Prediction of stent endflare, arterial stresses and flow patterns in a stenotic artery

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Abstract

Restenosis remains a significant problem in coronary intervention. While stent migrations, collapses, and positioning difficulties remain serious issues, it is the problem of restenosis that is the most common long term problem in treating atherosclerotic coronary arteries with stents. While much attention has focused on biocompatibility, thrombosis and neointimal pathology, less attention has been given to matching stents to the inflation balloon, artery and occlusion size.

Results from this study indicate a 100% increase in balloon overhang results in a 4% increase in maximum endflare and a 39% change in peak arterial stress. At the end of expansion, which is of the most clinical importance, the increase in maximum endflare is 2% and the increase in maximum arterial stress is 93% at the balloon point of contact and 45% at the point of contact with the far proximal and distal ends of the stent.

When comparing the results of calcified and cellular plaque, a maximum endflare of about 55% was observed for both the calcified and cellular plaque cases during expansion. At the end of expansion the increase in maximum endflare was 10% for the cellular plaque and 40% for the calcified plaque. The peak equivalent stress seen by the artery was about 100% larger in the cellular case than in the calcified plaque case.

Keywords: stent, vascular injury, balloon, restenosis, finite element analysis.

1 Introduction

Atherosclerotic stenosis and its ischemic complications necessitate arterial reconstruction. Current strategies to restore normal blood flow in stenotic coronary arteries include angioplasty, intracoronary stents, and coronary artery

bypass surgery. Depending on the method of treatment, the incidence of restenosis is high: up to 40% within six months after angioplasty [1], 25% after stenting [2], and 20% after bypass surgery [3]. Therefore, restenosis remains a significant problem in coronary intervention.

Advances in prosthetic science and engineering have spurred the rapid development of many new permanent implants such as arterial reinforcement grafts, venous filters, myocardial perforation-sealing clamshells, and stents that strengthen and scaffold the biliary duct, urethra, veins, and arteries. These devices are typically attached to a delivery catheter and threaded to the site of interest where they are expanded. The very nature of the remote delivery systems make the mechanical details of implantation difficult to ascertain, yet this is important to quantify since there may be a link between how the devices are emplaced and the body's acute and chronic response. Endovascular stents in particular are ideal devices to quantify these relationships because of the extreme levels of stress they impose and because of their ubiquity; more than one million are annually implanted in the U.S. alone [4].

These studies suggest an upper limit exists to the success of purely biomedical approaches for managing post-device implantation, and a return to examining the mechanical initiators of vascular injury that occur during implantation. A complete understanding of the manner that stents expand may thus lead to both a new understanding of the processes of vascular adaptation to implants and possibly to the design and development of less-injurious devices. Experimental data are indirect; stents are too thin to be fully radioopaque, and methods of bringing a camera to the stent, such as intravascular ultrasound, are blocked by the balloon that expands the stent during the critical moments of implantation. Post mortem examinations indicate that restenosis is paradoxically more severe in the parts of the artery immediately outside the stented region, and animal studies have shown an unusual pattern of endothelial cell denudation occurring at a regular pattern at the center of stent struts, a superficial injury that may be a marker for deep vascular injury [8]. These data are not explained by current finite element analyses of arterial stresses in a stent-expanded artery [9-12] because no finite element models included the expansion of a balloon catheter in the model of a plastically-deformed stent.

The inclusion of the balloon catheter in the stent expansion model is not trivial. The problem is highly non-linear and includes complex contact problems between the stent, balloon, and arterial wall. Additionally, the balloon properties change dramatically depending on whether it is fully or partially inflated.

2 Geometry

The finite element stress analysis was performed on a three-dimensional stent/balloon/plaque/artery geometry. In addition to the usual difficulties in modeling the mechanical behavior of soft tissue, the overall system response is highly nonlinear due to the large plastic/multilinear-elastic/hyperelastic deformations of the individual components. The component geometries and constitutive material models are described below.

2.1 Artery

The coronary artery modeled was 30mm in length, with an inside diameter of 2.8mm and thickness of 0.3mm. Average element size was about 0.5mm long with a thickness of 0.15mm. This configuration yielded a total of 7,680 elements.

The artery elements were defined by eight nodes capable of large deflections and hyperelasticity

2.2 Plaque

The plaque has a semi-parabolic profile and corresponds to percent blockage data presented by Lally *et al.* [12]. The plaque was 16mm in length with a maximum thickness of 0.48 mm. This configuration corresponds to a maximum percent blockage of about 60%. The characteristic shape of plaque can be seen in Figure 1.

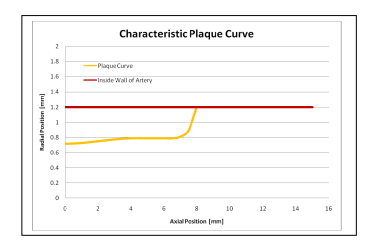


Figure 1: Characteristic plaque curve.

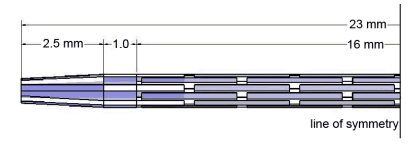


Figure 2: Balloon geometry, shown with mounted slotted tube stent.

2.3 Balloon

The balloon was modeled in its unfolded state, and already assumed to be in contact with the stent. The balloon dimensions are given at 0 Pa, before stent expansion occurs. Depending on the amount of balloon overhang, the overall length of the balloon can range from 22-23 mm. For 1mm balloon overhang, the total length of end balloon is 1mm, or 0.5 mm on each side of the inside balloon, yielding a total balloon length of 22 mm. For 2 mm balloon overhang, the total length of balloon overhang is 2 mm, or 1 mm on each side, yielding a total balloon length of 23 mm (see fig. 2).

The balloon was meshed using triangular shell elements with an average base size of 0.05 mm and an average side length of 0.05 mm. This yielded 54,456 elements for the 2 mm overhang case and 51,616 elements for the 1 mm overhang case. For finite element analysis, elements capable of modeling shell structures, large deflections and plasticity were used.

2.4 Stent

A three dimensional model of the slotted tube geometry intravascular stent was created. The stent is 16 mm in length (L), with an inside diameter (ID) of 1 mm, and a thickness (t) of 0.1 mm. The diamond-shaped stent consists of 5 slots in the longitudinal direction and 12 slots in the circumferential direction with a length of 2.88 mm. The slots were cut such that in a cross-section, the angle describing the slot was approximately 23 degrees, and the angle describing the metal between slots was 6.9 degrees (see figs. 3 and 4). These dimensions refer to the model in an unexpanded state [10, 12].

The stent was meshed using hexahedral elements. There are two elements through the thickness of the stent yielding a total of 12,036 elements. The stent was assigned an element type of solid45 for finite element analysis in ANSYS. These elements are defined by eight nodes and are capable of large deflections and plasticity.

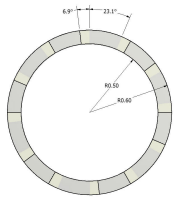


Figure 3: Medial slice of modeled slotted tube stent.

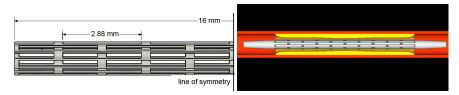


Figure 4: Side view: stent geometry.

Figure 5: Final model geometry.

A cross-section of the final model geometry with the stent/balloon/artery and plaque is shown in fig. 5.

3 Materials

3.1 Artery

The material properties of the artery are based on a previous study by Lally *et al.* [12]. This model describes the behavior of the artery using a five parameter, third-order, Mooney-Rivlin hyperelastic constitutive equation. This model has been found to be suitable for modeling an incompressible isotropic material [13].

Prendergast *et al.* developed the constants for this model by fitting the five-parameter Mooney-Rivlin expression to uniaxial and equibiaxial tension tests of human femoral arterial tissue data [14]. See Table 1.

3.2 Plaque

The material properties of the plaque are based on a previous study by Loree *et al.* [15]. Two histological classifications of plaques were modeled: cellular and calcified. The cellular and calcified specimen results were chosen to provide models of stent expansion dynamics with plaques whose stress-strain slopes differed significantly. This model describes the behavior of the plaque using a five parameter, third-order Mooney-Rivlin hyperelastic constitutive equation. This model for plaque behavior neglects the artery laminate compositions, tissue anisotropy as well as the residual strain and active smooth muscle stresses [6]. The final form of the strain density function used to model the artery is given in eqn. (1).

$$W = a_{10}(I_1 - 3) + a_{01}(I_2 - 3) + a_{20}(I_1 - 3)^2 + a_{11}(I_1 - 3)(I_2 - 3) + a_{30}(I_1 - 3)^3$$
 (1)

The constants were developed for this model by fitting the five-parameter Mooney-Rivlin expression to uniaxial tension tests of human aortic atherosclerotic tissue data [15]. The hyperelastic constants for the plaques are given in Table 2.

3.3 Balloon

To model the mechanical properties of the balloon without evaluating the balloon's behavior during unfolding, empirical data was used. The stress-strain

Table 1: Hyperelastic constants for the artery.

Hyperelastic Constants [Pa]		
a ₁₀ =	= 0.01890	
a ₀₁ =	= 0.00275	
a ₂₀ =	= 0.08572	
a ₁₁ =	= 0.59043	
	$a_{02} = 0$	

Table 2: Hyperelastic constants for plaque.

Hyperelastic Constants [Pa]	
Cellular Plaque	Calcified Plaque
$a_{10} = -0.088314$	$a_{10} = -3.0254$
$a_{01} = 0.10619$	$a_{01} = 3.1073$
$a_{20} = 0.11373$	$a_{20} = 107.39$
$a_{11} = 0.89382$	$a_{11} = -234.7$
$a_{02} = -0.96676$	$a_{02} = 137.22$

curve for the full expansion of the balloon produced a linear piecewise function. The first segment of the piecewise function is representative of the unfolding balloon, while the second is of the balloon expansion after unfolding.

3.4 Stent

The stent was modeled after the slotted tube geometry given by Migliavacca *et al.* [10]. This model assumes the stent to be made of 316LN stainless steel. The Poisson ratio is 0.3 and the Young Modulus is 200 GPa.

4 Boundary conditions

The artery, balloon, plaque, and stent were all constrained in the rotational directions allowing no rotation. The artery was constrained axially at the distal ends. The artery was at a minimum, 7mm longer than the stent on each side and 3.5mm longer than the end of the balloon on each side. This constraint on the artery did not affect the behavior of the artery at the point of contact with the stent or balloon because of the extra length of the artery on both sides. The same axial constraint was placed on the balloon. To model the expansion of the balloon the balloon was assigned a ramped internal pressure load.

5 Deformed geometry export and solution

In order to examine the blood flow patterns thorough the stented artery, a conversion module was written to export the final expanded stent/plaque/artery system for meshing and use within a finite volume computational fluid dynamics code (FLUENT/UNS). The deformed geometry was exported to STL format and the internal geometry (solid mirror of the stent/plaque/artery) was meshed using a hex dominant grid. Holes and seams between the various geometry parts were automatically filled within the meshing algorithms.

The inlet boundary condition was specified as a velocity inlet using the time dependent flow rate equation of Womersley found in [16]. The flow was

assumed to enter the simulation with a constant cross-sectional velocity. A noslip condition where the stent/plaque/artery surfaces were in contact with the blood was assumed. The Casson model was used to evaluate the viscosity of the blood since the artery was relatively large and inhomogeneities associated with Fahraeus effects were negligible.

6 Results and discussion

Figure 6 shows the endflare during stent expansion for the 1mm and 2mm balloon overhang cases when there is no plaque included in the models. Endflare is defined as the ratio of the stent diameter at the distal ends to the diameter at the stent centerline. There is a significant difference in the endflare both at the point of peak endflare and at the end of expansion, indicating that the amount of balloon overhang can have a significant impact on vascular injury.

Figure 7 shows the endflare during stent expansion for the 2mm balloon overhang case when calcified and cellular plaque are included in the models. The endflare with plaque present in the model is significantly higher than when it was not included. This is because the distal ends of the stent were located such that they did not contact the plaque. Therefore, the effective diameter against which the stent was expanding was significantly larger in this area leading to a lower expansion resistance. There is a significant difference in the endflare both at the point of peak endflare and at the end of expansion, indicating that the amount of balloon overhang can have a significant impact on vascular injury.

Figure 8 shows the arterial stresses at the end of stent expansion when calcified and cellular plaque was included in the model. At the end of expansion, the increase in maximum endflare for the cellular plaque geometry over the calcified plaque geometry is about 200%. The increase in maximum arterial stress is 200% at the point of stent contact at the proximal and distal ends.

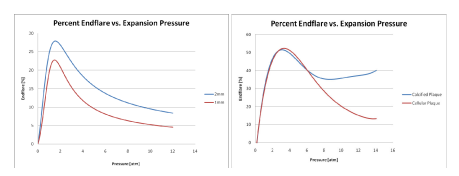


Figure 6: Endflare without plaque for 2mm (upper line) and 1mm (lower line) overhang.

Figure 7: Endflare with plaque for calcified (upper line) and cellular (lower line) plaque.

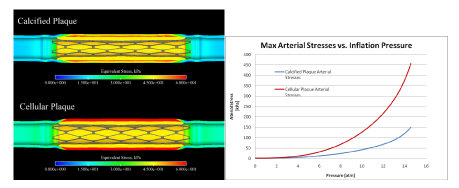


Figure 8: Arterial/plaque stresses for calcified (top) and cellular (lower) plaque.

Figure 9: Max arterial stresses for calcified (lower line) and cellular (upper line) plaque.

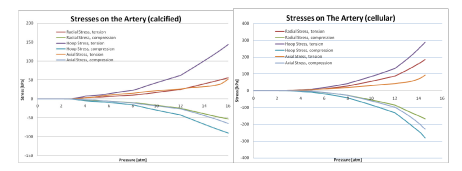


Figure 10: Arterial stresses (radial, hoop, and axial) for calcified plaque.

Figure 11: Arterial stresses (radial, hoop, and axial) for cellular plaque.

Figure 9 shows the max arterial stress during the expansion process for the cellular and calcified plaque cases. The cellular plaque case results in much higher stresses at a given balloon expansion pressure than the calcified plaque case.

Figures 10 and 11 show the directional stresses when calcified and cellular plaque are included in the model. As expected, the arterial stresses were significantly lower for the calcified plaque cases when compared to the cellular case. This is the result of the much higher rigidity of the cacified plaque layer over the celluar plaque. This is further supported by the higher stresses seen by the plaque in the calcified case as compared to the cellular case.

It should be mentioned that no plaque breakup model was included in the simulations. It would be expected that the calcified plaque would break up before the high stresses at the end of expansion predicted by the model.

As one would expect, the most significant compenent of stress is the hoop. The smallest compenent is the axial. The clinical significance of the directional stresses on vascular injury is unclear. Future experimental studies will be aimed at determining the most appropriate stress to analyze when evaluating vascular injury (von Misses, maximum principal, directional, etc.). The current model does not include the anisotropic behavior of the artery and plaque materials. Additionally, no distinction was made between the passive arterial medium and the active fibers, the orentations of which may vary from inner to the outer layers of the arterial wall. This is currently being included in a more sophisticated model which also includes arterial prestresses.

Finally, Figure 12 shows the preliminary results of the flow pattern within the final stented geometry. The development of Poiseuille flow is apparent as the blood moves from the model entrance along the artery and plaque surface (not show) and through the stented portion of the artery. The strain rates are quite interesting and show a decrease after the initial entry into the arterial section. However, they rapidly increase as the blood travels through the stented portion of the artery.

This CFD model is now being used to evaluate the diffusion process associated with drug eluting stents, as well as modeling neointima formation, thrombus formation mechanics, and blood flow patterns. It has been found that standing vortices and regions of stagnation are responsible for the rises in concentrations of platelet-activating agents within those regions. Platelets accumulate preferentially in the regions of large platelet-activating agent concentrations and low fluid velocities. Figure 13 shows a contour plot at 0.05 m/s. Regions of the domain near where the stent contacts the plaque exhibit

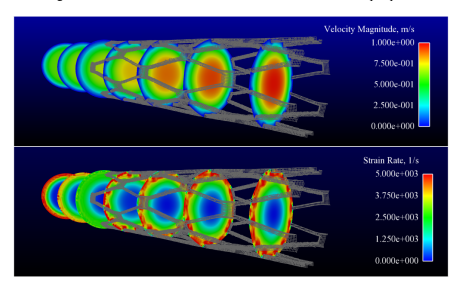


Figure 12: Velocity magnitudes and strain rates in stented artery.

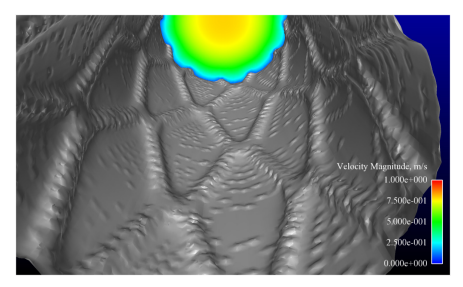


Figure 13: Iso-contour at 0.05 m/s on region near plaque/stent interface.

zones of low velocity. These regions are potential locations for thrombus formation. Additionally, the wall shear rates and blood strain rates at the wall can be evaluated to determine if they exceed a critical embolizing limit.

7 Clinical significance of report

Concerns that drug-eluting stents interfere with the process of reendothelization and thus may encourage long-term thrombosis have spurred interest in understanding the mechanisms causing acute deendothelization during the stenting procedure. This model aids in the prediction of regions of endothelial cell (EC) denudation during stent implantation. This is an important phenomenon since regions of EC denudation profoundly impact drug absorption/loading profiles of anti-proliferative agents in drug-eluting stents (DES). Additionally, anti-proliferative drugs are hypothesized to inhibit EC regrowth causing increased rates of long-term thrombosis, so predictive capability of regions of EC denudation during implantation provides the tool to reduce thrombosis rates of DES

The model developed also helps in the prediction of regions of high arterial stresses, which may cause vascular injury. Acute superficial and deep vascular injury has been found to be a strong predictor of chronic restenosis. This method provides a predictive tool to evaluate the degree of acute vascular injury of new stent geometries prior to *in-vivo* studies.

Finally, the ability to examine blood flow patterns in stented arteries allows for the prediction of standing vortices and high platelet-activating substances capable of trapping and stimulating platelets for aggregation. It also allows for the evaluation of embolizing stresses acting on a thrombus.

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