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Journal of Cardiovascular Computed Tomography

Review article

The Clinical Utility of Hybrid Imaging for the Identification

of Vulnerable Plaque and Vulnerable Patients

Short title: Hybrid imaging of vulnerable plaque

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1 ABSTRACT

2 Despite decades of research and major innovations in technology, cardiovascular 3 disease remains the leading cause of death globally. Our understanding of major 4 cardiovascular events and their prevention is centred around the atherosclerotic plaque and the processes that ultimately lead to acute plaque rupture. Recent 5 6 advances in hybrid imaging technology allow the combination of high spatial resolution and anatomical detail with molecular assessments of disease activity. This 7 8 provides the ability to identify vulnerable plaque characteristics and differentiate 9 active and quiescent disease, with the potential to improve patient risk stratification. Combined positron emission tomography and computed tomography is the 10 11 prototypical non-invasive hybrid imaging technique for coronary artery plaque 12 assessment. In this review we discuss the current state of play in the field of hybrid 13 coronary atherosclerosis imaging.

15 KEYWORDS

- 16 Vulnerable plaque
- 17 Atherosclerosis
- 18 Computed tomography coronary angiography
- 19 Myocardial perfusion
- 20 Positron emission tomography
- 21 Fractional flow reserve
- 22 Coronary physiology
- 23

24 **ABBREVIATIONS**

- 25 CMR Cardiac magnetic resonance
- 26 CT Computed tomography
- 27 CTCA Computed tomography coronary angiography
- 28 FFR Fractional flow reserve
- 29 FFRct Computed tomography coronary angiography-derived fractional flow
- 30 reserve
- 31 MRCA Magnetic resonance coronary angiography
- 32 PET Positron emission tomography

34 INTRODUCTION

Cardiovascular disease is the leading cause of death globally, despite advances in 35 risk stratification, diagnostic tools and preventative therapies (1). Consequently, 36 37 there remains major interest in refining our current methods of diagnosis and risk stratification to better individualise preventative therapies. Myocardial infarction is 38 most commonly caused by rupture of atherosclerotic plaque. Plaques that are prone 39 to rupture have certain common characteristics that together define the vulnerable 40 plaque. Vulnerable plaques have played an integral role in our understanding of 41 42 atherosclerosis and cardiovascular disease, with extensive research conducted to better characterise and identify these lesions (2). However, appreciation of the fact 43 that the majority of vulnerable plaques ruptures are clinically silent has led to a 44 45 paradigm shift in atherosclerotic plaque imaging; focus has shifted from the level of the individual plaque to the patient (3), and from invasive to non-invasive imaging 46 modalities. This change has coincided with advances in non-invasive imaging 47 48 techniques which now facilitate comprehensive assessments of plaque characteristics and disease activity across the coronary vasculature. Hybrid 49 50 cardiovascular imaging is at the frontier of clinical research in this field, although it has yet to become adopted for routine clinical use. 51

52

53 HYBRID IMAGING: RATIONALE AND CONCEPTS

The pathophysiology of atherosclerosis and the vulnerable plaque is well-described (2). There are hallmarks characteristics of high-risk plaque that have been identified on histology, invasive intracoronary imaging and computed tomography coronary angiography (CTCA) which serve as specific targets for hybrid coronary imaging. The prototypical thin-cap fibroatheroma features inflammation (predominantly

macrophage infiltration), a large lipid-rich necrotic core, a thin fibrous cap (<65 μ m), 59 superficial microcalcification and plague haemorrhage. Notably, these findings are 60 independent of stenosis severity; myocardial infarction is often due to plague rupture 61 62 in non-obstructive lesions (4, 5). As our ability to image these plaques has improved it has become clear that the majority of these lesions appear to either heal or rupture 63 sub-clinically with only very few leading to myocardial infarction. In the landmark 64 PROSPECT trial (4), 596 thin-cap fibroatheromas were identified on virtual-histology 65 66 intravascular ultrasound (IVUS) in a cohort of 697 patients with acute coronary syndrome. After 3 years' follow-up, only 31 myocardial infarctions, cardiac arrests or 67 cardiac deaths occurred. Of these, 14 were related to the original culprit lesion. The 68 69 recent Lipid Rich Plague study demonstrated that the lipid core burden index on near-infrared spectroscopy IVUS predicted both culprit and non-culprit major adverse 70 cardiovascular events at 24 months (6). The value of invasive assessments of 71 plaque morphology and plaque-directed therapies has therefore been questioned. 72 Vulnerable plaque assessment appears to be of greater value at the level of the 73 74 patient, using non-invasive imaging to interrogate the entire coronary vasculature. Those subjects in whom adverse plague characteristics are identified are at 75 increased risk of future events, although the originally identified lesion may not itself 76 77 result in a clinical event. As such, total atherosclerotic burden has yet to be superceded for prognostic purposes by any plaque-level imaging (3). 78

79

Hybrid imaging techniques combine two different modalities, taking advantage of
their individual strengths to provide a comprehensive dataset. A modality with high
temporal and spatial resolution is required to provide anatomical detail and
assessments of soft tissue composition. For the coronary arteries, the most common

84	of these is CTCA, although cardiovascular magnetic resonance (CMR) can also be
85	utilised. For hybrid coronary plaque assessment, this dataset is most commonly
86	fused with positron emission tomography (PET). This allows interrogation of plaque
87	biology and potentially any disease process but requires appropriately targeted
88	radiotracers. The ability of hybrid PET-CT and PET-MR to provide this breadth of
89	information about anatomy, plaque composition and disease activity make them
90	exciting techniques with which to study coronary atherosclerosis.

91

92 HYBRID IMAGING: PLAQUE CHARACTERISTICS

Non-invasive imaging of vulnerable plaque morphology has been extensively studied

94 with CT and MR as described below.

95

96 Computed tomography coronary angiography

CTCA has been at the forefront of coronary plaque characterisation for many years, 97 98 with studies demonstrating close correlation between CTCA and intravascular imaging findings of thin-cap fibroatheromas (7, 8). There are several classic CTCA 99 features of vulnerable plaque: low-attenuation (<30 Hounsfield Units), positive 100 101 remodelling (commonly defined as a remodelling index >1.1), spotty calcification and 102 the napkin-ring sign (low-attenuation plaque core with a rim of higher attenuation). 103 Early data from Motoyama et al described the increased prevalence of low attenuation plague (79% vs 9%), positive remodelling (87% vs 12%) and spotty 104 105 calcification (63% vs 21%) in culprit lesions compared to stable lesions (9). Further 106 prospective data demonstrated an increased rate of acute coronary syndromes in patients with high-risk plaque (16.3% vs 1.4% at mean follow-up of 3.9 ± 2.4 years, 107 108 hazard ratio [HR] 8.24 (95% confidence interval [CI] 5.26 - 12.96)) (10). There now

109 exists a large body of non-randomised evidence supporting these findings (5, 9-12). 110 Recent analyses from the two largest randomised trials of CTCA in symptomatic patients with suspected stable coronary artery – the Prospective Multicenter Imaging 111 112 Study for Evaluation of Chest Pain (PROMISE) and Scottish Computed Tomography of the Heart (SCOT-HEART) trials – have added further weight to the prognostic 113 power of CTCA assessments of vulnerable plaque. In PROMISE, 676 (15%) patients 114 had high-risk plaque, conferring a greater risk of major adverse events after 115 adjustment for significant stenoses and atherosclerotic cardiovascular disease risk 116 117 score (HR 1.72, 95% confidence interval [CI] 1.89 – 3.93) (13). Of note, this incremental prognostic power was seen only in patients with non-obstructive 118 disease. In SCOT-HEART, adverse plaque features, present in 608 (34%) patients 119 120 (40% of patients with non-obstructive plaque and 75% in those with obstructive plaque), conferred a 3-fold higher risk of coronary heart disease death or nonfatal 121 myocardial infarction (HR 3.01, 95% CI 1.61 – 5.63), an effect that was most 122 123 pronounced during short-term follow-up. The high prevalence of adverse plague features demonstrates the relatively low positive predictive value of these findings. 124 The prognostic power of adverse plaque features was not independent of coronary 125 artery calcium score, highlighting the importance of total atherosclerotic disease 126 burden. Additionally, approximately half of patients with subsequent adverse events 127 128 did not have obstructive coronary artery disease, while (14). It is important to note the clinical context of these trials, as vulnerable plaque features may perhaps be 129 more relevant in the acute setting (15) due to the dynamic nature of plague biology 130 131 and the increased use of long-term preventative therapies .

132

133 Cardiovascular magnetic resonance

134 CMR has become an imaging modality of major interest in recent years as a result of improvements in scanner technology and software. CMR offers a multiparametric 135 approach to cardiovascular imaging, providing unparalleled soft tissue 136 137 characterisation that is of particular value in the myocardium, alongside information regarding anatomy, function, perfusion and viability. CMR is also able to characterise 138 coronary atherosclerosis, utilising non-contrast T1-weighted (black blood) imaging to 139 detect methaemoglobin found in acute intraplaque haemorrhage or intraluminal 140 thrombosis. High-intensity coronary plagues correlate well with CTCA findings of low 141 attenuation and positive remodelling, demonstrate increased rates of in-situ 142 thrombus, and have been shown to reduce in intensity after statin therapy (16-20). 143 Meanwhile, novel targeted contrast agents have been developed, such as THI0567-144 targeted liposomal-gadolinium. This agent binds with high affinity to integrin $\alpha 4\beta 1$, a 145 key integrin involved in recruiting inflammatory cells to atherosclerotic plaques, and 146 is able to detect vulnerable aortic plaque in an animal model (21). However, as a 147 148 result of the inferior spatial resolution compared to CTCA, magnetic resonance coronary angiography (MRCA) is largely restricted to the proximal and mid-vessel 149 coronary segments. Sequences such as the Coronary Atherosclerosis T1-weighted 150 Characterization with integrated anatomical reference (CATCH) (22) have been 151 developed to overcome some of these limitations but remain exploratory at this point 152 in time. The majority of clinical research and histological validation has therefore 153 focused on larger, stationary vessels such as the carotid artery. Longer scan times in 154 comparison to CTCA, cost and accessibility are other potential barriers to wider 155 uptake of CMR for imaging of coronary atherosclerosis. 156

157

158 HYBRID IMAGING: DISEASE ACTIVITY

Molecular nuclear imaging techniques utilise targeted probes bound to radioactive 159 isotopes. An understanding of plaque biology and the various components of 160 vulnerable plaque are critical to determine suitable targets for molecular imaging. 161 162 PET has been studied for many years, primarily in other specialties such as oncology. Coronary PET imaging has previously been limited due to poor spatial 163 resolution, partial volume effects and cardiac motion. However, with improvements in 164 scanners and the development of advanced motion correction and co-registration 165 techniques, many of these limitations have been overcome. There is now major 166 research interest in coronary PET imaging for the assessment of disease activity 167 within atherosclerotic plagues. This interest had led to the advent of bespoke tracers 168 targeting specific aspects of plaque biology to complement the use of more 169 170 established radiotracers that have been re-purposed from other fields. 171

172 Positron emission tomography: 18F-fluorodeoxyglucose

18F-fluorodeoxyglucose (18F-FDG) PET has been used widely in oncology for many 173 years and was first utilised to image atherosclerosis in the carotid artery in 2002 (23). 174 As a glucose analogue, it is metabolised and accumulates intracellularly in tissues 175 with high metabolic activity via the glucose transporter protein system. 18F-FDG has 176 therefore been used as a non-specific marker of vascular inflammation in the aorta. 177 carotids and femoral arteries. 18F-FDG uptake has been shown to correlate with the 178 presence of atherosclerosis, features of plaque vulnerability, biomarkers of 179 inflammation (in particular macrophage burden) and clinical cardiovascular risk (24-180 26). 181

Although the 18F-FDG PET is excellent for myocardial viability assessment due to avid uptake in cardiomyocytes, coronary artery uptake is often obscured by the adjacent myocardial signal. This is the main limitation of 18F-FDG for coronary atherosclerosis imaging, even despite dietary restrictions prior to scanning (27-29). 187 18F-FDG may still prove of value in detecting plaque inflammation in the aorta and carotid arteries; large prospective outcome studies are awaited.

189

190 Positron emission tomography: 18F-sodium fluoride

Given 18F-FDG's lack of specificity, other radiotracers have been explored. 18F-191 sodium fluoride (18F-NaF) has been used as a bone tracer and for the detection of 192 bony metastases for many decades but has now found a potential application in 193 hybrid cardiac imaging. The ligand for 18F-NaF is hydroxyapatite, a key component 194 of early bone and vascular calcification. It preferentially binds to micro- rather than 195 macrocalcification due to the higher exposed surface area of hydroxyapatite (30). 196 197 Microcalcification is thought to be an early healing response to cell necrosis and inflammation that precedes the development of larger, macroscopic deposits of 198 calcium which can stabilise plaque. Microcalcification within a thin fibrous cap may 199 also increase local stress and destabilize the plaque, thereby increasing the chance 200 of rupture (31). Coronary microcalcification is therefore a key component of 201 202 vulnerable plaque and a biologically plausible target for imaging, with 18F-NaF providing different information to the more established, stable macrocalcification 203 identified on CT. 204

205

18F-NaF PET-CT was noted to identify aortic plaque in 2010 (32). Subsequent data
has shown that increased 18F-NaF activity can be identified in the coronary arteries,

208 localising to individual plaques and demonstrating excellent inter-observer 209 repeatability (33). This improved ability to detect discrete coronary artery uptake compared to 18F-FDG appears to be due to low 18F-NaF uptake in the adjacent 210 211 myocardium and very high affinity of the tracer for microcalcification (30, 34) Again, 18F-NaF appears to be providing different information to the presence of calcium on 212 CT; in one study, almost a half of patients with a calcium score >1000 Agatston 213 units did not have any coronary 18F-NaF uptake (33). In keeping with the hypothesis 214 215 that 18F-NaF uptake is associated with vulnerable plaque, several clinical studies 216 have demonstrated uptake to be associated with culprit and high-risk coronary plaque as defined by invasive angiography, intravascular ultrasound and CTCA. In 217 the first report, increased 18F-fluoride uptake was observed at the site of the culprit 218 219 coronary plaque in 37 of the 40 patients with recent myocardial infarction (29), a finding supported by two subsequent smaller studies (35, 36). 220

221

222 Recent technological advances have greatly improved the image guality of coronary 18F-NaF PET-CT imaging. These techniques have focused on optimizing image 223 reconstruction and correcting for cardiac, respiratory and gross patient motion, with 224 the important advantage that they can be applied retrospectively to PET datasets 225 (37-39). Newer data have also demonstrated the feasibility of fusing PET data with 226 227 previously acquired CTCA, expanding the potential practical application of this form of imaging (40). 18F-NaF PET-MR is also feasible with lower doses of ionising 228 radiation, but currently cannot provide the spatial resolution of CTCA for coronary 229 artery imaging (41). 230

231

232 **Positron emission tomography: other radiotracers**

Multiple alternative radiotracers, each with a specific target, have been studied for 233 234 use in atherosclerosis (Table 1) (42). 68Ga-DOTATATE, which targets the somatostatin receptor subtype 2 (SSTR2) on the surface of activated 235 236 proinflammatory M1 macrophages, has recently been investigated for coronary atherosclerosis imaging. The Vascular Inflammation imaging using Somatostatin 237 receptor positron emissION tomography (VISION) study utilised RNA sequencing, 238 autoradiology, histology and PET-CT in patients with stable and unstable 239 cardiovascular disease. The investigators elegantly demonstrated exclusive 240 241 expression of SSTR2 in M1 macrophages within atherosclerotic plaque, a strong correlation between SSTR2 expression and 68Ga-DOTATATE activity, and 242 improved discrimination of culprit and high-risk plaque in both the coronary and 243 carotid arteries compared to 18F-FDG (43). Further studies are keenly anticipated. 244 Other examples of alternative radiotracers include 11C-PK11195, which is a specific 245 ligand of the translocator protein that is highly expressed on activated phagocytes, 246 247 and the chemokine receptor CXCR4, which is upregulated in unstable plague and colocalizes with CD68 inflammatory cells. 11C-PK11195 is able to image intraplague 248 haemorrhage in recently symptomatic carotid plaques (44) as well as active disease 249 in large-vessel vasculitis, while CXCR4 has recently shown promise for the 250 distinguising culprit and non-culprit coronary plagues in ST-elevation myocardial 251 infarction (45). 252

253

254 HYBRID IMAGING: PHYSIOLOGY

In addition to measurements of plaque composition and disease activity, PET/CT
also allows for functional assessments of atherosclerotic lesions, whether using
myocardial perfusion studies or non-invasive fraction flow reserve (FFR). This is of

258 interest as recent data have suggested that vulnerable plaque characteristics are 259 associated with haemodynamically significant lesions, and that integrating luminal stenosis, adverse plaque characteristics and adverse haemodynamic characteristics 260 261 (comprised of CT-derived FFR (CT-FFR), delta CT-FFR across the vessel, wall shear stress and axial plaque stress) provides better identification of culprit lesions 262 than each individual parameter (46). Functional coronary assessments may 263 therefore act as both surrogates of adverse plaque features, as well as an additional 264 265 modality to add incremental prognostic information.

266

With invasive FFR now routinely used for decision making in interventional 267 cardiology, interest has grown in the potential for CT-FFR to enhance the role of 268 269 CTCA as a gatekeeper to invasive angiography. Although CT-FFR does not image vulnerable plaque directly, several studies have demonstrated that CTCA 270 assessments of plaque composition improve discrimination of ischaemic lesions as 271 defined by an invasive FFR \leq 0.80 or by decreased quantitative myocardial blood 272 flow (47-51). The number of adverse plaque features appear to increase as stenosis 273 severity increase, but the presence of high-risk plague also remains an independent 274 predictor of ischaemia regardless of stenosis severity, particularly positive 275 276 remodelling (47, 48, 51). The mechanisms for these findings are not clear but reflect the complex relationship between coronary atherosclerosis and ischemia. Positive 277 remodelling and the lipid-rich necrotic core of the vulnerable plaque may predispose 278 279 to local endothelial dysfunction and altered shear stress, thus altering impairing arterial vasomotor function. This hypothesis is supported by a recent exploratory 280 study in which high-risk plague characteristics were more strongly related to invasive 281 pressure measurements during hyperaemia than during rest (52). The adaptive 282

arterial remodelling response to the progression of atheroma – the Glagov
phenomenon – may also reach its limit with a certain volume of plaque, at which
point luminal encroachment, obstruction to flow and ischaemia may rapidly progress
(53).

287

PET provides the gold standard non-invasive assessment of myocardial perfusion. 288 Unlike FFR, PET is able to provide a quantitative assessment of absolute 289 hyperaemic blood flow and myocardial blood flow reserve, thus integrating the 290 291 combined effect of epicardial coronary arterial atherosclerosis as well as microvascular disease. The Prospective Comparison of Cardiac PET/CT, SPECT/CT 292 293 Perfusion Imaging and CT Coronary Angiography With Invasive Coronary Angiography (PACIFIC) study demonstrated PET to be superior to CTCA and single-294 photon emission computed tomography (SPECT) for the diagnosis of ischaemia 295 (based on invasive FFR) (54). Although this study showed only limited additional 296 297 diagnostic value with hybrid imaging, a recent meta-analysis confirmed incremental diagnostic performance with hybrid anatomy/perfusion imaging compared to CTCA 298 (55). Additionally, several retrospective studies consistently described an 299 incremental prognostic value of combining myocardial perfusion imaging and CTCA 300 (56, 57). Further data is now needed to investigate the association between adverse 301 plaque features and myocardial perfusion on PET, to assess whether the latter might 302 also provide a surrogate for unstable coronary plague phenotypes. 303 304

The future of non-invasive coronary anatomy/physiology imaging is therefore
extremely promising; there is great appeal in deriving a hybrid dataset assessing

307 coronary anatomy, plaque morphology, plaque burden and coronary flow at both a
308 lesion and vessel level. Randomized clinical trials will be highly anticipated.

309

310 CURRENT LIMITATIONS

Although the appeal of hybrid imaging is clear, there remain some limitations. Most 311 crucially, until recently there has been a lack of randomised data demonstrating the 312 ability of these imaging techniques to change outcomes. We now have this data 313 supporting the use of CTCA, but there is a major need for similar data demonstrating 314 315 the benefits of hybrid non-invasive ischaemia testing. Furthermore, access to scanners and integration of these imaging techniques into clinical workflows in 316 imaging departments must be considered. Costs is also an issue, particularly with 317 318 regards to the production of bespoke radiotracers for nuclear imaging. Consequently, which test to use in which patients in what setting must be carefully considered, 319 taking into account all of these factors. This is in addition to other clinical factors that 320 321 may influence the choice of test, such as patient age, likelihood of calcific disease, ability to achieve adequate heart rate control, comorbidities and acceptable radiation 322 dose. 323

324

325 CASE 1

A 57-year-old man with type 2 diabetes mellitus presented with a non-ST elevation myocardial infarction (NSTEMI) and underwent percutaneous coronary intervention with two drug-eluting stents to the proximal right coronary artery (RCA) and posterior left ventricular artery (PLV). Six months later, he re-presented with another NSTEMI despite appropriate preventative therapies. Invasive coronary angiography demonstrated severe in-stent restenosis in the proximal stent and a further severe

332 de novo mid-RCA (Figure 1A). Optical coherence tomography demonstrated plaque 333 rupture with red thrombus in the mid-vessel lesion and aggressive neointimal hyperplasia in the proximal lesion (Figure 1B-C). Two drug-eluting stents were 334 335 implanted with a good angiographic result (Figure 1D). The left system, particularly the left anterior descending artery (LAD), had diffuse plaque without obstructive 336 disease (Figure 1E). As part of a research study, the patient underwent 18F-NaF 337 PET-CT six weeks later. This demonstrated three discrete regions of focal uptake in 338 the proximal, mid and distal RCA (arrows). Although the mid-RCA lesion was the 339 340 culprit, the highest uptake was in the proximal restenotic lesion. In contrast to the RCA, the diffusely diseased LAD had non-obstructive calcific plague without high-341 risk features on CTCA and did not demonstrate any 18F-NaF uptake (Figure 1F). 342

343

344 CASE 2

A patient was referred three weeks after an episode of chest pain with a late 345 346 presentation myocardial infarction. He was not revascularized due to established infarction. Six months post-infarct, he underwent 18F-NaF PET-MR. Whole-heart, 3-347 dimensional, contrast-enhanced coronary MR angiography using a respiratory-348 navigated, electrocardiographically triggered, inversion-recovery fast spoiled 349 gradient-echo sequence demonstrated a severe culprit plague in the proximal LAD 350 351 (Figure 1A). Extensive near-transmural infarction in this territory was seen on late gadolinium enhancement (Figure 1B). Focal 18F-NaF uptake was noted in the culprit 352 lesion (Figure 1C-D, white arrowheads) as well as in the aorta and mitral annulus 353 (black arrows). Adapted from Robson et al (41). 354

355

356 CONCLUSIONS

357 The field of cardiovascular atherosclerosis imaging is burgeoning, with increasing 358 availability and uptake of CT and CMR in particular. Hybrid imaging platforms combine these modalities with PET, which together provide detailed information 359 about coronary anatomy, flow, plaque morphology and disease activity, potentially 360 361 expanding our pathophysiological understanding of atherosclerosis and improving risk stratification. Further studies are now required to investigate the clinical utility of 362 this approach and determine whether hybrid imaging of the vulnerable plaque can 363 improve patient outcomes. 364

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- 372

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FIGURE LEGENDS

Figure 1

Invasive angiography (A, D, E), optical coherence tomography (B, C) and positron emission tomography-computed tomography (F, G) for Case 1. Descriptions provided in case vignette.

Figure 2

Cardiac magnetic resonance for Case 2. Descriptions provided in case vignette.

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Table 1 Positron emission tomography radiotracers targeting vulnerable plaque				
Target	Ligand	Radiotracer	Current applications in atherosclerosis	
Macrophage activation	Glucose transporter protein system. Conversion to 18F- FDG-6-phosphate and intracellular accumulation	¹⁸ F-FDG	Prospective <i>in vivo</i> studies in extracardiac atherosclerosis. Correlation with atherosclerosis and cardiovascular risk. Myocardial signal spill-over limits coronary artery assessment (35, 37).	
	Somatostatin receptor subtype 2	68Ga-DOTATATE	Prospective <i>in vivo</i> studies in cardiac and extracardiac atherosclerosis. Correlation with culprit coronary lesions and high-risk features on CTCA (52).	
	Translocator protein 18-kDa	¹¹ C-PK11195	Prospective <i>in vivo</i> study in carotid atherosclerosis (67). Short half-life and variable metabolism.	
	Choline kinase phosphorylated to phosphatidylcholine	¹⁸ F-FCH	Retrospective <i>in vivo</i> study demonstrating correlation with large vessel atherosclerosis and an inverse relationship with calcification (68).	
Apoptosis	Phosphatidylserine	Annexin V	Prospective <i>in vivo</i> pilot data in carotid atherosclerosis (69).	
Нурохіа	Reduction to amine derivative in low oxygen environment	¹⁸ F-FMISO	Prospective <i>in vivo</i> pilot data in carotid atherosclerosis (70).	
	Reduction to amine derivative in low O ₂ environment	¹⁸ F-HX4	Prospective <i>in vivo</i> pilot data in carotid atherosclerosis (71).	
Microcalcification	Hydroxyapatite	¹⁸ F-NaF	Prospective <i>in vivo</i> studies in coronary and extracardiac atherosclerosis. Correlation with culprit coronary lesions and high-risk features on CTCA (37, 41, 44, 72).	
Angiogenesis	αVβ3 & αVβ5 integrin	¹⁸ F-Fluciclatide	Prospective <i>in vivo</i> pilot data in the aorta (73).	
	αVβ3 integrin	¹⁸ F-RGD-K5	<i>Ex vivo</i> study in carotid atherosclerosis (74).	
Thrombus	Glycoprotein IIb/IIIa platelet receptor	¹⁸ F-GP1	Prospective <i>in vivo</i> pilot data in arterial thromboembolic disease (75).	
18F-FDG: 18-fluor 18F-FMISO: 18F-f yl)propan-1-ol, 18F	odeoxyglucose, CTCA: comput luoromisonidazole, 18F-HX4: 18 F-NaF: 18F-sodium fluoride, 18F	ed tomography coron 3F-2-(4-((2-nitro-1H-i -RGD-K5: arginine-c	ary angiography, 18F-FCH: 18F-fluorocholine, midazol-1-yl)methyl)-1H-1,2,3-triazol-1- glycine-aspartate-K5.	













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