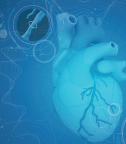


Identifying Appropriate Patients:

The Coronary CTA + HeartFlow FFR_{CT} Analysis Pathway



ACC/AHA Chest Pain Guidelines



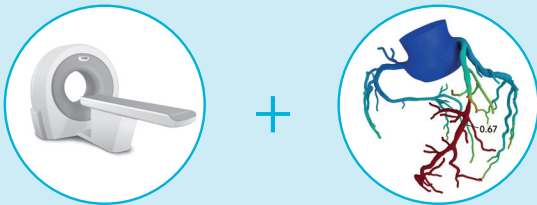
CCTA is the ONLY Class 1 non-invasive test with Level A evidence.



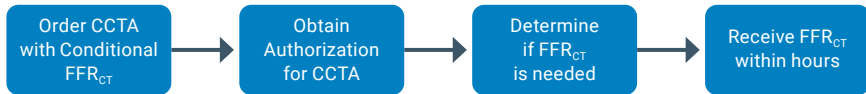
FFR_{CT} is designated as Class 2a with Level B evidence and provides actionable information across a broad range of patient populations.

Order coronary CTA + HeartFlow FFR_{CT} Analysis for patients with stable or acute chest pain with suspected or known CAD. Consider starting with these patients first:

- **Negative or equivocal** functional tests
- **Contraindications** to functional tests (e.g., arthritic, abnormal resting ECG, unable to exercise)
- **Pre-Test Probability** of CAD



To optimize workflows, we recommend submitting an order for CCTA with a conditional FFR_{CT} order and completing prior-authorizations as needed for both at the same time.



<4 hour median turnaround time as of March 2021. Data on file.

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