Molecular imaging of abdominal aortic aneurysms

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The candidate confirms that the work submitted is her own, except where work which has formed part of jointly authored publications has been included, and that appropriate credit has been given within the thesis where reference has been made to the work of others. The contributions of the candidate and other authors to this work is explicitly indicated in the Preface.

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Abstract

Abdominal aortic aneurysm (AAA) disease is characterised by an asymptomatic, permanent, focal dilatation of the abdominal aorta progressing towards rupture, which confers significant mortality. Patient management and surgical decisions currently rely on aortic diameter measurements via abdominal ultrasound screening. However, AAA rupture can occur at small diameters or may never occur at large diameters. Therefore, there is a need to develop molecular imaging-based biomarkers independent of aneurysm diameter that may help stratify patients with early-stage AAA to reduced surveillance. AAA uptake of $[^{18}F]$fluorodeoxyglucose on positron emission tomography (PET) has been demonstrated previously; however, its glucose-dependent uptake may overlook other key mechanisms. The cell proliferation marker $[^{18}F]$fluorothymidine ($[^{18}F]$FLT) is primarily used in tumour imaging. The aim of the overall study for this thesis was to explore the feasibility of $[^{18}F]$FLT PET / computed tomography (CT) to visualise and quantify AAA in the angiotensin II (AngII)-infused mouse model. The experiments presented in this thesis revealed increased uptake of $[^{18}F]$FLT in the 14-day AngII AAA model than in saline controls, followed by a decrease in this uptake at 28 days. Moreover, in line with the in vivo PET/CT findings, Western blotting of aortic tissue revealed increased levels of thymidine kinase-1 (the substrate of $[^{18}F]$FLT) and nucleoside transporters in the 14-day AngII AAA model than in saline controls, followed by decreased expression levels at 28 days. A pilot experiment further demonstrated that $[^{18}F]$FLT PET/CT could be used to detect an early therapeutic response to oral imatinib treatment in the AngII AAA model. Therefore, $[^{18}F]$FLT PET/CT may be a feasible modality to detect and quantify cell proliferation in the AngII AAA murine model. The findings of this thesis are encouraging for the application of $[^{18}F]$FLT PET/CT in patients with small AAA.
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List of Abbreviations

3D: three-dimensional
AAA: abdominal aortic aneurysm
AngII: angiotensin II
ApoE−/−: apolipoprotein E-knockout
CaCl₂: calcium chloride
CD: cluster of differentiation
CT: computed tomography
CNT: concentrative nucleoside transporter
CTA: computed tomography angiography
EVAR: endovascular aneurysm repair
ENT: equilibrative nucleoside transporter
ECM: extracellular matrix
[¹⁸F]FDG: [¹⁸F]fluorodeoxyglucose
[¹⁸F]FES: [¹⁸F]-fluoro-17b-estradiol
[¹⁸F]FLT: [¹⁸F]fluorothymidine
GLUT: glucose transporter
HPLC: high-performance liquid chromatography
ICER: incremental cost-effectiveness ratio
%ID/g: percentage of injected dose per gram of tissue
IVC: inferior vena cava
MRA: magnetic resonance angiography
MRI: magnetic resonance imaging
MMP: matrix metalloproteinase
MASS: Multicentre Aneurysm Screening Study
[¹⁸F]NaF: [¹⁸F]sodium fluoride
NHS: National Health Service
ER: oestrogen receptor
OAR: open aneurysm repair
PDGF: platelet-derived growth factor
PPE: porcine pancreatic elastase
PET: positron emission tomography
QALY: quality-adjusted life year
ROI: region of interest
SPECT: single-photon emission computed tomography
SMC: smooth muscle cell
SoFIA\textsuperscript{3}: Sodium Fluoride Imaging of AAA
SD: standard deviation
SEM: standard error of the mean
SUV: standardised uptake value
SUV\textsubscript{max}: maximum standardised uptake value
SUV\textsubscript{mean}: mean standardised uptake value
TBR: target-to-background ratio
TAA: thoracic aortic aneurysm
TK: thymidine kinase
TAC: time–activity curve
USS: ultrasound scanning
VEGF: vascular endothelial growth factor
VSMC: vascular smooth muscle cell
Preface

Historical context: The mystery that outsmarted Albert Einstein

Albert Einstein, one of the most influential scientists of the 20th century and father of the revolutionary Theory of Relativity, was faced with a puzzle that was daunting even for someone of his calibre. For years, he had suffered from attacks of upper abdominal pain every 2–3 months, each lasting for 2–3 days and often accompanied by bouts of vomiting. He was nearing 70 years of age, slightly overweight, and a regular pipe smoker; the rest of his history was unremarkable. In late 1948, he finally consulted with Dr Rudolph Nissen, a thoracic surgeon at the Brooklyn Jewish Hospital. Physical examination revealed a pulsating mass deep in his abdomen. Further assessment revealed a weakened, dilated span of the largest artery in the body: the aorta. Einstein, the brilliant theoretical physicist, was diagnosed with an abdominal aortic aneurysm (AAA).

Patients with AAA exhibit a focal ballooning of the abdominal portion of the aorta. Individuals who are male, are older than 60 years of age, are former or current smokers, have a positive familial history, and have high blood pressure fulfil the criteria for being at risk of developing AAA. Most cases of AAA remain clinically silent until they progress to a more critical stage. Aneurysms grow in size over time and eventually rupture, an event that causes profound internal bleeding and inevitable patient death (Sakalihasan et al., 2018). Nissen aimed to prevent Einstein’s aneurysm from rupturing by wrapping the aneurysm of arguably the most intelligent man in the world with a common kitchen item: cellophane. This was a ground-breaking procedure implemented in 1948, with the rationale that the tightly wrapped cellophane would elicit an immune response to the ‘foreign body’. Immune cells would tackle the perceived intruder and, by proximity, the cells within the wrapped aortic section, which would trigger a full-blown immune attack and eventually narrow the aneurysmal aorta. This pioneering measure was an ingenious approach at the time, as a bit of cellophane sufficed to keep Einstein symptom-free and able to continue with his physics research for roughly 7 years. Currently, the wrapping of cellophane has been replaced by the more sophisticated removal of the aneurysmal section and insertion of a specialised graft to conjoin the gap.

In 1955, however, Einstein developed severe abdominal and back pain, which are also common acute signs of worsening prognosis, as the
cellophane was no longer sufficient to restrict the AAA expansion. At this point, he was offered another exploratory surgery that might have added a few more years to his life. Aneurysms tend to exhibit prolonged courses of progression and can be detected well before they rupture. As the risk of AAA rupture and internal bleeding increases with aortic size, early AAA detection combined with the ability to stratify patients according to rupture risk is crucial. Einstein, who at 76 years of age was content with his long and gratifying life, refused the additional life-extending surgery, as he parted with the famous wise words, 'I have done my share, it is time to go. I will do it elegantly' (Zimmermann, 2018).

Motivation

AAA has long represented a significant clinical burden, owing to the lack of effective risk stratification methods and inability to confidently predict the incidence of post-surgical complications. The selection of patients to undergo surgical intervention is based on aneurysm size and growth rate revealed through abdominal ultrasound scanning (USS)-based screening, which is currently implemented in the UK for all men older than 65 years of age. However, the traditional anatomy-based screening of AAA fails to comprehensively elucidate the risk of aneurysm rupture and predisposition to surgery-related complications. Molecular imaging of biological events that are involved in AAA formation and rupture is likely to improve patient risk stratification and help distinguish between patients who would benefit from different therapies by dividing them into ‘watchful waiting’ and ‘imminent surgery’ groups, for example, eventually resulting in decreased AAA morbidity and mortality.

This thesis focusses on the novel application of positron emission tomography (PET) / computed tomography (CT) with a radiotracer that is predominantly used for tumour imaging: $^{18}$F]fluorothymidine ($^{18}$F]FLT). $^{18}$F]FLT is a marker of cell proliferation, thus being a suitable indicator of cancer progression. Meanwhile, there is emerging evidence of early-stage cell proliferation in the pathophysiology of AAA that proceeds to a late-stage process characterised by apoptosis and senescence; therefore, cell proliferation is an interesting molecular target for imaging. The overarching emphasis in this thesis is placed on assessing the feasibility of using $^{18}$F]FLT PET/CT to visualise and quantify cell proliferation in a classical murine model of AAA.

Thesis overview
Four main chapters comprise this thesis to convey the journey of implementing \(^{18}\text{F}\)FLT PET/CT in a AAA mouse model. An overview of the chapters is presented below.

**Chapter 1 (Background):** This chapter provides a comprehensive introduction to the basic concepts underlying AAA in terms of its pathophysiology and clinical issues. Clinical and preclinical imaging is discussed, followed by a particular focus on PET and radionuclide-based tracers. This then sets the stage for the introduction of the overall hypothesis, aim, and research objectives of the thesis.

**Chapter 2 (Methods):** The methods implemented for the experiments described in this thesis and their rationale are outlined in this chapter. Descriptions of the different mouse models used in the overall study are provided, following by details of the methodologies implemented, including histological staining, *in vivo* USS and PET/CT, *ex vivo* gamma counting and autoradiography, and Western blotting. The methods adopted for statistical analyses are also outlined.

**Chapter 3 (Results):** All the key findings of the overall research are systematically reported in this chapter. The presence of cell proliferation in different murine models of AAA was first investigated to prioritise a model to implement for the *in vivo* PET/CT experiments. The uptake of the gold-standard radiotracer \(^{18}\text{F}\)fluorodeoxyglucose (\(^{18}\text{F}\)FDG) was first tested, followed by \(^{18}\text{F}\)FLT uptake in healthy control mice, before finally assessing \(^{18}\text{F}\)FLT uptake in AAA models. The results of a pilot therapeutic-based PET/CT study are also presented in this chapter.

**Chapter 4 (Discussion):** A general discussion of the key results arising from the overall research is included in this chapter, starting with the histological evidence of cell proliferation in AAA and expression of key players in the \(^{18}\text{F}\)FLT mechanistic pathway. A comprehensive discussion of \(^{18}\text{F}\)FLT uptake in the AngII AAA model and PET image analysis is then provided, followed by a discussion of anti-proliferative treatment for AAA. The chapter then ends with potential future directions of AAA research and concluding thoughts.

**Key contributions**

The following are the specific achievements of this thesis:
(i) $[^{18}\text{F}]$FLT uptake on \textit{in vivo} PET/CT and \textit{ex vivo} gamma counting has been characterised for the first time in a classical murine model of AAA, revealing a peak in abdominal aortic uptake in the early stage of the disease;

(ii) the expression levels of key players in the thymidine mechanistic pathway were found to be increased in the early stage of AAA, mirroring the pattern of $[^{18}\text{F}]$FLT uptake;

(iii) $[^{18}\text{F}]$FLT PET/CT was identified as a feasible modality to evaluate the response to an anti-proliferative agent in a classical murine model of AAA, a finding that would benefit from further validation using large sample sizes; and

(iv) an optimised quantification approach was identified for PET image analysis, which may have implications for future analyses of PET data.

\textbf{Dissemination}

The following jointly authored publications and presentations are associated with the research presented in this thesis.

\textbf{Journal articles}


The candidate performed experiments and collected data, analysed data from experiments, wrote the manuscript, and obtained funding. The co-authors jointly performed experiments and collected data, edited the manuscript, produced critical reagents for experiments, provided intellectual input, obtained funding, conceived the idea, and had overall responsibility for the study.

In addition, attention is drawn to the following editorial article that focussed exclusively on (Gandhi et al., 2019):

This is the first study to confirm that cellular proliferation is indeed elevated within the aorta of AngII-infused ApoE−/− mice and that 18F-FLT can non-invasively quantify this process. Moreover, the timing of 18F-FLT uptake closely mirrored histological and proteomic analysis of aneurysmal tissue, with cellular proliferation appearing to be higher in the early stages of aneurysm development before later tailing off. This study has therefore provided novel insight into the cellular mechanisms observed in maturing aneurysmal tissue, and demonstrated 18F-FLT PET as a useful technique to detect these cellular changes (Syed et al., 2019).


The candidate conducted the systematic review, analysed data, wrote the manuscript, obtained funding, and conceived the idea. The co-author edited the manuscript, provided intellectual input, obtained funding, and had overall responsibility for the study.

**Conference presentations**


The candidate performed experiments and collected data, analysed data from experiments, obtained funding, plans to create the poster, and plans to present the poster at the conference.

- Bashair Alhummiany, **Richa Gandhi**, Stephen J. Archibald, Christopher Cawthorne, Marc A. Bailey, Charalampos Tsoumpas. Effect of different ROI definitions on the quantification of SUV. *European Molecular Imaging Meeting 2019*, Glasgow, Scotland, UK (Poster presentation)

The candidate supervised the MSc student (Bashair Alhummiany), conceived the idea for the project, collected data, provided intellectual input, edited the poster, and presented the poster at the conference. A copy of the poster is provided in Appendix A.
• Richa Gandhi, John D. Wright, Joanna Koch-Paszkowski, Stephen J. Archibald, Christopher Cawthorne, Marc A. Bailey, Charalampos Tsoumpas.\(^{[18]}\)F-fluorothymidine positron emission tomography/computed tomography of experimental abdominal aortic aneurysms in murine models. *European Molecular Imaging Meeting 2019*, Glasgow, Scotland, UK (Poster presentation)

The candidate performed experiments and collected data, analysed data from experiments, obtained funding, created the poster, and presented the poster at the conference. A copy of the poster is provided in Appendix A.

• Richa Gandhi, Michael Shires, Stephen J. Archibald, Marc A. Bailey Christopher Cawthorne & Charalampos Tsoumpas. Uptake of 18F-fluorothymidine in experimental abdominal aortic aneurysms of small rodents detected using PET/CT. *European Association of Nuclear Medicine 2018*, Dusseldorf, Germany (ePoster presentation)

The candidate performed experiments and collected data, analysed data from experiments, obtained funding, created the poster, and presented the poster at the conference. A copy of the poster is provided in Appendix A.

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Chapter 1 Background

This chapter presents a general introduction to aneurysms, before narrowing down to aortic aneurysms. A brief comparison between AAA and thoracic aortic aneurysms (TAA) is presented in terms of cellular biology and genetics. The spotlight then turns to AAA with a focus on important clinical aspects related to screening and diagnosis, along with pathophysiological and biomechanical characteristics that contribute to AAA formation. Next, the importance of preclinical imaging is highlighted with a brief overview of different imaging modalities in the context of AAA, followed by a discussion of PET and quantitative image analysis. Finally, $[{ }^{18} \text{F}]$-based PET radiotracers are discussed with a particular emphasis on $[{ }^{18} \text{F}]$FLT, in addition to literature focusing on recent studies investigating AAA using PET. Finally, the overall hypothesis, aim, and research objectives of the thesis are presented.

1.1 Aneurysms

The word ‘aneurysm’ is derived from the Greek word *aneurysma*, meaning dilation. An aneurysm is a focal dilation of a blood vessel to greater than 1.5 times its normal size, predominantly due to weakening of the arterial wall. Aneurysms can develop in various arteries throughout the body, with the most common sites being branches of the Circle of Willis in the brain and the aorta. Aneurysms that develop in the cerebral vasculature are referred to as intracranial aneurysms. Rupture of these aneurysms commonly causes subarachnoid haemorrhage, a severe type of stroke that involves bleeding into the area between the arachnoid membrane and pia mater surrounding the brain (Keedy, 2006). Meanwhile, the aorta is the largest artery in the body that originates in the left ventricle of the heart and passes through the chest and abdomen. AAA, which develops in the part of the aorta that passes through the abdominal cavity, is the most common type of aortic aneurysm. TAA develops in the part of the aorta that passes through the chest (Harris et al., 2016). Aneurysms can also occur in peripheral arteries, which are less likely to rupture than aortic aneurysms. Some examples include aneurysms of the popliteal artery in the leg behind the knee, carotid artery in the neck, splenic artery near the spleen, femoral artery in the groin, and mesenteric artery that supplies the intestines.

1.2 Aortic aneurysms
Aortic aneurysms can occur at different sites, which are named accordingly, such as ascending aortic aneurysms, aortic arch aneurysms, descending TAA, thoracoabdominal aortic aneurysms, and AAA (Figure 1.1). Increasing evidence suggests that TAA and AAA are unique pathophysiological entities with different characteristics (Ruddy et al., 2008, Kuivaniemi et al., 2015).

The aorta contains intimal, medial, and adventitial layers; however, these layers in the thoracic aorta are structurally different from those in the abdominal aorta. Essential components of the medial layer include lamellar units, which are unique fibromuscular layers of smooth muscle cells (SMCs) enveloped by structural proteins such as collagen, elastin, and proteoglycans that facilitate the dispersal of stress and supply elasticity. The media of the thoracic aorta contains approximately 60 lamellar units that are further distributed across avascular and vascular regions. Meanwhile, the abdominal aorta contains roughly 30 lamellar units and is exclusively avascular (Kuivaniemi et al., 2015). Variations in the delivery of oxygen, growth factors, and nutrients to cells of the thoracic and abdominal aortic media are likely to play a role in the variations observed in vascular remodelling between the two aortic regions. The smaller number of lamellar units, avascular form, and thinner wall of the abdominal aorta in comparison to the thoracic aorta may also contribute to the greater incidence of AAA than of TAA (Guo et al., 2006, Ruddy et al., 2008).
Figure 1.1 Anatomy of the aorta.
The role of genetics in TAA and AAA is well studied. Many cases of TAA are associated with genetic syndromes, such as Marfan syndrome due to mutations in fibrillin-1, Ehlers–Danlos syndrome caused by mutations in type III collagen, and Loeys–Dietz syndrome caused by mutations in transforming growth factor-β receptors 1 and 2 (Elefteriades and Pomianowski, 2013). TAA may also occur in individuals with genetic predispositions in the form of familial TAA and aortic dissection, the risk of which is also increased with mutations in vascular smooth muscle cell (VSMC) contractile proteins (Pannu et al., 2006, Guo et al., 2007). Meanwhile, specific mutations in relation to genetic predispositions in AAA are not entirely clear, although AAA is associated with a first-degree familial history of aneurysmal disease (Guo et al., 2006, Harris et al., 2016). A meta-genome-wide association study by Jones et al., to which Leeds contributed more than 500 patient samples, verified the roles of a number of risk loci that appear to exhibit particular specificity for AAA: **BCAR3, SORT1, NOTCH2, TDRD10, UBE2W, CDKN2B-AS1/ANRIL, LRP1, NAB2, FGF9, and PLTP** (Jones et al., 2017). The formation of both TAA and AAA are complex processes resulting from the interaction of various contributory factors. Whilst TAA is primarily a genetic condition with a single gene mutation resulting in aneurysm development in a specific patient, AAA is a more degenerative disease of older age with a polygenic footprint of an array of susceptibility genes. More studies are needed to build a more complete understanding of the differences in aetiologies and pathogeneses of aortic aneurysms. The remainder of this chapter lays the foundation for the AAA-centred experiments presented in this thesis, which were conducted to elucidate the molecular mechanisms of AAA.

### 1.3 Abdominal aortic aneurysms

#### 1.3.1 Clinical features

AAA disease is characterised by localised dilatation of the abdominal aorta from a normal diameter of approximately 10–20 mm to an aneurysmal diameter of 30 mm or greater. AAA is asymptomatic, yet simultaneously progressive towards rupture and profound internal bleeding, resulting in a high mortality rate in the range of 59–83% for patients with ruptured AAA who either cannot make it to the hospital in time or do not undergo surgery (Blanchard et al., 2000, National Institute for Health and Clinical Excellence, 2008, Kumar et al., 2017). From 2005 to 2012, 39,740 aneurysm-associated deaths occurred in England (Karthikesalingam et al., 2016). Ruptured
aneurysms manifest with severe abdominal or back pain, hypotension, and shock. Various environmental and genetic risk factors contribute to the development and progression of AAA; the predominant risk factors of AAA include increased age, male gender, cigarette smoking, and elevated diastolic blood pressure (Blanchard et al., 2000, Thompson et al., 2002, Sakalihasan et al., 2018). Meanwhile, the risk of aneurysm rupture may be enhanced by rapid aortic expansion, female gender, cigarette smoking, and hypertension. Although it is suggested that patients with AAA undergo interventions that lower cardiovascular risk, in addition to smoking cessation, there remains limited evidence that these approaches reduce AAA-associated morbidity and mortality (Toczek et al., 2016). Therefore, there is an unmet clinical need for interventions that specifically target AAA.

### 1.3.2 Screening and monitoring

Early-stage AAA does not manifest with any apparent signs or symptoms; therefore, it is usually diagnosed incidentally during unrelated medical check-ups or more recently through screening programmes, which have been formally launched in several countries including the US, the UK, and Sweden (Davis et al., 2013, Guirguis-Blake et al., 2014, Johansson et al., 2015, Zarrour et al., 2016). The National Health Service (NHS) AAA Screening Programme was fully implemented in 2013 in England to offer routine USS-based screening for all men aged older than 65 years. This programme was established based on the results of the UK Multicentre Aneurysm Screening Study (MASS), which demonstrated a nearly 50% reduction in AAA-related mortality risk after 13 years in men invited for screening compared with that in men who did not undergo screening (0.66% vs. 1.12%). The UK MASS further showed that this reduction was associated with an incremental cost-effectiveness ratio of £7600 per quality-adjusted life years (QALYs) gained at 10 years (Ashton et al., 2002, Thompson et al., 2012).

Cost-effectiveness is an important aspect to consider when implementing AAA screening programmes in the general population. If health benefits are assessed in terms of QALYs, then the cost-effectiveness of AAA screening may be evaluated based on the incremental cost-effectiveness ratio (ICER) of screening vs. no screening (Equation 1.1):

\[
ICER = \frac{\text{cost}_{\text{screening}} - \text{cost}_{\text{no_screening}}}{\text{QALYs}_{\text{screening}} - \text{QALYs}_{\text{no_screening}}}
\]

**Equation 1.1 Incremental cost-effectiveness ratio**
Based on economic modelling that incorporates AAA prevalence, size
distribution at initial screening, and screening attendance to determine the
ICER, AAA USS-based screening implemented in England for men aged 65
years and older remains highly cost-effective per QALYs gained (Glover et
al., 2014, Zarrouk et al., 2016). The gain in QALYs is attributed to screening
being used to detect when AAA reaches the intervention threshold, i.e. the
point at which the risk of intervention no longer outweighs the risks of
intervention.

1.3.3 Intervention thresholds

AAA is currently managed through careful monitoring of aortic diameter
based on USS; the frequency of surveillance in patients reflects AAA size.
Small AAA (30–49 mm in diameter) typically grow at a rate of 2–4 mm/year;
this growth rate increases to >7 mm/year beyond a diameter of 55 mm
(Vega de Céniga et al., 2006, Huang et al., 2019). Patients with small AAA
are regularly monitored (e.g. annually for AAA with diameters of 30–45 mm
and every 3 months for AAA with diameters of 45–55 mm) until the (a)
diameter exceeds 55 mm or (b) growth rate exceeds 10 mm/year; when
either of these criteria are fulfilled, patients are referred for surgery.
International guidelines, however, suggest a diameter threshold of 50 mm in
women, based on the observations that women have a three-fold higher risk
of AAA rupture than men and the mean aortic diameter preceding AAA
rupture is larger in men than in women (Powell et al., 1996, Brown and
Powell, 1999, Moxon et al., 2010). Aortic size has been shown to be
correlated with rupture risk. Oliver-Williams et al. recently reported rupture
risks of less than 0.5% for men with AAA less than 55 mm in diameter (Table
1.1) (Oliver-Williams et al., 2019).
Table 1.1 Estimated annual risk of AAA rupture. Sources: (Kent, 2014, Ullery et al., 2018)

<table>
<thead>
<tr>
<th>AAA Diameter (mm)</th>
<th>Rupture Risk (%/y)</th>
</tr>
</thead>
<tbody>
<tr>
<td>30–44</td>
<td>0.03</td>
</tr>
<tr>
<td>45–49</td>
<td>&lt;0.3</td>
</tr>
<tr>
<td>50–54</td>
<td>0.40</td>
</tr>
<tr>
<td>55–59</td>
<td>&lt;10</td>
</tr>
<tr>
<td>60–69</td>
<td>10–20</td>
</tr>
<tr>
<td>70–79</td>
<td>20–40</td>
</tr>
<tr>
<td>≥80</td>
<td>30–50</td>
</tr>
</tbody>
</table>
Furthermore, for each 5-mm increase in AAA diameter, the mean aortic growth rate increases by 0.59 mm/y, with an increase in the AAA rupture rate by a factor of 1.91. These values work out to more than 7 years of surveillance for a 30-mm AAA and about 8 months of surveillance for a 50-mm AAA to reduce the risk of AAA exceeding 55 mm in diameter to less than 10% in men (RESCAN Collaborators et al., 2013). These results indicate that a watchful waiting period of several years is suitable for patients with small AAA. Surgical intervention is offered once the aortic diameter exceeds 55 mm; adoption of this threshold in the clinic is supported by evidence from four different trials (the UKSAT, ADAM, CAESAR, and PIVOTAL studies) that demonstrates no significant advantage to early surgical intervention for small AAA (Filardo et al., 2015). Whether nonsurgical options may be effective for small AAA is currently unknown.

The important point to note here is that although there is extensive ongoing research to establish more reliable and informative risk markers of AAA, the only indicator that is used consistently in the clinic is aortic size. This is the simplest and most convenient parameter to extract using USS, but may not always accurately reflect prognosis, as small AAA have been reported to rupture, whilst large AAA have been reported as remaining stable (McGloughlin and Doyle, 2010). Some other proposed risk markers that are under ongoing validation are aortic wall stress, the finite element analysis rupture index, vessel asymmetry, intraluminal thrombus growth, and the rupture potential index; in addition, aortic wall stress and intraluminal thrombi have been demonstrated to be independent of aortic diameter, hence justifying the notion of utilising more than aortic size alone when evaluating AAA rupture risk (Stenbaek et al., 2000, Fillinger et al., 2003, Vande Geest et al., 2006, Doyle et al., 2009, Zhu et al., 2020). However, these potential markers require substantial investigation before being introduced into clinical practice.

1.3.4 Interventions

Surgical intervention for AAA is currently offered once the aortic diameter reaches 55 mm in maximal diameter. The exact timing of surgery is often difficult to determine due to the risks associated with AAA progression and surgery invasiveness; the 55-mm aortic diameter acts as a threshold at which point the annual risk of AAA rupture is similar to the mortality risk of surgery (Moxon et al., 2010, Cafueri et al., 2012). It is well established that the low rupture rate in small aneurysms and risks correlated with surgical
intervention do not sufficiently justify routine repair of small AAA (Filardo et al., 2015, Chuen and Theivendran, 2018).

The two main types of surgical intervention currently offered are open aneurysm repair (OAR) and endovascular aneurysm repair (EVAR). OAR has been performed since the mid-1950s and involves making a large abdominal incision to expose the abdominal aorta, followed by insertion of a synthetic graft that is sewn in place to replace the weakened portion of the aorta. OAR is a highly invasive procedure that confers considerable mortality and morbidity risks due to which many patients are denied for surgery. On the other hand, EVAR involves making an incision in the groin area, followed by insertion of a collapsed synthetic graft through the femoral artery that is opened inside the aneurysm with X-ray guidance and fixed in place with a stent. EVAR was introduced with the promise of being a solution to the pitfalls of OAR and comes with the advantages of requiring smaller incisions, conferring fewer perioperative complications, being less painful for the patient, being associated with lower mortality and morbidity rates, and requiring less hospitalisation in comparison to OAR (Greenhalgh, 2004, Prinssen et al., 2004, Kontopodis et al., 2020). However, EVAR is also reported to be associated with an increased frequency of readmissions due to the lack of long-term durability (Sandford et al., 2014). The short-term (i.e. 0–6 months following surgery) survival benefits of EVAR compared to those of OAR (4.6 vs. 10.0%/person-years) appear to be reduced beyond 8 years following surgery (1.3 vs. 0.2%/person-years) (Patel et al., 2016). The National Institute for Health and Care Excellence guidelines have previously suggested removing EVAR as an option from the NHS for elective cases (National Institute for Health and Clinical Excellence, 2018); however, these have since been updated to include EVAR in cases wherein OAR is contraindicated (National Institute for Health and Clinical Excellence, 2020). Nonetheless, the implementation of EVAR in AAA management remains controversial, and its effects on the long-term prognosis of patients with AAA remain under scrutiny to facilitate improvements in the procedure.

1.3.5 AAA development

The pathophysiology of AAA involves active and dynamic pathological remodelling of the connective tissue within the aortic wall; however, the exact sequence of events remains largely unclear (Davies, 1998, Kuivaniemi et al., 2015, Petsophonsakul et al., 2019). During early-stage AAA formation, changes in haemodynamic stress and aortic wall elasticity occur, likely in response to one or more of the causal risk factors described in section 1.3.1,
such as cigarette smoking and increased diastolic blood pressure. Macrophages also infiltrate the vessel wall due to increased oxidative stress, leading to protease secretion that deteriorates the vessel wall via extracellular matrix (ECM) degradation; the upregulation of pro-inflammatory cytokines and chemokines further contributes to compromised aortic wall integrity (Henderson et al., 1999, Thompson et al., 2002, Cafueri et al., 2012, Kuivaniemi et al., 2015, Sun et al., 2018). Although extensive research has focussed on the role of inflammation in AAA, inflammatory mechanisms only account for part of the overall story (Patel et al., 1996). This is further evidenced by the lack of definitive findings of anti-inflammatory therapy attenuating AAA growth, with some enhancing AAA growth (Yoshimura et al., 2018, Tedjawirja and de Waard, 2019).

It is increasingly recognised that the proliferation of VSMCs, which reside in the tunica media and are responsible for controlling blood pressure and flow distribution, contributes to AAA development (Owens et al., 2004, Chasman and Lawler, 2017). An early study in this field demonstrated that VSMC migration and proliferation are enhanced in AAA tissues compared to that in aortic occlusive disease and normal inferior mesenteric artery tissues (Patel et al., 1996). The same team also showed that these aortic SMCs produce increased levels of matrix metalloproteinases (MMPs) compared to those produced in control arterial tissues (Patel et al., 1996); these MMPs contribute to the degradation of elastin and various ECM proteins, progressively leading to degradation of the aortic wall. In the following year, another study revealed that the SMC density in the medial layer is significantly decreased in AAA tissues in association with SMC apoptosis, reflecting its role in aortic wall degradation (Lopez-Candales et al., 1997).

VSMCs usually exhibit a contractile, quiescent phenotype, characterised by the expression of proteins such as α-smooth muscle actin, smooth muscle myosin heavy chain, and smooth muscle-22α; however, when stimulated by injury or stress, they undergo a phenotypic switch to a proliferative and migratory state (Owens et al., 2004, Ailawadi et al., 2009, Michel et al., 2018, Petsophonsakul et al., 2019). In 2009, Ailawadi et al. demonstrated that this phenotypic switching is an early event in AAA, based on the downregulation of SMC marker genes and upregulation of MMPs (Ailawadi et al., 2009). One of the histopathological characteristics of AAA is extensive degradation of the aortic ECM, which is facilitated by elastolytic enzymes that are produced by VSMCs when they exhibit the proliferative phenotype. The ECM is a fundamental component of blood vessels that is produced by the cells located in vessel walls and contains fibrous proteins such as
collagen and elastin, proteoglycans such as versican, and glycoproteins such as thrombospondins and fibronectin. There are numerous kinds of ECM with varying compositions that each play different roles in facilitating different pathophysiological processes. Proteomics-based approaches have revealed distinct differences in the ECM composition between AAA and non-AAA tissues (Didangelos et al., 2011). Some proteins that have been identified to exhibit changes in AAA include collagen XII, which is found in tissues subject to high tensile forces; versican, a major extracellular proteoglycan; periostin and thrombospondins, which contribute to cellular adhesion and spreading; ACLP, which is known to be associated with collagen and fibrosis; and tenasin, which is correlated with macrophage accumulation (Satta et al., 1997, Layne et al., 1998, Theocharis et al., 2001, Didangelos et al., 2011). These ECM proteins significantly modulate VSMC phenotypic switching, and their reduction reflects the pathological remodelling of the ECM and VSMCs in the AAA disease course. Finally, calcification—the deposition of calcium phosphate crystals in the medial layer of the vessel wall—further contributes to vessel stiffening, and hence, VSMCs have been shown to contribute to vascular calcification (Proudfoot et al., 2000, Shanahan et al., 2011, Wu et al., 2013). Overall, synthetic VSMCs are associated with a loss of contractile proteins, greater production of elastolytic enzymes that degrade the ECM, the release of extracellular vesicles that promote inflammation and calcification, and presence of reactive oxygen species (Kapustin and Shanahan, 2016, Schurgers et al., 2018). These key biological processes thus collectively contribute to inducing vessel dilatation and aneurysm progression.
Figure 1.2 Flow diagram of the mechanisms contributing to AAA formation and progression.
Late-stage AAA is understood to involve an amalgamation of VSMC apoptosis, upregulated inflammatory cell infiltration, thinning of the aortic wall due to an imbalance between the concentrations of collagen and elastin, and calcification; these components collectively exacerbate focal aortic wall weakening, eventually resulting in rupture (MacSweeney et al., 1994, Wills et al., 1996, Lopez-Candales et al., 1997, Proudfoot et al., 2000, Rowe et al., 2000, Clement et al., 2019, Quintana and Taylor, 2019). The prediction of rupture risk is an important area of research, and in this context, mechanical properties of the aorta in relation to breaking stress and stiffness are well studied. Per the Law of Laplace, intraluminal pressure, wall thickness, and vessel diameter affect the wall tension necessary to counter a given internal fluid pressure, which in turn affects wall stress. Aneurysm rupture occurs when the wall stress exceeds wall tension (Vorp, 2007).

Studies of the biomechanical properties of aortic aneurysms typically involve computational modelling and finite element methods to apply an array of proposed criteria that may help predict rupture risk: expansion rate, wall tension, wall stiffness, peak wall stress, intraluminal thrombus thickness, and surface area (Hall et al., 2000, Hatakeyama et al., 2001, Metaxa et al., 2017, van Disseldorp et al., 2019). For example, Conlisk et al. demonstrated that sites of greater curvature exhibit increased stress, which then corresponds with a greater susceptibility of rupture (Conlisk et al., 2017). Importantly, some AAA that are smaller in diameter than the surgical intervention threshold exhibit high peak wall stress (van Disseldorp et al., 2019). Recent data have suggested that biomechanical assessment may be superior to using only the aortic diameter to evaluate rupture risk (Polzer et al., 2020). Basing clinical assessments on biochemical parameters in this way may be a promising adjunctive tool for precise diagnosis and disease course prediction, although their application in the clinic is likely to be hampered by the logistical difficulties of obtaining these measurements at repeated time points to suffice precise risk prediction as an aneurysm grows and remodels. In this case, more subtle stratification biomarkers based on underlying molecular mechanisms that come into play before physical appearance of the disease may be more feasible to implement in the clinic to predict whether patients have a high or low risk of rapidly progressive disease.

1.3.6 Clinical imaging of AAA

Current clinical interventions for AAA focus on physically excluding the aneurysm, which themselves are associated with significant risks. Therapies that instead inhibit the progressive deterioration and molecular dysfunction
of the aortic wall would decrease the necessity for surgery. Medical imaging plays a key role in better understanding the adverse molecular mechanisms underlying early AAA formation. Currently, USS is used in the clinic to screen for AAA and monitor disease progression over time. Whilst it is effective in revealing changes in aortic size and shape over time, the technique is not without its limitations. The most salient limitation is that USS is operator-dependent; therefore, results from USS depend on a sonographer’s experience and accuracy. This creates a two-variable problem: individual patient factors and sonographer accuracy. Implementing quality control is inevitably difficult, having to rely on standardised guidelines to reduce inter-patient and inter-operator variability. An increasingly common problem in Western societies is patient size. Larger patients are difficult to image via USS due to their greater mass and depth of tissue, which attenuates the sound waves reflecting from deeper structures in the body, such as the aorta, and back to the transducer. Ironically, it is overweight and obese patients who tend to carry a greater risk of developing AAA and would most benefit from screening and monitoring. Trapped air and gas can further obstruct the findings of USS; therefore, imaging of regions that are close to the air-filled bowel is not always ideal (Kornezos et al., 2010, Dobrucki and Sinusas, 2020). That being said, with a sensitivity of 95% and specificity of nearly 100%, USS is predominantly used in the clinic to screen for AAA because of its ease of operability, low costs, low post-imaging workload and instantaneous results, and widespread availability (Kumar et al., 2017).

Abdominal CT without contrast enhancement is equivalent to USS in terms of detecting AAA and is typically recommended in patients who cannot undergo USS. Meanwhile, CT with contrast enhancement can provide information on the presence of thrombus or dissection flaps, but may not provide as much information about branch vessel involvement and smooth three-dimensional (3D) rendering that CT angiography (CTA) may offer (Reis et al., 2017). CTA is another commonly used modality in the clinic for AAA. It is notably valuable in pre-intervention planning, such as in assessing aortic size, the presence of intraluminal thrombus, the involvement of visceral arteries, and extension to other parts of the aorta, and is even superior to USS in identifying and measuring the size of aneurysms (Kumar et al., 2017, Kyriakou et al., 2020). Use of CTA for maximum diameter measurements has been shown to be associated with high intra-observer reproducibility (Mora et al., 2014). However, CTA confers high radiation doses (Kumar et al., 2017).
Magnetic resonance imaging (MRI) has the ability to provide accurate anatomical information of AAA with excellent soft-tissue contrast and reduced exposure to ionising radiation compared to CT. MRI has previously been used to confirm the diagnosis of thrombus and assess blood flow in aortic diseases in patients with aortic aneurysms and dissection (Honda et al., 1999). Four-dimensional flow MRI has further been utilised to analyse changes in blood flow patterns in the abdominal aorta (Liu et al., 2018). Although long scan durations are a notable challenge with the use of MRI, advances in MRI-based approaches have revealed that 3D non-contrast black-blood MRI is associated with decreased scan durations with retention of image quality (Zhu et al., 2019). In a prospective study, this method was shown to reveal intraluminal thrombi in patients with AAA, with the finding that active changes in intraluminal thrombi are associated with large aortic diameters and rapid AAA growth (Zhu et al., 2019). Furthermore, magnetic resonance angiography (MRA) has a similar sensitivity to that of CTA and has also been shown to reveal endoleaks that could not be visualised using CT angiography (Wicky et al., 2003). MRA is particularly useful in cases wherein intravenous contrast is contraindicated, such as in cases of allergic responses or renal dysfunction, and is associated with a lack of ionising radiation (Kumar et al., 2017). However, the benefits of MRI and MRA are hampered by key disadvantages such as higher associated costs, reduced availability of scanners, motion artefacts and patient claustrophobia due to long scanning durations, and contraindications in patients harbouring pacemakers and metal clips.

Meanwhile, PET imaging has the potential to offer more precise functional information at the molecular level for risk stratification purposes as an adjunctive tool to USS-based screening and surveillance. As the primary focus of this thesis, section 1.5 is dedicated to a discussion of PET imaging in the context of AAA. The use of PET with a suitable radiotracer may be beneficial when applied at the time of AAA detection to help personalise a surveillance regimen or intervention threshold. Furthermore, there is currently no pharmacological treatment for AAA, and the indication for surgery is based solely on aortic size from USS. Establishing a pharmacological treatment that could slow or even reverse AAA progression is the ultimate aim of AAA research, but this necessitates a stratification imaging biomarker that could (a) help select patients who would benefit from such a treatment, (b) predict AAA growth and refine watchful waiting regimens, (c) determine patient-specific intervention thresholds, or (d) be applied in all of the aforementioned applications. Much of the dated
understanding of late-stage AAA is based on aortic tissue acquired during surgery; hence, theories derived from these samples may not accurately represent the mechanisms of early-stage disease, when a novel pharmacological therapy may be administered. Preclinical studies of AAA are the first step in achieving a more thorough understanding of early-stage AAA pathophysiology. Understanding the molecular workings of AAA in preclinical models can then lead to the development of stratification biomarkers that could be translated to the clinic. Following on from this, a preclinical model of AAA has been used to investigate a candidate radiotracerm for PET imaging in this thesis.

1.4 Preclinical imaging of AAA

In vivo imaging modalities are the central focus of preclinical research and confer the advantage of visualising biological processes in live animals non-invasively. The multifactorial contributors to the underlying AAA pathobiology each act as attractive molecular-level imaging targets to further our understanding of the early-stage changes that precede anatomical changes in aneurysmal aortae.

1.4.1 Preclinical models of AAA

For any disease, the use of clinically relevant animal models is essential to shed light on mechanisms of disease progression before clinical testing. That being said, no single model is a perfect representative of human AAA pathology; therefore, it is ideal to test hypotheses in multiple models. Rabbits, pigs, mice, and rats have been used previously to investigate AAA (Patelis et al., 2017). Mice are invaluable in studies of AAA for numerous reasons. They are small in size and thus convenient as model species; they can be easily housed and maintained, and they adapt well to changes in their surroundings. Mice are also relatively cost-effective, as large batches can be purchased from mass-producers that breed rodents specifically for research purposes (Daugherty and Cassis, 2004, Poulsen et al., 2016). Furthermore, the mouse genome is very similar to the human genome and can be easily modified to study genes of interest. Inbreeding of different mouse strains is a standard means to achieve gene knockout or knock-in models, the physiological and phenotypic implications of which can then be explored. For example, in the context of AAA, hyperlipidaemic mice are achieved via the knockout of apolipoprotein E or low-density lipoprotein receptors, and mice deficient in lysyl oxidase (resulting in a failure to crosslink elastin and collagen) and MMP genes have also been utilised to
better understand the role of the ECM (Daugherty and Cassis, 2004, van der Weyden et al., 2011). Herein, three commonly used murine models of AAA are described: angiotensin II (AngII) infusion in apolipoprotein E-knockout (ApoE−/−) mice, aortic application of calcium chloride (CaCl2), and aortic application of porcine pancreatic elastase (PPE).

1.4.1.1 AngII infusion model

The AngII model is the most commonly used AAA model in mice (Poulsen et al., 2016). The result of AngII infusion leading to the development of AAA was an unexpected finding of a study conducted to assess the effect of increased plasma concentrations of AngII on atherogenesis in ApoE−/− mice (Daugherty et al., 2000). AngII is a vasoconstrictor that mediates a variety of growth processes, including proliferation. The proliferative effects of AngII have long been established in adrenocortical cells (Gill et al., 1977), mesangial cells (Bakris and Re, 1993), endothelial cells (Wolf et al., 1996), hematopoietic stem cells (Kim et al., 2016), and VSMCs (Johnson et al., 1992, Yaghini et al., 2010), among other cell types. To study AAA, ApoE−/− mice are classically used for this model because of their inherent atherosclerotic susceptibility. AngII infusion in wildtype C57BL6/J mice will also produce aneurysms, but only in about 10% of animals compared to 80% of animals with an ApoE−/− background. Low-density lipoprotein receptor-deficient mice also exhibit an increased incidence of AAA following AngII infusion and are an alternative, as is a prolonged period of high-fat diet feeding. Therefore, although an underlying ApoE−/− condition is not necessary for AAA development, it appears to significantly enhance the incidence with which AAA develops in this model (Deng et al., 2003, Manning et al., 2003). In this model, AAA occurs spontaneously in the suprarenal abdominal aorta following 28 days of continuous AngII infusion at a dose of 500–1000 ng/kg/min via an osmotic mini-pump that has been subcutaneously implanted laterally into the flank through a midline neck incision without conferring an effect on the thoracic aorta (Rateri et al., 2011). Along with dilatation of the suprarenal aorta, the aneurysms in this model present with atherosclerosis, macrophage accumulation in the elastic lamina, leukocyte infiltration, and medial hypertrophy. There is also a predominance of AAA development in male mice, with roughly double the incidence than that in female mice. These features are similar to those observed in human AAA disease (Daugherty et al., 2000). This model is also popular because the AAA is progressive, i.e. it continues to grow as long as AngII continues to be infused (Tedjawirja and de Waard, 2019). The timeline of biological events in
this model is well established, initiating with macrophage infiltration into the medial layer of the susceptible aortic region, transmedial dissection that results in rapid luminal expansion within the first week of AngII infusion, and additional inflammatory mechanisms, such as the formation of an intramural thrombus, degradation of elastin, and significant vascular remodelling (Cao et al., 2010). It is important to acknowledge that this model is regarded by many as an aortic dissection and dissecting aneurysm model; this is because the aneurysms that form in this model have been found to be preceded by an intramural rather than intraluminal thrombus, medial tears at specific side branches instead of circumferential medial degradation, and aortic dissection, which are not typical features of human AAA (Trachet et al., 2017). This was the main model used for the experiments in this thesis for its procedural simplicity and characteristic similarities to AAA disease in humans. The most salient difficulty encountered in this model is the high AAA rupture rate (rates of 30–50% have been reported previously); most ruptures occur within the first week of AngII infusion, thus necessitating a greater starting number of animals for experiments involving this model (Cao et al., 2010, Nguyen et al., 2011, English et al., 2015).

1.4.1.2 PPE application model

The PPE perfusion model was first described by Anidjar et al. in 1990 and has been widely used as a murine model of AAA since then. This model entails in vivo isolation and cannulation of the abdominal aorta to administer elastase, followed by careful repair of the surgical incision. This model is technically demanding; murine aortic isolation from the adjacent inferior vena cava (IVC) and small lumbar branch vessels can prove to be difficult. In addition, there may be differences in the extent of vascular injury due to differences in perfusion pressure. Moreover, careful closure of the aortotomy whilst avoiding stenosis is imperative to restore antegrade blood flow (Anidjar et al., 1990).

More than two decades later, a modified version of this model wherein PPE is applied to the peri-adventitial aorta was introduced by Bhamidipati et al. to overcome the complexities associated with the PPE perfusion model. Their model can be achieved without individual vessel manipulation and generates aneurysms that involve greater accumulation of activated macrophages, elastic lamina degradation, decreased expression of smooth muscle protein, and enhanced matrix metalloproteinase activity, similar to that observed in other experimental AAA models (Bhamidipati et al., 2012). This modified
model was implemented for the research presented in this thesis for its technical simplicity.

The PPE model rarely if ever demonstrates rupture in comparison to the AngII model. Atherosclerosis is not a feature of the PPE model, whilst AAA in this model is associated with medial degeneration, leukocyte infiltration, and formation in the infra-renal abdominal aorta as the key features resembling human AAA. However, the PPE model is a non-progressive model of AAA, as the disease stabilises after a period of several days or weeks due to healing of the biological process (Senemaud et al., 2017).

1.4.1.3 CaCl\(_2\) application model

The CaCl\(_2\) model is another non-progressive model of AAA that involves peri-aortic application of CaCl\(_2\) to the infra-renal aorta, which induces AAA. In 1988, Gertz et al. first demonstrated that the application of CaCl\(_2\) to the adventitia of a carotid artery resulted in the formation of an aneurysm. This aneurysm was accompanied by a decrease in VSMCs, elastin calcification, and inflammatory cell infiltration (Gertz et al., 1988). In 1997, this approach was reported to induce AAA in New Zealand rabbits (Freestone et al., 1997). Finally, this model was first described in mice by Chiou et al. who reported a 110% increase in diameter 3 weeks after CaCl\(_2\) treatment. This model is based on the high affinity of calcium for elastin; disrupting this affinity weakens the vascular wall, activates inflammatory processes, and eventually leads to aneurysm development. Furthermore, AAA in this model is accompanied by the depletion of VSMCs, degradation of elastin, infiltration of inflammatory cells, and high circulating levels of inflammatory markers (Chiou et al., 2001, Wang et al., 2013). Disruption of elastic tissue by calcium depositions has been demonstrated in human atherosclerosis; in addition, calcification of the human aorta is observed in older adults, and approximately 80% of human AAA exhibit calcification of the aortic wall (Jayalath et al., 2005, Maier et al., 2010, Kamenskiy et al., 2018). These findings make the CaCl\(_2\) model a relevant model of human AAA. However, thrombus and rupture, which are observed in human AAA, are not features of CaCl\(_2\)-induced AAA (Wang et al., 2013).

1.4.2 USS

As the imaging modality of choice in the clinic, USS has also been used extensively in preclinical research to investigate morphological aspects of AAA. The feasibility of 3D micro-USS in monitoring aneurysm growth has been demonstrated in mice via comparisons with histological images
(Goldberg et al., 2007). High-frequency USS, which is better suited to visualise superficial body structures and can produce images of high axial resolution compared to low-frequency USS, has also been used extensively to detect AAA in mice based on measurements of luminal diameter and aortic wall thickness (Martin-McNulty et al., 2005, Barisone et al., 2006, Azuma et al., 2011). USS is a feasible modality to assess changes in aortic strain (Favreau et al., 2012). Moreover, preclinical USS combined with a semi-automatic image processing algorithm has been shown to reveal changes in arterial distension, which may prove to be informative as an early marker of arterial disease in animal models (Janus et al., 2018). USS can be challenging in mice with AAA; aortic rupture has been reported to occur during USS, implying that pressure from the probe on the abdominal region may be sufficient to induce rupture.

1.4.3 MRI

MRI uses strong magnetic fields and radiofrequency pulses to produce high-resolution images without the need for ionising radiation (Brangsch et al., 2017). MRI can be implemented with various probes that have different targets to elucidate AAA mechanisms and potentially act as distinct biomarkers supplemental to aortic diameter. Botnar et al. demonstrated the feasibility of using a fibrin-specific MRI probe to investigate fibrin content in the AngII AAA mouse model (Botnar et al., 2018). Bazeli et al. used contrast-enhanced MRI to detect MMPs in the rat elastase model of AAA, demonstrating a correlation between MMP activity and inflammation (Bazeli et al., 2010). Furthermore, a newly developed smart MRI nanoprobe has recently been demonstrated to be useful in detecting MMP activity in the AngII AAA mouse model and potentially correlating with severe progression or rupture (Yao et al., 2020). Brangsch et al. demonstrated the use of dual-probe MRI to evaluate ECM degradation and inflammatory activity in the AngII AAA murine model, which could then be associated with AAA rupture risk (Brangsch et al., 2019). Dysfunctional ECM remodelling has further been investigated using a gadolinium-based contrast agent specific to tropoelastin, the soluble precursor to elastin, which was then shown to correlate with accumulated tropoelastin in human aneurysmal tissue (Lavin et al., 2019). MRI has also been used to quantitatively analyse aortic motion and curvature, as well as changes in aortic size following an intervention in mice with AAA (Turner et al., 2008, Goergen et al., 2011). MRI has also successfully been used perioperatively to guide EVAR in a swine model of AAA, raising the possibility of its role in the clinic during EVAR in patients.
who are not suitable to undergo CTA (Raman et al., 2005). Despite its limitations associated with long scan durations and high costs, as a sensitive technique that confers excellent soft tissue contrast, MRI for AAA is a growing field of research.

### 1.4.4 CT

CT uses X-rays to generate three-dimensional images with or without contrast agents for target enhancement. Although CT is typically used as a supplement to functional imaging techniques, such as PET and SPECT, Wang et al. recently demonstrated that utilising gold nanoparticles as contrast agents with micro-CT can help predict elastin damage and rupture pressure in the AngII AAA mouse model (Wang et al., 2019). A CT-based method that targets phagocytosis to image vessel wall inflammation in the same model has also been developed (Toczek et al., 2018).

### 1.4.5 Single-photon emission computed tomography

Single-photon emission computed tomography (SPECT) is a modality that involves the detection of emitted gamma photons following the injection of radiotracers such as $^{99m}$Tc, $^{123}$I, and $^{111}$In. Golestani et al. used SPECT with RP805, a $^{99m}$Tc-labelle der tracer that targets matrix metalloproteinases (MMPs), to demonstrate that MMP-targeted SPECT may indicate aortic wall inflammation and help predict expansion or rupture in AAA (Golestani et al., 2015). Toczek et al. also used SPECT, but with a novel pan-MMP tracer called RYM1, to successfully demonstrate specific detection of MMP and inflammatory activity in AAA (Toczek et al., 2017). As a lower-cost option compared to PET, the application of SPECT to investigate AAA is promising, particularly as its spatial resolution is superior to that of preclinical PET (approximately 0.25 mm vs. 1 mm). The main obstacle lies in the translation of SPECT to the clinic, as clinical SPECT scanners have poorer resolutions than those of clinical PET scanners. SPECT is closely related to PET, which is the central focus of this thesis, thus warranting a more focussed separate discussion in the following section.

### 1.5 PET

Molecular imaging utilises biologically specific tracer molecules to target, characterise, and quantify cellular and subcellular pathways and processes (Fernandez-Friera et al., 2014, Golestani et al., 2016). PET is a molecular imaging modality that relies on radioactive tracers to generate three-dimensional images of functional processes within the body. The working
principle of PET involves the simultaneous detection of two gamma-ray photons emitted in opposite directions after the annihilation between an electron and tracer-emitted positron. Reconstructed images based on the localisation of pairs of photons then reveal the distribution of injected radioactive tracer throughout the body, effectively producing a three-dimensional functional map that is biologically specific to the injected tracer. These reconstructions may incorporate corrections that have been made for attenuation, scatter, dead time, and random photon coincidences, which further improve the spatial resolution of images (Ramaswamy et al., 2013, Tarkin et al., 2016).

1.5.1 Data analysis

Quantitative measurements are a defining feature of nuclear medicine techniques such as PET, allowing for the ability to precisely characterise physiological parameters of interest to inform the diagnosis, staging, and monitoring of various diseases. To provide anatomical context to supplement quantitative data, stand-alone PET imaging systems are often coupled with CT or MRI to compensate for the limited spatial resolution of PET; the integration of a function-dominant modality (PET) with readily available structure-dominant modalities (CT, MRI) improves the quantitative analysis of biological functions (Golestani et al., 2016).

PET/CT can counter some of the caveats associated with USS and, more importantly, supplement the information acquired from USS. Unlike USS, PET is operator-independent, as it relies on an automated scanner and a radiotracer, the activity distribution of which is independent of patient factors such as obesity; this feature makes PET an objective technique. Large patients, as long as they fit within the scanner, can undergo PET scans with minimal effects of tissue depth on the resultant PET signal. One concern involving PET is the possible inter-analytical variability depending on how the region of interest (ROI) is defined. There is ongoing research to standardise ROI definitions and analytical methods, and a significant advantage of PET is that data can be reviewed and corrected post-acquisition using analytic methods to correct for temporal and spatial resolution, something that USS that does not yet offer. Meanwhile, standardising the human error that contributes to operator-dependent variability in USS is difficult (Veronesi et al., 2015, Comelli et al., 2018).

PET image analysis often relies on the standardised uptake value (SUV) metric, a widely used radioactivity quantifier that takes the injected activity
and body weight of the subject into account. SUVs are calculated using the following equation (Equation 1.2):

\[
SUV = \frac{\text{Radioactivity concentration in a region of interest (kBq/mL)}}{\text{Injected activity (kBq)}} \times \frac{1}{\text{Body weight (g)}}
\]

**Equation 1.2 Standardised uptake value**

The resultant value is unitless (g/mL, where g and mL are equivalent), representing radiotracer accumulation in a given ROI. SUV thresholds are commonly used in clinical practice as a basis for disease classification and criterion for diagnoses, particularly in the cancer field; for example, regions of radioactive glucose uptake (further described in section 1.5.2) may indicate where a tumour is located and how aggressive and metabolically active the cancer might be (Kinahan and Fletcher, 2010, Nakajima et al., 2012, Hofman and Hicks, 2016). Use of the SUV to aid diagnosis is controversial based on the notions that (a) relative image appearance may suffice for diagnostic purposes and (b) calculated SUVs exhibit high variability due to non-standardised methods of image acquisition, reconstruction, and analysis. However, the SUV may be better used as a trend marker, especially in conservative surveillance, wherein clinicians might evaluate trends in SUV change to decide how to advance with treatment options in AAA; it may be useful to determine whether a ‘watch-and-wait’ strategy should be adopted to avoid performing invasive procedures that may not be necessary for AAA management (Gambhir, 2002, Boellaard et al., 2004, Kinahan and Fletcher, 2010).

Despite the widespread use of the SUV, its quantification may be largely affected by a number of physiological (e.g. body weight and composition, mode of radiotracer administration, time between radiotracer administration) and physical (e.g. scanner calibration, reconstruction parameters, ROI definition) factors. A variation of 10–25% in repeated SUV measurements can be expected in the same patients due to instrument-related and analytical factors (Fahey et al., 2010, Kinahan and Fletcher, 2010). The most common methods of calculating the SUV are based on the average or maximum SUV of all voxels within a predefined ROI (SUV$_{\text{mean}}$ and SUV$_{\text{max}}$, respectively). Although the SUV$_{\text{mean}}$ shows less sensitivity to image noise, it is subject to intra- and inter-observer variability, as it is highly dependent on the voxels that are included in the final calculation. Meanwhile, the SUV$_{\text{max}}$ is largely independent of the ROI definition, as it reflects the single ‘hottest’ voxel value; however, it may be adversely affected by image noise (Massaro
et al., 2009, Vanderhoek et al., 2013). In an effort to reduce the variability in SUVs, other methods of calculating the SUV have been explored, wherein the average SUV in a group of voxels surrounding the ‘hottest’ voxel value is determined. This is achieved by applying specific threshold values to calculate the SUV\textsubscript{mean} for voxels within predefined ROIs that are equal to or greater than the ‘hottest’ voxels. This approach may maintain the reproducibility of the SUV\textsubscript{max} with improved statistics to reduce noise (Nakamoto et al., 2002, Krak et al., 2005, Velasquez et al., 2009). Averaging the 10 ‘hottest’ voxels has been shown to reduce the variability in the SUV\textsubscript{max} by a factor of 2.7 (Burger et al., 2012). In similar studies, different threshold values have been investigated, ranging from 40% to 90% of the highest values, which evidently raises the need for a standardised percentage threshold that can be applied consistently to provide uniform SUV results (Boellaard, 2009, Wahl et al., 2009).

In addition to the quantitative threshold, the morphology of the ROI also contributes to SUV quantification. Manually drawing the ROI to precisely delineate the area of tracer uptake is the most intuitive and simple segmentation method. However, this method is time-consuming and highly subjective, leading to variability in the results (Velasquez et al., 2009, Vorwerk et al., 2009). Alternatively, implementing fixed-size ROIs permits semi-automatic delineation of areas of tracer uptake, hence requiring less time than that required for manual delineation (Nakamoto et al., 2002, Benz et al., 2008, Vanderhoek et al., 2013). Different sizes and shapes of fixed-size ROIs have been explored, including square, cylindrical, and spherical regions with side lengths ranging from 7 to 15 mm (Nahmias and Wahl, 2008, Weber et al., 2015). Moreover, advanced automatic image segmentation methods that utilise properties of image reconstruction algorithms, such as edge detection, region growing, and fuzzy locally adaptive Bayesian methods, are under continuous development (Day et al., 2009, Hatt et al., 2010); this domain has evolved to what is now called ‘radiomics’. Studies of the effects of different thresholds on SUV quantification have largely used data from the oncology field; however, the implications of the results are also useful for cardiovascular and other medical fields.

Another metric that is commonly implemented in PET image analysis is the target-to-background ratio (TBR). The TBR takes the SUV a step further, in dividing the target lesion’s SUV with the venous blood pool SUV, as follows (Chen and Dilsizian, 2015) (Equation 1.3):
\[ TBR = \frac{SUV_{\text{target}}}{SUV_{\text{background}}} \]

**Equation 1.3 Target-to-background ratio**

Doing so corrects the SUV for blood uptake of the tracer, enabling researchers and clinicians to differentiate between radiotracer uptake in a target lesion versus radioactivity due to radiotracer that has not been taken up into cells and has instead remained in the blood supply circulating around the target lesion. Nonetheless, similar to the use of the SUV, the use of the TBR is also controversial. The TBR depends on how the blood pool activity is measured, reflecting the denominator in the ratio. This measure could be acquired from the superior vena cava, IVC, right atrium, or jugular vein, a source that varies across different studies. Furthermore, the blood pool activity may vary because of biological (e.g. differential renal clearance) or analytical (e.g. differential ROI definitions or low spatial resolution) reasons (Chen and Dilsizian, 2015). Regarding this, Huet et al. implemented a mathematical model that summed vascular wall activity and blood activity with adjustments based on weighting factors considering activity spill-out. They suggested that differences in the TBR result from notable differences in vascular wall activity and in estimated blood pool activity, rendering the TBR less reproducible than the SUV (Huet et al., 2015). Improvements in spatial resolution and application of partial volume correction may enhance the precision and accuracy of methods of PET analysis in the future (Tsoumpas et al., 2016).

1.5.2 PET radiotracers

The most common radioactive tracer used in PET imaging is \([^{18}\text{F}]\text{FDG}\), an analogue of glucose. Uptake of \([^{18}\text{F}]\text{FDG}\) occurs in cells that metabolise glucose, after which it is phosphorylated by hexokinase into \([^{18}\text{F}]\text{FDG-6-phosphate}\). At this stage, it is then metabolically trapped, as it does not harbour the 2' hydroxyl group that is essential to proceed with glycolytic reactions (Figure 1.3). Therefore, the intracellular distribution of \([^{18}\text{F}]\text{FDG}\) is associated with the degree of metabolic activity and inflammation in regions of uptake (Tarkin et al., 2016). \([^{18}\text{F}]\text{FDG}\) uptake on PET has been demonstrated to be correlated with macrophage density in plaques, cardiovascular risk factors, the Framingham Risk Score, and various inflammatory and glycolysis-related biomarkers, such as glucose transporter (GLUT)-1, GLUT-3, and total lesion glycolysis (Zhao et al., 2002, Tarkin et al., 2014, Suzuki et al., 2018). A downside to the advantage of \([^{18}\text{F}]\text{FDG}\)
having the capacity to highlight all regions of active glucose metabolism is that, as a result, it is difficult to differentiate disease-specific activity. Moreover, the uptake of $[^{18}\text{F}]{\text{FDG}}$ can be influenced by conditions of disease microenvironments, such as hypoxia or increased myocardial muscle activity, or the efficiency with which the microcirculation distributes the radiotracer (Wykrzykowska et al., 2009, Folco et al., 2011, Taqueti et al., 2014). To help overcome these and similar limitations, novel radioactive tracers are continually under development to elucidate different pathobiological mechanisms in different diseases.
Figure 1.3 Schematic of the metabolic pathway of glucose and its analogue $^{18}$F-FDG. $^{18}$F-FDG, $^{18}$F-fluorodeoxyglucose; $^{18}$F-FDG-6P, $^{18}$F-fluorodeoxyglucose-6-phosphate; GLUT1, glucose transporter-1; HK, hexokinase; G6Pase, glucose-6-phosphatase; Glucose-6P, glucose-6-phosphate; TCA, tricarboxylic acid.
Primary research studies of AAA involving PET imaging that have been conducted in the last decade are presented in Table 1.2. There has been an extensive focus on markers of inflammatory activity, evidenced by the overwhelming predominance of $^{18}$F-FDG PET studies. Both preclinical and clinical studies of gold-standard $^{18}$F-FDG PET to assess glucose metabolism and inflammation in AAA have yielded varying results, with both increased and variable $^{18}$F-FDG uptake being demonstrated (Sakalihasan et al., 2002, Kotze et al., 2011, English et al., 2015, Huang et al., 2016, Nie et al., 2018, English et al., 2020). Some studies have shown that $^{18}$F-FDG PET may be useful to predict AAA expansion and/or progression (Reeps et al., 2008, Courtois et al., 2013, Nchimi et al., 2014), whilst other studies have contradicted this (Kotze et al., 2011, Barwick et al., 2014). For example, Reeps et al. showed that increased aortic uptake of $^{18}$F-FDG in patients was associated with a higher density of inflammatory markers, which may contribute to aortic expansion, whilst patients showing $^{18}$F-FDG uptake in AAA revealed no correlation with aortic expansion 12 months later in a study by Kotze et al (Reeps et al., 2008, Kotze et al., 2011). Moreover, Nchimi et al. demonstrated that $^{18}$F-FDG uptake correlated positively with wall stress and strength in AAA, whereas Barwick et al. found no significant difference in aortic uptake of $^{18}$F-FDG between patients with infra-renal AAA and patients without AAA (Barwick et al., 2014, Nchimi et al., 2014). Based on a study involving $^{18}$F-FDG PET and contrast-enhanced MRI, Kuzniar et al. further demonstrated that the hotspots of $^{18}$F-FDG uptake and late gadolinium enhancement rarely coincide in AAA, although both are associated with aneurysm growth, raising questions about the distribution of cellular activity in AAA (Kuzniar et al., 2019). Collectively, these findings lend to a complicated story of glucose metabolism in AAA.

Other PET radiotracers are also promising in the context of AAA. The Sodium Fluoride Imaging of AAA (SoFIA) trial has shown that uptake of $^{18}$F-labelled sodium fluoride (NaF), which reflects regions of microcalcification, in patients with asymptomatic AAA predicts AAA progression and rupture, providing proof-of-concept data for the feasibility of a non-$^{18}$F-FDG PET radiotracer for AAA stratification (Forsythe et al., 2018). A key point coming out of this study is that aneurysm size is not necessarily a predictor of rupture; AAA growth can be non-linear and influenced by biomechanical processes that may not exhibit a detectable pattern. The SoFIA trial demonstrated that $^{18}$F-NaF uptake is a positive predictor of aneurysm growth and clinical outcomes, which are independent but supplementary to classic clinical parameters such as aneurysm diameter.
(Forsythe et al., 2018). That being said, the findings of this trial must be considered in light of the confounding issue of spill-in contamination from the nearby bone into the aneurysm; thus, background correction techniques can provide more robust quantitative assessments of AAA (Akerele et al., 2019).

Alternative tracers that may prove to be useful include markers of other characteristics of AAA development, such as angiogenesis (Shi et al., 2015) and integrins (Kitagawa et al., 2013, Tegler et al., 2014). 

$^{18}$F]fluoromethylcholine, which is commonly implemented for staging prostate cancer, may also be useful to incidentally detect AAA in patients with prostate cancer (Ferda et al., 2019). In this way, PET will undoubtedly remain an attractive modality to advance our understanding of AAA pathophysiology, whilst novel molecular tracer agents are introduced and hybrid multimodality systems are improved. As PET is readily accessible and already being used in the clinic, continued research using PET has practical applications that are feasible for clinical translation.
Table 1.2 Studies using PET imaging to investigate AAA published in the most recent decade. The following Boolean operators were used as search terminology: (AAA OR abdominal aortic aneurysm*) AND (PET OR positron emission tomography). Only original research articles were included (i.e. reviews, case reports, conference proceedings, etc. were excluded).

<table>
<thead>
<tr>
<th>Reference</th>
<th>Radiotracer</th>
<th>Target (Species)</th>
<th>Key Conclusions</th>
</tr>
</thead>
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<tr>
<td>(Barwick et al., 2014)</td>
<td>$[^{18}F]$FDG</td>
<td>Glucose metabolism (humans)</td>
<td>Metabolic activity levels may not correlate with aortic size and may not differ between aneurysms vs. controls.</td>
</tr>
<tr>
<td>(Courtois et al., 2013)</td>
<td>$[^{18}F]$FDG</td>
<td>Correlation between inflammation and histological analysis (humans)</td>
<td>$[^{18}F]$FDG uptake in the aneurysmal wall may be associated with an active inflammatory process involving proliferating leukocytes and increased circulating C-reactive protein.</td>
</tr>
<tr>
<td>(Courtois et al., 2018)</td>
<td>$[^{18}F]$FDG</td>
<td>Circulating miRNAs (humans)</td>
<td>Specific miRNAs are significantly correlated with $[^{18}F]$FDG uptake in the aneurysmal wall and may be directly involved in AAA instability.</td>
</tr>
<tr>
<td>(Courtois et al., 2019)</td>
<td>$[^{18}F]$FDG</td>
<td>Prediction of complications after EVAR (humans)</td>
<td>Aortic $[^{18}F]$FDG uptake may be a predictor of post-EVAR complications.</td>
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<td>(English et al., 2014)</td>
<td>$[^{18}F]$FDG and $[^{11}C]$PBR28</td>
<td>Aortic wall inflammation (rats)</td>
<td>AAA wall inflammation can be detected using $[^{18}F]$FDG and $[^{11}C]$PBR28.</td>
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<tr>
<td>(English et al., 2015)</td>
<td>$[^{18}F]$FDG</td>
<td>Rupture prediction (rats)</td>
<td>Increased pre-rupture glucose uptake may be associated with increased inflammation in the ruptured AAA wall.</td>
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<tr>
<td>Reference</td>
<td>Tracer</td>
<td>Description</td>
<td>Result</td>
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<tr>
<td>(English et al., 2020)</td>
<td>$[^{64}\text{Cu}]\text{DOTA-ECL1i}$</td>
<td>Expression of chemokine receptor 2 (rats)</td>
<td>Chemokine receptor 2 may predict AAA rupture.</td>
</tr>
<tr>
<td>(Ferda et al., 2019)</td>
<td>$[^{18}\text{F}]\text{FCH}$</td>
<td>Incidental AAA detection in patients with prostate cancer (humans)</td>
<td>$[^{18}\text{F}]\text{FCH}$ PET/CT may be an effective approach for secondary prevention and stratification of AAA in patients with prostate cancer.</td>
</tr>
<tr>
<td>(Forsythe et al., 2018)</td>
<td>$[^{18}\text{F}]\text{NaF}$</td>
<td>AAA growth and clinical outcomes (humans)</td>
<td>$[^{18}\text{F}]\text{NaF}$ uptake may help identify advanced AAA and may be correlated with aneurysm growth and clinical AAA events that differ from established risk factors.</td>
</tr>
<tr>
<td>(Huang et al., 2016)</td>
<td>$[^{18}\text{F}]\text{FDG}$</td>
<td>Structural stress (humans)</td>
<td>Increased $[^{18}\text{F}]\text{FDG}$ is associated with high mechanical stress of thick intraluminal thrombus in AAA.</td>
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<td>(Kitagawa et al., 2013)</td>
<td>$[^{18}\text{F}]\text{FPPRGD}_2$</td>
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<td>Metabolic activity (humans)</td>
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<td>(Kotze et al., 2011)</td>
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</tr>
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<td>(Kotze et al., 2014)</td>
<td>$[^{18}\text{F}]\text{FDG}$</td>
<td>Correlation between CT texture analysis data and metabolism (humans)</td>
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</tr>
<tr>
<td>Study (Year)</td>
<td>Imaging Modality (FDG)</td>
<td>Main Findings</td>
<td>Additional Information</td>
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<tr>
<td>Kuzniar et al. (2019)</td>
<td>$[^{18}F]$FDG</td>
<td>Inflammation (humans)</td>
<td>$[^{18}F]$FDG PET/MRI may be used to assess inflammation in asymptomatic AAA, although the hotspots of $[^{18}F]$FDG uptake and late gadolinium enhancement are not always aligned.</td>
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<td>Lee et al. (2018)</td>
<td>$[^{18}F]$FDG</td>
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<td>Maier et al. (2012)</td>
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<td>Marini et al. (2012)</td>
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<td>Molacek et al. (2019)</td>
<td>$[^{18}F]$FDG</td>
<td>AAA progression (humans)</td>
<td>$[^{18}F]$FDG PET/CT or PET/MRI does not correlate with disease symptoms, AAA progression, or dissection.</td>
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<td>Morbelli et al. (2014)</td>
<td>$[^{18}F]$FDG</td>
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<td>Application</td>
<td>Result</td>
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<td>Morel et al. (2015)</td>
<td>[¹⁸F]FDG</td>
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<td>Metabolic changes in AAA may follow a cyclic pattern, similar to that observed with changes in maximal aortic diameter.</td>
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<td>Murakami et al. (2014)</td>
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<td>Infection (humans)</td>
<td>[¹⁸F]FDG PET is useful to diagnose infected AAA.</td>
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<td>Study</td>
<td>Tracer</td>
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<td>Sarda-Mantel et al., 2012</td>
<td>[^{18}\text{F}]\text{FDG},[^{18}\text{F}]\text{FCH},[^{18}\text{F}]\text{DPA714}\</td>
<td>Correlation between aortic wall inflammation and histopathological analysis (rats)</td>
<td>[^{18}\text{F}]\text{FDG} may have higher sensitivity than that of [^{18}\text{F}]\text{FCH} and [^{18}\text{F}]\text{DPA714} in detecting activated leukocytes in the aneurysmal wall.</td>
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<tr>
<td>Shi et al., 2015</td>
<td>[^{64}\text{Cu}]\text{NOTA-TRC105-Fab}\</td>
<td>Angiogenesis (mice)</td>
<td>[^{64}\text{Cu}]\text{NOTA-TRC105-Fab} uptake indicates regions of increased angiogenesis in AAA, based on CD105 expression.</td>
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<td>Tegler et al., 2012</td>
<td>[^{18}\text{F}]\text{FDG}\</td>
<td>Inflammation (humans)</td>
<td>[^{18}\text{F}]\text{FDG} PET cannot be used to detect chronic inflammation in asymptomatic aneurysms.</td>
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<td>Tegler et al., 2012</td>
<td>[^{11}\text{C}]\text{PK11195}\ and [^{11}\text{C}]\text{-d-deprenyl}\</td>
<td>Inflammation (humans)</td>
<td>Inflammation in AAA cannot be detected using [^{11}\text{C}]\text{PK11195}\ and [^{11}\text{C}]\text{-d-deprenyl}.</td>
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<td>Tegler et al., 2014</td>
<td>[^{18}\text{F}]\text{-fluciclatide}\</td>
<td>(\alpha_\text{v}\beta_3) expression (humans)</td>
<td>(\alpha_\text{v}\beta_3) integrin expression in AAA may be visualised using [^{18}\text{F}]\text{fluciclatide}</td>
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<td>Truijers et al., 2009</td>
<td>[^{18}\text{F}]\text{FDG}\</td>
<td>Aneurysm wall pathology (humans)</td>
<td>[^{18}\text{F}]\text{FDG PET/CT} may be useful in detecting concomitant malignancies in patients with AAA.</td>
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<td>Tsuruda et al., 2016</td>
<td>[^{18}\text{F}]\text{FDG}\</td>
<td>Aortic wall inflammation (humans)</td>
<td>Active aortic wall inflammation may contribute to AAA progression and rupture.</td>
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<tr>
<td>Xu et al., 2010</td>
<td>[^{18}\text{F}]\text{FDG}\</td>
<td>Wall stress and metabolic activity (humans)</td>
<td>High AAA wall stress and accelerated metabolism may be associated.</td>
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1.5.3 \[^{18}\text{F}]\text{FLT and thymidine}\n
A key idea underlying this thesis, as described in section 1.3.5, is that AAA formation involves cell proliferation, which may be a feasible target for \[^{18}\text{F}]\text{FLT PET.} \[^{18}\text{F}]\text{FLT is an example of a radioactive PET tracer that is useful to elucidate a non-glucose activity mechanism: cell proliferation.} \[^{18}\text{F}]\text{FLT has a similar structure to that of thymidine, with the exception of the fluorine atom that replaces the hydroxyl group at the 3’ position; this precise positioning prevents \[^{18}\text{F}]\text{FLT from proceeding along the complete mechanistic pathway of thymidine.} Once \[^{18}\text{F}]\text{FLT is actively transported from the blood into cells, it acts as a substrate for thymidine kinase (TK)-1, a cytosolic enzyme with peak expression during the synthesis phase of the cell cycle and very low expression in resting cells (He et al., 2004). At this stage, it is phosphorylated and metabolically trapped within the cell, such that it is restricted from incorporating into DNA (Figure 1.4). Few researchers have attempted kinetic analysis for \[^{18}\text{F}]\text{FLT quantification in PET imaging of non-neural regions. However, a mathematical model similar to that for \[^{18}\text{F}]\text{FDG has been established, which consists of (i) an extracellular blood pool compartment shared by \[^{18}\text{F}]\text{FLT and thymidine; (ii) an exchangeable tissue compartment, from which \[^{18}\text{F}]\text{FLT and thymidine exchange freely between the blood pool or continue to be metabolised intracellularly; and (iii) a compartment of phosphorylated (and thus trapped) \[^{18}\text{F}]\text{FLT nucleotides. Four rate constants are used to define the kinetic transfer rates between these two compartments and blood. The transfer from blood into tissue is represented by } K_1, \text{ with } k_2 \text{ representing the return of non-phosphorylated } \[^{18}\text{F}]\text{FLT from tissue. The phosphorylation of } \[^{18}\text{F}]\text{FLT is conveyed through } k_3, \text{ which is the rate-limiting step of } \[^{18}\text{F}]\text{FLT retention in tissue (Shields et al., 2002, Muzi et al., 2006). These phosphorylated products then show negligible dephosphorylation back to the extracellular compartment, represented by } k_4, \text{ as } \[^{18}\text{F}]\text{FLT is not a substrate for thymidine phosphorylase (Grierson et al., 2004).} \)
Figure 1.4 Schematic of the metabolic pathway of thymidine and its analogue $[^{18}\text{F}]\text{FLT}$. TK-1, thymidine kinase-1; dNT, deoxyribonucleotidase; DNA, deoxyribonucleic acid; $[^{18}\text{F}]\text{FLT}$, $[^{18}\text{F}]\text{fluorothymidine}$; $[^{18}\text{F}]$-FLT-MP, $[^{18}\text{F}]$fluorothymidine-monophosphate; Thymidine-MP, thymidine-monophosphate.
Based on the notion that TK-1 activity is directly associated with the concentration of $[^{18}\text{F}]$FLT in cells, $[^{18}\text{F}]$FLT signal and uptake are reflective of proliferative activity. Phosphorylated $[^{18}\text{F}]$FLT is unable to escape from cells and has been demonstrated to act as a substrate for TK-1 and not for mitochondrial TK-2, the latter being cell cycle-independent, making $[^{18}\text{F}]$FLT a tracer that exhibits high specificity for its biological target (i.e. proliferative cells) (Hannigan et al., 1993, Toyohara et al., 2002). Furthermore, the compartmental model of $[^{18}\text{F}]$FLT indicates that the metabolic flux parameter, $K_{\text{FLT}}$, which is a product of the rate constants, is strongly correlated with the Ki67 proliferative index as well as TK-1 expression (Muzi et al., 2005, Brockenbrough et al., 2011).

The utility of $[^{18}\text{F}]$FLT in cancer imaging has been extensively demonstrated because its biological target is one of the key hallmarks of cancer (Shields et al., 1998, Salskov et al., 2007, Yue et al., 2010, Hanahan and Weinberg, 2011, Viertl et al., 2011). Furthermore, Ye et al. demonstrated that $[^{18}\text{F}]$FLT PET could be used to visualise proliferating macrophages, hematopoietic stem cells, and progenitor cells in preclinical and clinical models of atherosclerosis (Ye et al., 2015). Their findings highlight the promise of $[^{18}\text{F}]$FLT PET for non-cancer applications; however, its usefulness for other cardiovascular applications is unclear.

### 1.6 Research hypotheses, aims, and objectives

The hypothesis to be tested in this thesis is that there is an active period of cell proliferation in AAA that can be detected using $[^{18}\text{F}]$FLT PET/CT and modulated in response to anti-proliferative drugs. To test this hypothesis, the aim of this thesis was to determine the feasibility of $[^{18}\text{F}]$FLT PET/CT to visualise and quantify cell proliferation in the AngII AAA murine model. The key objectives of this study were to

(i) investigate if there is significant $[^{18}\text{F}]$FLT uptake in the aneurysms of the AngII and PPE AAA mouse models;

(ii) evaluate the expression of key proteins involved in the thymidine proliferative pathway;

(iii) investigate a variety of image quantification metrics by assessing the effects of different ROI definitions; and

(iii) determine if $[^{18}\text{F}]$FLT PET/CT can be used to assess a therapeutic response in the AngII AAA model.
Chapter 2 Methods

This chapter provides an overview of the methods implemented to conduct the experiments presented in this thesis. The rationale for each method chosen is also described.

2.1 Animal models

All animal work was conducted in accordance with the UK Home Office, Animals (Scientific Procedures) Act 1986 under Project Licence P606320FB. Male Jax™ ApoE−/− mice (B6.129P2-Apoetm1Unc/J; Charles River, UK) underwent surgery at 14 weeks of age. Male C57BL6/J mice (Charles River, UK) were used at 8 weeks of age for baseline biodistribution studies. Male mice were chosen for the overall study as AAA predominantly occurs in male patients (Bloomer et al., 2012). All animals were part of the Jackson Laboratories Genetic Stability Programme to limit cumulative genetic drift. Mice were maintained at 21°C with a 12-hour light / dark cycle and 50–70% humidity in GM500 individually ventilated cages (Techniplast, Italy) with a maximum of 5 animals per cage and fed a standard RM1 chow pellet diet (Special Diet Services) and triple-filtered water via Hydropac pouches ad libitum. All mice were provided with a housing dome and two chew sticks as environmental enrichment. All mice were checked daily by an animal technician.

The traditional murine AAA models were generated as described below. All surgical procedures were performed under isoflurane anaesthesia via inhalation (to allow for tight control of anaesthetic depth with rapid recovery) in a purpose-built murine operating facility under sterile surgical conditions using an OPMI Pico operating microscope (Zeiss) in accordance with the guiding principles of the laboratory animal science association (www.lasa.co.uk) by Dr Marc Bailey. Mice were maintained at 37°C during surgery and recovery using a heated operating table and recovery platform. Wounds were closed using continuous Vicryl (Ethicon) sutures, 6-0 for the peritoneum and 4-0 for the skin. All animals received suitable buprenorphine analgesia via intraperitoneal injection (100 μL of 0.1 mg/mL solution with additional 50-μL injections as needed). Mice were identified by ear notching.

The AngII AAA model was the main model used for the experiments presented in this thesis because of (i) the favourable location of the AAA distant from the bladder, which would impart partial volume effects, and (ii) its surgical ease, as these mice needed to undergo surgery at the University
of Hull, where the surgical facilities were limited. For this model, 14-week-old male ApoE−/− mice (Jackson Laboratories) received human AngII (Sigma A9525) infusions at 750 ng/kg-min via Alzet® 1002 (14 days) or 1004 (28 days) osmotic mini-pumps that were implanted subcutaneously by posterior neck incision under recovery isoflurane anaesthesia, as described previously (Bridge et al., 2017). Matched male ApoE−/− mice infused with a saline solution were used as controls for this model.

For the PPE model, 12-week-old male C57BL6/J mice were subjected to laparotomy, and the abdominal aorta was exposed by blunt dissection. PPE (10 µL, Sigma E1250) was applied to the adventitia of the infra-renal aorta for 5 min. For the CaCl2 model, the approach was identical to that of the PPE model, but CaCl2 (0.5 M) was applied to the adventitia of the infra-renal aorta for 14 min on disassociated cotton bud tips. Sham-operated matched male C57BL6/J mice were used as controls for both the PPE and CaCl2 models; these animals underwent identical laparotomy and aortic exposure, but saline washout only.

For all mice used for experiments at the University of Hull, the presence of AAA was determined based on visual inspection at post-imaging necropsy (i.e. the presence or lack of AAA) because of the limited resources available at this site; moreover, it was difficult to grade the severity of the AAA without access to a microscope. For all mice used for experiments at the University of Leeds, the presence of AAA was determined using USS, which also provided measurements of size.

### 2.2 Histological staining

For the initial experiment investigating cell proliferation in murine models of AAA, AAA was first induced in mice by three methods: AngII infusion to ApoE−/− mice or peri-adventitial application of CaCl2 or PPE to the aorta in C57BL6/J mice, as described in section 2.1. In each case, AAA tissues were compared with equivalent tissues from appropriate and matched sham control animals. The aortic tissues were harvested at 28 days post-induction of AngII and CaCl2 AAA and at 14 days post-induction of PPE AAA (per the standard time points of the models) and used for 3,3′-diaminobenzidine staining for proliferative cells using an antibody raised against the cell proliferation marker Ki67. The Ki67 index is defined by the proportion of cells in a specified ROI with positive immunohistochemical staining and has been demonstrated to be particularly useful in cancer research to monitor the proliferative rate of tumour cells (Scholzen and Gerdes, 2000). Ki67 protein
expression peaks in cells throughout the active G1, S, G2, and M cell cycle phases, with drastically reduced expression in resting cells (i.e. G0) (Miller et al., 2018). Therefore, Ki67 was chosen for the initial experiments to explore the detectability of cell proliferation in murine AAA models.

To perform Ki67 staining, the mice were first terminally exsanguinated via the IVC. Murine aortae were each fixed in situ by perfuse fixation with 10 mL of phosphate-buffered saline followed by 5 mL of 4% paraformaldehyde in terminally anaesthetised animals via cardiac puncture with an outflow tract through the severed pulmonary arteries. Aortae were then fixed at 4°C for 48 hours before they were embedded in Cellwax (Cellpath Ltd., UK) using the Leica EG1150H embedding station. Four-µm thickness sections were cut onto Plus Frost slides (Solmedia, UK) using a Leica RM2235 microtome, and the slides were dried overnight at 37°C. Proliferating cells were stained with rabbit anti-mouse Ki67 antibody (ab15580, Abcam; 1:750 dilution) for 1 hour and Menapath Polymer HRP secondary antibody (Menarini Diagnostics Ltd., Winnersh, UK) for 30 minutes. The specificity of the antibody was first confirmed in positive control (spleen) tissue with and without primary antibody. Following a final wash, all sections were treated with Menapath diaminobenzidine peroxidase for 5 minutes and counterstained with Mayer’s Haematoxylin for 2 minutes, dehydrated, cleared in xylene, and mounted in dibutyl phthalate xylene. The slides were then imaged using the Aperio® AT2 (Leica Biosystems, Wetzlar, Germany) digital pathology slide scanner with x20 maximal magnification. Images were stored on the secure Leeds Institute of Cancer and Pathology digital pathology server and accessed remotely using Aperio® Image Scope software (Leica Biosystems). All quantitative analyses were performed using ImageJ 1.51k software (Schneider et al., 2012). For quantification of the Ki67 staining, the proportions of Ki67-positive nuclei were determined in three 50-µm² ROIs and averaged for each animal. Finally, aortae from control models were compared against aortae from the three different aneurysm models.

2.3 In vivo USS

To evaluate aortic volumes and diameters, in vivo USS was performed in two cohorts: (i) the animals used for Ki67 analysis and (ii) those that were subsequently used for [18F]FLT gamma counting analysis. USS was performed using the Vevo2100 high-resolution (30 µm), high-frequency preclinical µUSS system (VisualSonics, FUJIFILM VisualSonics, Inc., Toronto, ON, Canada) with an MS-550D transducer at a 40-MHz frequency on a heated platform maintaining a core body temperature of 35°C to 37°C,
monitored via a rectal probe. Recovery anaesthesia was induced in the mice at 5% isoflurane and maintained at 2%. Prior to imaging, the abdomen area was shaved and hair removed using depilatory cream. Imaging was performed using Aquasonic® clear gel (Parker Labs). Transverse imaging was acquired using a motor along an 11.96-mm ROI from the right renal artery in the cranial direction with 157 frames at 0.076-mm intervals gated for respiration with electrocardiographic triggering 50 ms after the r-wave. Images were reconstructed and measured using Vevo Lab v1.7.0 (VisualSonics, FUJIFILM VisualSonics, Inc., Toronto, ON, Canada) to generate three-dimensional lumen volume measurements. For the co-registration analysis, bony landmarks on CT and USS images were aligned and displayed using ImageJ 1.51k (Schneider et al., 2012).

2.4 In vivo [18F]FLT PET/CT

[18F]FLT was prepared from [18F]fluoride produced by a radiochemist at the University of Hull using an on-site 7.5-MeV ABT Biomarker Generator cyclotron and purified using an in-house developed microfluidic electrochemical cell for electrode trapping, through which irradiated target water (0.5 mL, [18O]H2O) containing ca. 1 GBq of [18F]fluoride was pumped at 0.2 mL/min whilst applying a 20 V potential, and the cell was flushed with 2 mL of MeCN at 1 mL/min with no potential applied. Subsequently, 0.4 mL of a solution containing KHCO3 (30 mM) and K222 (37 mM) in MeCN was pumped through the cell at 40°C at 0.1 mL/min. Ten mg of fluorothymidine precursor 3’-N-Boc-5’-dimethoxytrityl-3’-O-nosyl-thymidine was added to the released solution to perform a radio-labeling reaction at 100°C for 10 min. After the fluorination, the unreacted [18F]fluoride was trapped on the neutral alumina cartridge (light) and reaction mixture treated for 5 min at room temperature with an equivalent volume of 2 N HCl solution. The mixture was then neutralised with a stoichiometric amount of 8 N NaOH solution and purified by semi-preparative high-performance liquid chromatography (HPLC) on an ACE 5 C18 10×250 5A column eluted with 35% acetonitrile in water (both 0.1% trifluoroacetic acid) (flow rate=4.7 mL/min, Rf=12 min). The HPLC fraction containing partially protected product was diluted 2–3-fold with water and passed over an HBL Oasis C18 cartridge. Trapped product was eluted with 0.5 mL of ethanol and 3 mL of diethyl ether and dried at 60°C under an inert gas stream. The heat applied for drying generated the fully de-protected [18F]FLT (confirmed by analytical HPLC with a cold [19F]FLT standard), which was re-dissolved in 10% ethanol/PBS solution, filtered through a 0.22-µm filter for sterility, and delivered for animal
administration. Tracer preparation started with 1.0–1.3 GBq of cyclotron-produced $^{18}$Ffluoride and yielded 40±8 MBq (n.d.c.) $^{18}$FFLT intravenous-injectable formulation in 136±15 min ($n$=5) (RCY (decay-corrected)=8±2%).

Prior to imaging for the AngII AAA study, mice were induced with 5% isoflurane/oxygen (v/v) anaesthesia before maintenance at 2% at 1 L/min. Mice were cannulated in the tail vein using a bespoke catheter before being placed into an imaging cell where temperature and respiration were monitored (Minerve, France). $^{18}$FFLT was injected with the following means ± standard deviations (SDs) of activity in 200 µL of 0.9% saline solution (Aqupharm No1, Animalcare Ltd., York, UK) through the pre-cannulated lateral tail vein at the beginning of a 90-minute dynamic imaging sequence: 7.27 ± 2.89 MBq (0.20 ± 0.08 mCi) (14-day scans; $n$=12) and 10.08 ± 1.77 MBq (0.27 ± 0.05 mCi) (28-day scans; $n$=7). Images were acquired using the Super Argus (Sedecal) small-animal PET/CT scanner installed at the University of Hull, with animals placed prone and the field-of-view centred on the abdominal aorta. Mice were maintained at 1% anaesthesia during scanning, with temperature and respiration monitored throughout. Following the 90-minute dynamic PET scan, a CT image was acquired for anatomic co-registration (40 kV, 140 µA, 360 projections, 8 shots). PET images were histogrammed into 15 2-second, 2 15-second, 4 60-second, 1 300-second, and 8 600-second frames and reconstructed using the 3D ordered subsets expectation maximisation algorithm with 2 iterations and 16 subsets with attenuation correction (as standard protocol at the University of Hull), yielding voxel dimensions of 0.39 × 0.39 × 0.78 mm$^3$.

Prior to imaging for the PPE model and imatinib studies, mice were induced with 5% isoflurane/oxygen (v/v) anaesthesia before maintenance at 2% at 1 L/min. $^{18}$FFLT was injected at mean ± SD doses of 9.3±0.7 MBq and 9.4±0.1 MBq in 100 µL of 0.9% saline solution (Aqupharm No1, Animalcare Ltd., York, UK), respectively. After 80 and 90 min to allow for biodistribution of the radiotracer, respectively, images were acquired using the Albira Si (Bruker) small-animal PET/SPECT/CT scanner at the University of Leeds, with animals placed prone and the field-of-view centred on the abdominal aorta. Mice were maintained at 1% anaesthesia during scanning, with temperature and respiration monitored throughout (Bruker). Following the 20-min static and 90-min dynamic PET scans, respectively, a CT image was acquired for anatomic co-registration. PET/CT images were reconstructed using the maximum likelihood estimation maximisation algorithm (25 iterations), yielding voxel dimensions of 0.5 mm$^3$. 


A cohort of animals received imatinib via oral gavage on days 14–15 following mini-pump implantation. Matched controls received tap water, the solvent for imatinib.

The imaging protocols implemented in the present study were based not only on the standard protocols at the University of Hull, but also on a number of well-established notions, as follows. In a study that involved toxicological evaluation of \(^{18}\text{F}\)FLT in patients, blood-derived TACs showed a significant peak in activity within 5 min, followed by a gradual decrease to 110 min after radiotracer injection (Turcotte et al., 2007). Another study further showed a gradual decrease in the blood biodistribution of \(^{18}\text{F}\)FLT in patients with cancer (Cysouw et al., 2017). Moreover, \(^{18}\text{F}\)FLT uptake in proliferative lesions has been found to progressively increase over time to 80–100 min after radiotracer injection until eventual renal clearance (Lovinfosse et al., 2019). Evaluation of the compartmental model of \(^{18}\text{F}\)FLT kinetics, which describes the radiotracer’s uptake and retention over 120 min, has revealed that restricting PET data acquisition to 60 min leads to a loss in the accuracy of ROI-specific information. This study also revealed that curves corresponding to \(k_3\), which reflects \(^{18}\text{F}\)FLT retention in tissues, are fairly stable between 90 and 120 min, with data acquisition beyond 90 min adding minimal value (Muzi et al., 2005). High repeatability of SUV quantification has also been demonstrated to occur when PET imaging is initiated at least 60 min after radiotracer administration (Lodge et al., 2017). Keeping with these findings, dynamic PET imaging protocols commonly include short timeframes that progress to increasingly longer timeframes by the end of a 90–120-min scan (Ukon et al., 2016). For these reasons, each dynamic scan was 90 min in duration, and short timeframes were implemented at the beginning of each dynamic scan in the present study to capture the peak activity of and any interesting fluctuations in the radiotracer immediately post-injection.

2.4.1 PET/CT image analysis

Animals with failed or paravenous radiotracer injection were excluded from the analyses. Three-dimensional isocontour ROIs incorporating the supra-renal abdominal aortic area (between the renal arteries and diaphragm) were manually drawn on the reconstructed images using AMIDE v1.0.4 software (Loening and Gambhir, 2003). The aortic ROI was constructed between the kidneys and anterior to the anterior border of the vertebral column based on the CT images including the most prominent PET signal in the 80- to 90-minute timeframe. For saline controls, the aortic ROI was
constructed between the kidneys, superior to the bladder, and anterior to the anterior border of the vertebral column based on the CT images. These ROIs had dimensions of 1.08 mm$^3$ and contained 15 voxels. Incorporating mouse weights and injected radioactivity doses, the output statistics revealed the SUV$_{\text{max}}$ and SUV$_{\text{mean}}$ across all frames. These values were used to generate time–activity curves (TACs) for ROIs in the abdominal aorta, spleen, kidneys, and urinary bladder. The SUV$_{\text{max}}$ was the metric of choice for quantification of the aortic ROI to avoid partial volume effects of the adjacent kidneys and bladder, where high [$^{18}\text{F}$]FLT signals were observed consistently.

2.4.2 Generation of different ROI definitions for SUV analyses

For the experiments conducted to assess the effect of different ROI definitions on the resulting SUVs, analyses were conducted based on two types of ROI definitions: (i) shape (manually drawn vs. fixed-size ROIs) and (ii) voxel threshold (the proportion of voxels included in the final SUV calculation).

To address (i), the ellipsoid ROI-drawing tool was used to draw 4-mm ellipsoid ROIs in the abdominal aortic regions on images from both saline control and AngII AAA mice; these were the fixed-size ROIs. The size of the fixed-size ROI was chosen based on the maximum observed aneurysmal diameter of the aortae (approx. 3 mm). Therefore, the fixed-size ROIs were sufficiently large to contain the aneurysmal uptake without a deliberate attempt to align precisely with its contours, whilst also avoiding the inclusion of adjacent organs, such as the kidneys and urinary bladder. Manually drawn ROIs were generated using the 3D isocontour ROI-drawing tool to include as much of the abdominal aortic uptake as possible without including edges that may be subject to partial volume effects. For saline control images (i.e. the images that did not reveal abdominal aortic uptake of [$^{18}\text{F}$]FLT), the aortic ROI was constructed between the kidneys, superior to the bladder, and anterior to the anterior border of the vertebral column based on the CT images, a similar approach to that described in section 2.4.1. To test the efficiency of each segmentation method, the time spent drawing each ROI was recorded.

To address (ii), the following values were then determined for all fixed-size and manually drawn ROIs: SUV$_{\text{mean}}$, SUV$_{40}$, SUV$_{50}$, SUV$_{70}$, and SUV$_{90}$, and SUV$_{\text{max}}$. The calculations for SUV$_{40}$, SUV$_{50}$, SUV$_{70}$, and SUV$_{90}$ included voxels that were equal to or greater than 40%, 50%, 70%, and 90% of the
highest value voxels, respectively, as defined in the AMIDE software manual (Leoning, 2014). For clinical relevance, the threshold values were chosen based on those implemented in previous studies (Dutour et al., 2009, Kahraman et al., 2011).

2.5 Ex vivo [$^{18}$F]FLT gamma counting

Animals were anaesthetised with 5% isoflurane with an oxygen flow rate of 2 L/min. [$^{18}$F]FLT in 100 µL of 0.9% saline solution (Aquapharm No1, Animalcare Ltd., York, UK) was injected into the tail vein at the following doses (mean ± SD): C57BL6/J study, 6.07 ± 2.57 MBq (0.16 ± 0.07 mCi); AngII AAA study, 0.25 ± 0.32 MBq (0.01 ± 0.01 mCi). A significant difference in the injection doses is noted here. For the C57BL6/J study, the mice underwent in vivo PET imaging prior to their organs being used for ex vivo gamma counting; the PET imaging necessitated a sufficiently high radiotracer dose for the signals to be detected on the resultant images. For the AngII AAA study, the mice did not undergo prior in vivo PET imaging; thus, the injected dose was reduced to allow more animals to be included in a single experiment with a single delivery of [$^{18}$F]FLT. This was possible owing to the high sensitivity of the Hidex gamma counter; furthermore, the reduced dose helped avoid oversaturation of the gamma ray detectors. Animals were recovered for 90 minutes and subsequently humanely culled by cervical dislocation under Schedule 1 of the UK (Scientific Procedures) Act 1986. Samples (blood, plasma, heart, spleen, kidneys, small and large intestines, supra-renal abdominal aorta, bone, and tail) were collected, weighed, and measured for radioactivity using the Hidex gamma counter. Importantly, blood was removed from the aortic lumen before it was placed in the gamma counter. All ex vivo biodistribution data were decay-corrected, and radioactivity counts were expressed as the percentage of injected dose per gram of tissue (%ID/g). The following equation was used for decay corrections (Multi-Agency Radiological Laboratory Analytical Protocols, 2004) (Equation 2.1):

\[
\frac{\%ID}{g} = \left( \frac{C_G - C_B}{\lambda \Delta t_c} \times e^{-\lambda \Delta t_i} \right) \times 100
\]

\[
A_i = \frac{C_G - C_B}{\lambda \Delta t_c} \times e^{-\lambda \Delta t_i}
\]

\[
M_S
\]
Equation 2.1 Normalised percentage of injected dose per gram of sample (%ID/g). \( C_0 \), counting rate of sample (cps); \( C_B \), counting rate of blank/background (cps); \( \lambda \), decay constant; \( \Delta t_c \), total counting time (s), \( \Delta t_i \), time from injection to start of counting (s); \( A_i \), injected activity (MBq); \( M_S \), mass of sample (g)

2.6 Ex vivo \(^{18}\text{F}\)FLT autoradiography

Following a 90-minute delay for biodistribution (corresponding to the peak radiotracer uptake observed on in vivo PET images) and gamma counting, spleen, control aorta, and 14-day AngII AAA whole-organ samples were placed on a phosphor screen (PerkinElmer) covered with clear cellophane wrap. After overnight exposure, the imaging plates were scanned using the Cyclone Plus imager (PerkinElmer) at a resolution of 300 DPI. Images were processed using OptiQuant software (PerkinElmer).

2.7 Western blotting

Aortic tissue (supra-renal abdominal aortic segment only) was harvested under terminal isoflurane anaesthesia, snap-frozen in liquid nitrogen after phosphate-buffered saline perfusion, and stored at −80°C until use. Proteins were extracted using cell extraction buffer (FNN0011, Invitrogen) supplemented with Protease Inhibitor Cocktail (P8340, Sigma). The DC Protein assay kit (5000112, Bio-Rad) was used to quantify isolated proteins. The average yield was more than 3 mg of protein per aorta. Western blotting was performed for TK-1, a key enzyme expressed during DNA synthesis and the substrate for \(^{18}\text{F}\)FLT, and the specific carriers responsible for the transport of both \(^{18}\text{F}\)FLT and thymidine across the cell membrane: equilibrative nucleoside transporter (ENT)-1 and -2 and concentrative nucleoside transporter (CNT)-1 and -3. Each protein’s expression was compared to the expression of \( \beta \)-actin, a housekeeping protein expressed in all cells that is commonly used for data normalisation and as a loading control; this ensures that each well of the agarose gel contains consistent amounts of protein to be tested. For each protein, blots were conducted as per the manufacturer’s suggestions using 100 µg of protein per well. To create the running samples, \( \beta \)-mercaptoethanol (a reducing agent added to reduce disulphide bonds to irreversibly denature RNases) was added to laemmli sample buffer at a ratio of 1:4, and 3 parts of the target protein sample were then diluted into 1 part of this mixture. Antibodies and dilutions used were anti-TK-1, 1:500 (GTX113281; GeneTex, CA, USA); ENT-1, 1:1000 (ab135756); ENT-2, 1:1000 (ab181192); CNT-1, 1:500 (ab192438); CNT-3, 1:500 (ab223085); and \( \beta \)-actin, 1:1000 (Ab8226; all Abcam,
Cambridge, UK). Proteins were detected using SuperSignal West Femto Maximum Sensitivity enhanced chemiluminescent substrate (ThermoFisher Scientific, Loughborough, UK). ImageJ 1.51k software was used to quantify the optical intensity of the bands relative to that of corresponding β-actin bands (Schneider et al., 2012).

2.8 Statistical analyses

All statistical analyses were performed using Origin 2017 software (OriginLab Corporation, Northampton, MA, USA). All graphs show the mean ± the standard error of the mean (SEM) unless otherwise stated. There are two types of statistical tests: parametric and non-parametric. Parametric tests are typically used when the data to be analysed are normally distributed (i.e. the mean is a more accurate representation of the centre of the data) and sample sizes are large. Meanwhile, non-parametric tests are typically used when the data to be analysed are ‘distribution-free’ (i.e. the median more accurately represents the centre of the data) and when sample sizes are small. The Shapiro–Wilk test revealed that the data populations were normally distributed, unless indicated otherwise in the subsequent chapter; thus, parametric statistical tests were conducted to analyse all data. However, given the small sample sizes, non-parametric tests may be an appropriate alternative for statistical analysis, the results of which are presented in Appendix B. Histology, USS, and some gamma counting results were analysed using two-sampled t tests with Welch’s correction. Welch’s correction was implemented rather than Student’s t test as the former tends to be more reliable when sample sizes and variances are unequal (Glen, 2015). PET/CT, western blotting, and some gamma counting results were analysed using one-way analysis of variance with post hoc Bonferroni–Holm correction. Funding limitations generally restrict the number of animals that can be used; thus, it is important to implement a statistical test with maximum statistical power. Bonferroni–Holm correction is a uniformly more robust correction method than the Bonferroni method, as the former controls for the probability of type I error adjustments and the false discovery rate (Aickin and Gensler, 1996). The threshold for statistical significance was set at \( p < 0.05 \).
Chapter 3
Results

This chapter describes the outcomes of a series of experiments performed to answer critical questions about AAA-associated cell proliferation. Central to this thesis, PET/CT was undertaken with the aim of assessing the feasibility of using $[^{18}\text{F}]\text{FLT}$ to visualise and quantify cell proliferation in the AngII AAA model. This was followed up with Western blotting to directly test for biochemical markers of $[^{18}\text{F}]\text{FLT}$ activity, including the substrate and transporters of $[^{18}\text{F}]\text{FLT}$. An attempt was also made to explore $[^{18}\text{F}]\text{FLT}$ PET/CT in a second model of AAA, and the findings of a pilot proof-of-concept therapeutic study are also described in this chapter. The overall experimental flow of all batches of ApoE$^{-/-}$ mice is illustrated in Figure 3.1. These ApoE$^{-/-}$ mice were used to obtain the results presented in sections 3.1, 3.3, 3.4, 3.5, and 3.7.
Figure 3.1 Different cohorts of ApoE−/− mice were used for different experiments. The age (weeks) of the mice at each stage is presented across the top of the figure.
3.1 Cell proliferation is detectable in murine AAA models

AAA was first induced in mice by three methods: AngII infusion in ApoE<sup>−/−</sup> mice or CaCl<sub>2</sub> or PPE application to the aorta in C57BL6/J mice (as described in section 2.1). The aortic tissues were harvested at 28 days post-induction of AAA for the AngII and CaCl<sub>2</sub> models and 14 days post-induction of AAA for the PPE model (per the standard time points of the models). The tissues were then used for immunohistochemical staining for proliferative cells using an antibody raised against the cell proliferation marker Ki67, as described in section 2.2. Histological staining results from mouse aortic samples were analysed for the average proportion of Ki67-positive cells in three randomly selected regions containing proliferative cells (i.e. within the aortic wall), which was reflective of the proliferative activity. Aortae from control animals were compared against aortae from the three different aneurysm models. The abdominal aortic tissue from all three AAA models showed greater Ki67 staining compared to the control aortic tissue (Figure 3.2).
Aneurysm tissues show greater Ki67 staining than control aortic tissue. Representative histological images of Ki67-stained aortic tissue from the three AAA models: AngII, angiotensin II (blue); PPE, porcine pancreatic elastase (yellow); and CaCl₂, calcium chloride (green). A single example control aorta is shown in red. Ki67 positivity is indicated by the brown colour, and a blue haematoxylin counterstain has been applied to aid visualisation.
On quantification, the proportions of Ki67-positive nuclei (mean±SEM) for the PPE-, AngII-, and CaCl₂-induced AAA and saline control aortic tissues were 81.4±1.1, 78.5±2.2, 68.7±1.0, and 0.11±0.01, respectively. Tissue from all the AAA models revealed a significant increase in the proportion of Ki67-positive nuclei in the aortic wall compared to that in the saline controls (all \( p<0.001 \)) (Figure 3.3). The greatest proportions of Ki67-positive nuclei were observed in the PPE and AngII AAA models (both with significantly greater proportions compared to that in the CaCl₂ model, \( p<0.001 \)), whilst almost no proliferating cells were observed in the control aortic tissue. Overall, these data support the idea that cell proliferation occurs in the three major murine models of AAA. As the Ki67 staining was most pronounced in the AngII and PPE models, these were prioritised for use in the rest of the thesis.

3.2 AngII AAA volume correlates with the proportion of Ki67-positive nuclei

*In vivo* USS imaging was performed in the AngII AAA animals eventually used for the Ki67 analysis. Evaluation by 3D USS revealed significant increases in aortic diameter and volume following 14-day AngII infusion compared to those in saline-infused controls. The aortic diameters (mean±SEM) were 1.39±0.06 mm and 1.10±0.03 mm (\( p<0.001 \)) and aortic volumes (mean±SEM) were 15.34±0.80 mm³ and 11.16±0.73 mm³ (\( p<0.01 \)) in the 14-day AngII AAA model (\( n=9 \)) and saline controls (\( n=4 \)), respectively (Figure 3.4). The proportion of Ki67-positive nuclei showed a positive correlation with aortic volume (\( p<0.05 \), Pearson’s \( r=0.85 \)) (Figure 3.5). The results of the non-parametric statistical tests for these figures are presented in Appendix B.
Figure 3.3 PPE and AngII AAA tissue reveal the greatest proportions of Ki67-positive nuclei. Proportion of Ki67-positive nuclei in saline controls ($n=3$) and in the AngII ($n=6$), PPE ($n=7$), and CaCl$_2$ ($n=4$) AAA models. *** $p<0.001$, ns: not significant on one-way ANOVA with post-hoc Bonferroni–Holm correction.
Figure 3.4 Aortic volumes and diameters are larger in the AngII AAA model than in saline controls. Three-dimensional USS aortic reconstructions, volumes, and diameters in 14-day AngII AAA (n=9) vs. saline control aortae (n=4). ** $p<0.01$, *** $p<0.001$ on two-sample t test.
Figure 3.5 AngII AAA volume and the proportion of Ki67-positive nuclei are positively correlated. AngII AAA (n=6; blue). Pearson’s $r=0.85$, $p<0.05$. 
3.3 Cell proliferation can be visualised and quantified using $[^{18}\text{F}]$FLT PET/CT in the AngII AAA model

Although the PPE and AngII models showed no significant difference in the proportion of Ki67-positive nuclei, the AngII model was chosen to implement for further experiments described in this thesis for two main reasons: (i) the surgery for AngII AAA is much simpler than that for PPE AAA and the experiments were performed at the University of Hull, where only basic surgical facilities were available; and (ii) AngII AAA consistently develops in the suprarenal region of the abdominal aorta, whereas PPE AAA develops in the infra-renal portion (Daugherty and Cassis, 2004). As $[^{18}\text{F}]$-based PET radiotracers are excreted via the bladder, the signal in AngII AAA was anticipated to be further away from the signal due to bladder excretion than the signal in PPE AAA would be; hence, the AngII AAA was predicted to be easier to visualise and analyse.

3.3.1 Animal survival decreased over the course of the experiment

Twenty-two male ApoE$^{-/-}$ mice were enrolled in the saline vs. AngII PET/CT experiment performed at the University of Hull. Fourteen mice died at various time points during the experiment, as outlined in Figure 3.6. Images were only excluded from the subsequent SUV analysis in cases wherein the radiotracer injection was suboptimal, based on (i) judgment by the individual administering the injection and (ii) a lack of radiotracer signal in the inferior vena cava within the first 15 seconds of the scan. Meanwhile, Figure 3.7 presents Kaplan–Meier curves illustrating the proportion of surviving animals over the course of the experiment. All mice were harvested by terminal anaesthesia following the 28-day PET scan. There was a greater proportion of surviving AngII AAA mice than surviving saline controls (Kaplan–Meier survival log rank $p<0.01$); all mice were then sacrificed in week 18.
Figure 3.6 Number of surviving animals decreased from 13 to 18 weeks of age. Flow of surviving animals that produced images for analysis through the experiment.
Figure 3.7 Number of surviving animals decreased from 13 to 18 weeks of the study period. Kaplan–Meier survival curves illustrate the proportions of surviving animals due to (A) all causes of death, (B) aortic rupture, and (C) PET-related deaths. All mice were sacrificed after the final PET/CT scan in week 18.
3.3.2 $^{18}$F]FDG uptake differs between saline controls and AngII AAA mice

$^{18}$F]FDG is the most well-established radiotracer used in the clinic, particularly in the field of oncology, and is normally the first radiotracer to be considered for diagnostic imaging. Therefore, $^{18}$F]FDG was used for the initial experiment, the first preclinical PET study in collaboration with and performed at the University of Hull PET Imaging Centre. Starting with $^{18}$F]FDG allowed us to become familiar with and resolve any difficulties associated with the imaging protocol without any issues being attributed to $^{18}$F]FLT, a more novel radiotracer.

For this initial experiment, ApoE−/− mice were divided into two treatment groups: saline or AngII infusions for 28 days (saline controls and AngII AAA mice, respectively). Dynamic PET/CT was performed on day 28 following implantation of the osmotic mini-pumps in the ApoE−/− mice. Radiotracer uptake was assessed in the IVC, myocardium, paraspinal region, and abdominal aorta. $^{18}$F]FDG uptake in the IVC was visualised within the first 15 s of each scan, reflecting the bolus radiotracer injection. This early signal confirmed successful intravenous radiotracer delivery and uptake after administration (Figure 3.8).

$^{18}$F]FDG uptake was visualised in the myocardium of all mice, reflecting the typical basal myocardial energetics and confirming the radiotracer’s expected biodistribution. The myocardial $^{18}$F]FDG signal in the AngII AAA model was noticeably greater (Figure 3.9). Three-dimensional isocontour ROIs were manually drawn on the acquired images to extract SUV$_{\text{max}}$ information over time, revealing the following values for myocardial ROIs (mean±SEM SUV$_{\text{max}}$ 80–90 min post-radiotracer injection, consistent with all other SUV calculations):

1.91±0.00 (healthy controls, n=3); 5.06±0.20 (28-day AngII AAA, n=4) vs. 1.92±0.02 (28-day saline control, n=6) (two-sample t test, p<0.001) (Figure 3.10). Myocardial uptake of $^{18}$F]FDG in the 28-day AngII AAA model was greater than that in 28-day saline controls, as AngII infusion is also a model system for heart failure (Figure 3.10).
Figure 3.8 $[^{18}\text{F}]$FDG uptake was visualised in the IVC at the beginning of each scan. Representative coronal-view PET images of dynamic IVC uptake of $[^{18}\text{F}]$FDG from 4 to 14 s post-radiotracer injection. Colour scale bar indicates SUV thresholding to aid visualisation.
Figure 3.9 [$^{18}$F]FDG uptake was visualised in the myocardium of all mice. Representative coronal-view PET/CT image of static myocardial uptake 80–90 min post-radiotracer injection. Arrows indicate the myocardium. Colour scale bar indicates SUV thresholding to aid visualisation.
Figure 3.10 $[^{18}F]$FDG uptake in the myocardium increased from 0 to 90 min post-$[^{18}F]$FDG injection. (A) Time–activity curves of SUV$_{\text{max}}$ and (B) absolute values of SUV$_{\text{max}}$. (C) Time–activity curves of SUV$_{\text{mean}}$ and (D) absolute values of SUV$_{\text{mean}}$. SUV$_{\text{max}}$ and SUV$_{\text{mean}}$ in (B) and (D) represent 28-day $[^{18}F]$FDG uptake in myocardial ROIs 80–90 min post-radiotracer injection of $[^{18}F]$FDG in 28-day saline controls ($n=6$) and the 28-day AngII AAA model ($n=4$).
Uptake of $^{18}$FFDG was also visualised in the paraspinal region in several mice. This was likely in response to standard animal handling practices and resultant stress induced in the animals (Fueger et al., 2006) (Figure 3.11).

$^{18}$FFDG uptake in the abdominal aortic region in the 28-day AngII AAA model was variable, with only two mice showing clear uptake in AAA (Figure 3.12). The TACs indicated that the radiotracer distributed to the abdominal aortic region at later time points of the scan in the mice that exhibited AAA uptake (Figure 3.13). Thus, 3D isocontour ROIs were drawn on images from the final time frame, which revealed the following values (mean±SEM 80–90 min post-radiotracer injection): SUV$_{\text{mean}}$, 28-day saline controls ($n=6$) 0.07±0.004 vs. 28-day AngII AAA ($n=4$) 0.17±0.06 (two-sample t test, $p=0.06$); SUV$_{\text{max}}$, 28-day saline controls ($n=6$) 0.09±0.01 vs. 28-day AngII AAA ($n=4$) 0.24±0.08 (two-sample t test, $p<0.05$). A significant difference was noted in $^{18}$FFDG abdominal aortic SUV$_{\text{max}}$ between saline controls and 28-day AngII AAA mice (Figure 3.13).
Figure 3.11 $[^{18}\text{F}]$FDG uptake was visualised in the paraspinal region of most mice. Representative sagittal-view PET/CT image of static paraspinal uptake 80–90 min post-radiotracer injection. Arrow indicates the paraspinal region. Colour scale bar indicates SUV thresholding to aid visualisation.
Figure 3.12 $^{18}$F]FDG uptake in the abdominal aorta was greater but variable in the 28-day AngII AAA model than in 28-day saline controls. Coronal-view PET/CT images of static 28-day abdominal aortic uptake of $^{18}$F]FDG 80–90 min post-radiotracer injection in all animals. Arrows indicate the abdominal aorta. Colour scale bars indicate SUV thresholding to aid visualisation.
Figure 3.13 $[^{18}\text{F}]$FDG uptake in the abdominal aorta was variable in the 28-day AngII AAA model than in saline controls. (A) Time–activity curves of SUV$_{\text{max}}$ and (B) absolute values of SUV$_{\text{max}}$. (C) Time–activity curves of SUV$_{\text{mean}}$ and (D) absolute values of SUV$_{\text{mean}}$. SUV$_{\text{max}}$ and SUV$_{\text{mean}}$ in (B) and (D) represent 28-day $[^{18}\text{F}]$FDG uptake in abdominal aortic ROIs 80–90 min post-radiotracer injection in 28-day saline controls ($n=6$) and the 28-day AngII AAA model ($n=4$).
3.3.3 \(1^{\text{8}}\text{F}\)FLT localises to expected regions of cell proliferation in control mice

To establish reference data, healthy control mice were used to assess aortic \(1^{\text{8}}\text{F}\)FLT uptake and validate the biodistribution of \(1^{\text{8}}\text{F}\)FLT for the overall study. First, 90-min dynamic PET/CT revealed \(1^{\text{8}}\text{F}\)FLT uptake in the IVC within the first 15 s of each scan, reflecting the bolus radiotracer injection. This early signal confirmed successful intravenous radiotracer delivery and uptake after administration (Figure 3.14).

\(1^{\text{8}}\text{F}\)FLT uptake was then visualised in the spleen of all mice, reflecting the role of the spleen as a reservoir of proliferative cells and confirming the radiotracer's expected biodistribution (Figure 3.15). Three-dimensional isocontour ROIs were manually drawn on the acquired images to extract SUV\(_{\text{max}}\) information over time, revealing a mean±SEM SUV\(_{\text{max}}\) 80–90 min post-radiotracer injection of 0.47±0.03 (n=3) in the spleen (Figure 3.16). These results were further confirmed by greater ex vivo uptake of \(1^{\text{8}}\text{F}\)FLT uptake in the spleen relative to the heart by gamma counting (mean±SEM: 186.99±9.99 vs. 86.82±17.18 %ID/g, \(p<0.001\)), whilst the aorta exhibited comparable uptake to the heart (mean±SEM: 76.23±24.98 vs. 86.82±17.18 %ID/g, \(p=0.69\)), as expected (Figure 3.17). Hepatic uptake of \(1^{\text{8}}\text{F}\)FLT was noted in some animals, but this was less consistently observed across all the animals than splenic uptake. Moreover, the low signal in the bone marrow was difficult to detect as a distinct region that could be easily delineated for further ROI analyses without significantly increasing the SUV threshold on the image visualisation software (AMIDE), which resulted in saturated signal across the image and reduced visibility of other ROIs. The heart and spleen were therefore used as negative and positive control organs, respectively, for all experiments involving \(1^{\text{8}}\text{F}\)FLT thereafter.
Figure 3.14 $^{[18F]}$FLT uptake was visualised in the IVC at the beginning of each PET scan. Representative coronal-view PET images of dynamic IVC uptake of $^{[18F]}$FLT from 4 to 14 s post-radiotracer injection. Colour scale bar indicates SUV thresholding to aid visualisation.
Figure 3.15 $[^{18}\text{F}]$FLT uptake was visualised in the spleen of all healthy controls. Coronal-view PET/CT images of static splenic uptake of $[^{18}\text{F}]$FLT 80–90 min post-radiotracer injection. Arrows indicate the spleen. Colour scale bar indicates SUV thresholding to aid visualisation.
Figure 3.16 $[^{18}\text{F}]$FLT uptake in the spleen increased from 0 to 90 min post-$[^{18}\text{F}]$FLT injection. Time–activity curves representing splenic uptake of $[^{18}\text{F}]$FLT in healthy control mice ($n=3$).
Figure 3.17 *Ex vivo* $[^{18}F]$FLT uptake in the spleen was greater than that in the heart and abdominal aorta in healthy control mice. Decay-corrected *ex vivo* $[^{18}F]$FLT counts per mass units ($n=3$). *** $p<0.001$ on one-way ANOVA with post-hoc Bonferroni–Holm correction.
3.3.4 \([^{18}\text{F}]\text{FLT uptake is observed in regions of cell proliferation in the AngII model of AAA}\)

For the AngII AAA experiments, dynamic PET/CT was performed on days 14 and 28 following implantation of the osmotic mini-pumps. \([^{18}\text{F}]\text{FLT uptake in the IVC was visualised within the first 15 s of each scan, reflecting the bolus radiotracer injection. Observation of this peak was facilitated by the short 2-s timeframes, which effectively revealed the early fluctuations in activity. The appearance of this signal at such an early time point of the scan confirmed successful intravenous radiotracer delivery and uptake after administration in the saline controls and AngII AAA mice (Figure 3.18).}

\([^{18}\text{F}]\text{FLT uptake was consistently visualised in the spleen of all mice, as expected based on the baseline \([^{18}\text{F}]\text{FLT experiment in healthy control mice (Figure 3.19). Three-dimensional isocontour ROIs were manually drawn on the acquired images to extract SUV}_{\text{max}}\) information over time, revealing the following values for splenic ROIs (mean±SEM SUV}_{\text{max}}\) 80–90 min post-radiotracer injection): 0.52±0.01 (14-day AngII AAA, \(n=5\)) vs. 0.52±0.00 (14-day saline control, \(n=5\)) (two-sample \(t\) test, \(p=0.48\)); 0.37±0.07 (28-day AngII AAA, \(n=3\)) vs. 0.52 (28-day saline control, \(n=1\)) (Figure 3.20). Splenic uptake confirmed that \([^{18}\text{F}]\text{FLT distributed to expected regions, as the spleen exhibits high basal proliferative activity. Interestingly, splenic uptake in the 28-day AngII AAA model was variable.}\)
Figure 3.18 \([^{18}\text{F}]\text{FLT}\) uptake was visualised in the IVC at the beginning of each PET scan. Representative coronal-view PET images of dynamic IVC uptake of \([^{18}\text{F}]\text{FLT}\) from 4 to 14 s post-radiotracer injection. Colour scale bar indicates SUV thresholding to aid visualisation.
Figure 3.19 [\textsuperscript{18}F]FLT uptake was visualised in the spleen of all mice. Representative coronal-view PET/CT image of static splenic uptake of [\textsuperscript{18}F]FLT 80–90 min post-radiotracer injection. Arrow indicates the spleen. Colour scale bar indicates SUV thresholding to aid visualisation.
Figure 3.20 [\textsuperscript{18}F]FLT uptake in the spleen increased from 0 to 90 min post-[\textsuperscript{18}F]FLT injection. Time–activity curves representing splenic uptake of [\textsuperscript{18}F]FLT in (A) 14-day saline controls (n=5) and the 14-day AngII AAA model (n=5) and (B) 28-day saline control (n=1) and the 28-day AngII AAA model (n=3).
3.3.5 \([^{18}\text{F}]\text{FLT}\) uptake is observed in AngII AAA

The dynamic PET data revealed that \([^{18}\text{F}]\text{FLT}\) distributed to the abdominal aortic region in the AngII AAA model in the later time points of the scan (Figure 3.21). Static PET/CT images revealed that \([^{18}\text{F}]\text{FLT}\) uptake in the 14-day AngII AAA model in the final timeframe (80–90 min post-injection of \([^{18}\text{F}]\text{FLT}\)) was consistently enhanced compared to that in the saline controls (Figure 3.22).

Three-dimensional isocontour ROIs were manually drawn between the renal arteries and diaphragm to incorporate the suprarenal abdominal aortic area in the final timeframe using AMIDE v1.0.4 software. The abdominal aortic ROI was constructed between the kidneys and anterior to the anterior border of the vertebral column based on the CT images, including the most prominent PET signal in the 80–90-min timeframe. Efforts were made to avoid including the edges of the region of \([^{18}\text{F}]\text{FLT}\) uptake to minimise partial volume effects due to signal spill-over from adjacent organs, notably the bladder. For saline controls, the aortic ROI was constructed between the kidneys, superior to the bladder, and anterior to the anterior border of the vertebral column based on the CT images; these ROIs had dimensions of 1.08 mm\(^3\) and contained 15 voxels (Figure 3.23).

Incorporating mouse weights and injected radioactivity doses, output statistics revealed the SUV\(_{\text{max}}\) and SUV\(_{\text{mean}}\) across all timeframes. These values were used to generate TACs for ROIs in the abdominal aorta. The late uptake of \([^{18}\text{F}]\text{FLT}\) in the 14-day AngII AAA model was also reflected in these corresponding TACs (Figure 3.24). As the \([^{18}\text{F}]\text{FLT}\) signal was the most prominent in the final timeframe, the SUVs were extracted from the final timeframe for analysis. The following values were obtained from the data of the 14-day scan (mean±SEM 80–90 min post-radiotracer injection): SUV\(_{\text{max}}\), 0.010±0.002 (14-day saline controls, \(n=5\)) vs. 0.33±0.02 (14-day AngII AAA, \(n=5\)) (two-sample \(t\) test, \(p<0.001\)); SUV\(_{\text{mean}}\), 0.004±0.002 (14-day saline controls, \(n=5\)) vs. 0.25±0.02 (14-day AngII AAA, \(n=5\)) (two-sample \(t\) test, \(p<0.001\)) (Figure 3.24). The Shapiro–Wilk test revealed that the data of the 14-day saline control group were not normally distributed. The results of the non-parametric statistical test are presented in Appendix B.
Figure 3.21 [$^{18}$F]FLT uptake in the abdominal aorta was visualised late in the 90-min scan. Representative coronal-view PET images of dynamic AAA uptake of [$^{18}$F]FLT from 50 to 90 min post-radiotracer injection. Colour scale bar indicates SUV thresholding to aid visualisation. Arrows indicate the AAA.
Figure 3.22 $[^{18}\text{F}]$FLT uptake in the abdominal aorta was greater in the 14-day AngII AAA model than in saline controls. Coronal-view PET/CT images of static 14-day abdominal aortic uptake of $[^{18}\text{F}]$FLT 80–90 min post-radiotracer injection in 14-day saline controls ($n=5$) and the 14-day AngII AAA model ($n=5$). Arrows indicate the abdominal aorta. Colour scale bars indicate SUV thresholding to aid visualisation.
Figure 3.23 Abdominal aortic ROIs for SUV analysis were constructed avoiding adjacent organs. Representative transverse-, coronal-, and sagittal-view static PET images and corresponding abdominal aortic ROI generation in (A) saline controls (yellow) and (B) 14-day AngII AAA mice (blue).
Figure 3.24 [\(^{18}\text{F}\)]FLT uptake in the abdominal aorta was consistently greater in the 14-day AngII AAA model than in saline controls. (A) Time–activity curves of SUV\(_{\text{max}}\) and (B) absolute values of SUV\(_{\text{max}}\). (C) Time–activity curves of SUV\(_{\text{mean}}\) and (D) absolute values of SUV\(_{\text{mean}}\). SUV\(_{\text{max}}\) and SUV\(_{\text{mean}}\) in (B) and (D) represent 14-day [\(^{18}\text{F}\)]FLT uptake in abdominal aortic ROIs 80–90 min post-radiotracer injection in 14-day saline controls (\(n=5\)) and the 14-day AngII AAA model (\(n=5\)). *** \(p<0.001\) on two-sample \(t\) test.
Of all mice with AngII AAA, [\(^{18}\)F]FLT scans could be conducted at both the 14- and 28-day stages in only 3 mice, allowing for a comparison between early- and late-stage AngII AAA uptake of [\(^{18}\)F]FLT. Data from the other animals were not available due to episodes of aneurysm rupture or unsuccessful radiotracer injection. All acquired static images of [\(^{18}\)F]FLT uptake in the abdominal aortic regions of saline controls and 28-day AngII AAA mice in the 80–90-min timeframe are presented in Figure 3.25. The TACs for abdominal aortic ROIs following the 28-day PET/CT scans revealed greater [\(^{18}\)F]FLT uptake in the 28-day AngII AAA model than in the 28-day saline control (Figure 3.26). Similar to the analysis of the 14-day scan data, the SUVs were extracted from the final timeframe for analysis. Compared to [\(^{18}\)F]FLT uptake in 14-day AngII AAA, reduced uptake was noted in 28-day AngII AAA (n=3) with a mean±SEM (80–90 min post-radiotracer injection) SUV\(_{\text{max}}\) of 0.15±0.01 and SUV\(_{\text{mean}}\) of 0.13±0.02. The single 28-day saline control mouse from which suitable data were retrieved presented with an SUV\(_{\text{max}}\) of 0.03 and SUV\(_{\text{mean}}\) of 0.02, which were similar to the results of the 14-day saline controls (Figure 3.26); however, these data were not used for further analysis as they were insufficient for tests of statistical rigour.
Figure 3.25 [18F]FLT uptake in the abdominal aorta was greater in the 28-day AngII AAA model than in the saline control. Coronal-view PET/CT images of static 28-day abdominal aortic uptake of [18F]FLT 80–90 min post-radiotracer injection in 28-day saline controls (n=1) and the 28-day AngII AAA model (n=3). Arrows indicate the abdominal aorta. Colour scale bars indicate SUV thresholding to aid visualisation.
Figure 3.26 [18F]FLT uptake in the abdominal aorta was consistently greater in the 28-day AngII AAA model than in the saline control. (A) Time–activity curves of SUV$_{\text{max}}$ and (B) absolute values of SUV$_{\text{max}}$. (C) Time–activity curves of SUV$_{\text{mean}}$ and (D) absolute values of SUV$_{\text{mean}}$. SUV$_{\text{max}}$ and SUV$_{\text{mean}}$ in (B) and (D) represent 28-day [18F]FLT uptake in abdominal aortic ROIs 80–90 min post-radiotracer injection in 28-day saline controls ($n=1$) and the 28-day AngII AAA model ($n=3$).
3.3.6 Different ROI definitions may affect resultant SUVs

Amide, the image visualisation and analysis software used for the PET/CT analysis in this study, includes parameters related to the ROI that can be modified by the user. A pilot investigation was conducted to determine whether changing these parameters in the software would influence the PET image analysis results. Fixed-size and manually drawn ROIs were generated and SUV\text{mean}, SUV_{40}, SUV_{50}, SUV_{70}, SUV_{90}, and SUV_{\text{max}} were determined, as described in section 2.4.2. The mean±SEM SUV\text{mean}, SUV_{40}, SUV_{50}, SUV_{70}, SUV_{90}, and SUV_{\text{max}} for both fixed-size and manually drawn abdominal aortic ROIs on all images are presented in Figure 3.27. In all images, the fixed-size and manually drawn abdominal aortic ROIs revealed the following mean±SEM values, respectively: SUV\text{mean}, 0.14±0.01 and 0.16±0.02; SUV_{40}, 0.14±0.01 and 0.16±0.02; SUV_{50}, 0.14±0.02 and 0.16±0.02; SUV_{70}, 0.16±0.02 and 0.16±0.02; SUV_{90}, 0.19±0.02 and 0.17±0.02; and SUV_{\text{max}}, 0.20±0.02 and 0.18±0.03. Following a Shapiro–Wilk test that confirmed that the data were normally distributed (all \( p<0.05 \)), paired \( t \) tests were conducted for all fixed-size and manually drawn ROI pairs of SUV thresholds, which revealed significant differences at all SUV thresholds (\( p<0.01 \)) except at SUV_{70} (\( p=0.73 \)). This indicated that the SUVs were correlated between fixed-size and manually drawn ROIs only when a 70% threshold was applied to the voxels included in the given ROI.

The average times spent delineating the abdominal aortic regions were 0.49±0.67 min and 1.54±0.54 min when implementing the fixed-size and manually drawn approaches, respectively; thus, fixed-size ROIs are less time-consuming to generate, a finding that is likely important when tasked with analysing large datasets containing many images.
Figure 3.27 SUV\textsubscript{70} corresponding to both fixed-size and manually drawn aortic ROIs exhibited no significant differences. The SUV\textsubscript{mean}, SUV\textsubscript{40}, SUV\textsubscript{50}, SUV\textsubscript{90}, and SUV\textsubscript{max} were significantly different between fixed-size and manually drawn aortic ROIs. ** p<0.01, ns: not significant on paired t test. FS, fixed-size; MD, manually drawn.
3.3.7 $[^{18}\text{F}]$FLT uptake in the AngII AAA model may be localised to the aortic wall

Using ImageJ 1.51k software, USS and PET/CT images were manually co-registered. For the co-registration analysis, bony landmarks on CT and USS were aligned, which revealed $[^{18}\text{F}]$FLT PET signal in the aortic wall in the AngII AAA model (Figure 3.28).
Figure 3.28 [¹⁸F]FLT uptake in the AngII AAA model may occur in the aortic wall. Representative transverse-view ultrasound, [¹⁸F]FLT PET/CT, and fused ultrasound–PET/CT images. Short arrow, aortic lumen; long arrow, remodelling aortic wall.
3.4 *Ex vivo* $[^{18}\text{F}]\text{FLT}$ uptake is greater in AngII AAA than in saline control aortae

As described previously in section 3.3.3, the baseline $[^{18}\text{F}]\text{FLT}$ biodistribution study demonstrated high uptake in the spleen and low uptake in the aorta and heart; the aorta exhibited comparable uptake to the heart, whilst the spleen exhibited greater uptake relative to the heart. Therefore, the heart and spleen were used as negative and positive control reference tissues, respectively, going forward. To confirm the tissue-specific origin of the $[^{18}\text{F}]\text{FLT}$ signal observed on PET/CT in the abdominal aortic ROI, *ex vivo* gamma counting was performed using whole organs from 14-day AngII AAA mice and saline controls. Uptake in the aorta and spleen were then normalised to uptake in the heart to control for mouse-to-mouse variability. $[^{18}\text{F}]\text{FLT}$ counts in the abdominal aorta relative to the heart in the 14-day AngII-infused mice ($n=9$) were significantly increased compared to those in saline controls ($n=4$) (mean±SEM: 3.30±1.01 vs. 0.37±0.14 %ID/g, $p<0.05$) (Figure 3.29). The Shapiro–Wilk test revealed that the data of the 14-day AngII AAA group were not normally distributed. The results of the non-parametric statistical test are presented in Appendix B.

Moreover, a pilot attempt was made to perform *ex vivo* autoradiography of $[^{18}\text{F}]\text{FLT}$ uptake in whole-organ samples. Although the number of samples was small, greater $[^{18}\text{F}]\text{FLT}$ uptake in the aneurysmal aorta compared to that in the control aorta was observed, as expected based on the *ex vivo* biodistribution results.

3.4.1 Aortic $[^{18}\text{F}]\text{FLT}$ uptake is correlated with aortic volume

A significantly positive correlation was noted between $[^{18}\text{F}]\text{FLT}$ uptake on *ex vivo* gamma counting and aortic volume on *in vivo* USS ($p<0.05$, Pearson’s $r=0.67$) (Figure 3.30).
Figure 3.29 *Ex vivo* $[^{18}\text{F}]\text{FLT}$ uptake in the abdominal aorta was greater in the 14-day AngII AAA model than in 14-day saline controls. Decay-corrected *ex vivo* $[^{18}\text{F}]\text{FLT}$ counts per mass units. (A) All organs. (B) Uptake in the spleen and abdominal aorta normalised to uptake in the heart in the 14-day AngII AAA model ($n=9$) and 14-day saline controls ($n=4$). * $p<0.05$ on two-sampled $t$ test with Welch’s correction.
Figure 3.30 Aortic volume is positively correlated with $[^{18}\text{F}]\text{FLT}$ uptake. Correlation between aortic volume and ex vivo $[^{18}\text{F}]\text{FLT}$ counts in AngII AAA (blue, $n=9$). Pearson’s $r=0.67$, $p<0.05$. 
3.5 Expression of key molecules involved in [¹⁸F]FLT metabolism are upregulated in the AngII AAA model

The [¹⁸F]FLT PET/CT data presented in section 3.3 suggest increased abdominal aortic proliferative activity in the 14-day AngII AAA model compared to that in control animals, which diminishes at the 28-day time point. To further investigate the change in [¹⁸F]FLT signal observed between 14- and 28-day AngII AAA, an independent cohort of mice was divided into 4 groups, and aortae were harvested: baseline group (n=3), containing ApoE⁻/⁻ mice; saline control group (n=3), containing ApoE⁻/⁻ mice that received saline infusions for 14 days; 14-day AngII AAA group (n=3), containing ApoE⁻/⁻ mice that received AngII infusions for 14 days; and 28-day AngII AAA (n=3), containing ApoE⁻/⁻ mice that received AngII infusions for 28 days. This was followed by western blotting for (i) TK-1, the key substrate of [¹⁸F]FLT, and (ii) the nucleoside transporters that carry [¹⁸F]FLT into the cell: ENT-1, ENT-2, CNT-1, and CNT-3. The response of these proteins to AngII-induced AAA is unknown.

3.5.1 Expression of the substrate of [¹⁸F]FLT is increased in the AngII AAA model

Consistent with [¹⁸F]FLT uptake observed using PET/CT, the TK-1 band intensity was increased after 14-day AngII infusion compared to that after saline control and 28-day AngII infusion (mean±SD TK-1:β-actin band intensities: 2.45±0.24, 0.34±0.02, and 1.07±0.07, respectively; all p<0.001) (Figure 3.31). The results of the non-parametric statistical test are presented in Appendix B.
Figure 3.31 TK-1 expression was the greatest in 14-day AngII AAA. (A) Representative TK-1 western blots. (B) Quantitative analysis. TK-1 band intensity was normalised to β-actin band intensity. *** p<0.001, ns: not significant on one-way analysis of variance with post-hoc Bonferroni–Holm correction.
3.5.2 Expression of [$^{18}$F]FLT transporters is increased in the AngII AAA model

The ENT-1 band intensity was increased after 14-day AngII infusion compared to that after saline control and 28-day AngII infusion (mean±SD ENT-1:β-actin band intensities: 2.74±0.09, 0.10±0.06, and 1.45±0.11, respectively; all $p<0.001$) (Figure 3.32). The Shapiro–Wilk test revealed that the data of the saline control group were not normally distributed. The intensity of the ENT-2 band was also increased after 14-day AngII infusion compared to that after saline control and 28-day AngII infusion (mean±SD ENT-2:β-actin band intensities: 2.66±0.13, 0.29±0.00, and 1.68±0.04, respectively; all $p<0.001$) (Figure 3.33). Keeping with a similar pattern, the CNT-1 band intensity was increased following 14-day AngII infusion compared to that following saline control and 28-day AngII infusion (mean±SD CNT-1:β-actin band intensities: 4.06±0.45, 0.24±0.02, and 1.53±0.09, respectively; all $p<0.001$) (Figure 3.34). Finally, the intensity of the CNT-3 band was increased after 14-day AngII infusion compared to that after saline control and 28-day AngII infusion (mean±SD CNT-3:β-actin band intensities: 3.69±0.70, 0.40±0.21, and 2.03±0.44, respectively; all $p<0.001$) (Figure 3.35). Overall, the key proteins tested that play a role in [$^{18}$F]FLT metabolism were upregulated in 14-day AngII AAA tissue compared to 28-day AngII AAA and saline control tissue. For reference, all the tested proteins were found to be expressed in tissue from a saline control spleen, which was expected as the spleen contains proliferative cells. The differences in protein expression were insignificant between baseline and saline control aortae for all the tested proteins. The results of all non-parametric statistical tests corresponding to these findings are presented in Appendix B.
Figure 3.32 ENT-1 expression was the greatest in 14-day AngII AAA. (A) Representative ENT-1 western blots. (B) Quantitative analysis. ENT-1 band intensity was normalised to β-actin band intensity. *** $p<0.001$, ns: not significant on one-way analysis of variance with post-hoc Bonferroni–Holm correction.
Figure 3.33 ENT-2 expression was the greatest in 14-day AngII AAA. (A) Representative ENT-2 western blots. (B) Quantitative analysis. ENT-2 band intensity was normalised to β-actin band intensity. *** p<0.001, ns: not significant on one-way analysis of variance with post-hoc Bonferroni–Holm correction.
Figure 3.34 CNT-1 expression was the greatest in 14-day AngII AAA. (A) Representative CNT-1 western blots. (B) Quantitative analysis. CNT-1 band intensity was normalised to β-actin band intensity. *** \( p < 0.001 \), ns: not significant on one-way analysis of variance with post-hoc Bonferroni–Holm correction.
Figure 3.35 CNT-3 expression was the greatest in 14-day AngII AAA. (A) Representative CNT-3 western blots. (B) Quantitative analysis. CNT-3 band intensity was normalised to β-actin band intensity. *** $p<0.001$, ns: not significant on one-way analysis of variance with post-hoc Bonferroni–Holm correction.
3.6 Cell proliferation is difficult to visualise and quantify using $[^{18}\text{F}]$FLT PET/CT in the PPE AAA model

The PPE model of AAA is a well-established non-progressive disease model that is less likely to rupture than the AngII infusion model (Senemaud et al., 2017). As one of the commonly studied murine models of AAA and the model that showed the greatest proportion of Ki67-positive nuclei (reported in section 3.1), aortic volumes and $[^{18}\text{F}]$FLT uptake were explored in the PPE AAA model to investigate if the findings from the AngII model could be replicated in a second model.

3.6.1 Aortic volumes and diameters in the PPE AAA model are larger than those in sham controls

Evaluation by 3D USS revealed the expected significant increases in aortic diameter ($p<0.05$) and volume ($p<0.05$) 14 days post-PPE application compared to baseline and sham-operated controls. The aortic diameters were 1.66±0.53 mm (mean±SEM) and 0.52 mm and aortic volumes were 18.72±5.47 mm$^3$ (mean±SEM) and 2.70 mm$^3$ in the 14-day PPE model ($n=3$) and 14-day sham control ($n=1$), respectively (Figure 3.36).
Figure 3.36 Aortic volume and AP diameters were greater in the 14-day PPE AAA model than in sham controls. Aortic volumes and AP diameters based on 3D USS of 14-day PPE AAA (n=3) vs. 14-day sham control aortae (n=1). ns: not significant on two-sample t test.
3.6.2 *In vivo* $[^{18}\text{F}]$FLT uptake is variable in the PPE AAA model

Static PET/CT images revealed that $[^{18}\text{F}]$FLT uptake in the 14-day PPE AAA model 80–95 min post-injection of $[^{18}\text{F}]$FLT was variable and noisy due to uptake in regions surrounding the abdominal aorta, such as areas of the bowel, leading to inconclusive evidence of $[^{18}\text{F}]$FLT uptake in PPE AAA (Figure 3.37). Additionally, multiple hotspots of diffuse radiotracer uptake were noted on the PET images of the abdominal region, making it difficult to confidently delineate the abdominal aorta. This background uptake was not observed in the AngII AAA model, leading to the suspicion that these hotspots in the abdominal and bowel areas resulted from the method of PPE application. For this method, the mice are laparotomised, and 10 µL of PPE is applied to the abdominal aorta using a syringe. The $[^{18}\text{F}]$FLT signal may relate to post-surgical changes in the abdomen involving the formation of adhesions, which can occur following any laparotomy.

3.6.3 *Ex vivo* $[^{18}\text{F}]$FLT uptake in PPE AAA is similar to that in sham control aortae

*Ex vivo* gamma counting was then performed, as per section 2.5. The experiment revealed no significant difference in $[^{18}\text{F}]$FLT counts in the abdominal aorta relative to the heart between the 14-day PPE AAA and 14-day sham control (Figure 3.38). These results suggested that there was no significant uptake in the AAA.
Figure 3.37 $^{18}$FFLT uptake in the abdominal aorta was difficult to visualise in the 14-day PPE AAA model. Coronal-view PET/CT images of static 14-day abdominal aortic uptake of $^{18}$FFLT 80–95 min post-radiotracer injection in all animals. Arrows indicate the abdominal aortic region. Colour scale bars indicate SUV thresholding to aid visualisation.
Figure 3.38 *Ex vivo* [18F]FLT uptake in the abdominal aorta was similar in PPE AAA and sham control aortae. Decay-corrected *ex vivo* [18F]FLT counts per mass units (A) All organs. (B) Uptake in the spleen and abdominal aorta normalised to uptake in the heart in PPE AAA (*n*=3) and sham control (*n*=1) aortae.
3.7 [¹⁸F]FLT PET/CT might be useful to assess a therapeutic response in the AngII AAA model

Finally, [¹⁸F]FLT PET/CT was used in a proof-of-concept exploratory experiment to determine if the response to an anti-proliferative tyrosine kinase inhibitor commonly used as a cancer therapeutic (imatinib) could be detected in the AngII AAA model. Considering the high rupture rate of the AngII AAA model and economic feasibility, 15 mice received AngII infusions. Aortic rupture occurred in 5 mice within 12 days of AngII treatment. Three-dimensional USS after 12-day AngII infusion revealed that aneurysms developed in all surviving mice; 6 of the 10 mice presented with aneurysms that were neither dissecting nor ectatic, which were categorised as ‘high priority’ for PET/CT imaging. The remaining 4 mice with smaller aneurysms were categorised as ‘low priority’ for PET/CT imaging.

Pre-treatment [¹⁸F]FLT PET/CT was performed on day 13 of AngII infusion, during which 2 of the high-priority mice and 2 of the low-priority mice died following [¹⁸F]FLT injection. The PET/CT data acquired from 3 of the 4 mice that survived to produce follow-up data revealed notable [¹⁸F]FLT uptake in the AAA region. Three-dimensional isocontour ROIs revealed a mean±SEM (n=3) SUV\textsubscript{max} of 0.19±0.01 and SUV\textsubscript{mean} of 0.18±0.01. All mice were then administered 3 doses of imatinib or tap water (as the vehicle treatment) via oral gavage before undergoing follow-up PET/CT on day 15 of AngII infusion. Post-imatinib PET data revealed a mean±SEM (n=2) SUV\textsubscript{max} of 0.02±0.0 and SUV\textsubscript{mean} of 0.02±0.0; these were significantly decreased compared to the pre-treatment values (p<0.001 and p<0.001, respectively). [¹⁸F]FLT uptake in AAA decreased after 3 doses of imatinib, reflecting an anti-proliferative response that was detectable in the early stage of AAA progression using PET/CT. Meanwhile, matched tap water treatment resulted in no significant change in [¹⁸F]FLT uptake, lending support to the imatinib data (n=1: SUV\textsubscript{max} of 0.23 and SUV\textsubscript{mean} of 0.21, respectively, compared to pre-treatment) (Figure 3.39).

Ex vivo gamma counting was then performed. [¹⁸F]FLT counts in the abdominal aorta relative to the heart in the AngII AAA mice that received imatinib were decreased compared to that in the AngII AAA mouse that received vehicle treatment (mean±SEM: 23.89±8.57 vs. 51.55 %ID/g) (Figure 3.40).
Figure 3.39 \([^{18}F]\)FLT uptake in AngII AAA decreased following imatinib treatment. Coronal-view PET/CT images of static abdominal aortic uptake of \([^{18}F]\)FLT 80–100 min post-radiotracer injection pre- and post-treatment with imatinib or vehicle (tap water). Arrows indicate the abdominal aorta. Colour scale bars indicate SUV thresholding to aid visualisation.
Figure 3.40 Ex vivo $[^{18}\text{F}]$FLT uptake in AngII AAA was reduced following imatinib treatment than that following vehicle treatment. Ex vivo $[^{18}\text{F}]$FLT uptake in the spleen and abdominal aorta normalised to uptake in the heart following vehicle ($n=1$) and imatinib ($n=2$) treatment.
3.8 Summary and conclusions

This chapter presented critical experiments conducted to explore if $^{[18F]}$FLT uptake could be observed in experimental murine models of AAA. The proportion of Ki67-positive nuclei was shown to be the greatest in the PPE and AngII AAA models and is correlated with aortic volume in the AngII AAA model. Uptake of $^{[18F]}$FDG was investigated in saline controls and the AngII AAA model, revealing variable aneurysmal uptake. Myocardial $^{[18F]}$FDG uptake was observed in all animals, with significantly greater uptake in AngII AAA mice. Furthermore, the uptake of $^{[18F]}$FLT was explored in healthy controls to establish control data and validate the use of $^{[18F]}$FLT as a cell proliferation marker in a non-disease control model. Finally, and most importantly, $^{[18F]}$FLT PET/CT appears to be a feasible modality to visualise and quantify cell proliferation in the AngII AAA model. This was further validated by ex vivo $^{[18F]}$FLT uptake in AAA, which was correlated with aortic volume 14 days post-induction of AAA. The expression levels of TK-1, the substrate of $^{[18F]}$FLT, and the transporters that carry $^{[18F]}$FLT into cells are also increased in the 14-day AngII AAA model, mirroring the in vivo $^{[18F]}$FLT PET data. The $^{[18F]}$FLT results in the AngII model could not be positively replicated in the PPE model. An exploratory study demonstrated that $^{[18F]}$FLT PET/CT may be used to visualise a decrease in the proliferative signal in the AngII AAA model following 3 doses of imatinib; however, these results must be interpreted with caution owing to the small sample sizes.
Chapter 4 Discussion

The main objective of this thesis was to assess the feasibility of using $^{18}$FFLT PET/CT to visualise and quantify cell proliferation in the AngII AAA mouse model. In 14-day AngII-induced AAA compared to 28-day AngII-induced AAA and saline-infused controls, the $^{18}$FFLT PET signal was found to be significantly increased, along with greater $^{18}$FFLT counts in excised aortic tissue, which correlate with aortic volume. The expression of the $^{18}$FFLT substrate TK-1 and $^{18}$FFLT transporters ENT-1, ENT-2, CNT-1, and CNT-3 are also increased in 14-day AngII AAA than in 28-day AngII AAA and saline controls. These findings collectively suggest an early period of cell proliferation in the AngII AAA murine model, which is detectable using PET/CT. Some key findings are discussed in this chapter, including the evidence of cell proliferation and an early active growth phase, the correlation between aortic volume and cell proliferation, and features of PET/CT image analysis that may be improved to generate even more robust results. These key aspects are further discussed in the context of clinical applications to highlight the long-term translatability of the findings of this thesis. This chapter concludes with a discussion of areas for future research and a brief summary.

It is well known that the AngII AAA mouse model exhibits a number of features that pose a challenge to the type of experiments described in this thesis, wherein large numbers of subjects are required to optimally utilise both the expensive bespoke radiotracer production and time on the PET/CT scanner: (i) the AngII model has a high rate of mortality due to aneurysm rupture, with most deaths occurring within the first 7 days following minipump implantation (Cao et al., 2010); (ii) a proportion of animals will not respond to AngII infusion to form an aneurysm (i.e. non-responders); and (iii) tail vein injection for radiotracer delivery in the AngII model was more challenging than in wildtype C57BL6/J mice, and some animals needed to be excluded because of failure of radiotracer delivery via the fragile and poorly accessible tail vein. To overcome (i–iii), the sample sizes at the start of the experiments were overestimated to take account of estimated losses due to aortic rupture, non-response, and technical problems. However, to achieve this, ApoE$^{-/-}$ mice had to be bought from commercial suppliers, as sufficient numbers could not be guaranteed from the University’s internal breeding programme. During the project, the main supplier discontinued breeding in the UK, and the animals had to be imported from the EU, posing
a significant economic challenge. Therefore, the sample sizes in this study were relatively small.

4.1 Histological evidence of cell proliferation in AAA

The first finding in the present study was evidence of a significantly increased proportion of cells with Ki67-positive nuclei in the suprarenal abdominal aortic wall from AngII-infused ApoE⁻/⁻ mice compared to those from saline-infused controls with a relative difference of approx. 200%, which was positively correlated with aortic volume. This result indicated that cell proliferation is indeed upregulated in the AngII AAA model. This is an important finding to justify the use of an expensive tool like $^{18}$FFLT PET/CT later in the study to test its feasibility in visualising and quantifying the observed cell proliferation. Ki67 staining is a well-established method to identify proliferative cells, and $^{18}$FFLT is a marker of cell proliferation; in line with this, the correlation between $^{18}$FFLT uptake and Ki67 positivity has been demonstrated in several tumour models (Grierson and Shields, 2000, Salskov et al., 2007). Ki67 is an extensively utilised marker for cell proliferation in laboratory and clinical studies, and its utility has been demonstrated in detecting proliferative activity in numerous cell types in various diseases. In the present study, significantly greater proportions of Ki67-positive nuclei were observed in tissues from three different AAA murine models than that from saline control tissue. This is in line with reports of increased Ki67 positivity in patients with AAA, reflecting VSMCs, endothelial cells, and macrophages based on combined staining for anti-α-smooth muscle actin, cluster of differentiation (CD)-31, and CD68, respectively (Cafueri et al., 2012, Ryer et al., 2015). In terms of other non-aneurysmal diseases, positive Ki67 expression is observed in systemic lupus erythematosus in association with disease activity (Hudspeth et al., 2019), and positive Ki67 staining has also been noted in human and ApoE⁻/⁻ murine atherosclerotic lesions, wherein the proliferating cells are suggested to be rich in macrophages and inflammatory cell infiltrates in the active growth phase of lesion development, eventually leading to localised apoptosis; however, the exact cell types contributing to the disease process remain mostly uncharacterised, similar to that in AAA (Lhoták et al., 2016). The heightened Ki67 positivity observed in the present study is also similar to that observed in human prostate carcinoma (Keshgegian et al., 1998, Claudio et al., 2002), follicular thyroid tumours (Mu et al., 2018), severe cervical cancer lesions (Silva et al., 2017), and breast cancer (Qiu et al.,
Therefore, the positive Ki67 data in the AngII AAA mouse samples suggest that there is indeed a proliferative phase in the AngII AAA model.

4.2 Expression of key players in the $[^{18}\text{F}]$FLT mechanistic pathway

In line with the Ki67 data, the level of TK-1 expression was significantly increased in 14-day AngII AAA tissue than in saline control tissue with a relative difference of approximately 151%, followed by a decrease to a 78% relative difference in 28-day AngII AAA tissues, supporting the notion that cell proliferation is an event that occurs during the progressive phase of AAA in this model. $[^{18}\text{F}]$FLT is a substrate for cytoplasmic TK-1, which has been demonstrated as a proliferation biomarker in leukaemia, Hodgkin’s and non-Hodgkin’s lymphoma, lung carcinoma, and breast cancer, amongst other cancers (Zhou et al., 2013). Furthermore, studies of TK-1 expression in cancer have revealed its associations with tumour stage, histological grade, metastasis, size, and distant and local recurrence (He et al., 2000, Mao et al., 2002, He et al., 2006, Aufderklamm et al., 2012, Nisman et al., 2013). The (i) overexpression of TK-1 and (ii) its correlation with the Ki67 proliferation index have also been demonstrated in various cancers (Mao et al., 2005, Chen et al., 2013, Bagegni et al., 2017).

In addition to TK-1, the expression levels of the equilibrative and concentrative transporters ENT-1, ENT-2, CNT-1, and CNT-3 were found to be significantly increased in 14-day AngII AAA tissue compared to those in saline control tissues with relative differences of 160–185%, followed by a decrease to 45–90% relative differences in 28-day AngII AAA tissues. These transporters contribute to nucleoside homeostasis, and as they are responsible for the cellular uptake of some nucleoside-based drugs, they may be valuable in nucleoside-derived therapy (Molina-Arcas et al., 2009, Jiraskova et al., 2018). These nucleoside transporters are also reported to influence $[^{18}\text{F}]$FLT uptake; Paproski et al. previously characterised $[^{18}\text{F}]$FLT transport by these transporters in cancer cell lines and demonstrated that they contribute to $[^{18}\text{F}]$FLT uptake, with ENT-1 showing the most significant expression (Paproski et al., 2008, Paproski et al., 2010). There is an established association between $[^{18}\text{F}]$FLT uptake and proliferation- or thymidine-associated marker expression in various cancers, some of which include Ki67 expression in breast cancer, CD8 and Ki67 expression in metastatic prostate cancer, Ki67 and TK-1 expression in lung cancer, and proliferating cell nuclear antigen and TK-1 expression in fibrosarcoma; these
associations are interesting to draw parallels with correlations amongst $^{18}$F-FLT uptake and the expression of TK-1, ENT-1, ENT-2, CNT-1, and CNT-3 observed in AAA (Barthel et al., 2003, Brockenbrough et al., 2011, Woolf et al., 2014, Scarpelli et al., 2019). Furthermore, concomitant malignancies have been reported in up to 14% of AAA cases, of which most involve colorectal cancer (Jibawi et al., 2011). Although it might be ambitious to propose that cancers and AAA involve similar proliferative mechanisms, further studies are warranted to determine whether significant commonalities exist between their pathogeneses. Considering our current understanding of $^{18}$F-FLT in cancer progression and proliferation, the findings of these proteins being upregulated and expressed in AngII AAA tissue support the notion of proliferation occurring at an early stage of AAA development.

4.3 $^{18}$F-FLT uptake in the AngII AAA model

Given the significantly high proportion of Ki67-positive nuclei at the end of the period of AngII infusion (28 days post-implantation of the mini-pump), $^{18}$F-FLT PET/CT was planned to be performed on day 28. As this experiment was complex and expensive, the decision was made to perform an additional scan at a second time point (i.e. day 14) to identify if there was a change in radiotracer uptake over time in the AngII AAA disease course. In this study, the uptake of $^{18}$F-FLT in AAA was successfully demonstrated for the first time. Interestingly, the uptake significantly decreased by a relative difference of 55% between days 14 and 28 of the model, suggesting that there may be a decrease in proliferative activity in the late stage of AAA. Furthermore, the alignment of $^{18}$F-FLT results with Ki67 staining data and supporting evidence from Western blotting for key players in the $^{18}$F-FLT mechanistic pathway support the PET/CT observations and raise the potential of using $^{18}$F-FLT as a tool to observe proliferative activity in in vivo model systems or indeed in humans.

As described in section 1.3.5, AAA formation and progression rely on multiple contributing factors. These include changes in the mechanical properties of the vessel wall, such as wall stress and elasticity; inflammatory cell infiltration of the aortic wall; increased autoimmunity; enhanced oxidative stress; vascular remodelling; degradation of the ECM; and microcalcification. This multifactorial nature corresponds to many cells contributing to the AAA pathophysiology, such as lymphocytes, mast cells, macrophages, ECM proteins, VSMCs, and endothelial cells, amongst others (Wang et al., 2014, Kuivaniemi et al., 2015, Sun et al., 2018). Reports of the precise cell types that may contribute to the early-stage proliferative signal are largely
inconclusive, owing to the wide variety of implicated cell types and difficulties in accurately identifying them in vivo using antibody staining. Although the experiments presented in this thesis do not concretely prove that the proliferative signal originates from VSMCs, the findings are in line with reports of VSMC apoptosis, medial wall thinning, and degeneration in late-stage AAA, which are critical for aortic dilatation and rupture (Lopez-Candales et al., 1997, Henderson et al., 1999, Ailawadi et al., 2009, Riches et al., 2013, Salmon et al., 2013, Clement et al., 2019, Quintana and Taylor, 2019). These events are associated with VSMC phenotypic switching early in AAA development. The reduced density of VSMCs in late-stage AAA is suggested to be the result of apoptosis, evidenced by the observation of apoptotic VSMCs in the medial layer of AAA in humans (Rowe et al., 2000, Kuivaniemi et al., 2015). When the rate of VSMC apoptosis exceeds the rate of VSMC proliferation, the number of SMC layers in the aortic wall decreases, eventually leading to rupture. When VSMC proliferation is promoted via anti-inflammatory treatment with interleukin-10, the degradation of SMCs is inhibited, leading to a delay in the development of AAA in rabbits (Zhu et al., 2019). Aneurysm growth is also inhibited in ApoE−/− mice following treatment with the xanthine derivative KMUP-3, which inhibits AAA phenotypic switching and apoptosis (Lai et al., 2020). Moreover, studies have demonstrated VSMC proliferation and the role of dedifferentiated medial VSMCs in neo-intimal development following vascular injury (Herring et al., 2014, Roostalu et al., 2018). These findings propose an important role for VSMCs in AAA development; however, further experiments are warranted to determine the precise contributions of other cell types and regardless of the implicated cell types, it is suggested that the pathobiology differs between early- and late-stage AAA.

The overall findings of an increase in proliferation-associated biomarkers at 14 days followed by a decrease at 28 days point to an active period of cell proliferation early in the AngII-infused AAA disease course that then leads to replicative senescence and reduced proliferative activity late in the disease course. ‘Repli- cative senescence’ was first described in the context of human fibroblasts in culture and reflects the process that limits the proliferative activity of cells, as cells have a finite life span in which division occurs (Campisi, 1997). Cell senescence has been reported in patients with AAA and patients manifesting risk factors of AAA (Liao et al., 2000, Gacchina et al., 2011), and senescence can also progress to apoptosis (Thompson et al., 1997). In addition, accelerated replicative senescence comprises a distinct phenotype of VSMCs in human AAA compared to that of VSMCs in the non-
aneurysmal inferior mesenteric artery, indicating that senescence may play a role in the VSMC reduction that is specifically observed in AAA (Liao et al., 2000, Riches et al., 2018). Furthermore, this late-stage VSMC reduction may reflect the increased accumulation of ECM degradative proteins, the production of which is stimulated by VSMCs, as well as endothelial cells, adventitial fibroblasts, and inflammatory cells that are adherent to the ECM. In AAA, the amount of proteins that confer aortic wall integrity, such as collagen and elastin, is reduced compared to that in normal aortae (Lin et al., 2018, Quintana and Taylor, 2019). An imbalance between proteolytic enzymes, such as MMPs, and their inhibitors contributes to this reduction; correspondingly, various MMPs have been shown to be produced by VSMCs, and a marked increase in MMP expression has then been shown to be associated with a reduction in VSMCs with AAA disease progression (Knox et al., 1997, Mao et al., 1999, Kadoglou and Liapis, 2004, Fanjul-Fernández et al., 2010, Courtois et al., 2013, Lin et al., 2018, Quintana and Taylor, 2019). This imbalance leads to a disruption in the equilibrium between ECM synthesis and degradation, which then influences the course of expansion and rupture of AAA. For example, slow-growing aneurysms may reflect ECM synthesis mechanisms counterbalancing the ECM degradation mechanisms. Acceleration of aneurysm growth and rupture in the AAA disease course may result from this equilibrium shifting towards degradation. Therefore, it is encouraging that the data presented in this thesis suggest a period of decreased proliferative activity late in the AAA disease course, which may suggest that VSMCs may partially contribute to the proliferative signal being detected.

*Ex vivo* gamma counting of whole organs, which provided definitive evidence of the $^{[18]}$F FLT hotspots noted on the PET/CT images, revealed 160% greater $^{[18]}$F FLT uptake in 14-day AngII AAA compared to saline control aortae. All the counted organs also demonstrated greater uptake of $^{[18]}$F FLT compared to that in the saline control organs (Figure 3.29). This observation may be explained by the effects of AngII on widespread cell proliferation in the AngII-infused model of AAA. As introduced in section 1.4.1.1, AngII mediates growth processes and has been shown to induce the activity of other vasoactive factors, such as endothelin, which further confer growth-altering effects in cells of the kidneys, lungs, and intestines, among other organs (Johnson et al., 1992, Wolf and Wenzel, 2004, Slice et al., 2005, Wang et al., 2015). Regardless of the effects of AngII in other organs, it is encouraging and convincing to observe such a significant increase in proliferative activity as measured by $^{[18]}$F FLT uptake in the aneurysmal aorta
of the classical AngII-infused model of AAA. In addition to the abdominal aorta, the spleen may be another interesting region for analysis. As expected, $[^{18}\text{F}]$FLT uptake was observed in the spleen of all animals. Significant differences in splenic SUVs were not observed between saline control and 14-day AngII AAA mice. However, in 28-day AngII AAA mice, the splenic SUVs showed a 33% relative difference and a slightly greater SEM. Further investigation of this splenic uptake in a greater number of 28-day AngII AAA mice is warranted to better understand if long-term AAA disease is associated with changes in proliferative activity in the spleen. The spleen is a major repository of proliferative cells, as monocytes differentiate into dendritic cells and macrophages in tissue healing and repair processes (Drutman et al., 2012); thus, $[^{18}\text{F}]$FLT uptake in the spleen is expected. Splenic uptake of $[^{18}\text{F}]$FLT was also noted by Ye et al. in atherosclerotic mice, although with a difference in splenic SUVs between wildtype and ApoE$^{-/-}$ mice (Ye et al., 2015). Furthermore, in male patients with AAA, maximal aortic diameter and spleen volume exhibit a strong positive correlation (Li et al., 2017); however, more data are needed to suggest that spleen enlargement may play a role as an indicator of AAA progression and rupture. The variations in splenic SUVs observed in this study are in line with these previous findings, indicating that further research on the association between splenic activity and AAA progression is warranted.

Finally, a positive correlation was observed between ex vivo $[^{18}\text{F}]$FLT uptake and aortic volume at 14 days. Given the limitations of this study, there was no evidence to suggest that aortic volume remains positively correlated specifically with $[^{18}\text{F}]$FLT uptake in later stages of AAA development, while other mechanisms may contribute to changes in aortic volume with further AAA progression. Nonetheless, this result is important in the context of early-stage AAA management post-US$S$ screening, as $[^{18}\text{F}]$FLT uptake may provide an additional dimension of information about AAA progression: proliferative activity in relation to early aortic volume changes. The utility of this additional dimension has considerable potential in risk assessment; for example, a small but highly active AAA may warrant treatment, whereas an indolent large AAA may not. Similar to the way in which SUV thresholds of $[^{18}\text{F}]$FDG uptake are currently implemented in the cancer field to classify tumour grade/aggressiveness and guide patient intervention, there may thus be a role for $[^{18}\text{F}]$FLT uptake thresholds in AAA management. Following from the previous example, a patient exhibiting greater aneurysmal $[^{18}\text{F}]$FLT uptake (and thus harbouring early-stage AAA) might benefit more from an anti-proliferative drug therapy, compared to a patient exhibiting lower
[18F]FLT uptake (and thus harbouring late-stage AAA) who would benefit from surgical repair. Therapy for AAA currently involves patient health optimisation and surgical intervention. An added layer of information may pave the way for trialling medical therapies as opposed to implementing surgery-only options based on physical aneurysm characteristics. However, to confirm the correlation between [18F]FLT uptake in the abdominal aorta and AAA severity, further longitudinal preclinical investigations of aneurysmal [18F]FLT uptake and its associations with aortic size at baseline, 14 days, and 28 days in the AngII AAA model are needed, a key limiting aspect in the current study due to the high rates of mortality, non-response to AngII, and failure of intravenous radiotracer administration, as well as the significantly high costs of the AngII AAA mouse model. Using the same cohort of mice from study initiation (i.e. day of mini-pump implantation) to completion (i.e. day 28 of AngII infusion) would yield robust results, although large numbers of animals would be needed to mitigate the aforementioned challenges. In terms of clinical research, a large-scale analysis of patient data to precisely determine the relationship between aortic sizes based on USS screening and [18F]FLT uptake on PET would facilitate a clearer understanding of whether [18F]FLT SUVs might be used as informative indicators of size and/or AAA stage.

4.4 PET image analysis

It is worth noting here that the abdominal aorta and spleen are located in close proximity to the urinary bladder and kidneys; the two latter organs readily take up [18F]FLT (as all [18F]-based radiotracers are renally excreted), which consequently raises the possibility of partial volume effects due to signal spill-over into the aortic ROI. Moreover, the combination of the PET scanner’s spatial resolution (1.55 mm) (Sanchez et al., 2013) and diameter of the aneurysmal aorta (1.5–3 mm) made it difficult to definitively study [18F]FLT uptake within specific regions of the aortic wall. Nonetheless, to validate that the PET/CT signal was indeed originating from the aortic wall, ex vivo [18F]FLT gamma counting of whole aortae was performed following PET scanning, and the [18F]FLT PET/CT and 3D USS datasets were manually co-registered. To mitigate the risk of introducing significant partial volume effects in the image analysis results, SUV_{max} was the parameter of choice when reporting PET data in presentations or publications, for example. The SUV_{max} is prone to being affected by image noise, whilst SUV_{mean} has high inter-observer variability (Büyükereli et al., 2016); therefore, applying a percentage threshold of the maximum voxel values
within preselected ROIs is an approach that may help exclude regions of low radiotracer uptake. Determining optimal quantification metrics to assess PET images of AAA is an ongoing effort (Akerele et al., 2020).

A pilot investigation of the effects of applying different SUV thresholds and ROI definitions on the resultant SUVs in the present study revealed that the SUV\text{mean}, SUV\text{40}, SUV\text{50}, SUV\text{90}, and SUV\text{max} of abdominal aortic uptake of [\text{18F}]FLT in both saline controls and AngII AAA mice were significantly different between fixed-size and manually drawn ROIs; meanwhile, the SUV\text{70} was not significantly different between the two types of ROIs. A potential explanation of these findings may lie in the presumption that applying a 70% SUV threshold may be an optimal balance between the SUV\text{max}, which is predominantly affected by noise, and SUV\text{mean}, which is predominantly affected by partial volume effects in a given ROI. Furthermore, these findings are in line with supporting evidence to suggest that applying threshold values of 70–75% may yield consistent quantitative outcomes of lesion uptake throughout a longitudinal study period, regardless of changes in lesion morphology (Krak et al., 2005). However, conversely, other studies have found that thresholding values of 40–43% may be adequate to segment lesions (Ford et al., 2006, Sher et al., 2016), implicating the need for further studies to achieve conclusive results of optimal thresholds. In the present study, the resultant SUVs corresponding to the manually drawn ROIs showed less variability compared to those corresponding to the fixed-size ROIs, which may have been attributed to the fact that the manually drawn ROIs were generated uniquely to each dataset that i) avoided the edges of the region of uptake to minimise partial volume effects, ii) were in accordance with the morphology of the aneurysms, and iii) could be adapted to individual differences in the anatomy. On the other hand, the fixed-size ROIs did not account for these factors, as they were simply sized according to the maximum aortic diameter acquired in the study model without individual considerations of ROI edges, leading to more variable resultant SUVs, because of the included voxels being independent of the lesion size and shape. In addition, implementing fixed-size ROIs was significantly more time-efficient and a more objective approach than manually drawing ROIs. As a result, future research on ROI definitions will need to resolve the issues that manually drawn ROIs are currently able to overcome.

The overall findings are likely to have clinical implications when analysing large datasets and images containing regions of low uptake; in these cases,
it may be more beneficial to determine the SUV$_{70}$ for consistency between different approaches of ROI generation. Delineating lesions using fixed-size ROIs is a simple, relatively quick, semi-automatic approach that is correlated with less sensitivity to partial volume effects. Meanwhile, a potential problem associated with using the fixed-size method in the context of AAA is the variability in the size and shape of aneurysms across different patients, reflected in the variability in radiotracer uptake, in turn leading to differences in the proportion of an aneurysm contained within a fixed-size ROI. This may particularly cause difficulties in longitudinal studies, wherein the size and shape of aneurysms are likely to alter over time. Although studies have been conducted to determine the size of a fixed-size ROI that would result in the least deviation from the actual radioactivity concentration (Nahmias and Wahl, 2008, Weber et al., 2015), these issues are minimised when implementing manually drawn ROIs.

4.5 Anti-proliferative treatment for AAA

Currently, no targeted pharmacological treatments for AAA are used in clinical practice; EVAR and OAR remain the mainstay of current interventions. The evidence of a period of active cell proliferation in the AngII AAA model is encouraging to test the effectiveness of anti-proliferative therapies in targeting AAA, which was the original motivation for investigating cell proliferation in the present study. Imatinib, also known as Gleevec®, is a tyrosine kinase inhibitor that is suggested to play a role in the inhibition of T lymphocyte proliferation and mast cell activation, both of which contribute to AAA development (Juurikivi et al., 2005, Seggewiss et al., 2005, Shimizu et al., 2006, Sun et al., 2007). Imatinib has also been demonstrated to inhibit endothelial cell population growth, platelet-derived growth factor (PDGF) receptor activation, and the proliferation and migration of VSMCs, which are key players in AAA formation (Vrekoussis et al., 2006, Hacker et al., 2007, Ballinger et al., 2010). Based on these findings, imatinib as a treatment for AAA was tested by Vorkapic et al. using the AngII AAA murine model and human samples of AAA tissue. They demonstrated that imatinib attenuates AAA development by inhibiting the recruitment of T lymphocytes and mast cells and SMC-mediated processes (Vorkapic et al., 2016). Drawing inspiration from this study but with a focus on early changes in cell proliferation post-treatment, the present pilot study demonstrated increased abdominal aortic [$^{18}$F]FLT uptake in the AngII AAA model at 14 days, which was then shown to decrease by 89% following only 3 doses of imatinib. An important limitation in the present results was the small sample
size; the high rate of deaths immediately after radiotracer injection and unusually high bone uptake in this cohort led to the suspicion that the radiotracer stock might have been supplied from an unstable batch, resulting in unintended interactions once injected. Anti-proliferative treatments, such as imatinib, might be a promising therapeutic option for early-stage AAA, although further preclinical studies with larger numbers are needed to justify and validate this. The fundamental aim of treating AAA is to prevent aortic rupture and its corresponding high rate of mortality. Given that the only treatments currently offered (EVAR and OAR) are plagued with considerable postoperative complications, such as the need for re-repair, occurrence of endoleaks, and postoperative morbidity and mortality, there is a precedence for identifying less invasive means to treat AAA. In this context, a prevailing issue with the prospect of imatinib as a treatment option for patients with AAA is the reduced ability to detect AAA in patients during the early, proliferative growth stage of the disease. Furthermore, as a chemotherapeutic agent, imatinib is likely to confer toxic effects on non-aneurysmal proliferative cells, such as cells of the spleen and bone marrow, and also lead to adverse side effects, such as gastrointestinal disruption and hair loss. Nevertheless, the present findings provide further validation of the increased \(^{[18]F}\)FLT signal observed at the 14-day time point in this model and demonstrate that \(^{[18]F}\)FLT PET/CT can be used to detect changes in cell proliferation that are induced by short-term anti-proliferative treatment.

Regardless of whether imatinib becomes a treatment option for AAA, it may be beneficial to adopt oncologic clinical approaches, such as using \(^{[18]F}\)FLT PET/CT as a tool to stratify patients in the pursuit of personalised medicine and to evaluate the response to a less toxic equivalent of chemotherapy.

It is important to recognise that proliferation in AAA may not be solely detrimental, and thus, the use of anti-proliferative agents may not be a ‘one-off’ treatment option. As outlined in sections 1.3.5 and 4.3, accumulating evidence suggests that there is an active proliferative phase early in the AAA disease course. Halting this activity via anti-proliferative agents may indeed slow down or delay AAA progression. However, when also considering other factors contributing to aortic wall stress, if the aortic wall integrity has already been compromised, then it may be beneficial to leave proliferation unhindered in an attempt to stabilise the aortic wall and prevent further degradation. Thus, anti-proliferative therapy could ideally be determined on a patient-to-patient basis, according to AAA progression and aortic wall stability in those who show abdominal aortic \(^{[18]F}\)FLT uptake on PET imaging. With insignificant aortic expansion and low wall stress, anti-
proliferative therapy may not be as effective because the proliferative activity may protect the wall from further degradation. However, in cases of rapid aortic expansion and increased wall stress, patients may benefit from anti-proliferative therapy to inhibit further wall weakening and degradation leading to rupture. In this way, with an imminent need for pharmacological treatment options for AAA, patient-specific therapy decisions incorporating different biomarkers hold significant potential.

4.6 Future directions

AAA remains a significant cause of mortality in adults as a result of aortic rupture. USS provides information regarding anatomical changes in AAA size and shape; however, there remain gaps in our knowledge of the molecular changes that precede physical manifestation of the disease and in our ability to visualise these pathological mechanisms at the molecular level. A better understanding of these functional changes in early-stage AAA may help stratify patients based on rupture risk early in the disease course and guide appropriate treatment selection. Despite the large number of AAA studies conducted, our knowledge of AAA pathobiology is incomplete, and there remains a lack of concrete data to translate preclinical findings to clinical practice. Although several risk factors and potential contributors to the disease process have been identified, these are not yet sufficiently predictive when applied to individual patients.

As we move forward, a shift away from aortic diameter as the main indicator of AAA repair is perhaps required. There is a need to investigate the feasibility of alternative markers of AAA rupture that may be more reliable, such as circulating biomarkers, MRI findings, and haemodynamic parameters. In this context, the pathophysiology of AAA in humans remains partially clear and represents an ever-growing area in cardiovascular research. This is largely due to the fact that studies of AAA in humans are predominantly limited to using late-stage aortic tissue obtained during surgery, owing to the asymptomatic presentation of early-stage AAA. A wide array of imaging techniques have been and are currently being explored for applications in AAA diagnosis and management. Anatomical modalities have reigned over the past few decades in both preclinical and clinical imaging of AAA. Meanwhile, functional imaging is gaining increasing importance for AAA, and several new functional imaging studies of AAA are anticipated in forthcoming years. Non-invasive molecular imaging of AAA is significantly promising for clinical translation, particularly in facilitating patient risk stratification. It is important to note that anatomical and molecular imaging
techniques are not competitive; rather, they are complementary in their usefulness for AAA management. Thus, non-invasive imaging modalities such as PET offer a means to detect and evaluate components of potentially contributing pathways in patients to better understand the development of AAA beyond morphological features. The overall findings of the present study open up several avenues for further research to elucidate the pathobiology of AAA beyond the scope of this thesis.

4.6.1 Considerations of the AngII AAA mouse model

Aortic tissues cannot be obtained from patients at early stages of the AAA disease course; this creates an evident need for appropriate laboratory models to study early-stage mechanisms. The evidence of heightened cell proliferation in the 14-day AngII AAA mouse model is encouraging to investigate changes in the proliferative activity at more time points in the disease course using $[^{18}\text{F}]\text{FLT}$ PET/CT. With less stringent economic restrictions, it would be ideal to image mice of this model at serial time points, i.e. weekly, to track even slight fluctuations in cell proliferation until the aneurysm ruptures. Moreover, the cellular composition of early-stage AAA remains largely unclear owing to the lack of access to early-stage human tissue. Histological staining of early-stage AAA tissues from mice could be performed using markers such as alpha-smooth muscle actin to identify SMCs, FLK1 to identify endothelial cells, collagen, and macrophage markers (e.g., CD45, CD68). The use of mice also facilitates genetic studies, wherein the effects of knocking out or overexpressing a gene can be investigated; some candidate genes to study in the setting of early-stage AAA to further elucidate the cellular composition include those specific for SMCs, such as SM22α; MMPs; collagen, such as COL3A1; or elastin, such as ELN (Lin et al., 2018, Quintana and Taylor, 2019).

Although it is clear that preclinical models are invaluable in the progression to clinical studies, whether animal models of AAA accurately represent human AAA remains a point of discussion. With the widespread use of rodent models of AAA, several pathophysiological characteristics of human AAA are mimicked, such as ECM degradation, SMC loss, inflammation, calcification, and neovascularisation (Daugherty and Cassis, 2004). However, when translating the findings of this study to clinical practice, it is important to consider that a notable difference between AAA in humans and AAA in the AngII mouse model is the localisation of the disease in the infrarenal and suprarenal regions of the abdominal aorta, respectively. The suprarenal localisation of AAA is consistently observed in the AngII AAA
murine model (Cao et al., 2010, Rateri et al., 2011, Lu et al., 2015). Meanwhile, regional differences in collagen and elastin content are proposed to contribute to the infra-renal localisation of AAA in humans (Halloran et al., 1995). These points highlight the importance of considering the distinguishing characteristics of different models and species when tailoring AAA investigations to a specific aim. However, because of the lack of data regarding AAA formation in humans due to research being limited to late-stage tissues from patients with confirmed AAA, it remains difficult to assess the comparability of molecular mechanisms between animal models and humans.

Whilst the findings of this study are encouraging, it is important to acknowledge that cell proliferation in patients with AAA likely occurs at a much slower rate than that observed in mouse models of 14-day AngII-induced AAA. This is based on the reported mean rate of AAA growth amongst patients included in the UK Small Aneurysm Trial: 2.6 mm/year (Brady et al., 2004). Another study of AAA growth rates in the Chinese population found that the mean growth rates are 2.8 mm/year in patients with small AAA (aortic diameters 30–49 mm) and 7.5 mm/year in patients with large AAA (aortic diameters ≥50 mm) (Huang et al., 2019). The typical spatial resolutions of clinical PET scanners (4 mm) may be insufficient to detect slight changes in [18F]FLT uptake based on slow changes in cell proliferation, like that reported in small AAA (Khalil et al., 2011). Therefore, [18F]FLT PET/CT shows greater potential in stratifying patients, guiding therapeutic trials, and aiding the prediction of AAA-targeted drug response after AAA has been diagnosed, rather than in initial disease detection.

4.6.2 Future studies of calcification

Identifying key pathways in the pathogenesis of AAA warrants the application of additional radiotracers to visualise and quantify other contributing factors. The role of calcification in AAA formation is one active area of research. Calcification, which entails calcium phosphate crystal deposition in the medial vessel wall layer, contributes to mechanical stiffening of the vessel wall and decreased vascular compliance (Wu et al., 2013). Through these structural changes, calcification has been demonstrated to contribute to AAA rupture risk and is increased in patients who exhibit AAA symptoms (Buijs et al., 2013, O'Leary et al., 2015, Chowdhury et al., 2018, Forsythe et al., 2018). Furthermore, apoptosis—as reported in late-stage AAA—involves the production of apoptotic bodies, which play a role in promoting calcification in AAA (Proudfoot et al., 2000).
CT is a useful tool to generate images with detailed anatomical information, particularly of hard tissues such as bone. Thus, calcification is an ideal target for CT imaging of AAA, as evidenced by the use of calcium scoring in the clinic to indicate the narrowing or blockage of coronary arteries due to plaque build-up, which in turn helps assess the severity of coronary artery disease. A retrospective study of patients with TAA or AAA who underwent CT-based calcium scoring revealed that macrocalcification is correlated with greater cardiac and all-cause mortality (Chowdhury et al., 2018). Furthermore, vascular calcification is enhanced in symptomatic AAA and augments AAA rupture risk (Buijs et al., 2013, O'Leary et al., 2015). The most convincing evidence of calcification in AAA is reflected in the SoFIA³ trial, the findings of which suggest that $[^{18}\text{F}]{\text{NaF}}$ PET/CT is an effective tool to identify AAA disease activity in patients, based on the localisation of the radiotracer in regions of microcalcification, which in turn, is a susceptibility marker of aneurysm expansion and rupture early after AAA formation (Forsythe et al., 2018). These findings collectively suggest that calcification can be used as a prognostic marker in patients with AAA to help stratify high-risk patients for risk-reducing interventions. It is then also worth considering the addition of a vascular contrast-enhanced CT or CTA scan, which would yield more detailed anatomical information than plain CT alone, following $[^{18}\text{F}]\text{FLT}$ PET/CT. This combination would offer insight into two key molecular processes that contribute to the AAA disease course (i.e. cell proliferation and calcification), thus conferring a more comprehensive assessment of patient prognosis. Additionally, it would be beneficial to perform $[^{18}\text{F}]{\text{NaF}}$ PET/CT of mice with AAA to acquire data that could serve as the preclinical version of the SoFIA³ trial. Preclinical data of $[^{18}\text{F}]{\text{NaF}}$ uptake in AAA have not yet been published, but a study using animal models would be useful to further elucidate the timing of microcalcification in AAA formation and progression.

### 4.6.3 Prospective studies based on genetic pathways

$[^{18}\text{F}]{\text{FDG}}$, $[^{18}\text{F}]{\text{NaF}}$, and $[^{18}\text{F}]{\text{FLT}}$ are some commonly used radiotracers in the clinic for various diseases, largely due to their ease of access and availability. In addition to the array of radiotracers used to investigate AAA as outlined in Table 1.2, there is scope for the development and testing of radiotracers related to pathways of genes identified to play a role in AAA formation and progression. For example, the Arg95 genetic variant of factor XIII-B is associated with an increased risk of AAA (Macrae et al., 2014), suggesting the role of platelets and coagulation factors in AAA
pathogenesis, which represent an important field of study. Radiotracers that are selective for coagulation factors may prove to be useful in elucidating AAA pathogenesis, much like the radiotracer $^{18}$FENC2015, which is specific for factor XIIa and has been used to investigate arterial thrombi in rats (Andrews et al., 2019). The rs6511720 single-nucleotide polymorphism in the low-density lipoprotein receptor gene has also been shown to be a risk factor of AAA, implicating the correlation between AAA and cholesterol homeostasis (Bradley et al., 2013); thus, cholesterol-targeted radiotracers may be informative in the context of AAA. Furthermore, a meta-genome-wide association study by Jones et al. confirmed the roles of several risk loci in the following genes that seem to exhibit specificity for AAA than for other cardiovascular diseases: BCAR3, SORT1, NOTCH2, TDRD10, UBE2W, CDKN2B-AS1/ANRIL, LRP1, NAB2, FGF9, and PLTP (Jones et al., 2017). The specificities of these genes are promising for the development of radiotracers that specially target AAA-associated gene transcription. For example, BCAR3 is known to play a role in the development of anti-oestrogen resistance in oestrogen receptor (ER)-positive breast cancer (Wallez et al., 2014). The radiotracer 16a-$^{18}$F-fluoro-17b-estradiol ($^{18}$FES) has recently been demonstrated to be an effective PET radiotracer to detect ER-positive breast cancer (Liu et al., 2019); thus, based on the genetic link, $^{18}$FES may also have PET imaging applications in patients with AAA harbouring upregulated BCAR3 expression. Moreover, determining the correlation between the expression of these susceptibility genes and $^{18}$FFLT uptake in AAA may be informative. For example, CDKN2B-AS1/ANRIL is a tumour suppressor that is also consistently associated with cardiovascular diseases and has been shown to influence cell proliferation, senescence, and apoptosis via epigenetic mechanisms (Congrains et al., 2013). A patient that exhibits both upregulated CDKN2B-AS1/ANRIL expression and increased $^{18}$FFLT uptake may have a high-risk AAA that would benefit from immediate therapeutic intervention as opposed to a patient with only one of these risk factors.

4.6.4 Potential studies related to angiogenic and immune mechanisms in AAA

Angiogenesis is another mechanistic area that provides a basis for PET imaging targets. AAA is associated with a marked angiogenic response and enhanced medial neovascularisation (Thompson et al., 1996, Choke et al., 2006). Various pro-angiogenic factors have been implicated in the formation and progression of AAA. In particular, vascular endothelial growth factor
VEGF has been shown to promote AngII-induced AAA formation in ApoE<sup>−/−</sup> mice (Kobayashi et al., 2002). A VEGF-specific PET radiotracer, <sup>89</sup>Zr-ranibizumab, has been investigated in human xenograft tumour models and shown to correlate with VEGF status and angiogenesis (Nagengast et al., 2011). In addition, <sup>64</sup>Cu-DOTA-VEGF<sub>121</sub> has been demonstrated to reveal VEGF receptor levels in small-animal tumour models (Chen et al., 2009). Another pro-angiogenic factor that has been reported to contribute to AAA is PDGF, the receptors of which are strongly expressed in VSMCs in AAA, and reducing PDGF receptor activation via imatinib has been shown to attenuate AAA formation (Kanazawa et al., 2005, Vorkapic et al., 2016). Meanwhile, the tracer <sup>111</sup>In-DOTA-Z09591 has been demonstrated to accumulate in regions with increased levels of PDGF receptor expression (Tolmachev et al., 2014). Radiotracers specific for the microRNA-195 family may also be useful to study AAA development, as microRNA-195, which modulates angiogenesis, has been demonstrated to regulate the aortic ECM in murine AAA (Zampetaki et al., 2014). Exploring the usefulness of angiogenesis-specific tracers that have been validated in other disease models in visualising sites of angiogenesis in AAA models may be interesting avenues in preclinical research.

Immune cells, such as lymphocytes, cytokines, and antigen-presenting cells are observed in AAA (Kuivaniemi et al., 2008); however, no dedicated PET studies have been conducted previously to specifically investigate these cell types in AAA. The spleen plays a key role in immunologic functions, being a site of immune cell subsets (Lewis et al., 2019). Thus, the variations in splenic uptake of [<sup>18</sup>F]FLT demonstrated in this study may correspond to the distinct varying roles of immune cells at different stages of the AAA disease course. The radiotracer <sup>64</sup>Cu-DOTA-ipilimumab has been investigated in models of non-small cell lung cancer (Ehlerding et al., 2017). This tracer targets cytotoxic T lymphocyte-associated protein 4, the levels of which are reported to be increased in AAA (Sakthivel et al., 2007), thus making it a potential biomarker for AAA. Additionally, <sup>68</sup>Ga-pentixafor has been studied in atherosclerotic plaques to image the cytokine receptor CXCR4, which may also be informative in AAA (Hyafil et al., 2017). Immune mechanisms thus remain an active area of AAA research.

### 4.6.5 Clinical translation

It is well established that PET can be used to evaluate treatment response, in addition to its uses in disease management. Its ability to provide detailed longitudinal information at the molecular level makes it ideal to stratify
patients according to disease risk. In the context of AAA, $[^{18}\text{F}]\text{FLT PET}$ may be useful as a management indicator in high-risk patients who have not yet developed AAA. For example, if a patient exhibits many risk factors (such as a man aged 65 years or older with heavy smoking habits and a familial history of AAA) but does not yet physically manifest AAA, then $[^{18}\text{F}]\text{FLT PET}$ may be used as a tool to determine if there is evidence of abnormal cell proliferation based on the radiotracer’s uptake in the aortic region. Cases of positive $[^{18}\text{F}]\text{FLT}$ uptake suggest a role for prophylactic interventions, such as early placement of anti-proliferative drug-eluting aortic stents. Additionally, $[^{18}\text{F}]\text{FLT}$ may be utilised as a predictor of prognosis in patients who are already diagnosed with AAA. For example, if a patient with USS-established AAA does not reveal $[^{18}\text{F}]\text{FLT}$ uptake on PET, then he/she may be stratified to a more advanced disease stage, possibly requiring surgical intervention. Conversely, if a patient with established AAA reveals $[^{18}\text{F}]\text{FLT}$ uptake on PET, then he/she may be stratified with an earlier disease stage, requiring a less surgically intensive intervention. In this way, $[^{18}\text{F}]\text{FLT}$ PET may then play a more significant role in personalised medicine to aid disease risk stratification as opposed to population-level screening.

The clinical translation of radionuclide-based tracers is crucial to maximally benefit patients with AAA. PET imaging is immensely useful in comparison to USS and optical imaging methods; however, much of the preclinical PET research on AAA never proceeds to the clinic. This is largely due to hurdles such as the overwhelming cost of introduction in the clinic, a lack of understanding between preclinical scientists and clinicians, and long-standing regulations. Cooperative partnerships amongst researchers, clinicians, and pharmaceutical chemists are imperative to identify and verify novel imaging markers and more clinically applicable preclinical models, as well as to generate and test tracers, followed by smooth transition for application to patient management. Ultimately, the optimal imaging marker will reveal key aspects of AAA formation in humans, be used for patient risk stratification, and help assess treatment efficacy in novel clinical trials.

4.7 Conclusion

Overall, in this thesis, $[^{18}\text{F}]\text{FLT PET/CT}$ was demonstrated to be a feasible modality to visualise and quantify cell proliferation in a classical preclinical model of AAA, the AngII AAA murine model. This method shows great potential as a clinical risk stratification biomarker and in monitoring pharmacological treatment response in patients with AAA. Future studies of PET image analysis would benefit all clinical fields by establishing
quantitative metrics that could yield reliable and consistent results across varying ROI definitions. Prospective studies of PET in the context of AAA would benefit from focussing on the development of PET radiotracers that target alternative molecular pathways involved in the development of AAA disease before its physical manifestation. Until then, 65 years on, two issues remain unresolved: Einstein’s unified field theory and how best to manage early-stage AAA.
References


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Appendix A

Copies of all conference posters associated with this thesis are provided here.

Figure A.1 Poster for the European Molecular Imaging Meeting 2018
Effect of different ROI definitions on the quantification of SUV

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Background

• The standardized uptake value (SUV) is commonly used in clinical and preclinical PET applications to analyze tracer accumulation within a specified region of interest (ROI)1,2.

• The most common ways of calculating SUV are based on the average (SUV$_{\text{mean}}$) or maximum (SUV$_{\text{max}}$) of all voxels within an ROI.

• It has been suggested that applying a threshold to estimate the average SUV within a group of voxels can potentially reduce limitations in the SUV$_{\text{max}}$ or SUV$_{\text{mean}}$.

Aims

1. To identify a reproducible threshold value using two different segmentation methods.

2. To assess the applicability of using these methods in preclinical PET-CT.

Methods

1. Image acquisition and reconstruction

Fourteen mice (10 with abdominal aortic aneurysms (AAA), 4 controls) underwent 90-min dynamic $^{[18]}$F-FLT PET-CT acquisitions using the Sedecal Super Argus PET-CT scanner. Images were reconstructed with the 3D OSEM algorithm (2 iterations, 16 subsets).

2. Image analysis using Amide software

• Manual ROIs were drawn using the 3D isocontour tool (Figure 1 A and B).

• Fixed-size ROIs were 4-mm spheres (Figure 1 C and D).

3. Statistical analysis

Paired sample t-tests were conducted to determine whether a difference existed between pairs of SUV measurements.

Results

• SUV$_{60}$, SUV$_{90}$, and SUV$_{90}$ for manually drawn vs. fixed-size ROIs showed significant differences (all $P < 0.05$), while SUV$_{70}$ showed no statistically significant difference between the segmentation techniques ($P = 0.72$).

• The average volumes of the fixed-size ROIs were relatively larger than those of the manual ROIs, and the volumes of both ROIs decreased with the application of higher threshold values.

• The choice of segmentation tool for ROI definitions in PET image analysis has a direct impact on SUV measurements.

• The implementation of fixed-size ROIs with a 70% threshold appears to provide the most stable SUV measurements in the analysis of aortic lesions in preclinical PET-CT images of mice.

References


Figure A.2 Poster for the European Molecular Imaging Meeting 2018
Aim:
- Abdominal aortic aneurysm (AAA) disease is characterised by a localised dilatation of the abdominal aorta that affects approx. 4% of men aged 65–74 years in the UK.
- AAA risk factors include age, male gender, former/current smoking habits, positive familial history, and high diastolic blood pressure.
- AAA is currently managed with routine ultrasound screening, which does not elucidate the molecular mechanisms that precede physical disease manifestation.
- The aim of this study was to investigate the feasibility of using positron emission tomography/computed tomography (PET/CT) with $[^{18}F]$Fluorothymidine ($[^{18}F]$FLT) to visualise AAA in murine models.

Materials & Methods:
- Fourteen-week-old apolipoprotein E-knockout (ApoE$^{-/-}$) mice were administered either saline or angiotensin II (AngII) infusions via subcutaneously implanted osmotic mini-pumps for 28 days.
- Ninety-minute dynamic PET/CT under recovery isoflurane was performed on days 14 and 28.
- Radiotracers were administered via intravenous tail vein injections
  - $[^{18}F]$Fluorodeoxyglucose ($[^{18}F]$FDG): 9.76±2.10 MBq
  - $[^{18}F]$FLT: 7.63±3.02 MBq (day 14), 9.61±1.34 MBq (day 28)
- Images were reconstructed using the three-dimensional (3D) ordered subsets expectation maximisation (OSEM) algorithm.
- 3D isocontour regions of interest (ROIs) were manually drawn in all abdominal aortic regions.
- Maximum standardised uptake values (SUVs) were calculated over all voxels for each ROI.

Results:
- $[^{18}F]$FDG and $[^{18}F]$FLT uptake in the inferior vena cava were observed in the first 15 seconds of each scan, indicating successful intravenous tracer delivery.
- Expected tracer biodistribution was observed in control and AAA models:
  - $[^{18}F]$FDG: myocardium (Figure 1)
  - $[^{18}F]$FLT: spleen (Figure 2)
- Significantly greater $[^{18}F]$FLT uptake was observed in aneurysmal aorta than in control aorta.
  - SUV % differences: 4466.80 (day 14; p=0.001) (Figure 3), 430.35 (day 26) (Figure 4)
- $[^{18}F]$FLT uptake was more consistent than $[^{18}F]$FDG (Figure 5)
  - Standard deviations: $[^{18}F]$FDG, 0.14, $[^{18}F]$FLT, 0.04 (day 14) and 0.02 (day 28)
- Aneurysmal aorta showed greater $[^{18}F]$FLT positivity than control aorta (Figure 6)
  - % difference: 4669.19, p=0.001

Conclusions:
- $[^{18}F]$FLT uptake is consistently enhanced in AAA models than in control models in comparison to $[^{18}F]$FDG uptake.
- To our best knowledge, this is the first study to demonstrate the potential utility of $[^{18}F]$FLT PET/CT in evaluating AAA-associated cell proliferation.
- Further validation work to elucidate the time course and molecular mechanisms of AAA is ongoing.

Figure A.3 ePoster contents for the Annual Congress of the European Association of Nuclear Medicine 2018
The results of non-parametric statistical tests are presented here. The Kruskal–Wallis test was used for the data presented in Figures B.1, B.5, B.6, B.7, B.8, and B.9. The Mann–Whitney $U$ test was used for the data presented in Figures B.2, B.3, and B.4. All significant results according to parametric tests remain statistically significant following non-parametric analysis.

Figure B.1 PPE and AngII AAA tissue reveal the greatest proportions of Ki67-positive nuclei. Proportion of Ki67-positive nuclei in saline controls ($n=3$) and in the AngII ($n=6$), PPE ($n=7$), and CaCl$_2$ ($n=4$) AAA models. * $p<0.05$, ns: not significant on Kruskal–Wallis test.
Figure B.2 Aortic volumes and diameters are larger in the AngII AAA model than in saline controls. Aortic volumes and diameters in 14-day AngII AAA (n=9) vs. saline control aortae (n=4). * p<0.05, ** p<0.01 on Mann–Whitney U test.
Figure B.3 \([^{18}F]\)FLT uptake in the abdominal aorta was greater in the 14-day AngII AAA model than in saline controls. Absolute values of \(SUV_{\text{max}}\) and \(SUV_{\text{mean}}\) representing 14-day \([^{18}F]\)FLT uptake in abdominal aortic ROIs 80–90 min post-radiotracer injection in 14-day saline controls (\(n=5\)) and the 14-day AngII AAA model (\(n=5\)). ** \(p<0.01\) on Mann–Whitney \(U\) test.
Figure B.4 *Ex vivo* $[^{18}F]$FLT uptake in the abdominal aorta was greater in the 14-day AngII AAA model than in 14-day saline controls. Decay-corrected *ex vivo* $[^{18}F]$FLT counts per mass units. Uptake in the spleen and abdominal aorta normalised to uptake in the heart in the 14-day AngII AAA model ($n=9$) and 14-day saline controls ($n=4$). * $p<0.05$ on Mann–Whitney $U$ test.
Figure B.5 TK-1 expression was the greatest in 14-day AngII AAA. Quantitative analysis wherein TK-1 band intensity was normalised to β-actin band intensity. * $p<0.05$, ns: not significant on Kruskal–Wallis test.
Figure B.6 ENT-1 expression was the greatest in 14-day AngII AAA. Quantitative analysis wherein ENT-1 band intensity was normalised to β-actin band intensity. * $p<0.05$, ns: not significant on Kruskal–Wallis test.
Figure B.7 ENT-2 expression was the greatest in 14-day AngII AAA. Quantitative analysis wherein ENT-2 band intensity was normalised to β-actin band intensity. * $p<0.05$, ns: not significant on Kruskal–Wallis test.
Figure B.8 CNT-1 expression was the greatest in 14-day AngII AAA. Quantitative analysis wherein CNT-1 band intensity was normalised to β-actin band intensity. * $p<0.05$, ns: not significant on Kruskal–Wallis test.
Figure B.9 CNT-3 expression was the greatest in 14-day AngII AAA.
Quantitative analysis wherein CNT-3 band intensity was normalised to β-actin band intensity. * $p<0.05$, ns: not significant on Kruskal–Wallis test.