

Computational Methods for the Analysis of Ascending Aortic Aneurysms

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Overview



Introduction

Aortic Aneurysms





Aortic Aneurysms







5-10 cases per 100,000 person/year 22% of patients with ruptured aneurysm die before reaching a hospital

Linked to age, sex, hypertension, genetic conditions

Clinical problem

Current practice: Surgery is determined by **diameter**.



Problem:

- It's too generic
- Unpredicted aneurysm rupture
- Unnecessary intervention

Post-operative complications:

- ► Hemorrhage
- Infection
- Cardiac fatigue.



Clinical need to gain insight of the patient's HEMODYNAMICS & WALL DETERIORATION for accurate personalized treatment

Surgical decision



Surgical decision



Aortic Aneurysm

Patient specific:

- ► Aorta Shape
- ► Valve morphology
- Valve pathology
- ► Hemodynamic BCs
- Aortic wall



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Aortic Aneurysm

Patient specific:

- Aorta Shape
- Valve morphology
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Healthy



Aneurysm



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Computational tools for personalized treatment



Final Conclusions

Computational Methods

Section I

Computational methods for accurate turbulence and viscosity modelling

Introduction

No standardized methodology exists for the computation of cardiovascular flows



Introduction Computational Methods CFD Biomarkers Patient Specific FSI Final Conclu

Introduction

Viscosity

- Blood is a mixture of plasma and red blood cells with a shear-thinning behaviour.
- Eddy development and near-wall flow is influence by this property [1].
- It is argued that, under the high shear-rates present in the aorta, the variations in viscosity are negligible and constant viscosity can be assumed.



Turbulence

- Turbulence causes bursts of shear stress, damaging endothelial cells [2].
- Turbulence generates additional stresses on aneurysm wall leading to wall vibration and increases the rate of wall dilation [2].
- Pulsatile flow with a low averaged Reynolds number, averaged Reynolds suggests laminar flow.
- Flow deceleration during diastole favours turbulence generation.

LAMINAR FLOW	TURBULENT FLOW
	3,26,55

[2] Tan et al. "Analysis of flow patterns in a patient-specific thoracic aortic aneurysm model," *Computers and Structures* 87 (2009)

[1] Wyk et al., "Non-Newtonian perspectives on pulsatile blood-analog
flows in a 180° curved artery model", Physics of Fluids 27 (2015)

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Previous work

Viscosity

Newtonian model causes:

- Underestimation of WSS and hemolysis
- Growth and decay of eddies
- Premature turbulent transition

Turbulence

Laminar model causes:

- WSS underestimated between 0-6% (depending on author)
- Platelet activation and hemolysis
- Underestimated TKE



Karimi et al. Journal of Non-Newtonian Fluid Mechanics 207 (2014)

No publication exists on the combined effect of viscosity and turbulence models



Bozzi et al. *Journal of Biomechanics 128* (2021)

Introduction

Objective



Understand the interaction between models and the importance of the model choices

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Scope

Viscosity

- Newtonian: $\mu(\dot{\gamma}) = \mu_{\infty}$
- Non-Newtonian: Carreau viscosity (CV)

$$\mu(\dot{\gamma}) = \mu_{\infty} + (\mu_0 - \mu_{\infty}) \left[1 + (\lambda \dot{\gamma})^2 \right]^{\frac{n-1}{2}}$$

 $\mu_{\infty} = 3.5 \text{ mPa} \cdot \text{s}$ $\mu_0 = 56 \text{ mPa} \cdot \text{s}$ $\lambda = 3.313 \text{ s}$ n = 0.3568

Turbulence

- No model: Laminar flow model (LFM)
- Turbulent: LES

$$\begin{aligned} \frac{\partial \bar{u}_i}{\partial x_i} &= 0 ,\\ \frac{\partial \bar{u}_i}{\partial t} + \bar{u}_j \frac{\partial \bar{u}_i}{\partial x_j} &= -\frac{1}{\rho} \frac{\partial \bar{p}}{\partial x_i} + \nu \frac{\partial}{\partial x_j} \left(\frac{\partial \bar{u}_i}{\partial x_j} \right) - \frac{\partial \tau_{ij}}{\partial x_j} \\ \tau_{ij} - \frac{1}{3} \tau_{kk} \delta_{ij} &= -2\nu_{sgs} \bar{S}_{ij} \\ \nu_{sgs} &= (C_S \Delta)^2 |\bar{S}| \end{aligned}$$
Dynamic Smagorinsky-Lilly (DSL) subgrid-scale turbulence model

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Model setup







Windkessel outlets





Stenotic



10 days per scenario

20 heart beats32 cores



Introduction

Patient Specific FSI

Vortex structure is influenced by the turbulence model.

- Non-Newtonian viscosity has greater impact (2.9-5.0%) on wall shear stress than Large Eddy Simulation turbulence modelling (0.1-1.4%).
- Wall shear stress is underestimated when considering Newtonian viscosity by 2.9-5.0%.
- The contribution of non-Newtonian viscosity is amplified when combined with a LES model.
- Cycle-to-cycle variability can impact the results as much as the numerical model if insufficient cycles are performed.



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CFD Biomarkers

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Martinez et al., "Effect of Turbulence and Viscosity Models on Wall Shear Stress Derived Biomarkers for Aorta Simulations," *Computers in Biology* and Medicine, 167 (2023)

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Future works

Additional viscous models: Power law, Casson, Cross

Realistic aortic jet shapes

FSI effects

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Section II

CFD biomarkers for aneurysm growth prediction

Computational tools for personalized treatment



Introduction

- Hemodynamics conditions influence the biomechanical processes in the arterial wall:
 - ► Endothelial damage.
 - Elastin and smooth muscle cell damage.
 - Extra cellular matrix dysregulation.
- A debate exists on whether genetic conditions or hemodynamics are responsible for the development of aneurysms.

Elevated WSS Fbr: for tors cellagen Matrix

Guzzardi et al, "Valve-Related Hemodynamics Mediate Human Bicuspid Aortopathy: Insights From Wall Shear Stress Mapping," J. Am Coll Cardiol. 66 (2015)

IN THIS SECTION: The correlation between fluid biomarkers and aneurysm growth will be assessed.

CFD Biomarkers

Patient Specific FSI

Dataset



- Echocardiography: 20 patients
- MRI flow: 5 patients
- No data: 8 patients



Introduction

CFD Biomarkers

Patient Specific FSI

Growth analysis



Growth rate = Diameter change per year [mm/year]

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Fluid Biomarkers





FLOW ANALYSIS

Introduction

CFD Biomarkers

Patient Specific FSI

Fluid Biomarkers Wall Shear

Time-average WSS

Oscilating Shear Index









$$TAWSS = \frac{1}{T} \int_0^T |\mathbf{WSS}(t)| dt \qquad OSI = 0.5 \left(1 - \frac{\left| \int_0^T \mathbf{WSS}(t) dt \right|}{\int_0^T |\mathbf{WSS}(t)| dt} \right) \qquad SA = \frac{2}{\pi} \arctan\left(\frac{WSS_{Axial}}{WSS_{Circ}} \right)$$

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Fluid Biomarkers Wall Shear



Fluid Biomarkers Flow







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Fluid Biomarkers Flow

Flow Asymmetry:

Offset of flux centroid. Normalized by mean radius.

$$FA = \frac{\|P_{Center} - P_{FMC}\|}{R_{mean}}$$

Angle: Between flow and plane

Flow Asymmetry - Bounded:

Offset of bounded fast-moving region centroid Normalized by mean radius.

$$FA_{20\%} = \frac{\left\| P_{Center} - P_{FMC_{20\%}} \right\|}{R_{mean}}$$

Flow Dispersion:

$$FD = \frac{A_{20\%}}{A_{Total}}$$



- 0.8 - 0.7 - 0.6

0.5 0.4 0.3

0.2

0.1

-0.1



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Results: Growth



on PC = 0.25 for TAV and on PC = 0.40 for BAV.

Initial diameter does not correlate with *GR* (R= 0.04)

Results: Growth



(R= 0.04)

Results: Growth



The maximum diameter was located, on average, on PC = 0.25 for TAV and on PC = 0.40 for BAV. Initial diameter does not correlate with *GR* (R= 0.04)

Results: Fluid biomarkers

6 days per patient







CFD Biomarkers

Patient Specific FSI

Results: Correlations

PEAK SYSTOLE SHEAR ANGLE

- External wall of BAV patients.
- Weak correlation with GR_D and GR_L .
- Suggest reversed and rotating flow are linked to wall degeneration.
- Agrees with previous works:
- ► FSI on Marfan syndrome patients Pons et al., Royal Society Open Science 7 (2020)
- MRI flow on BAV patients Minderhoud et al., European Heart Journal – Cardiovascular Imaging 23 (2022)

- Only 17 BAV patients \rightarrow Statistical relevance is debatable.
- Largest CFD study on aneurysm growth up to date.





$SA = \frac{2}{2}$ spatan	(WSS_{Axial})
$SA = -\arctan \pi$	$\left(\overline{\mathrm{WSS}_{\mathrm{Circ}}} \right)$

		TAV				B	AV		
Biomarker	Measure	GR_D		GR_L		GR_D		GR_L	
		R	р	R	р	R	р	R	р
TAWSS	Max	-0.223	0.407	-0.274	0.304	-0.160	0.541	-0.256	0.321
	Mean	-0.054	0.843	-0.190	0.480	-0.128	0.623	-0.209	0.421
PSWSS	Max	-0.132	0.626	-0.162	0.549	-0.053	0.841	-0.148	0.570
	Mean	-0.178	0.510	-0.282	0.291	-0.095	0.717	-0.213	0.411
OSI	Mean	-0.030	0.911	0.108	0.692	-0.089	0.734	0.002	0.995
SA	TA-Mean	0.061	0.823	-0.048	0.860	0.255	0.324	0.274	0.287
	PS-Mean	0.004	0.987	-0.048	0.859	-0.482	0.050	-0.481	0.051
RFR	TA	0.034	0.899	0.073	0.787	-0.266	0.303	-0.306	0.232
	\mathbf{PS}	0.048	0.859	0.072	0.792	0.243	0.347	0.275	0.286

Future Works

Larger time window

Reduce the error in the growth rate measurements. Follow the evolution during the initial phase.

MRI 4D calibrated aortic jet

The spatio-temporal velocity profile of the aortic jet will severely determine the flow structure throughout the cardiac cycle, hence, the biomarkers.

Topological WSS skeleton analysis

Evaluated the topological shear variation index (TSVI) and fixed-point relative residence time ($RT\nabla$).

CFD Biomarkers







Section III

Patient-specific FSI models

Computational tools for personalized treatment



Computational tools for personalized treatment

Personalized hemodynamic conditions

- Aortic jet derived from MRI 4D flow
- Windkessel outlets calibrated with patient's data

Fluid biomarkers

WSS, Flow



Personalized aorta wall

- Thickness
- Elasticity

Structural biomarkers

Stress



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Methods: Aortic jet

MRI 4D Flow



Velocity extraction on aortic valve plane

Transfer onto the fluid model

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Methods: Aortic jet

Resampling and filtering:

- Finer grid (x3) using modified Akima interpolation: reduced undulations and over-flattening.
- Gaussian 2-D filter was applied to smooth each of the three velocity components. Smoothing kernel with standard deviation 2.5.



Methods: Aortic jet



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Patient Specific FSI

Methods: Windkessel

Measure	Value	Unit
Q_{max}	456.2	ml/s
Q_{min}	15.3	ml/s
Q_{mean}	6.89	l/min
Q_{DA}	3.48	l/min
P_{Sys}	60	mmHg
P_{Dias}	0.0	mmHg
Δt	0.1	s
A_{BT}	185.2	mm^2
A_{LCC}	20.4	mm^2
A_{LS}	67.3	mm^2

$R_T = \frac{P_{mean}}{Q_{mean}},$	
$R_i = R_T / f_i ,$	
$C_T = \frac{Q_{max} - Q_{min}}{P_{Sys} - P_{Dias}} \Delta t ,$	
$C_T^* = C_T - C_{As}^{3D},$	
$C_i = f_i C_T^* \frac{R_i}{R_{d_i}} = C_T^* \frac{R_T}{R_{d_i}}.$	

RESULTS		
Component	Value	
R _{p_{BT}}	3.858×10^{6}	
$R_{d_{BT}}$	6.504×10^7	
C_{BT}	1.587×10^{-9}	
$R_{p_{LCC}}$	$3.569{\times}10^7$	
$R_{d_{LCC}}$	6.016×10^{8}	
C_{LCC}	1.715×10^{-10}	
$R_{p_{LS}}$	1.065×10^{7}	
$R_{d_{LS}}$	1.796×10^{8}	
C_{LS}	5.746×10^{-10}	
$R_{p_{DA}}$	$2.575{\times}10^6$	
$R_{d_{DA}}$	4.340×10^{7}	
C^*_{DA}	9.407×10^{-10}	

Patient Specific FSI

Methods: Aortic wall - Clinical data

4 sections: Anterior, lateral, posterior and medial



Equi-biaxial tensile test performed in the University Hospital of Dijon.

S. Lin, "Biomechanics of human ascending aorta and aneurysm rupture risk assessment", PhD Thesis, 2021.

CFD Biomarkers

Patient Specific FSI

0.4

0.45

Methods: Aortic wall - Hyperelastic material

Ascending aorta: Third order Yeoh material model.

$$W = \sum_{i=1}^{3} C_{i0} (\bar{I}_1 - 3)^i.$$

The model coefficients for each quadrant were obtained after performing a curve fitting via minimization of normalized error of the circumferential strain-stress curves.



Supra-aortic vessels and DA: Second order Yeoh material model derived from estimated pulse wave velocity (PWV).

$$PWV = \frac{\alpha}{(2 \times 10^3 r_v)^{\beta}} \qquad E_{inc} = \frac{2r_v \rho}{T_v} PWV^2$$

Methods: Aortic wall - Model definition

Spatially varying material properties

Ascending aorta: 2 node interpolation

$$T_n = T_{s1} \frac{D_{n,s1}}{D_{n,s1} + D_{n,s2}} + T_{s2} \frac{D_{n,s2}}{D_{n,s1} + D_{n,s2}}$$

Aortic arch: 3 node interpolation

$$D_{n,v}^{\text{mod}} = D_{n,v}^* \frac{D_{\text{Lim}}}{D_{\text{Lim}} - D_{n,v}^*}$$
$$T_n = T_n^* \frac{D_{n,s}^{\text{Min}}}{D_{n,s}^{\text{Min}} + D_{n,v}^{\text{mod}}} + T_v \frac{D_{n,v}^{\text{mod}}}{D_{n,s}^{\text{Min}} + D_{n,v}^{\text{mod}}}$$

DA: Constant properties



0.2

Introduction

Methods: Aortic wall - Model definition



Initial

Introduction

Methods: Aortic wall - Model definition



Smooth

Introd	luction
III CO G	action

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Patient Specific FSI

Methods: Aortic wall - Boundary conditions

- Radial displacement on outlets
- Viscoelastic support on wall

$K_{n_i} =$	$= (K_{Soft})$	$+ W_d V$	$V_i K_{\text{Spine}}$	$A_n e_{n_i}$
103	1. 0010		J spine,	in neg /

Coefficient	Value
$K_{ m Soft}$	$1.5 \times 10^4 \text{ Pa/m}$
$K_{ m Spine}$	$10^6 \ Pa/m$
W_d	0.53
W_x	0.60
W_y	0.02
W_z	0.04



Geronzi et al., "Calibration of the Mechanical Boundary Conditions for a Patient-Specific Thoracic Aorta Model Including the Heart Motion Effect," *IEEE Trans Biomed Eng.* 70-11 (2023)

Introd	luction
muou	uction

Methods: Aortic wall - Zero pressure

Augmented Sellier's Inverse Method

- Inverse problem: loads and final deformation are known, initial geometry is to be computed.
- The zero-stress state will be approximated by the zero pressure state.





Results: Stress - Growth



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Conclusions



Clinical outcomes: one patient only, it is not possible to hypothesise on the relationship between growth and stress.



A large cohort should be analysed, considering both healthy, stable and dilating aneurysms.



A model combining patient specific hemodynamics and aorta wall has been presented. Further improvements will enable an accurate estimation of risk of rupture.



- Non-Newtonian viscosity is necessary.
- LES is optional, but computational requirement is negligible.

- Aneurysm growth could be related to:
 - BAV: Peak systole shear angle.
- Larger cohort with MRI flow data is needed.



- Hemodynamic personalization requires MRI 4D flow data.
- Aorta wall definition requires spatially varying thickness and elastic properties.
- Accurate risk of rupture estimation requires high fidelity models.











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