INTRODUCTION

Originally, computational fluid dynamics (CFD) was a mechanical engineering method for comprehensive analysis of fluid flow, heat transfer, and associated phenomena with the use of computer-based simulations. Recently, it has become a widely adopted methodology for solving complex problems in many modern engineering fields. The CFD-based approach is used for understanding the relationship between structure and flow by solving systems of partial differential equations to simulate fluid flow. Advances in medical imaging and computational methods provide an opportunity to apply the CFD-based approach in clinical medicine. Cardiovascular imaging is one of the most common medical applications of CFD modeling. Although CFD modeling is essentially studied by applied mathematicians and engineers who are specialists in fluid mechanics, as CFD modeling continues to spread into clinical medicine it is very important for clinicians to understand the basic principles, benefits, limitations, and pitfalls of CFD modeling. In this paper, we present an overview of CFD modeling and its application to aortic diseases, and primarily to aortic aneurysms.

FRAMEWORK OF CFD MODELING

CFD modeling is performed by solving differential equations to simulate fluid flow. The construction and solution of a CFD model involves several steps.

Geometrical discretization for CFD modeling

The first step of CFD modeling is to define the geometry of the region of interest to simulate the fluid flow. Geometrical extraction converts medical images into numerical geometries that define the physical boundaries of the model region of interest. A variety of medical imaging techniques can be used for geometrical extraction, including ultrasound, CT, MRI, and X-ray angiography. Spatial discretization, or “meshing,” divides the geometry into a number of discrete volumetric elements or cells. Temporal discretization divides the system into discrete time steps [1]. As the accuracy of a solution of a CFD model is essentially dependent on geometry extraction—namely, the spatial resolution and image quality of medical imaging—medical imaging must provide sufficient anatomical resolution to enable subsequent segmentation and data extraction. Most of the time spent on CFD modeling is devoted to this geometrical extraction process [2]. In cardiovascular systems, computational imaging tools may confer grid generation information, but there are limitations in that the resolution of current imaging tools is still low and the geometry varies according to the cardiac cycle [2]. Additionally, the time required for the computational solution of a CFD system depends on the resolution of the grid. Recently, high-performance computing systems have addressed many of the challenges facing huge, high-resolution CFD systems incorporating detailed geometrical data, thereby relieving CFD modeling of the long times required to solve complicated systems.
Definition of fluid properties

Prior to performing a CFD simulation of a cardiovascular system, the essential fluid properties of blood need to be clearly defined. Most non-biological fluids approximate Newtonian fluids, which result in a constant viscosity. The various components of blood, including plasma, blood cells, and other carried material, tend to cause blood to behave in a non-Newtonian manner in various degrees depending on the biological condition and anatomical scale of vessels, where the blood's viscosity changes with the shear rate of the flow [1]. Thus, the correct viscosity model, consisting of a mathematical equation, should be selected based on the range of shear rates in the system. For example, when the shear rate is sufficiently high, such as in an aorta or in large vessels, blood flow exhibits Newtonian flow behavior, but the non-Newtonian behavior of the fluid cannot be ignored in small vessels [1]. The rate of change in velocity from along the vessel wall to the center of the vessel is proportional to the wall shear stress (WSS) [1].

Boundary conditions

Because it is impossible to discretize the entire cardiovascular system in clinical situations, the region to be analyzed will have at least one inlet and one outlet of flow. To enable CFD modeling, boundary conditions of the flow, namely the physiological conditions at the inlet and outlet boundaries, must be specified [1]. Boundary conditions are a set of applied physiological parameters that may vary over time, such as such as blood pressure, blood flow velocity, and temperature. Boundary conditions may be based on patient-specific data, population data, physical models, or assumptions [1,2].

Numerical solution

After the pre-processing steps discussed above, the next step in CFD modeling is finding a numerical solution. Almost all CFD analyses solve the Navier-Stokes equations and continuity equations that govern fluid motion. A variety of numerical solution techniques are available, such as finite difference, finite element, finite volume, and spectral methods [2]. In a typical three dimensional (3D) cardiovascular simulation, millions of non-linear partial differential equations are solved simultaneously, and repeatedly, over all elements, at all time-steps. Once the numerical solution is found, the CFD model describes the condition of the flow, such as the pressure and velocity fields, over all elements and at each time-step.

Post-processing

The generated results of a numerical solution of a CFD model by a computer cannot be assessed directly. Also, some specific CFD parameters that are associated with the histopathological condition of the disease are of interest in the medical simulation. Therefore, some post-processing is required to extract and visualize relevant data. After this process, the researcher can easily understand the simulation results. Many visualization tools have been developed, including domain geometry and grid display, vector plots, line and shaded contour plots, two-dimensional and 3D surface plots, particle tracking, and color postscript outputs [2]. Various parameters of CFD modeling that are relevant to cardiovascular medicine are discussed in a later section.

CLINICAL SIGNIFICANCE OF CFD MODELING FOR AORTIC ANEURYSMS

For many years, the size of the aorta has been the principle decision making criteria for intervention of an aortic aneurysm. Many guidelines for the treatment of aortic aneurysms advocate a maximal aortic diameter and clinical risk factors as the determination of the timing for intervention [3]. Many studies have suggested that some small aortic aneurysms may rupture prior to reaching 55 mm of maximum short diameter, which is the suggested threshold size for surgical intervention [4], whereas as many aneurysms larger than the threshold never rupture. Therefore, appropriate predictors for the risk of rupture and the progression of an aneurysm, other than aortic size, would improve current treatment strategies by providing appropriate timing for intervention, while also reducing the expense and risk of unnecessary intervention.

One goal of CFD modeling in cardiovascular medicine is to understand the relationships between anatomical geometry (i.e., vascular anatomy), local hemodynamics (blood flow), fluid-structure interaction (FSI) including stress and pressure on the vessel wall, and the vascular pathology (development and progression of atherosclerosis, aneurysm, dissection, etc.) (Fig. 1). Several studies have suggested that hemodynamic forces may provide one of the most significant biomechanical effects underlying aneurysm development [5]. As measurement of in vivo hemodynamics assessment can be difficult and invasive, researchers are gradually focusing on the assessment of hemodynamics, using non-invasive CFD modeling to predict the risk of future rupture for aneurysms.

HEMODYNAMIC FORCES RELEVANT TO AORTIC PATHOLOGY

Clinical and experimental observations have indicated that various biomechanical conditions interactively influence the progression of an aortic aneurysm [6,7]. Fig. 2 illustrates the various hemodynamic forces that are relevant to biomechanical conditions in aortic pathologies.
Wall shear stress

WSS, which is the tangential force of blood flowing on the endothelial surface of blood vessels, is the most significant of the hemodynamic forces in a blood vessel. The interactions of pulsatile blood flow with arterial structures generate complex hemodynamic forces on the vessel wall that exhibit spatial and temporal variation (Fig. 3) [8]. Endothelial cellular responses to these physical stimuli influence vessel wall homeostasis. Constant laminar blood flow with tangential WSS protects the vascular endothelium [9]. After fluctuations of WSS promote changes in biochemical signals, the biophysical characters of vessel wall starts to degenerate. The degeneration of vessel wall will directly lead to the initiation and progression of some cardiovascular diseases, including aortic aneurysm [10] and coronary artery disease [11].

Pitfalls in the assessment of various WSS parameters

Although WSS is one of the most frequently studied hemodynamic parameters in cardiovascular diseases, careful attention should be paid to the interpretation of CFD modeling that is used in articles or in studies. As the pulsatile nature of blood flow produces complicated temporal and spatial WSS patterns during the cardiac cycle, various WSS parameters have been linked with cardiovascular complications, such as WSS at peak systole, time-averaged WSS, and time-averaged spatial WSS gradient. In addition, most studies rely on measures of WSS magnitude while little attention has been paid to the vectoral behavior of WSS [12]. One study suggests that the transverse WSS perpendicular to the flow direction is a more significant WSS parameter rather than the WSS magnitude [13].

A typical pitfall in WSS assessment is the fact that “the time-average of magnitude of WSS” and “the magnitude of time-averaged WSS” are different because WSS is a vector-valued quantity. The difference in values of “the time-average of magnitude of WSS” and “the magnitude of time-averaged WSS” becomes significant in flows with fluctuation where the directions of WSS vector changes with time. This pitfall is also discussed in the oscillatory shear index (OSI) section below.

Oscillatory shear index

OSI is another significant hemodynamic parameter used to assess WSS in that it has the time-averaged magnitude of WSS in...
the denominator. OSI quantifies the fluctuations of WSS from the primary flow direction during the cardiac cycle (Fig. 4). For example, blood flow with fluctuations possesses a high OSI even though its time-averaged WSS is low. Fluctuations of WSS disturb the vessel wall homeostasis and may induce aortic pathologies [10].

**Fig. 2.** Various hemodynamic forces of blood flow in the aorta. Schematic illustration of various hemodynamic forces of blood flow in the aorta. Through FSI, pulsatile aortic flow generates WSS (the tangential force exerted by moving blood along the vessel wall) and tensile stress (the perpendicular forces acting on the vascular wall) on the aortic wall during the cardiac cycle. FSI: fluid-structure interaction, WSS: wall shear stress.

**Fig. 3.** Time course change of flow vector and WSS. The time course change of the flow vector of blood flow (upper row) and WSS (lower row). Note that the pulsatile nature of blood flow produces complicated temporal and spatial WSS patterns. WSS: wall shear stress.
Vessel wall strain and distensibility

Recently, several researchers investigated other biomechanical forces aside from WSS for predicting the risk of rupture. Di Martino et al. [14] reported that maximum tissue stiffness was inversely correlated with wall strength, suggesting lower stiffness as a possible predictor of aneurysm rupture. Also, Wilson et al. suggested that distensibility at baseline and the change of the aorta during follow-up were correlated with the risk of rupture, in addition to the diastolic pressure and larger parameter [15]. Several ultrasound studies reported assessment of local wall strain of the whole abdominal aortic aneurysm in vivo [16]. Satriano et al. [17] proposed a 3D image-based approach to compute aortic wall strain maps in vivo, which was used for various imaging modalities. In terms of CFD, Stevens et al. [18] used a CFD method to assess how a change of flow-related biomechanical properties affects the wall stress and the related wall strain. Although further clinical studies are required, such hypotheses regarding vessel wall strain and wall distensibility would add more insight in predicting the precise risk for aneurysm rupture.

Fig. 4 illustrates the interrelationship between flow, WSS, OSI, and pressure on the vessel wall (i.e., distention force).

REGIONAL HEMODYNAMIC DIFFERENCES RELEVANT TO AORTIC ANEURYSMS

Few studies have investigated regional pathogenic risks of aortic pathologies. Even in different regions in the aorta that have the same diameter, the behavior of aortic dilatation differs depending on the location in the aorta. Usually, the abdominal aorta has a marked predilection for aneurysmal dilatation when compared to the thoracic aorta, where the infrarenal aorta is the most common site of aortic aneurysm formation. Different hemodynamic influences present along the length of the aorta may work in concert with other regional factors to explain this preferential distribution [6].

Region-specific structural differences are well recognized along the aorta. For example, the elastin-collagen ratio declines along the length of the aorta, reducing elasticity and wall motion [19]. Reduced distal aortic elasticity, in combination with augmented pressure due to pulse wave reflections from aortic bifurcation and other downstream arteries, may increase wall strain and aneurysm susceptibility [20].

Most relevant to aortic disease pathophysiology, and its predilection for the distal aortic segment, is the marked difference between aortic WSS in the proximal and distal aorta. In proximal aortic segments, flow is antegrade throughout the cardiac cycle, providing continuous antegrade laminar WSS [6]. As discussed above, constant laminar blood flow with normal WSS regulates homeostasis of the vascular endothelium [9]. In general, the time-averaged WSS is high, while the OSI is low in the distal thoracic aortic arch because of uniform laminar flow. In contrast, many experimental and numerical studies have indicated that multiple secondary reverse flows with vortex formation were observed in the late systolic and diastolic phases during the cardiac cycle in the distal abdominal aorta, while they were not observed in the thoracic and proximal aorta [6,11,21]. As a result, the time-averaged WSS is low and the OSI is high in the distal abdominal aorta when compared with proximal aorta. These distinct regional differences in hemodynamic influences may account for some component of the differential aneurysm risk noted between the thoracic and abdominal aortic segments.

Furthermore, there is individual variation in the location of aneurysmal formation among patients. Although the infrarenal aorta is the most common site of aortic aneurysm formation, some patients have an aneurysm in the distal aortic arch or in the thoraco-abdominal junction. Differences of anatomical geometry (shape of the aorta) can generate patient-specific flow patterns and FSIs during the cardiac cycle [22]. CFD modeling improves our understanding of variation in the location of an-
eurysmal formation among individual patients (Fig. 5).

LIMITATIONS AND PITFALLS OF CFD MODELING

In spite of the many advantages of the clinical application of CFD modeling, and the variety of useful visualizations provided by CFD modeling, simulated CFD data should be thoroughly validated. The accuracy of the model is determined by the model's design and the quality of the input data. The validity of CFD modeling results depends significantly on the selection of appropriate geometry discretization, boundary conditions, fluid properties, and the numerical solution methodology [23]. Due to the many simplifications and assumptions taken into account in CFD modeling, the degree of accuracy of the results needs to be assessed prior to application in clinical practice [23].

In this sense, doctors should learn how to assess a CFD model and understand its inherent limitations and pitfalls. Particularly, doctors should make critical judgments about the computed results from the perspective of medical validity [2]. Some kind of clinical variation should be attempted in clinical trials.

CONCLUSIONS

The advancement of CFD modeling has presented a unique opportunity to provide new insights into vascular hemodynamics in the assessment of cardiovascular diseases. Patient-specific CFD modeling has the potential to provide a comprehensive understanding of the interaction between vascular morphology, blood flow, and FSI. Information about biomechanical forces in aortic pathologies may help to predict the risk of aortic aneurysm and to select appropriate treatment.

Conflicts of Interest

The authors declare that they have no conflict of interest.

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