 cm should prompt follow-up imaging with MRI (Fig. 1). Therefore, for most incidental lesions in our algorithm, radiologists should seek to identify definitively benign features to prevent unnecessary follow-up imaging.

The most commonly encountered benign hepatic lesions fall into four major categories: hepatic cysts, perfusional changes, hemangiomas, and focal nodular hyperplasias (FNHs) [39-41]. Hepatic cysts, particularly if ≥1 cm, can generally be characterized by their low attenuation (discussed further in the “Reporting Considerations” section). If <1 cm, an accurate density measurement may not be attainable, a circumstance that is addressed in our algorithm by considering the patient’s underlying risk for malignancy (Fig. 1). Per fusional changes, including areas of fatty sparing of the liver, have characteristic locations and enhancement features, which generally enable their definitive characterization without further follow-up (also discussed further in the “Reporting Considerations” section) [42,43].

**Table 1. Patient risk factors**

<table>
<thead>
<tr>
<th>Low-risk patients†</th>
<th>High-risk patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>No known malignancy</td>
<td>Known malignancy with a propensity to metastasize to the liver</td>
</tr>
<tr>
<td>No hepatic dysfunction</td>
<td>Cirrhosis</td>
</tr>
<tr>
<td>No hepatic risk factors‡</td>
<td>Presence of hepatic risk factors‡</td>
</tr>
</tbody>
</table>

*Within the low-risk category, older patients (>40 years of age) are at higher risk than younger patients for malignancy. †Hepatic risk factors: hepatitis, nonalcoholic steatohepatitis, alcoholism, sclerosing cholangitis, primary biliary cirrhosis, choledochal cysts, hemochromatosis and other hereditary hepatic conditions, and anabolic steroid use.
Hepatic hemangiomas have an appearance that varies on the basis of the presence of contrast material and the available phase(s) of postcontrast imaging. On unenhanced CT, hemangiomas are similar in density to the blood pool. After the administration of contrast material, in the arterial phase, hemangiomas demonstrate peripheral, nodular enhancement, with progressive fill-in at later phases [41,44-46]. As a consequence, in the portal venous phase and thereafter, they are usually isodense or hyperdense relative to the normal liver [41,44-46]. Importantly, large hemangiomas may not enhance centrally on any postcontrast phase because of cystic degeneration, thrombosis, and/or fibrosis [41,44-46].

In the arterial phase, some smaller hemangiomas will uniformly enhance, rather than adhering to the aforementioned pattern of enhancement, an observation described as “flash-filling.” For this reason, if only the arterial phase (including late arterial/early portal venous phase) is available for evaluation, it can be challenging to distinguish hemangiomas from hypervascular neoplasms (metastases or hepatocellular carcinomas). Unlike hemangiomas, malignant neoplasms usually become hypodense relative to the normal liver in the portal venous phase [41,44-46]. Therefore, the availability of additional postcontrast imaging phases can help discern hemangiomas from hypervascular malignant neoplasms.

Enhancement characteristics of hemangiomas on MRI are analogous to those described for CT [41,44-46]. On unenhanced T1-weighted sequences, hemangiomas are low in signal intensity. On T2-weighted sequences, they are typically bright; internal fibrotic areas are dark [41,44-46].

FNH also has a CT appearance that varies on the basis of the presence of contrast material and the available phase(s) of postcontrast imaging. FNH is usually hypodense or isodense relative to the normal liver on unenhanced CT. A hypodense central scar is seen in one-third of cases [41,45,46]. After the administration of contrast material, FNH avidly enhances in the arterial phase. In the portal venous phase and thereafter, FNH becomes isodense; if present, the central scar enhances more gradually, and may appear hyperdense on delayed phases [41,44-46].

Enhancement characteristics of FNH on MRI are again similar to those described for CT. On unenhanced T1-weighted sequences, FNH is isointense relative to normal liver; on T2-weighted sequences, FNH is slightly hyperintense to isointense. The central scar, if present, is dark on T1-weighted sequences and bright on T2-weighted sequences [41,44-46].

Distinguishing FNH and hepatocellular adenoma (HCA) is important; HCAs can hemorrhage or transform into hepatocellular carcinoma, as described later. This is particularly important if the lesion is larger than 3 cm and subcapsular. For such patients, use of MRI with gadoxetate disodium is helpful because FNHs, unlike HCAs, typically demonstrate its uptake on hepatobiliary-phase imaging [47-52].

HCA
There are three primary subtypes of HCAs. From most to least frequent, they are inflammatory HCA, hepatocyte nuclear factor–1α inactivated HCA, and β-catenin-activated HCA [53-56]. Inflammatory HCA carries the greatest risk for hemorrhage, whereas β-catenin-activated HCA carries the greatest risk for malignancy [53-56]. All subtypes are associated with arterial-phase enhancement; however, enhancement characteristics in subsequent phases vary [53-56].

Comparisons With Prior Studies
Comparisons with prior imaging are critically important to determine if a hepatic lesion is new or growing. Importantly, review of prior ultrasound, chest CT, PET/CT and PET/MRI, and/or spinal CT or MRI examinations can also help establish stability. Our algorithm does not explicitly incorporate growth; this is because thresholds for intervention may vary substantially across patient risk categories and imaging features. However, in general, absence of growth over a 1-year time period favors the presence of a benign lesion.

Biopsy of Liver Lesions
Liver biopsy is commonly used when follow-up MRI cannot confirm the presence of liver metastases or a primary hepatic malignancy and when such knowledge would affect subsequent management decisions. Importantly, the risks for morbidity (about 0.5%) and mortality (about 0.05%) should be considered when making a decision about the need for, and technique of, biopsy (core versus fine-needle aspiration) [26,29,57]. Characteristic imaging features of hepatocellular carcinoma may obviate the need for biopsy in many patients [57]. However, personalization of treatment may require tissue to be obtained.

REPORTING CONSIDERATIONS
To optimize lesion characterization and management recommendations, the following elements should be considered when reporting an incidental liver lesion detected on CT. Although each element does not require...
mention for every incidental hepatic lesion (eg, a simple cyst), radiologists are strongly encouraged to address those elements that guide specific follow-up recommendations.

1. Lesion size
2. Lesion attenuation
3. Lesion homogeneity versus complexity
4. Lesion enhancement pattern
5. Lesion margin
6. Lesion multiplicity
7. Lesion growth pattern
8. Lesion location

Lesion Size
Most liver lesions ≤ 1 cm are benign [41,58,59]. Even in high-risk patients, such lesions are commonly benign; however, a potentially new lesion in a high-risk patient warrants follow-up (Fig. 1) [30-36].

Lesion Attenuation
In low-risk patients, lesions with the following characteristics are considered cysts: –10 to +20 Hounsfield units (HU), homogeneous, sharply marginated and without enhancement, mural thickening or nodularity, or septations [39,41]. Importantly, liver metastases can be cystic (eg, in ovarian cancer and gastrointestinal stromal tumors). Therefore, low density is not definitive for a simple cyst in certain patients with cancer [40].

Lesion Homogeneity Versus Complexity
To evaluate the homogeneity versus complexity (heterogeneity) of an incidental hepatic lesion on CT, multiple regions of interest should be placed throughout the lesion, including the highest density areas [41]. Wall thickening or peripheral enhancement, mural nodules, and thick septa raise the likelihood of malignancy [41]. It should be recognized that in the appropriate clinical setting, these features could also indicate an abscess.

Lesion Enhancement Pattern
Hepatic cysts should enhance ≤ 20 HU after the administration of contrast material. However, in small lesions, attenuation measurements can be inaccurate; moreover, it is uncommon to have both unenhanced and postcontrast CT available to evaluate an incidental liver lesion. To verify the presence of a cyst, MRI is superior to CT; ascertainment of no enhancement versus enhancement is more reliable, and additional T2-weighted sequences and diffusion-weighted imaging are helpful for confirmation [45]. Hemangiomas and FNH have characteristic enhancement features, described earlier.

Importantly, the “flash-filling” property of a lesion—uniform enhancement on arterial-phase imaging (including late arterial/early portal venous-phase imaging)—should be reported, particularly when it is the sole detectable feature of the lesion. In our algorithm, such lesions are managed separately in instances in which additional multiphasic imaging is not available to enable their further, definitive characterization (eg, as hemangiomas or hepatocellular carcinomas) (Fig. 1).

Lesion Margin
Benign lesions usually have smooth margins; malignant lesions may have smooth, irregular, or ill-defined margins [41].

Lesion Multiplicity
The presence of multiple liver lesions in patients with cancer often raises suspicion for metastatic disease. However, benign entities such as multiple biliary hamartomas are similarly associated with multiplicity [39,41]. If multiple lesions are present, index lesions that are largest in size and/or demonstrate the most concerning features should be identified to guide follow-up.

Lesion Growth Pattern
Growth of a hepatic lesion raises concern for malignancy, but benign and malignant lesions can grow over time [59-61]. Although our algorithm does not explicitly incorporate growth, in general, absence of growth over a 1-year time period strongly favors the presence of a benign lesion.

Lesion Location
Specific regions of the liver are susceptible to effects of perfusional changes and fatty infiltration or sparing; such effects may mimic liver lesions [62-65]. Peripherally, so-called THADs (transient hepatic attenuation differences, seen on CT) and THIDs (transient hepatic intensity differences, seen on MRI) reflect changes in enhancement of the parenchyma due to relative differences in hepatic arterial versus portal venous supply. Near the falciform ligament and the gallbladder fossa, alterations in venous drainage can result in focal fatty deposition or sparing [42,43,62-65].
INCLUSION AND EXCLUSION CRITERIA FOR USE OF THE ALGORITHM

The algorithm should only be applied to incidental liver lesions in asymptomatic adult patients (≥18 years of age) for whom CT was requested for an unrelated reason. As described earlier, the algorithm is designed for use in patients with varied underlying risk levels (low versus high) for a malignant hepatic lesion. However, the algorithm should not be applied when index CT (ie, that which demonstrates the incidental lesion) was requested to evaluate a known or suspected liver lesion or hepatic abnormality. There are some hepatic lesions that present with associated vascular invasion, biliary dilation, or abnormality. There are some hepatic lesions that present with associated vascular invasion, biliary dilation, or adenopathy. Patients with these associated findings should be referred directly for oncologic evaluation.

IMPLICATIONS OF IMAGING AND CLINICAL FEATURES

Five Basic Principles of the Algorithm

1. In a low-risk patient, an incidental hepatic lesion <1 cm generally does not require further workup and can be considered benign. Radiologists should feel comfortable deviating from this recommendation in instances in which such lesions have suspicious features (ie, ill-defined margins, heterogeneous density, mural thickening or nodularity, or thick septa). In these instances, MRI should be considered.

2. Incidental hepatic lesions that are ≥1 cm and have distinctly benign imaging features do not require follow-up. Such features include sharp margin, homogeneous low attenuation (≤20 HU) on non-contrast or portal venous-phase imaging, or characteristic features of hemangiomas, FNH, or perfusional changes (including focal fatty sparing or deposition). If pseudoenhancement is present, then a benign cyst may measure >20 HU; radiologists’ discretion is necessary.

3. Incidental hepatic lesions that are ≥1 cm and have suspicious imaging features require further workup with prompt MRI or biopsy, depending on the lesion’s size and features and the patient’s risk level. Suspicious imaging features include ill-defined margins, heterogeneous density, mural thickening or nodularity, thick septa, and intermediate to high attenuation on portal venous-phase imaging (>20 HU, in the absence of pseudoenhancement). If pre- and postcontrast CT is available, enhancement >20 HU is a suspicious feature.

4. In this algorithm, “flash-filling” lesions are classified separately as incidental liver lesions that are characterized by uniform enhancement on arterial-phase imaging (including late arterial/early portal venous-phase imaging), and for which multiphasic imaging is not available to enable definitive characterization. In low-risk patients, they are generally benign; MRI is suggested for follow-up only when they exceed 1.5 cm. In high-risk patients, MRI is advised routinely because of a higher probability of malignancy. Nevertheless, even in patients with cirrhosis with small, wedge-shaped hypervascular lesions, the vast majority of such lesions are benign [66].

5. If inadequate imaging is available to ascertain the presence of benign versus suspicious features in a ≥1 cm lesion (eg, a homogeneous 3-cm, 40-HU incidental hepatic lesion with a sharp margin on a non-contrast CT), prompt MRI should be considered for complete characterization of the lesion.

OVERVIEW OF THE ALGORITHM

Low-Risk Patients (Fig. 1)

Incidental Liver Lesion ≤1.5 cm. In low-risk patients, incidental liver lesions less than 1 cm generally do not require further workup and can be considered benign. Incidental liver lesions that are 1.0 to 1.5 cm and have benign or flash-filling features also do not require further workup. Prompt MRI is advised for lesions with suspicious features that are 1.0 to 1.5 cm.

Incidental Liver Lesion >1.5 cm. In low-risk patients, for incidental liver lesions that are greater than 1.5 cm and have benign imaging features, no further workup is necessary. If the lesion has suspicious or flash-filling features, prompt MRI is advised.

High-Risk Patients (Fig. 1)

Incidental Liver Lesion ≤1.5 cm. In high-risk patients with incidental liver lesions less than 1 cm, MRI is advised in 3 to 6 months to both characterize the lesion and document the presence or absence of growth. For lesions that are 1.0 to 1.5 cm and have benign features, no further workup with MRI is necessary; for lesions of this size with suspicious or flash-filling features, we recommend prompt MRI.

Incidental Liver Lesion >1.5 cm. In high-risk patients, for incidental liver lesions that are greater than 1.5 cm and do not have benign imaging features, at minimum, we recommend prompt MRI. For large and highly suspicious lesions (eg, a 3-cm lesion that is likely to be a solitary colorectal metastasis), direct referral to biopsy
may be considered, depending on the clinical scenario. In
general, core biopsy is preferred over fine-needle aspira-
tion and is often necessary for the accurate diagnosis of a
primary hepatocellular neoplasm.

**IMAGING PROTOCOL OPTIMIZATION**

**CT and MRI**

When performed for liver lesion evaluation, a CT pro-
tocol may include multiple phases: unenhanced imaging
and late arterial, portal venous, and delayed-phase post-
contrast imaging. If a dual-energy CT (DECT) exami-
nation is performed, the unenhanced CT phase should
be eliminated [67-69]. In general, the unenhanced CT
phase should be eliminated whenever possible, as it
does not provide additional information in many
scenarios.

We favor MRI over CT for the characterization of an
incidental liver lesion. In general, gadolinium blood pool
agents should be used rather than hepatobiliary agents;
however, to distinguish FNH and HCA, gadoxetate
disodium is recommended, as described earlier. In most
cases, MRI enables better characterization of a lesion’s
internal features, and ascertainment of enhancement is
more reliable relative to CT. In addition, radiation
exposure is avoided.

**DECT**

Depending on the method used to process images, if a
DECT examination is performed on a DECT-capable unit,
it may identify the presence and even quantity of iodine
within a lesion. Confirming iodine content demonstrates
that a lesion has blood perfusion, rather than simply being a
hyperdense lesion from another cause, such as protein-
aceous material, calcium, or iron. A color-coded iodine map
may be generated to localize foci of enhancement. Iodine
can also be detected by comparing the density on different
simulated monoenergetic images. On virtual unenhanced
series, an iodine-containing lesion would be low attenua-
tion, and dense lesions from other causes would remain
higher in attenuation. The ability to generate virtual
unenhanced series from a postcontrast examination may
eliminate the need to perform a conventional unenhanced
series. Using this technology, various types of lesions can be
characterized as nonenhancing, including hyperattenuating
hepatic cysts and bile duct hamartomas, as opposed to
enhancing lesions such as metastases and other malig-
nancies [67,68]. However, if an indeterminate lesion is
found on conventional CT, MRI may be preferred over
DECT [67-71].

**Reduced-Dose CT Scanning**

We recommend use of dose-reduction techniques that are
responsible and tailored to the clinical question at hand.
Detection and characterization of small liver lesions can be
a challenging task when aggressive dose reduction
techniques have been used. As such, particularly in high-
risk patients, we emphasize the need to maintain
diagnostic-quality imaging [72-74].

**PET/CT and PET/MR Evaluation**

In larger hepatic lesions (>1 cm), PET/CT and PET/MR
have precluded the need for biopsy in some patients
[75-77].

**CONCLUSIONS**

Hepatic incidental findings are a common problem on
CT; we provide an algorithm for their management that
is tailored to the patient’s risk for malignancy and the
lesion’s specific imaging features.

Four recommendations that define our updated
algorithm include (1) to forgo workup of incidental he-
patic lesions that are less than 1 cm in low-risk patients,
(2) to forgo workup of lesions with distinctly benign
features (regardless of patient risk level), (3) to pursue
workup of lesions that are ≥1 cm and without distinctly
benign features in high-risk patients, and (4) to use MRI
for further workup.

We hope that this update provided by the Liver
Subcommittee of the IFC will help accurately characterize
most incidental hepatic lesions that are detected on CT
and minimize the frequency of unnecessary patient
workup.

**TAKE-HOME POINTS**

- Forgo workup of incidental hepatic lesions less than
  1 cm in low-risk patients.
- Forgo workup of incidental hepatic lesions with
  distinctly benign features regardless of risk level.
- Pursue workup of incidental hepatic lesions that are
  ≥1 cm and without distinctly benign features in
  high-risk patients.

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