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Xu, H., Li, Z., Dong, H. et al. (6 more authors) (2017) Hemodynamic parameters that may predict false-lumen growth in type-B aortic dissection after endovascular repair: A preliminary study on long-term multiple follow-ups. Medical Engineering & Physics, 50. pp. 12-21. ISSN 1350-4533

https://doi.org/10.1016/j.medengphy.2017.08.011

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# Hemodynamic Parameters That May Predict False-lumen Growth in Type-B Aortic Dissection after Endovascular Repair: A Preliminary Study on Long-term Multiple Follow-ups

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#### 1 ABSTRACT

Thoracic endovascular aortic repair (TEVAR) is commonly applied in type-B aortic 2 dissection. For patients with dissection affects descending aorta and extends downward to 3 involve abdominal aorta and possibly iliac arteries, false lumen (FL) expansion might occur 4 5 post-TEVAR. Predictions of dissection development may assist in medical decision on re-intervention or surgery. In this study, two patients are selected with similar morphological 6 features at initial presentation but with different long-term FL development post-TEVAR 7 (stable and enlarged FL). Patient-specific models are established for each of the follow-ups. 8 Flow boundaries and computational validations are obtained from Doppler ultrasound 9 velocimetry. By analyzing the hemodynamic parameters, the false-to-true luminal pressure 10 difference (PDiff) and particle relative residence time (RRT) are found related to FL 11 remodeling. It is found that (i) the position of the first FL flow entry is the watershed of 12 negative-and-positive PDiff and, in long-term follow-ups, and the position of largest PDiff is 13 consistent with that of the greatest increase of FL width; (ii) high RRT occurs at the FL 14 proximal tip and similar magnitude of RRT is found in both stable and enlarged cases; (iii) 15 comparing to the RRT at 7days post-TEVAR, an increase of RRT afterwards in short-term is 16 found in the stable case while a slight decrease of this parameter is found in the enlarged case, 17 indicating that the variation of RRT in short-term post-TEVAR might be potential to predict 18 long-term FL remodeling. 19

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- 21 Key Words: aortic dissection; hemodynamics; endovascular procedures.

#### 22 INTRODUCTION

Aortic dissection (AoD) is a severe cardiovascular disease, where a surge of blood 23 flowing into the aortic wall via an initial tear or damage of the intima and splitting the 24 single aortic lumen into a true and false lumen (TL and FL). Stanford type-B AoD 25 26 indicates those with the dissection begins distal to the supraaortic branches. Interventional treatment of Stanford type-B AoD commonly involves thoracic 27 endovascular aortic repair (TEVAR)[1]. In a number of patients, FL expansion is 28 found post-TEVAR, especially in the infrarenal aorta. Recent study confirmed that 29 abdominal aortic expansion can be frequently found after TEVAR and is independent 30 from thoracic FL thrombosis[2]. Prediction of FL growth may contribute to early 31 decision-making of re-intervention or surgery. The post-TEVAR development of 32 dissection is highly dependent on local hemodynamics[3]. Medical imaging tools such 33 34 as Doppler ultrasound[4] and phase-contrast MR (pcMR)[5] are able to capture the flow velocity within aorta. However, the former provides velocity information at a 35 certain position of the vessel, and the latter reveals flow movement with relatively low 36 spatial and temporal resolution[6, 7]. On the other hand, the uptake of 37 <sup>18</sup>F-fluorodeoxyglucose in PET-CT can indicate complications in AoD[8] and positive 38 correlation between the uptake and wall shear stress is found in aortic aneurysm 39 study[9]; however, PET-CT is relatively expensive and the flow information cannot be 40 directly reported. Thus, computational simulations that can provide hemodynamic 41 parameters, such as flow, pressure and shear stress distributions, may enrich analysis. 42

Previous computational works focusing on type-B AoD include investigations on 43 hemodynamic features[10-14], luminal flow exchange[12, 15], post-TEVAR flow 44 45 effects[16-18], tear-induced flow effects[19, 20] and fluid-structure interaction studies[21, 22]. Besides, 4D pcMR[5, 23] and phantom[24, 25] measurements have been 46 47 conducted to compare with or validate the computed results. In this study, we investigate the flow-driven dissection development after TEVAR based on long-term 48 multiple follow-ups. Flow conditions in patients with stable and enlarged FL are 49 compared and key hemodynamic parameters that are related to dissection growth in 50 abdominal aorta are proposed, facilitating medical decision-making on post-TEVAR 51 treatment. 52

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#### 55 **METHODS**

#### 56 Image Acquisition and Model Reconstruction

This study was approved by the institutional review board of the Chinese PLA 57 General Hospital. Written informed consent was obtained from the patients involved 58 in this study. Two male patients (PI and PII) with subacute Stanford type-B AoD 59 underwent arterial-phase CT angiography (CTA) at initial presentation and during the 60 follow-up examinations after TEVAR via a dual-source CT scanner (SOMATOM 61 Definition Flash, SIEMENS, Germany). Details of the CTA scan and the patient 62 information are described in S1, Supporting Document. Image segmentation and 63 surface reconstruction of AoD were conducted through Mimics (Materialise, 64 65 Belgium). The cross-sectional contours of the reconstructed geometries were mapped back to CTA images to ensure that the 3D models present the actual outline of the 66 vessel lumen. Detailed views of the models are shown in Fig.1, where PI/II-1 67 indicates the models pre-TEVAR while PI/II-2 and others are models post-TEVAR. 68 69 After TEVAR, PI and PII have experienced six- (7days~53months) and four-times (7days~35months) CTA scans. The FL in PI was in stable condition (PI-2~7, Fig.1b) 70 71 while that in PII was expanding (PII-2~5, Fig.1d). The models were meshed in ICEM (ANSYS Inc, Canonsburg, USA) with tetrahedral elements in the core region and 72 73 prismatic cells (10 layers) in the boundary layer near the aortic wall. The grid resolution varies from 2,564,019 to 3,153,829 cells. 74

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# 76 Doppler Ultrasound and Boundary Conditions

77 Time-variant velocities at ascending aorta (AAo), brachiocephalic trunk (BT), left common carotid artery (LCCA), left subclavian artery (LSA) were measured via 78 79 Doppler ultrasound of the patients, and velocity variation at the distal thoracic aorta (DTAo, about 5cm above celiac trunk) was measured in the final examination of PI to 80 provide validation of the computational results. This is because the true lumen (TL) 81 remodeling at this position in PI is sufficient, so that relatively organized flow is 82 found and the central line of vessel can be accurately identified. The velocity of AAo 83 was measured through the apical 5-chamber view and the suprasternal long axis view 84 85 of aortic arch. The two results were compared to ensure the maximum velocity at AAo could be captured. For other arteries (BT, LCCA, LSA and DTAo), Doppler 86

velocimetry has been conducted at the proximal and distal sites of the targeted measurement vessel. Mean velocity over a cardiac cycle at the two sites for each particular vessel was then calculated and compared. When the difference between them is less than 5%, the measured velocity is considered effective. Details of the measurement are described in S2, Supporting Document.

The upper edge of the velocity sonogram was extracted (Fig.1e) as the variation of 92 the maximum velocity at the measured site. The flow rates at AAo, BT, LCCA and 93 LSA, as the velocity boundary conditions, can then be calculated based on the 94 95 measured time-variant maximum velocity and the assumed flat flow profile for AAo and parabolic flow profile for the others. The velocity boundary conditions of the 96 models is shown in Fig.S1a, Supporting Document. Pulsatile waveforms of the 97 pressure at celiac artery (CA), superiormesenteric artery (SMA), renal arteries and the 98 outlets at common iliac arteries were obtained from previous study[26] (Fig.S1b, 99 Supporting Document). As shown in Fig.1, two models in PII (PII-1 and PII-4) were 100 cropped below the iliac bifurcation due to the relatively shorter CT scanning range. To 101 eliminate the outlet effects, time-variant pressure distribution at the cropping plane 102 has been calculated in PII-2~3 and PII-5. The averaged pressure information at this 103 104 cropping plane was mapped to PII-1 and PII-4, serving as the pulsatile pressure outlets. 105

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## 107 Numerical Models

The vessel wall was assumed as no-slip and rigid, due to low distensibility in 108 long-term follow-ups[27]. The blood was treated as Newtonian and incompressible 109 with density of 1044kg/m<sup>3</sup> and dynamic viscosity of 0.00365kg·m<sup>-1</sup>·s<sup>-1</sup>[28]. The 110 average Reynolds number over a cardiac cycle, calculated based on the equivalent 111 diameter  $(D_e = 2\sqrt{Area/\pi})$  and velocity at the inlet of the ascending aorta in PI and 112 PII, were between 2066-2197 and 2844-2960, respectively. Our previous study 113 confirmed that laminar simulations with adequately fine mesh resolutions, especially 114 refined near the walls, can capture flow patterns as turbulence model[15]. To further 115 confirm this, we solved the flow in the first follow-up cases (PI-2 and PII-2) by both 116 laminar and k-w SST turbulence models, where the flow in the abdominal aortic 117 region is the fastest during the follow-ups. Similar flow patterns were found and the 118 discrepancy of the maximum velocity and wall shear stress (WSS) in the abdominal 119

aorta was 4.2% and 5.2% respectively (the laminar and turbulent results of PII-2 were 120 shown in Fig.S2, Supporting Document), ensuring the rationality of laminar model to 121 be applied in the current problem. A finite volume solver, CFD-ACE+ (ESI Group, 122 France) was employed. The heart-beat cycle of the patients was measured at each 123 clinical examination. The averaged cardiac cycle for PI and PII were 71 and 124 69beat/min respectively. Temporal discretization of numerical models was assigned as 125 45step/cycle. Simulation was carried out for 5 cardiac cycles to achieve a periodic 126 solution and results of the final cycle were presented. Grid and temporal 127 independency analyses on finer grids and finer temporal discretizations were 128 conducted to ensure the base resolution with the base time step settings are adequate 129 (S3, Supporting Document). 130

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## 132 **RESULTS**

#### 133 Aortic Remodeling

Aortic remodeling was assessed by: (i) size and numbers of the aortic tears (AoT), (ii) 134 change of luminal volume, and (iii) growth of aortic diameter. There are four major 135 136 AoTs along the aorta for both patients at initial presentation. The primary entry (AoT-1) in both patients is located at the proximal region of descending aorta and the 137 locations of other AoTs (AoT-2~4) are displayed in Fig.1. After TEVAR, the primary 138 entry was sealed and after the first follow-up of PI, AoT-2 was disappeared. 139 140 Considering the position and local aortic curvature of the tears, AoT-4 is the exit of the flow from FL to TL, while the function of AoT-2 and AoT-3 is uncertain. Since 141 only the flow entries towards the FL is able to bring mechanical impact into the 142 dissection, size measurement was only performed on AoT-2 and AoT-3 (Table 1). 143 Post-TEVAR, both AoT-2 and AoT-3 in PI and PII are enlarged, probably due to the 144 greater flow impact on these sites after AoT-1 was sealed. 145

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Tear	Geometry	PI-1	PI-2	PI-3	PI-4	PI-5	PI-6	PI-7
AoT-2	H[mm]	7.87	7.15	-	-	-	-	-
	W[mm]	8.97	10.76	-	-	-	-	-
	$A[mm^2]$	64.61	70.71	-	-	-	-	-
AoT-3	H[mm]	13.86	18.79	17.93	15.31	18.23	8.81	8.57
	W[mm]	10.04	9.38	6.03	8.90	7.30	6.72	5.75
	$A[mm^2]$	116.13	150.82	106.40	119.19	128.82	93.49	44.09
Tear	Geometry	PII-1	PII-2	PII-3	PII-4	PII-5		
AoT-2	H[mm]	8.17	8.41	5.74	11.04	11.20	-	
	W[mm]	9.34	9.96	7.36	9.59	6.03		
	$A[mm^2]$	62.61	70.47	37.33	89.50	54.05		
AoT-3	H[mm]	9.91	9.55	7.77	6.25	6.19	-	
	W[mm]	10.01	11.47	9.34	8.04	9.90		
	$A[mm^2]$	85.27	99.32	61.51	40.35	53.32	_	
		TT TT	ai alet. W	Width	A A # 2 0		-	

Table 1. The size of the aortic tears in the patients with long-term follow-ups

H-Height; W-Width; A-Area.

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The change of luminal volume is shown in Fig.2. The volume of TL  $(V_{TL})$  involves 147 the initial dissection-affected TL region, while V<sub>TL-Part</sub> indicates the 148 dissection-affected TL region during the follow-ups. Variations of V<sub>TL</sub>, volume of FL 149 150  $(V_{FL})$  and the ratio between  $V_{TL-Part}$  and  $V_{FL}$  are displayed in Fig.2b.  $V_{FL}$  in the stable case (PI) reduces gradually post-TEVAR, except for the final examination, where a 151 trivial increase of  $V_{FL}$  is found (131.6mm<sup>3</sup> in 20months). However,  $V_{FL}$  in the 152 expanding case (PII) reduces in the first two follow-ups but enlarges afterwards, 153 where the V<sub>FL</sub> growths in PII-4 and PII-5 are 2,138mm<sup>3</sup> and 34,989mm<sup>3</sup> respectively, 154 155 the durations of which are both 17months (Fig.2b). For both cases, the TL-to-FL volume ratio (V<sub>TL-Part</sub>/V<sub>FL</sub>) keeps increasing, except for PI-6 and PII-3 (pointed by 156 arrows). This is because significant FL regression occurs at the proximal region (blue 157 squares in Fig.1b,1d), which shortens the comparison region. Since TL remodeling is 158 insufficient in this region, value of V<sub>TL-Part</sub>/V<sub>FL</sub> reduces. The averaged increase rates of 159 V<sub>TL-Part</sub>/V<sub>FL</sub> in the follow-ups of PI and PII are 1.8% and 0.3% per month, 160 respectively, indicating a significant luminal remodeling difference. 161

The aortic diameter (D) is measured in each model, taking into account the width of 162 TL and FL ( $W_{TL}$  and  $W_{FL}$ ) and the thickness of flap. As shown in Fig.1f, 163 measurements have been conducted in the axial plane of the CTA datasets, along a 164 line that crosses the centre of the flap and is perpendicular to the flap. Four 165 measurement positions are selected in the abdominal aorta. As shown in Fig.2a, they 166 are 1cm below diaphragm (L1), 1cm above CA (L2), 3.5cm below SMA (L3) and 9cm 167 below SMA (L4). Fig.2c~e display the variation of D, W<sub>TL</sub> and W<sub>FL</sub> respectively. 168 Positive TL remodeling is found in PI and PII in general, except for L3 and L4 in PI-3 169

and L3 in PII-4~5.  $W_{FL}$  at the proximal region of PI (L1 and L2) reduces greatly during the follow-up, while in the distal region (L3 and L4), variation of  $W_{FL}$  is stable. In PII,  $W_{FL}$  increases sharply since the second follow-up (PII-3). Comparing the variation of  $W_{TL}$ ,  $W_{FL}$  and D, it can be found that the variation pattern of D is mainly determined by the variation of  $W_{FL}$ .

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# 176 Flow Pattern, Validation and Luminal Flow Exchange

Fig.3 shows the flow streamlines at systolic peak where the color map of velocity 177 magnitude is restricted to a certain range in PI ( $0 \sim 2.51$  m/s) and PII ( $0 \sim 3.78$  m/s) to 178 assist visualization of the longitudinal velocity variation. At systolic peak, fast and 179 180 organized flow is found in TL; while vortical and relatively slow flow presents in FL. Highest velocity presents in the short-term follow-ups (PI-2~3, 4months for PI; 181 PII-2~3, 1months for PII) at the region below stentgraft and above AoT-2 and -3, 182 where TL remodeling is insufficient and the blood has not been diverted. In long-term 183 follow-ups (PI-4~7 and PII-4~5), the flow in the proximal FL is generally slow 184 (<0.5mm/s); while below AoT-2 (the first re-entry), where blood perfusion occurs, 185 faster flow up to 1.53m/s in PI and 2.22m/s in PII are found in the FL with helical 186 feature (Fig.3c). 187

To validate the computational results, Doppler ultrasound velocimetry was 188 performed at distal thoracic aorta (5cm above CA) along the centerline of TL. This 189 was only conducted at the final examination of PI, because its TL in the distal thoracic 190 aorta is fully remodeled and the local flow is therefore organized (Fig.3a). The 191 measured and computed velocities at the centre of the cross-section of the vessel, 5cm 192 above CA, are 1.58m/s and 1.66m/s respectively, indicating a difference of 5.1%. 193 Moreover, the computed velocity variation pattern is similar to the measured one 194 (Fig.S3, Supporting Document), ensuring the rationality of computational results. 195

Fig.4 displays the luminal flow exchange via the primary entry for initial presentation and that via AoT-2 and AoT-3 during the follow-ups. As abovementioned, AoT-4 functions as the outlet of FL throughout the cardiac cycle, thus it is not displayed in Fig.4. As shown in Fig.4a (positive values indicates flow enters FL), before treatment, the TL-to-FL flow exchange in PI and PII presents similar pattern: the primary entry serves as the main flow inlet of FL; AoT-2 functions as the outlet of FL in the initial part of systole and behaves as inlet in the rest part of

cardiac cycle; and AoT-3 mainly serves as the outlet of FL. There are 31.48% and 203 40.00% of the total flow diverted to FL over a cardiac cycle for PI and PII 204 respectively, at initial presentation. After TEVAR, the function of the tears changes. In 205 PI, AoT-2 becomes the inlet of FL in the entire heart-beat cycle (Fig.4b), while AoT-3 206 functions as the inlet of FL during systole and serves as the outlet in diastole (Fig.4c). 207 Since the second follow-up of PI, AoT-2 is closed and AoT-3 gradually becomes the 208 inlet of FL throughout the entire cardiac cycle. The flow entering FL are 16.67%, 209 6.98%, 8.24%, 9.59%, 5.43%, 3.09% of the total flow for case PI-2~7 respectively, 210 211 the variation of which is consistent to the variation of tear size (Table 1) and it is generally reducing. In PII, both of AoT-2 and AoT-3 are existed during the follow-ups. 212 AoT-2 mainly serves as the inlet of FL; especially in model PII-4 and PII-5, positive 213 values of mass flow rate are found in the entire cardiac cycle (Fig.4b). AoT-3 diverts a 214 small amount of flow into the FL during systole and mainly behaves as the outlet of 215 FL (Fig.4c). There are 5.22%, 3.06%, 14.76% and 10.23% of the total flow diverted 216 into FL in case PII-2~5 respectively, the variation of which is also consistent with the 217 variation of tear size. 218

219

# 220 Loading Distribution along the Aorta

Fig.5a~b display the pressure drop at systolic peak (PDsys, the difference between the 221 local pressure and the pressure at the bottom of the model). In TL, PDsys reduces 222 from the inlet of AAo to the outlets of common iliac arteries. At initial presentation 223 (PI-1 and PII-1), FL diverts part of the total flow (31.48% for PI and 40.00% for PII), 224 maintaining PDsys in relatively low values. After TEVAR, in short-term follow-ups 225 (PI-2~3 and PII-2~3), proximal descending aorta is supported by stentgraft and 226 reshaped; however, below the endograft, TL remains collapsed, inducing larger 227 resistance and resulting in higher velocity and regional lower pressure in this region 228 (indicated by arrows in Fig.5a~b). In long-term follow-ups (PI-4~7 and PII-4~5), TL 229 remodeling in distal thoracic aorta is improved; PDsys thus gradually reduces along 230 aorta. In contrary, PDsys in FL is relatively uniform, which is probably due to the 231 232 higher energy exchange induced by the vortical flow. Fig.5d shows the variation of the maximum PDsys during the follow-ups. For both patients, the maximum PDsys 233 increases greatly soon after TEVAR (7days); along with the progress of TL 234 remodeling, the maximum PDsys reduces. The PDsys in normal aorta model (Fig.5c) 235

based on the same boundary conditions as PI and PII is also computed and the
averaged maximum PDsys is shown by the dash line in Fig.5d. Since the growth of
FL in the two patients shows obvious difference, the similar variation pattern of
PDsys indicates that the pressure drop is mainly affected by TL remodeling.

The mean arterial pressure drop (PDmean) from AAo to the common iliac arteries during a cardiac cycle is also calculated. It is 6.29mmHg for the initial model of PI and 4.70mmHg for the final model (53months post-TEVAR), which is slightly higher than that in normal aorta for the studied segment (4mmHg[29]). The PDmean of PII at the initial and final (35months post-TEVAR) models are 6.53mmHg and 6.07mmHg, respectively, consistent with its insufficient TL remodeling.

In the final examination of PI (PI-7), V<sub>FL</sub> is slightly increased, D and W<sub>FL</sub> also 246 increase at L3 (Fig.2); however, the flow exchange and tear size at this stage are both 247 reduced. Fig.5e displays the pressure distribution at systolic peak (Psys) on a slice of 248 FL for PI-6~7. It shows that the smaller tear size reduces the flow entering FL, 249 however, it increases the velocity and induces higher pressure impact on the outer 250 wall of FL (indicated by arrows in Fig.5e). The highest pressure of the impact region 251 in PI and PII at final examination are 114.37 and 115.41mmHg respectively. Besides, 252 253 by blanking the region with Psys<109.15mmHg, which is the 95% of the averaged maximum pressure in PI and PII at final examination, Fig.5f shows that the high 254 255 pressure region (in red) in the FL of PII is much larger than that in PI. In fact, the FL growth rates of PI and PII at the final examination are 0.22mm<sup>3</sup>/day and 256 68.61mm<sup>3</sup>/day respectively. 257

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## 259 Luminal Pressure Difference, Wall Shear Stress and Relative Residence Time

To investigate the pressure difference (PDiff) between TL and FL, a series of slices 260 that are perpendicular to the centerline of TL are extracted and the net pressures in TL 261 (P<sub>TL</sub>) and FL (P<sub>FL</sub>) on each slice over a cardiac cycle are calculated. Fig.6 displays 262 PDiff (PDiff=P<sub>FL</sub>-P<sub>TL</sub>) in each model. In both patients, pre-TEVAR (Fig.6a,6e), P<sub>FL</sub> is 263 smaller than P<sub>TL</sub> at the level above AoT-2; while, below AoT-2, P<sub>FL</sub> is larger than P<sub>TL</sub>, 264 265 pushing the FL towards TL. Post- TEVAR, the primary entry is closed. In PI, negative values of PDiff are found above AoT-3 (Fig.6b~c), indicating higher P<sub>TL</sub> presents in 266 this region, which supports the aortic wall and assists in TL expansion. However, in 267 the region below AoT-3 in PI, due to the blood perfusion into the FL, positive values 268

of PDiff present, indicating FL propulsion on the TL. In the final follow-up of PI 269 (Fig.6d), the maximum value of PDiff is 0.22mmHg. This maximum PDiff occurs 270 close to L3. At this position,  $W_{FL}$  and  $W_{TL}$  increase by 1.02mm and 0.46mm 271 respectively (Fig.2), indicating this small value of PDiff (0.22mmHg) is not large 272 enough to squeeze TL but the local pressure in FL (Fig.5e) is able to induce further 273 FL expansion. On the other hand, in PII, post-TEVAR, AoT-2 becomes the main inlet 274 of the flow into FL. Higher above this tear, P<sub>TL</sub> is larger than P<sub>FL</sub>, supporting TL 275 expansion (Fig.6e~f). In the near region above AoT-2, TL remodeling is insufficient 276 277 for the short-term follow-ups but PFL gradually increases; thus, regional positive PDiff is found in PII-3 above AoT-2 (arrow indicated in Fig.6f). Below AoT-2, positive 278 values of PDiff dominate and the maximum PDiff in the final examination of PII is 279 280 1.39mmHg, more than 6-times as high as that in PI, occurring close to L3. Taking account of that, in PII-5, W<sub>TL</sub> decreases yet W<sub>FL</sub> increases at L3 (Fig.2d~e), this high 281 282 PDiff propels TL collapse and the local pressure pushes FL growth.

Fig.7 shows the wall shear stress (WSS) distribution on the flap at systolic peak. Its 283 284 variation over a cardiac cycle is shown in Video S1-6, Supporting Document. In both patients, the WSS on the flap at the FL side (WSS<sub>FL</sub>) is significantly smaller than that 285 286 on the TL side (WSS<sub>TL</sub>); and the WSS<sub>TL</sub> in PII is generally higher than that in PI. At initial presentation, the maximum WSS<sub>TL</sub> occurs at the edge of primary entry; while 287 the maximum WSS<sub>FL</sub> occurs at the proximal descending aorta along the side opposite 288 to the primary entry (indicated by arrows in Fig.7b). In follow-ups, the maximum 289 WSS<sub>TL</sub> occurs at the region where TL presents most collapse; while the maximum 290 WSS<sub>FL</sub> occurs at the edge of the tears. AoT-3 and AoT-2 are the main FL flow entries 291 for PI and PII in follow-ups, which induces helical flow in the downstream and high 292 WSS<sub>FL</sub> on the side opposite to the tears (indicated by hollow arrows in Fig.7b). To 293 further visualize the variation of WSS<sub>FL</sub> at different regions along the flap on the FL 294 side, the color map is assigned to 0~5Pa (Fig.7c) and 0~0.5Pa (Fig.7d) to show the 295 distal and proximal region respectively. In PI, WSS<sub>FL</sub> at the region below AoT-3 296 reduces obviously from PI-4 to PI-6; however, in the final examination, it is slightly 297 increased, although the maximum WSS<sub>FL</sub> in this region is still smaller than 1Pa. In 298 PII, WSS<sub>FL</sub> presents obvious increase in PII-4 and reduces slightly in PII-5; the 299 maximum WSS<sub>FL</sub> at the region between AoT-2 and AoT-3 in the final examination is 300 5.21Pa. 301

Particle residence time is proposed to be related to thrombosis establishment[14].

The relative residence time (RRT), which is based on the time-averaged WSS 303 (TAWSS) and oscillatory shear index (OSI), [(1-2·OSI)·TAWSS]<sup>-1</sup>, reflects particle 304 residence time and thus may correspond with the region of thrombosis[14, 30]. 305 Fig.8a~b show the RRT distributions (normalized by the maximum RRT value) in PI 306 and PII post-TEVAR, respectively. In PI, the highest RRT occurs at the FL proximal 307 tip in PI-2~5; while in PI-6~7, RRT is greatly decreased and the highest RRT no 308 longer occurs at the FL tip. In PII, the highest RRT also occurs at the FL proximal tip. 309 Fig.8d displays the variation of the maximum RRT (normalized by the maximum 310 RRT at the first follow-up, i.e. PI-2 and PII-2 for PI and PII respectively). The RRT 311 variation in PI and PII shows significant difference during short-term follow-up: the 312 normalized maximum RRT in PI increases greatly in PI-3 (from 1 to 5.516) and 313 gradually decreases afterwards to 0.00405 in PI-7; while in PII, it decreases in PII-3 314 (from 1 to 0.025) and then maintains between 0.00324-0.319. The difference in the 315 maximum RRT's variation pattern shows potential to predict different FL remodeling 316 in the two patients. 317

318

#### 319 **DISCUSSION**

Thoracic endografts, aiming to seal the primary entry and diverting blood flow into 320 TL, are increasingly used in treating type-B aortic dissection[31]. Ideally, this 321 approach would lead to thrombosis establishment in the FL and morphologic change 322 in TL, to stabilize the aorta and consequently reduce aorta-related death. Previous 323 studies confirmed the favorable results of TEVAR; however, also reported FL 324 expansion on the segment distal to endografts, usually in the infrarenal aorta[32, 33]. 325 This is related to the patency of FL[33] or, in other words, it is related to the flow field 326 and hemodynamic conditions in AoD. Current literatures provide little information 327 concerning the fate of the abdominal aorta post-TEVAR and, to our knowledge, 328 computational studies on long-term multiple follow-up cases that are able to report 329 the change of hemodynamic parameters have been lacking. Therefore, in the current 330 study, we investigate two patients showing similar physical and hemodynamic 331 features at initial presentation but presenting different FL development (stable in PI 332 and expanded in PII) during the follow-ups. We preliminarily identify the possible 333 hemodynamic parameters that could help to evaluate/predict FL enlargement and 334 thrombosis formation. In this study, the variation trends of the hemodynamic 335

parameters are more important than their absolute values.

The mechanical load acting on the FL boundary, which includes the outer wall of 337 FL and the flap, induces FL enlargement and at the same time restricts TL 338 remodeling. The pressure that is normally applied on the FL wall plays a key role in 339 luminal remodeling[34] and the local flow directly relates to its distribution and 340 magnitude. The flow can be assessed by its amount and the velocity. The first is 341 mainly related to the size of tears. By comparing Fig.4b~c and Table 1, it can be 342 found that the absolute flow mass passes each tear per second has positive relationship 343 344 with the size of the tear. This can be shown on AoT-3 of PI and AoT-2~3 of PII, since they have multiple follow-up data, and it is consistent with previous report[14]. The 345 second, velocity, is determined by both the flow rate as well as the tear size; a smaller 346 tear size is correspondent to smaller flow rate yet accelerates the flow at the same 347 time. For instance, in the final two examinations of PI (PI-6 and PI-7), the amount of 348 the blood entering FL via AoT-3 decreases from 5.43% to 3.09% of the total flow and 349 the tear size decreases from 93.49mm<sup>2</sup> to 44.09mm<sup>2</sup>. However, the flow passing this 350 tear has been accelerated and induces stronger impact on the FL outer wall (Fig.5e). If 351 studying a longer period (PI-4 $\sim$ 7), the variation of W<sub>FL</sub> at L3 (Fig.2e), where just 352 353 below AoT-3 in PI, is similar to the variation of the ratio between the diverted flow amount and the tear size. 354

Along the aorta, the pressure in TL decreases generally but that in FL, due to the 355 vortical flow and its higher energy exchange, it does not present significant spatial 356 difference. The AoTs, functioning as the bridge between TL and FL, transport blood 357 flow and also pressure gradient. This induces similar pressure in the TL and FL near 358 the tears. Because of the relatively uniform  $P_{FL}$  and its connection with  $P_{TL}$  at the 359 tears, in general, P<sub>FL</sub> is smaller than P<sub>TL</sub> in the proximal region (above the AoTs) and 360 higher than P<sub>TL</sub> in the downstream. This general distribution feature is shown in Fig.5 361 in all of the post-interventional cases at the moment of systolic peak and similar 362 patterns can be found in other time steps of the cardiac cycle. 363

At the flap,  $P_{FL}$  and  $P_{TL}$  conflict each other; the difference between them (PDiff) may be associated with subsequent luminal remodeling[14]. Luminal remodeling is a long-term effect; investigation of PDiff in short-term follow-ups may show the variation trends of lumen remodeling, while PDiff in long-term follow-ups may be consistent to lumen remodeling results. Indeed, taking L3 as an example, in short-term (PI-2~3, PII-2~3), PD in PII increases from 1.44 to 2.93mmHg (7days-1month);

while, PDiff in PI remains about 0.55mmHg (7days-4month). This, earlier than 370 luminal change (Fig.2), shows the potential of FL enlargement for PII. In long-term 371 follow-ups (PI-4~7, PII-4~5), AoT-3 becomes the main FL flow entry in PI and 372 AoT-2 is the main entry in PII. PDiff increases from negative to positive slightly 373 below the position of AoT-3 in PI and AoT-2 in PII. Moreover, in both patients, in the 374 final examination, the maximum PDiff occurs near L3, where W<sub>FL</sub> increases the most 375 (among the compared four positions) and W<sub>TL</sub> in PII decreases. The abovementioned 376 indicates: (i) in short-term follow-ups, great increase of PDiff may imply potential FL 377 expansion; (ii) while, in long-term follow-up, the position of the first flow entry of FL 378 is the negative-to-positive watershed of PDiff, the position of the maximum PDiff is 379 consistent with the greatest W<sub>FL</sub> increase, and when the maximum PDiff is small 380 (0.22mmHg in PI-7), the pressure induces slight FL expansion without restricted acts 381 on TL, but when it is relatively large (1.39mmHg in PII-5), both FL expansion and TL 382 collapse are found. In clinical examinations, monitoring PDiff at early-stage and 383 identifying the position of FL entries and the position of largest PDiff may assist in 384 385 wisely control of the untreated aorta segment.

The WSS is related to the formation of thrombosis. Previous studies suggested that 386 the tearing of the aortic wall and high WSS in the near region of the tears could 387 promote initial activation of platelets as well as the formation of platelet aggregates; 388 while, the highly vortical flow pattern in the FL corresponding with low WSS 389 promotes platelet aggregation and deposition, so that leads to surface thrombosis [35, 390 36]. In other words, lower WSS may induce surface thrombus and thus lead to 391 constructive FL remodeling [37, 38]. In the long-term follow-ups (PI-4~5 and 392 PII-4~5), complete thrombosis is found at the proximal region of dissection and 393 partial thrombosis remains above the re-entries. The partial thrombosis in both PI and 394 PII is aligned with the intimal flap; thus, WSS on the flap along the TL and FL sides 395 are compared. The WSS<sub>FL</sub> is significantly lower than WSS<sub>TL</sub> throughout the cardiac 396 cycle; the low  $WSS_{FL}$  possibly induces surface thrombus along the flap in FL while 397 the high WSS<sub>TL</sub> can keep the TL patent. The lowest WSS<sub>FL</sub> (<0.25Pa) occurs at the 398 proximal region of FL (Fig.7d), implying potential thrombosis in these regions. 399 Indeed, partial thrombosis in PI-4~5 with very low WSS<sub>FL</sub> turns to complete 400 thrombosis in PI-6~7 (indicated by arrow inFig.7d). Moreover, in PII, the WSS<sub>FL</sub> at 401 the tip of the flap in PII-3 is small (<0.25Pa). In its next follow-up (PII-4), growth of 402 partial thrombosis can be found (indicated by arrow in Fig.7d). However, slightly 403

404 higher  $WSS_{FL}$  is shown in PII-4 at the proximal tip, and in its next follow-up (PII-5), 405 the partial thrombosis is slightly reduced (indicated by hollow arrow in Fig.7d). This 406 indicates the surface thrombosis is possibly very sensitive to WSS, and during the 407 long-term recovery, FL regression/growth could be repeatedly occurred at the 408 proximal tip of the FL.

The derived parameter of shear stress - RRT is related to particle residence and may 409 reflect thrombosis establishment as well<sup>[14]</sup>. To identify the location of high RRT, 410 Fig.8a~b draw the distribution of the normalized RRT to its maximum value in each 411 412 model. It shows that high RRT corresponds to the region with highly vortical and low velocity flow. This occurs at the FL proximal tip for both the patients (PI-2~5, 413 PII-2~5). Moreover, Fig.8ccompares the magnitude of RRT in PI-5 and PII-4, in 414 which the follow-up periods are similar (21 and 18months for PI and PII 415 respectively). The maximum RRT in PI-5 and PII-4 are 68.94 and 70.66Pa<sup>-1</sup> 416 respectively. The similar distribution pattern and magnitude of RRT in PI and PII 417 indicates that the values of RRT alone may not be able to predict FL remodeling. To 418 further investigate this, the variation of the maximum RRT is studied (Fig.8d). The 419 maximum RRTs in PI-3~7 and PII-3~5 are normalized to the correspondent first 420 421 follow-up results (PI-2 and PII-2, 7days post-TEVAR). The variation patterns of this normalized maximum RRT show significant difference between PI and PII. This 422 423 implicates that, post-TEVAR, variation of RRT in short- to middle-term follow-up (PI-2~5, PII-2~4) may play a key role in thrombosis establishment: an increase of 424 425 RRT after TEVAR and maintaining the relative normalized maximum RRT value to be above 1.0 (Fig.8d) may lead to positive FL remodeling. 426

Common morphological predictors for re-intervention or surgery after TEVAR 427 include aortic diameter >55mm and growth rate >10mm/year [39]. Hemodynamic 428 429 condition of the dissected aorta plays an important role in driving TL and FL remodeling. In other words, hemodynamic parameters may have the potential to 430 predict the dissection development earlier than morphological change. However, 431 hemodynamic markers that can possibly predict FL development post-TEVAR have 432 not been proposed yet, which would require long-term multiple follow-up analyses. 433 The current study investigated the correlation of hemodynamic parameters to the 434 development of post-TEVAR dissection. It preliminarily proposed the parameters that 435 are potential to differentiate the enlarged and stable FL in an early stage post-TEVAR. 436 Although this study was based on a limited number of patient cases and thus no 437

clinical conclusion can be drawn at this stage, it is the basis to future studies on a
larger amount of patient cases and would contribute to the research regarding early
decision-making on re-intervention or surgery for AoD after TEVAR.

441

## 442 LIMITATIONS

This study, based on long-term multiple follow-up data of two patients, preliminarily 443 shows the relationship of the variations between hemodynamic parameters and 444 luminal remodeling. However, critical values of these parameters should be better 445 determined by involving a greater number of patient cases. Besides, more detailed 446 mechanical analysis should involve the fluid-structure interaction analysis, which 447 448 does not only provide the stress information in the aortic wall but also offer more accurate results on the WSS. However, due to the complex geometry and the lack of 449 the actual material properties, the existed fluid-structure interaction studies on AoD 450 often generate the aortic/dissection wall with arbitrary thickness and assume the 451 mechanical properties of the aortic and dissection wall similar to the properties of 452 aortic aneurysms. More accurate simulations are highly dependent on accurate model 453 establishment and material property measurements, which are currently carried on in 454 our laboratory. 455

456

## 457 ACKNOWLEDGEMENTS

This study was supported by National Natural Science Foundation of China (81471752, 81353265), National Science and Technology Pillar Program of China (2015BAI04B03), and National Key Research and Development Program of China (2017YFC0107900). PW was partially supported by UK EPSRC (EP/N014642/1).

462

## 463 CONFLICTS OF INTEREST

464 No

465

## 466 ETHICAL APPROVAL

467 This study was approved by the Institutional Review Board of Chinese PLA General468 Hospital (ref no. 20110903.V1.1)

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#### **FIGURE LEGENDS**

- Fig.1 (a)~(d) display the reconstructed models of AoD (D-days, M-months); (e) is a sonogram of Doppler ultrasound velocimetry, the upper edge of which is shown in green; (f) shows an axial slice of CTA scan at initial presentation of PI, in which the segmented lumen boundary is shown in yellow.
- Fig.2 The luminal remodeling. (a) displays the measured axial positions (L1-4) and the regions to calculate luminal volume ( $V_{TL}$ ,  $V_{TL}$ -Part and  $V_{FL}$ ); (b)~(e) show the variation of luminal volume, aortic diameter, width of TL and FL respectively.
- **Fig.3** Flow patterns of AoD. (a)~(b) display the flow streamlines at systolic peak; (c) shows the streamlines at the proximal region of FL in the final model of PII.
- Fig.4 Flow exchange between the true and false lumen. (a) displays the variation of the mass flow rate towards the FL via the primary entry, AoT-2 and AoT-3 over a cardiac cycle at initial presentation; (b) and(c) respectively show the mass flow rate variation at AoT-2 and AoT-3 during the longitudinal follow-ups.
- Fig.5 Pressure distribution along the dissected aorta. (a) and (b) display the PDsys distribution in PI and PII respectively; (c) shows the PDsys in a normal aorta;
  (d) displays the variation of the maximum PDsys during the follow-up; (e) shows the Psys at a slice crossing the FL in the final two examinations of PI and in the final examination of PII; and (f) shows the region with Psys≥ 09.15mmHg (in red).
- **Fig.6** Pressure difference between FL and TL for PI (a)~(d) and PII (e)~(f).
- **Fig.7** WSS distribution along the flap. (a) and (b) display the WSS distribution at systolic peak on the flap of TL and FL side respectively; (c) and (d) show the WSS distribution on the flap of FL side during the long-term follow-ups in different WSS ranges.
- **Fig.8** RRT distribution and variation. (a) and (b) show the normalized RRT to its maximum value post-TEVAR in PI and PII respectively; (c) shows the RRT distribution at PI-5 and PII-4, where the follow-up periods in PI and PII are similar; and (d) displays the variation of the normalized maximum RRT to its value in the first follow-up (PI-2 for PI and PII-2 for PII).

# Hemodynamic Parameters That May Predict False-lumen Growth in Type-B Aortic Dissection after Endovascular Repair: A Preliminary Study on Long-term Multiple Follow-ups

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#### 1 ABSTRACT

Thoracic endovascular aortic repair (TEVAR) is commonly applied in type-B aortic 2 dissection. For patients with dissection affects descending aorta and extends downward to 3 involve abdominal aorta and possibly iliac arteries, false lumen (FL) expansion might occur 4 5 post-TEVAR. Predictions of dissection development may assist in medical decision on re-intervention or surgery. In this study, two patients are selected with similar morphological 6 features at initial presentation but with different long-term FL development post-TEVAR 7 (stable and enlarged FL). Patient-specific models are established for each of the follow-ups. 8 Flow boundaries and computational validations are obtained from Doppler ultrasound 9 velocimetry. By analyzing the hemodynamic parameters, the false-to-true luminal pressure 10 difference (PDiff) and particle relative residence time (RRT) are found related to FL 11 remodeling. It is found that (i) the position of the first FL flow entry is the watershed of 12 negative-and-positive PDiff and, in long-term follow-ups, and the position of largest PDiff is 13 consistent with that of the greatest increase of FL width; (ii) high RRT occurs at the FL 14 proximal tip and similar magnitude of RRT is found in both stable and enlarged cases; (iii) 15 comparing to the RRT at 7days post-TEVAR, an increase of RRT afterwards in short-term is 16 found in the stable case while a slight decrease of this parameter is found in the enlarged case, 17 indicating that the variation of RRT in short-term post-TEVAR might be potential to predict 18 long-term FL remodeling. 19

- 20
- 21 Key Words: aortic dissection; hemodynamics; endovascular procedures.

#### 22 INTRODUCTION

Aortic dissection (AoD) is a severe cardiovascular disease, where a surge of blood 23 flowing into the aortic wall via an initial tear or damage of the intima and splitting the 24 single aortic lumen into a true and false lumen (TL and FL). Stanford type-B AoD 25 26 indicates those with the dissection begins distal to the supraaortic branches. Interventional treatment of Stanford type-B AoD commonly involves thoracic 27 endovascular aortic repair (TEVAR)[1]. In a number of patients, FL expansion is 28 found post-TEVAR, especially in the infrarenal aorta. Recent study confirmed that 29 abdominal aortic expansion can be frequently found after TEVAR and is independent 30 from thoracic FL thrombosis[2]. Prediction of FL growth may contribute to early 31 decision-making of re-intervention or surgery. The post-TEVAR development of 32 dissection is highly dependent on local hemodynamics[3]. Medical imaging tools such 33 34 as Doppler ultrasound[4] and phase-contrast MR (pcMR)[5] are able to capture the flow velocity within aorta. However, the former provides velocity information at a 35 certain position of the vessel, and the latter reveals flow movement with relatively low 36 spatial and temporal resolution[6, 7]. On the other hand, the uptake of 37 <sup>18</sup>F-fluorodeoxyglucose in PET-CT can indicate complications in AoD[8] and positive 38 correlation between the uptake and wall shear stress is found in aortic aneurysm 39 study[9]; however, PET-CT is relatively expensive and the flow information cannot be 40 directly reported. Thus, computational simulations that can provide hemodynamic 41 parameters, such as flow, pressure and shear stress distributions, may enrich analysis. 42

Previous computational works focusing on type-B AoD include investigations on 43 hemodynamic features[10-14], luminal flow exchange[12, 15], post-TEVAR flow 44 45 effects[16-18], tear-induced flow effects[19, 20] and fluid-structure interaction studies[21, 22]. Besides, 4D pcMR[5, 23] and phantom[24, 25] measurements have been 46 47 conducted to compare with or validate the computed results. In this study, we investigate the flow-driven dissection development after TEVAR based on long-term 48 multiple follow-ups. Flow conditions in patients with stable and enlarged FL are 49 compared and key hemodynamic parameters that are related to dissection growth in 50 abdominal aorta are proposed, facilitating medical decision-making on post-TEVAR 51 treatment. 52

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#### 55 **METHODS**

#### 56 Image Acquisition and Model Reconstruction

This study was approved by the institutional review board of the Chinese PLA 57 General Hospital. Written informed consent was obtained from the patients involved 58 in this study. Two male patients (PI and PII) with subacute Stanford type-B AoD 59 underwent arterial-phase CT angiography (CTA) at initial presentation and during the 60 follow-up examinations after TEVAR via a dual-source CT scanner (SOMATOM 61 Definition Flash, SIEMENS, Germany). Details of the CTA scan and the patient 62 information are described in S1, Supporting Document. Image segmentation and 63 surface reconstruction of AoD were conducted through Mimics (Materialise, 64 65 Belgium). The cross-sectional contours of the reconstructed geometries were mapped back to CTA images to ensure that the 3D models present the actual outline of the 66 vessel lumen. Detailed views of the models are shown in Fig.1, where PI/II-1 67 indicates the models pre-TEVAR while PI/II-2 and others are models post-TEVAR. 68 69 After TEVAR, PI and PII have experienced six- (7days~53months) and four-times (7days~35months) CTA scans. The FL in PI was in stable condition (PI-2~7, Fig.1b) 70 71 while that in PII was expanding (PII-2~5, Fig.1d). The models were meshed in ICEM (ANSYS Inc, Canonsburg, USA) with tetrahedral elements in the core region and 72 73 prismatic cells (10 layers) in the boundary layer near the aortic wall. The grid resolution varies from 2,564,019 to 3,153,829 cells. 74

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# 76 Doppler Ultrasound and Boundary Conditions

77 Time-variant velocities at ascending aorta (AAo), brachiocephalic trunk (BT), left common carotid artery (LCCA), left subclavian artery (LSA) were measured via 78 79 Doppler ultrasound of the patients, and velocity variation at the distal thoracic aorta (DTAo, about 5cm above celiac trunk) was measured in the final examination of PI to 80 provide validation of the computational results. This is because the true lumen (TL) 81 remodeling at this position in PI is sufficient, so that relatively organized flow is 82 found and the central line of vessel can be accurately identified. The velocity of AAo 83 was measured through the apical 5-chamber view and the suprasternal long axis view 84 85 of aortic arch. The two results were compared to ensure the maximum velocity at AAo could be captured. For other arteries (BT, LCCA, LSA and DTAo), Doppler 86

velocimetry has been conducted at the proximal and distal sites of the targeted measurement vessel. Mean velocity over a cardiac cycle at the two sites for each particular vessel was then calculated and compared. When the difference between them is less than 5%, the measured velocity is considered effective. Details of the measurement are described in S2, Supporting Document.

The upper edge of the velocity sonogram was extracted (Fig.1e) as the variation of 92 the maximum velocity at the measured site. The flow rates at AAo, BT, LCCA and 93 LSA, as the velocity boundary conditions, can then be calculated based on the 94 95 measured time-variant maximum velocity and the assumed flat flow profile for AAo and parabolic flow profile for the others. The velocity boundary conditions of the 96 models is shown in Fig.S1a, Supporting Document. Pulsatile waveforms of the 97 pressure at celiac artery (CA), superiormesenteric artery (SMA), renal arteries and the 98 outlets at common iliac arteries were obtained from previous study[26] (Fig.S1b, 99 Supporting Document). As shown in Fig.1, two models in PII (PII-1 and PII-4) were 100 cropped below the iliac bifurcation due to the relatively shorter CT scanning range. To 101 eliminate the outlet effects, time-variant pressure distribution at the cropping plane 102 has been calculated in PII-2~3 and PII-5. The averaged pressure information at this 103 104 cropping plane was mapped to PII-1 and PII-4, serving as the pulsatile pressure outlets. 105

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## 107 Numerical Models

The vessel wall was assumed as no-slip and rigid, due to low distensibility in 108 long-term follow-ups[27]. The blood was treated as Newtonian and incompressible 109 with density of 1044kg/m<sup>3</sup> and dynamic viscosity of 0.00365kg·m<sup>-1</sup>·s<sup>-1</sup>[28]. The 110 average Reynolds number over a cardiac cycle, calculated based on the equivalent 111 diameter  $(D_e = 2\sqrt{Area/\pi})$  and velocity at the inlet of the ascending aorta in PI and 112 PII, were between 2066-2197 and 2844-2960, respectively. Our previous study 113 confirmed that laminar simulations with adequately fine mesh resolutions, especially 114 refined near the walls, can capture flow patterns as turbulence model[15]. To further 115 confirm this, we solved the flow in the first follow-up cases (PI-2 and PII-2) by both 116 laminar and k-w SST turbulence models, where the flow in the abdominal aortic 117 region is the fastest during the follow-ups. Similar flow patterns were found and the 118 discrepancy of the maximum velocity and wall shear stress (WSS) in the abdominal 119

aorta was 4.2% and 5.2% respectively (the laminar and turbulent results of PII-2 were 120 shown in Fig.S2, Supporting Document), ensuring the rationality of laminar model to 121 be applied in the current problem. A finite volume solver, CFD-ACE+ (ESI Group, 122 France) was employed. The heart-beat cycle of the patients was measured at each 123 clinical examination. The averaged cardiac cycle for PI and PII were 71 and 124 69beat/min respectively. Temporal discretization of numerical models was assigned as 125 45step/cycle. Simulation was carried out for 5 cardiac cycles to achieve a periodic 126 solution and results of the final cycle were presented. Grid and temporal 127 independency analyses on finer grids and finer temporal discretizations were 128 conducted to ensure the base resolution with the base time step settings are adequate 129 (S3, Supporting Document). 130

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## 132 **RESULTS**

#### 133 Aortic Remodeling

Aortic remodeling was assessed by: (i) size and numbers of the aortic tears (AoT), (ii) 134 change of luminal volume, and (iii) growth of aortic diameter. There are four major 135 136 AoTs along the aorta for both patients at initial presentation. The primary entry (AoT-1) in both patients is located at the proximal region of descending aorta and the 137 locations of other AoTs (AoT-2~4) are displayed in Fig.1. After TEVAR, the primary 138 entry was sealed and after the first follow-up of PI, AoT-2 was disappeared. 139 140 Considering the position and local aortic curvature of the tears, AoT-4 is the exit of the flow from FL to TL, while the function of AoT-2 and AoT-3 is uncertain. Since 141 only the flow entries towards the FL is able to bring mechanical impact into the 142 dissection, size measurement was only performed on AoT-2 and AoT-3 (Table 1). 143 Post-TEVAR, both AoT-2 and AoT-3 in PI and PII are enlarged, probably due to the 144 greater flow impact on these sites after AoT-1 was sealed. 145

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Tear	Geometry	PI-1	PI-2	PI-3	PI-4	PI-5	PI-6	PI-7
AoT-2	H[mm]	7.87	7.15	-	-	-	-	-
	W[mm]	8.97	10.76	-	-	-	-	-
	$A[mm^2]$	64.61	70.71	-	-	-	-	-
AoT-3	H[mm]	13.86	18.79	17.93	15.31	18.23	8.81	8.57
	W[mm]	10.04	9.38	6.03	8.90	7.30	6.72	5.75
	$A[mm^2]$	116.13	150.82	106.40	119.19	128.82	93.49	44.09
Tear	Geometry	PII-1	PII-2	PII-3	PII-4	PII-5		
AoT-2	H[mm]	8.17	8.41	5.74	11.04	11.20	-	
	W[mm]	9.34	9.96	7.36	9.59	6.03		
	$A[mm^2]$	62.61	70.47	37.33	89.50	54.05		
AoT-3	H[mm]	9.91	9.55	7.77	6.25	6.19	-	
	W[mm]	10.01	11.47	9.34	8.04	9.90		
	$A[mm^2]$	85.27	99.32	61.51	40.35	53.32	_	
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Table 1. The size of the aortic tears in the patients with long-term follow-ups

H-Height; W-Width; A-Area.

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The change of luminal volume is shown in Fig.2. The volume of TL  $(V_{TL})$  involves 147 the initial dissection-affected TL region, while V<sub>TL-Part</sub> indicates the 148 dissection-affected TL region during the follow-ups. Variations of V<sub>TL</sub>, volume of FL 149 150  $(V_{FL})$  and the ratio between  $V_{TL-Part}$  and  $V_{FL}$  are displayed in Fig.2b.  $V_{FL}$  in the stable case (PI) reduces gradually post-TEVAR, except for the final examination, where a 151 trivial increase of  $V_{FL}$  is found (131.6mm<sup>3</sup> in 20months). However,  $V_{FL}$  in the 152 expanding case (PII) reduces in the first two follow-ups but enlarges afterwards, 153 where the V<sub>FL</sub> growths in PII-4 and PII-5 are 2,138mm<sup>3</sup> and 34,989mm<sup>3</sup> respectively, 154 155 the durations of which are both 17months (Fig.2b). For both cases, the TL-to-FL volume ratio (V<sub>TL-Part</sub>/V<sub>FL</sub>) keeps increasing, except for PI-6 and PII-3 (pointed by 156 arrows). This is because significant FL regression occurs at the proximal region (blue 157 squares in Fig.1b,1d), which shortens the comparison region. Since TL remodeling is 158 insufficient in this region, value of V<sub>TL-Part</sub>/V<sub>FL</sub> reduces. The averaged increase rates of 159 V<sub>TL-Part</sub>/V<sub>FL</sub> in the follow-ups of PI and PII are 1.8% and 0.3% per month, 160 respectively, indicating a significant luminal remodeling difference. 161

The aortic diameter (D) is measured in each model, taking into account the width of 162 TL and FL ( $W_{TL}$  and  $W_{FL}$ ) and the thickness of flap. As shown in Fig.1f, 163 measurements have been conducted in the axial plane of the CTA datasets, along a 164 line that crosses the centre of the flap and is perpendicular to the flap. Four 165 measurement positions are selected in the abdominal aorta. As shown in Fig.2a, they 166 are 1cm below diaphragm (L1), 1cm above CA (L2), 3.5cm below SMA (L3) and 9cm 167 below SMA (L4). Fig.2c~e display the variation of D, W<sub>TL</sub> and W<sub>FL</sub> respectively. 168 Positive TL remodeling is found in PI and PII in general, except for L3 and L4 in PI-3 169

and L3 in PII-4~5.  $W_{FL}$  at the proximal region of PI (L1 and L2) reduces greatly during the follow-up, while in the distal region (L3 and L4), variation of  $W_{FL}$  is stable. In PII,  $W_{FL}$  increases sharply since the second follow-up (PII-3). Comparing the variation of  $W_{TL}$ ,  $W_{FL}$  and D, it can be found that the variation pattern of D is mainly determined by the variation of  $W_{FL}$ .

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# 176 Flow Pattern, Validation and Luminal Flow Exchange

Fig.3 shows the flow streamlines at systolic peak where the color map of velocity 177 magnitude is restricted to a certain range in PI ( $0 \sim 2.51$  m/s) and PII ( $0 \sim 3.78$  m/s) to 178 assist visualization of the longitudinal velocity variation. At systolic peak, fast and 179 180 organized flow is found in TL; while vortical and relatively slow flow presents in FL. Highest velocity presents in the short-term follow-ups (PI-2~3, 4months for PI; 181 PII-2~3, 1months for PII) at the region below stentgraft and above AoT-2 and -3, 182 where TL remodeling is insufficient and the blood has not been diverted. In long-term 183 follow-ups (PI-4~7 and PII-4~5), the flow in the proximal FL is generally slow 184 (<0.5mm/s); while below AoT-2 (the first re-entry), where blood perfusion occurs, 185 faster flow up to 1.53m/s in PI and 2.22m/s in PII are found in the FL with helical 186 feature (Fig.3c). 187

To validate the computational results, Doppler ultrasound velocimetry was 188 performed at distal thoracic aorta (5cm above CA) along the centerline of TL. This 189 was only conducted at the final examination of PI, because its TL in the distal thoracic 190 aorta is fully remodeled and the local flow is therefore organized (Fig.3a). The 191 measured and computed velocities at the centre of the cross-section of the vessel, 5cm 192 above CA, are 1.58m/s and 1.66m/s respectively, indicating a difference of 5.1%. 193 Moreover, the computed velocity variation pattern is similar to the measured one 194 (Fig.S3, Supporting Document), ensuring the rationality of computational results. 195

Fig.4 displays the luminal flow exchange via the primary entry for initial presentation and that via AoT-2 and AoT-3 during the follow-ups. As abovementioned, AoT-4 functions as the outlet of FL throughout the cardiac cycle, thus it is not displayed in Fig.4. As shown in Fig.4a (positive values indicates flow enters FL), before treatment, the TL-to-FL flow exchange in PI and PII presents similar pattern: the primary entry serves as the main flow inlet of FL; AoT-2 functions as the outlet of FL in the initial part of systole and behaves as inlet in the rest part of

cardiac cycle; and AoT-3 mainly serves as the outlet of FL. There are 31.48% and 203 40.00% of the total flow diverted to FL over a cardiac cycle for PI and PII 204 respectively, at initial presentation. After TEVAR, the function of the tears changes. In 205 PI, AoT-2 becomes the inlet of FL in the entire heart-beat cycle (Fig.4b), while AoT-3 206 functions as the inlet of FL during systole and serves as the outlet in diastole (Fig.4c). 207 Since the second follow-up of PI, AoT-2 is closed and AoT-3 gradually becomes the 208 inlet of FL throughout the entire cardiac cycle. The flow entering FL are 16.67%, 209 6.98%, 8.24%, 9.59%, 5.43%, 3.09% of the total flow for case PI-2~7 respectively, 210 211 the variation of which is consistent to the variation of tear size (Table 1) and it is generally reducing. In PII, both of AoT-2 and AoT-3 are existed during the follow-ups. 212 AoT-2 mainly serves as the inlet of FL; especially in model PII-4 and PII-5, positive 213 values of mass flow rate are found in the entire cardiac cycle (Fig.4b). AoT-3 diverts a 214 small amount of flow into the FL during systole and mainly behaves as the outlet of 215 FL (Fig.4c). There are 5.22%, 3.06%, 14.76% and 10.23% of the total flow diverted 216 into FL in case PII-2~5 respectively, the variation of which is also consistent with the 217 variation of tear size. 218

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# 220 Loading Distribution along the Aorta

Fig.5a~b display the pressure drop at systolic peak (PDsys, the difference between the 221 local pressure and the pressure at the bottom of the model). In TL, PDsys reduces 222 from the inlet of AAo to the outlets of common iliac arteries. At initial presentation 223 (PI-1 and PII-1), FL diverts part of the total flow (31.48% for PI and 40.00% for PII), 224 maintaining PDsys in relatively low values. After TEVAR, in short-term follow-ups 225 (PI-2~3 and PII-2~3), proximal descending aorta is supported by stentgraft and 226 reshaped; however, below the endograft, TL remains collapsed, inducing larger 227 resistance and resulting in higher velocity and regional lower pressure in this region 228 (indicated by arrows in Fig.5a~b). In long-term follow-ups (PI-4~7 and PII-4~5), TL 229 remodeling in distal thoracic aorta is improved; PDsys thus gradually reduces along 230 aorta. In contrary, PDsys in FL is relatively uniform, which is probably due to the 231 232 higher energy exchange induced by the vortical flow. Fig.5d shows the variation of the maximum PDsys during the follow-ups. For both patients, the maximum PDsys 233 increases greatly soon after TEVAR (7days); along with the progress of TL 234 remodeling, the maximum PDsys reduces. The PDsys in normal aorta model (Fig.5c) 235

based on the same boundary conditions as PI and PII is also computed and the
averaged maximum PDsys is shown by the dash line in Fig.5d. Since the growth of
FL in the two patients shows obvious difference, the similar variation pattern of
PDsys indicates that the pressure drop is mainly affected by TL remodeling.

The mean arterial pressure drop (PDmean) from AAo to the common iliac arteries during a cardiac cycle is also calculated. It is 6.29mmHg for the initial model of PI and 4.70mmHg for the final model (53months post-TEVAR), which is slightly higher than that in normal aorta for the studied segment (4mmHg[29]). The PDmean of PII at the initial and final (35months post-TEVAR) models are 6.53mmHg and 6.07mmHg, respectively, consistent with its insufficient TL remodeling.

In the final examination of PI (PI-7), V<sub>FL</sub> is slightly increased, D and W<sub>FL</sub> also 246 increase at L3 (Fig.2); however, the flow exchange and tear size at this stage are both 247 reduced. Fig.5e displays the pressure distribution at systolic peak (Psys) on a slice of 248 FL for PI-6~7. It shows that the smaller tear size reduces the flow entering FL, 249 however, it increases the velocity and induces higher pressure impact on the outer 250 wall of FL (indicated by arrows in Fig.5e). The highest pressure of the impact region 251 in PI and PII at final examination are 114.37 and 115.41mmHg respectively. Besides, 252 253 by blanking the region with Psys<109.15mmHg, which is the 95% of the averaged maximum pressure in PI and PII at final examination, Fig.5f shows that the high 254 255 pressure region (in red) in the FL of PII is much larger than that in PI. In fact, the FL growth rates of PI and PII at the final examination are 0.22mm<sup>3</sup>/day and 256 68.61mm<sup>3</sup>/day respectively. 257

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## 259 Luminal Pressure Difference, Wall Shear Stress and Relative Residence Time

To investigate the pressure difference (PDiff) between TL and FL, a series of slices 260 that are perpendicular to the centerline of TL are extracted and the net pressures in TL 261 (P<sub>TL</sub>) and FL (P<sub>FL</sub>) on each slice over a cardiac cycle are calculated. Fig.6 displays 262 PDiff (PDiff=P<sub>FL</sub>-P<sub>TL</sub>) in each model. In both patients, pre-TEVAR (Fig.6a,6e), P<sub>FL</sub> is 263 smaller than P<sub>TL</sub> at the level above AoT-2; while, below AoT-2, P<sub>FL</sub> is larger than P<sub>TL</sub>, 264 265 pushing the FL towards TL. Post- TEVAR, the primary entry is closed. In PI, negative values of PDiff are found above AoT-3 (Fig.6b~c), indicating higher P<sub>TL</sub> presents in 266 this region, which supports the aortic wall and assists in TL expansion. However, in 267 the region below AoT-3 in PI, due to the blood perfusion into the FL, positive values 268

of PDiff present, indicating FL propulsion on the TL. In the final follow-up of PI 269 (Fig.6d), the maximum value of PDiff is 0.22mmHg. This maximum PDiff occurs 270 close to L3. At this position,  $W_{FL}$  and  $W_{TL}$  increase by 1.02mm and 0.46mm 271 respectively (Fig.2), indicating this small value of PDiff (0.22mmHg) is not large 272 enough to squeeze TL but the local pressure in FL (Fig.5e) is able to induce further 273 FL expansion. On the other hand, in PII, post-TEVAR, AoT-2 becomes the main inlet 274 of the flow into FL. Higher above this tear, P<sub>TL</sub> is larger than P<sub>FL</sub>, supporting TL 275 expansion (Fig.6e~f). In the near region above AoT-2, TL remodeling is insufficient 276 277 for the short-term follow-ups but PFL gradually increases; thus, regional positive PDiff is found in PII-3 above AoT-2 (arrow indicated in Fig.6f). Below AoT-2, positive 278 values of PDiff dominate and the maximum PDiff in the final examination of PII is 279 280 1.39mmHg, more than 6-times as high as that in PI, occurring close to L3. Taking account of that, in PII-5, W<sub>TL</sub> decreases yet W<sub>FL</sub> increases at L3 (Fig.2d~e), this high 281 282 PDiff propels TL collapse and the local pressure pushes FL growth.

Fig.7 shows the wall shear stress (WSS) distribution on the flap at systolic peak. Its 283 284 variation over a cardiac cycle is shown in Video S1-6, Supporting Document. In both patients, the WSS on the flap at the FL side (WSS<sub>FL</sub>) is significantly smaller than that 285 286 on the TL side (WSS<sub>TL</sub>); and the WSS<sub>TL</sub> in PII is generally higher than that in PI. At initial presentation, the maximum WSS<sub>TL</sub> occurs at the edge of primary entry; while 287 the maximum WSS<sub>FL</sub> occurs at the proximal descending aorta along the side opposite 288 to the primary entry (indicated by arrows in Fig.7b). In follow-ups, the maximum 289 WSS<sub>TL</sub> occurs at the region where TL presents most collapse; while the maximum 290 WSS<sub>FL</sub> occurs at the edge of the tears. AoT-3 and AoT-2 are the main FL flow entries 291 for PI and PII in follow-ups, which induces helical flow in the downstream and high 292 WSS<sub>FL</sub> on the side opposite to the tears (indicated by hollow arrows in Fig.7b). To 293 further visualize the variation of WSS<sub>FL</sub> at different regions along the flap on the FL 294 side, the color map is assigned to 0~5Pa (Fig.7c) and 0~0.5Pa (Fig.7d) to show the 295 distal and proximal region respectively. In PI, WSS<sub>FL</sub> at the region below AoT-3 296 reduces obviously from PI-4 to PI-6; however, in the final examination, it is slightly 297 increased, although the maximum WSS<sub>FL</sub> in this region is still smaller than 1Pa. In 298 PII, WSS<sub>FL</sub> presents obvious increase in PII-4 and reduces slightly in PII-5; the 299 maximum WSS<sub>FL</sub> at the region between AoT-2 and AoT-3 in the final examination is 300 5.21Pa. 301

Particle residence time is proposed to be related to thrombosis establishment[14].

The relative residence time (RRT), which is based on the time-averaged WSS 303 (TAWSS) and oscillatory shear index (OSI), [(1-2·OSI)·TAWSS]<sup>-1</sup>, reflects particle 304 residence time and thus may correspond with the region of thrombosis[14, 30]. 305 Fig.8a~b show the RRT distributions (normalized by the maximum RRT value) in PI 306 and PII post-TEVAR, respectively. In PI, the highest RRT occurs at the FL proximal 307 tip in PI-2~5; while in PI-6~7, RRT is greatly decreased and the highest RRT no 308 longer occurs at the FL tip. In PII, the highest RRT also occurs at the FL proximal tip. 309 Fig.8d displays the variation of the maximum RRT (normalized by the maximum 310 RRT at the first follow-up, i.e. PI-2 and PII-2 for PI and PII respectively). The RRT 311 variation in PI and PII shows significant difference during short-term follow-up: the 312 normalized maximum RRT in PI increases greatly in PI-3 (from 1 to 5.516) and 313 gradually decreases afterwards to 0.00405 in PI-7; while in PII, it decreases in PII-3 314 (from 1 to 0.025) and then maintains between 0.00324-0.319. The difference in the 315 maximum RRT's variation pattern shows potential to predict different FL remodeling 316 in the two patients. 317

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#### 319 **DISCUSSION**

Thoracic endografts, aiming to seal the primary entry and diverting blood flow into 320 TL, are increasingly used in treating type-B aortic dissection[31]. Ideally, this 321 approach would lead to thrombosis establishment in the FL and morphologic change 322 in TL, to stabilize the aorta and consequently reduce aorta-related death. Previous 323 studies confirmed the favorable results of TEVAR; however, also reported FL 324 expansion on the segment distal to endografts, usually in the infrarenal aorta[32, 33]. 325 This is related to the patency of FL[33] or, in other words, it is related to the flow field 326 and hemodynamic conditions in AoD. Current literatures provide little information 327 concerning the fate of the abdominal aorta post-TEVAR and, to our knowledge, 328 computational studies on long-term multiple follow-up cases that are able to report 329 the change of hemodynamic parameters have been lacking. Therefore, in the current 330 study, we investigate two patients showing similar physical and hemodynamic 331 features at initial presentation but presenting different FL development (stable in PI 332 and expanded in PII) during the follow-ups. We preliminarily identify the possible 333 hemodynamic parameters that could help to evaluate/predict FL enlargement and 334 thrombosis formation. In this study, the variation trends of the hemodynamic 335

parameters are more important than their absolute values.

The mechanical load acting on the FL boundary, which includes the outer wall of 337 FL and the flap, induces FL enlargement and at the same time restricts TL 338 remodeling. The pressure that is normally applied on the FL wall plays a key role in 339 luminal remodeling[34] and the local flow directly relates to its distribution and 340 magnitude. The flow can be assessed by its amount and the velocity. The first is 341 mainly related to the size of tears. By comparing Fig.4b~c and Table 1, it can be 342 found that the absolute flow mass passes each tear per second has positive relationship 343 344 with the size of the tear. This can be shown on AoT-3 of PI and AoT-2~3 of PII, since they have multiple follow-up data, and it is consistent with previous report[14]. The 345 second, velocity, is determined by both the flow rate as well as the tear size; a smaller 346 tear size is correspondent to smaller flow rate yet accelerates the flow at the same 347 time. For instance, in the final two examinations of PI (PI-6 and PI-7), the amount of 348 the blood entering FL via AoT-3 decreases from 5.43% to 3.09% of the total flow and 349 the tear size decreases from 93.49mm<sup>2</sup> to 44.09mm<sup>2</sup>. However, the flow passing this 350 tear has been accelerated and induces stronger impact on the FL outer wall (Fig.5e). If 351 studying a longer period (PI-4 $\sim$ 7), the variation of W<sub>FL</sub> at L3 (Fig.2e), where just 352 353 below AoT-3 in PI, is similar to the variation of the ratio between the diverted flow amount and the tear size. 354

Along the aorta, the pressure in TL decreases generally but that in FL, due to the 355 vortical flow and its higher energy exchange, it does not present significant spatial 356 difference. The AoTs, functioning as the bridge between TL and FL, transport blood 357 flow and also pressure gradient. This induces similar pressure in the TL and FL near 358 the tears. Because of the relatively uniform  $P_{FL}$  and its connection with  $P_{TL}$  at the 359 tears, in general, P<sub>FL</sub> is smaller than P<sub>TL</sub> in the proximal region (above the AoTs) and 360 higher than P<sub>TL</sub> in the downstream. This general distribution feature is shown in Fig.5 361 in all of the post-interventional cases at the moment of systolic peak and similar 362 patterns can be found in other time steps of the cardiac cycle. 363

At the flap,  $P_{FL}$  and  $P_{TL}$  conflict each other; the difference between them (PDiff) may be associated with subsequent luminal remodeling[14]. Luminal remodeling is a long-term effect; investigation of PDiff in short-term follow-ups may show the variation trends of lumen remodeling, while PDiff in long-term follow-ups may be consistent to lumen remodeling results. Indeed, taking L3 as an example, in short-term (PI-2~3, PII-2~3), PD in PII increases from 1.44 to 2.93mmHg (7days-1month);

while, PDiff in PI remains about 0.55mmHg (7days-4month). This, earlier than 370 luminal change (Fig.2), shows the potential of FL enlargement for PII. In long-term 371 follow-ups (PI-4~7, PII-4~5), AoT-3 becomes the main FL flow entry in PI and 372 AoT-2 is the main entry in PII. PDiff increases from negative to positive slightly 373 below the position of AoT-3 in PI and AoT-2 in PII. Moreover, in both patients, in the 374 final examination, the maximum PDiff occurs near L3, where W<sub>FL</sub> increases the most 375 (among the compared four positions) and W<sub>TL</sub> in PII decreases. The abovementioned 376 indicates: (i) in short-term follow-ups, great increase of PDiff may imply potential FL 377 expansion; (ii) while, in long-term follow-up, the position of the first flow entry of FL 378 is the negative-to-positive watershed of PDiff, the position of the maximum PDiff is 379 consistent with the greatest W<sub>FL</sub> increase, and when the maximum PDiff is small 380 (0.22mmHg in PI-7), the pressure induces slight FL expansion without restricted acts 381 on TL, but when it is relatively large (1.39mmHg in PII-5), both FL expansion and TL 382 collapse are found. In clinical examinations, monitoring PDiff at early-stage and 383 identifying the position of FL entries and the position of largest PDiff may assist in 384 385 wisely control of the untreated aorta segment.

The WSS is related to the formation of thrombosis. Previous studies suggested that 386 the tearing of the aortic wall and high WSS in the near region of the tears could 387 promote initial activation of platelets as well as the formation of platelet aggregates; 388 while, the highly vortical flow pattern in the FL corresponding with low WSS 389 promotes platelet aggregation and deposition, so that leads to surface thrombosis [35, 390 36]. In other words, lower WSS may induce surface thrombus and thus lead to 391 constructive FL remodeling [37, 38]. In the long-term follow-ups (PI-4~5 and 392 PII-4~5), complete thrombosis is found at the proximal region of dissection and 393 partial thrombosis remains above the re-entries. The partial thrombosis in both PI and 394 PII is aligned with the intimal flap; thus, WSS on the flap along the TL and FL sides 395 are compared. The WSS<sub>FL</sub> is significantly lower than WSS<sub>TL</sub> throughout the cardiac 396 cycle; the low  $WSS_{FL}$  possibly induces surface thrombus along the flap in FL while 397 the high WSS<sub>TL</sub> can keep the TL patent. The lowest WSS<sub>FL</sub> (<0.25Pa) occurs at the 398 proximal region of FL (Fig.7d), implying potential thrombosis in these regions. 399 Indeed, partial thrombosis in PI-4~5 with very low WSS<sub>FL</sub> turns to complete 400 thrombosis in PI-6~7 (indicated by arrow inFig.7d). Moreover, in PII, the WSS<sub>FL</sub> at 401 the tip of the flap in PII-3 is small (<0.25Pa). In its next follow-up (PII-4), growth of 402 partial thrombosis can be found (indicated by arrow in Fig.7d). However, slightly 403

higher WSS<sub>FL</sub> is shown in PII-4 at the proximal tip, and in its next follow-up (PII-5),
the partial thrombosis is slightly reduced (indicated by hollow arrow in Fig.7d). This
indicates the surface thrombosis is possibly very sensitive to WSS, and during the
long-term recovery, FL regression/growth could be repeatedly occurred at the
proximal tip of the FL.

The derived parameter of shear stress - RRT is related to particle residence and may 409 reflect thrombosis establishment as well<sup>[14]</sup>. To identify the location of high RRT, 410 Fig.8a~b draw the distribution of the normalized RRT to its maximum value in each 411 412 model. It shows that high RRT corresponds to the region with highly vortical and low velocity flow. This occurs at the FL proximal tip for both the patients (PI-2~5, 413 PII-2~5). Moreover, Fig.8ccompares the magnitude of RRT in PI-5 and PII-4, in 414 which the follow-up periods are similar (21 and 18months for PI and PII 415 respectively). The maximum RRT in PI-5 and PII-4 are 68.94 and 70.66Pa<sup>-1</sup> 416 respectively. The similar distribution pattern and magnitude of RRT in PI and PII 417 indicates that the values of RRT alone may not be able to predict FL remodeling. To 418 further investigate this, the variation of the maximum RRT is studied (Fig.8d). The 419 maximum RRTs in PI-3~7 and PII-3~5 are normalized to the correspondent first 420 421 follow-up results (PI-2 and PII-2, 7days post-TEVAR). The variation patterns of this normalized maximum RRT show significant difference between PI and PII. This 422 423 implicates that, post-TEVAR, variation of RRT in short- to middle-term follow-up (PI-2~5, PII-2~4) may play a key role in thrombosis establishment: an increase of 424 425 RRT after TEVAR and maintaining the relative normalized maximum RRT value to be above 1.0 (Fig.8d) may lead to positive FL remodeling. 426

427 Common morphological predictors for re-intervention or surgery after TEVAR include aortic diameter >55mm and growth rate >10mm/year [39]. Hemodynamic 428 429 condition of the dissected aorta plays an important role in driving TL and FL remodeling. In other words, hemodynamic parameters may have the potential to 430 predict the dissection development earlier than morphological change. However, 431 hemodynamic markers that can possibly predict FL development post-TEVAR have 432 not been proposed yet, which would require long-term multiple follow-up analyses. 433 The current study investigated the correlation of hemodynamic parameters to the 434 development of post-TEVAR dissection. It preliminarily proposed the parameters that 435 are potential to differentiate the enlarged and stable FL in an early stage post-TEVAR. 436 Although this study was based on a limited number of patient cases and thus no 437

clinical conclusion can be drawn at this stage, it is the basis to future studies on a
larger amount of patient cases and would contribute to the research regarding early
decision-making on re-intervention or surgery for AoD after TEVAR.

441

## 442 LIMITATIONS

This study, based on long-term multiple follow-up data of two patients, preliminarily 443 shows the relationship of the variations between hemodynamic parameters and 444 luminal remodeling. However, critical values of these parameters should be better 445 determined by involving a greater number of patient cases. Besides, more detailed 446 mechanical analysis should involve the fluid-structure interaction analysis, which 447 448 does not only provide the stress information in the aortic wall but also offer more accurate results on the WSS. However, due to the complex geometry and the lack of 449 the actual material properties, the existed fluid-structure interaction studies on AoD 450 often generate the aortic/dissection wall with arbitrary thickness and assume the 451 mechanical properties of the aortic and dissection wall similar to the properties of 452 aortic aneurysms. More accurate simulations are highly dependent on accurate model 453 establishment and material property measurements, which are currently carried on in 454 our laboratory. 455

456

## 457 ACKNOWLEDGEMENTS

This study was supported by National Natural Science Foundation of China (81471752, 81353265), National Science and Technology Pillar Program of China (2015BAI04B03), and National Key Research and Development Program of China (2017YFC0107900). PW was partially supported by UK EPSRC (EP/N014642/1).

462

## 463 CONFLICTS OF INTEREST

464 No

465

#### 466 ETHICAL APPROVAL

467 This study was approved by the Institutional Review Board of Chinese PLA General468 Hospital (ref no. 20110903.V1.1)

469

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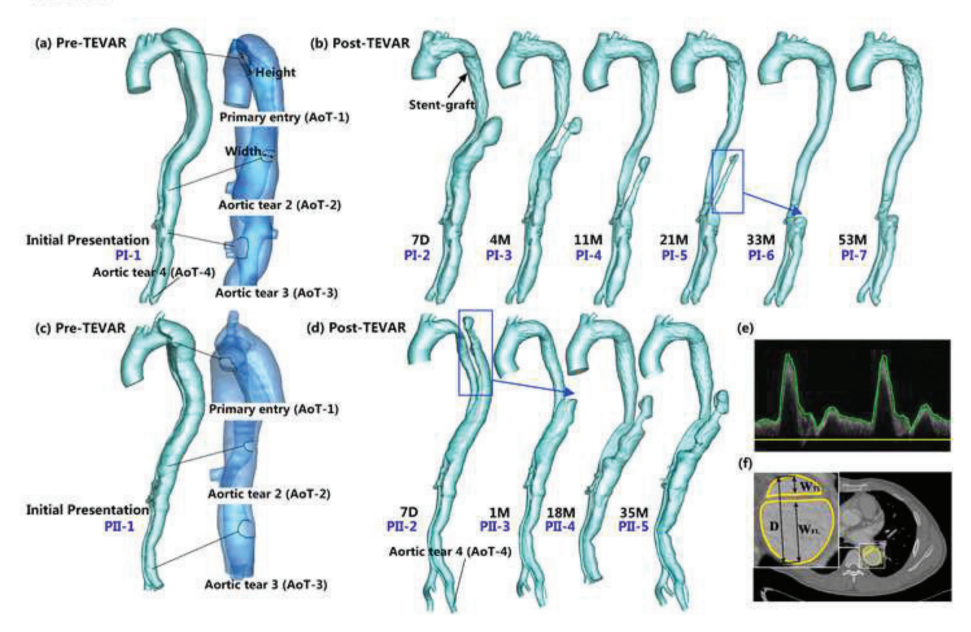
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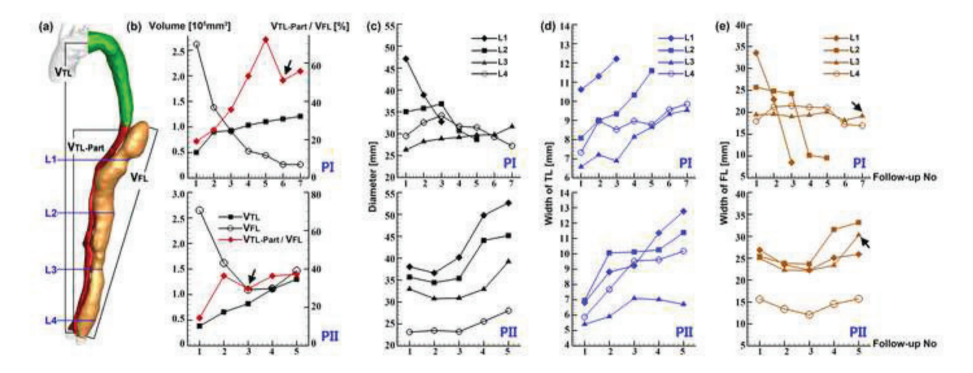
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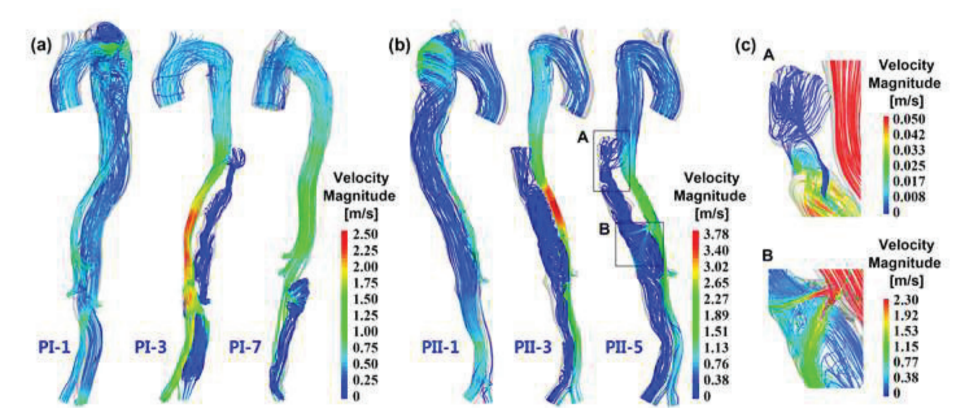
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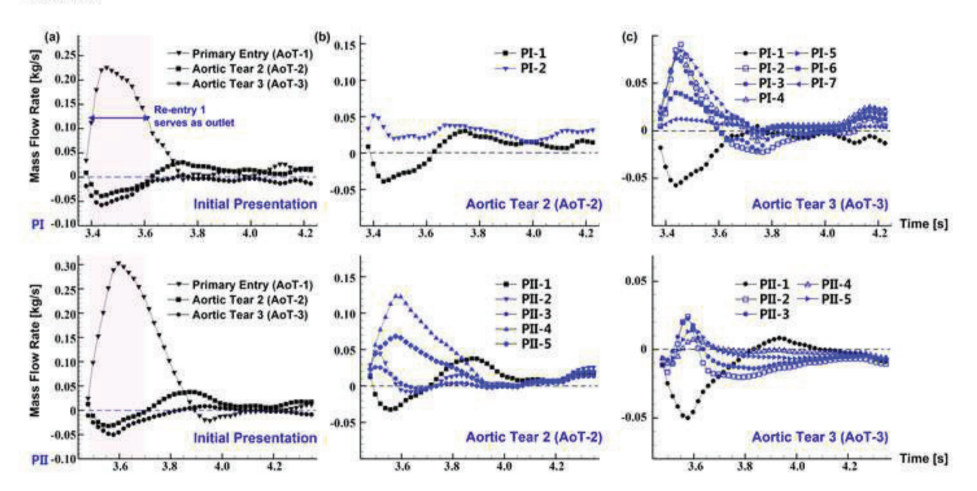
#### **FIGURE LEGENDS**

- Fig.1 (a)~(d) display the reconstructed models of AoD (D-days, M-months); (e) is a sonogram of Doppler ultrasound velocimetry, the upper edge of which is shown in green; (f) shows an axial slice of CTA scan at initial presentation of PI, in which the segmented lumen boundary is shown in yellow.
- Fig.2 The luminal remodeling. (a) displays the measured axial positions (L1-4) and the regions to calculate luminal volume ( $V_{TL}$ ,  $V_{TL}$ -Part and  $V_{FL}$ ); (b)~(e) show the variation of luminal volume, aortic diameter, width of TL and FL respectively.
- **Fig.3** Flow patterns of AoD. (a)~(b) display the flow streamlines at systolic peak; (c) shows the streamlines at the proximal region of FL in the final model of PII.
- Fig.4 Flow exchange between the true and false lumen. (a) displays the variation of the mass flow rate towards the FL via the primary entry, AoT-2 and AoT-3 over a cardiac cycle at initial presentation; (b) and(c) respectively show the mass flow rate variation at AoT-2 and AoT-3 during the longitudinal follow-ups.
- Fig.5 Pressure distribution along the dissected aorta. (a) and (b) display the PDsys distribution in PI and PII respectively; (c) shows the PDsys in a normal aorta;
  (d) displays the variation of the maximum PDsys during the follow-up; (e) shows the Psys at a slice crossing the FL in the final two examinations of PI and in the final examination of PII; and (f) shows the region with Psys≥ 09.15mmHg (in red).
- **Fig.6** Pressure difference between FL and TL for PI (a)~(d) and PII (e)~(f).
- **Fig.7** WSS distribution along the flap. (a) and (b) display the WSS distribution at systolic peak on the flap of TL and FL side respectively; (c) and (d) show the WSS distribution on the flap of FL side during the long-term follow-ups in different WSS ranges.
- **Fig.8** RRT distribution and variation. (a) and (b) show the normalized RRT to its maximum value post-TEVAR in PI and PII respectively; (c) shows the RRT distribution at PI-5 and PII-4, where the follow-up periods in PI and PII are similar; and (d) displays the variation of the normalized maximum RRT to its value in the first follow-up (PI-2 for PI and PII-2 for PII).

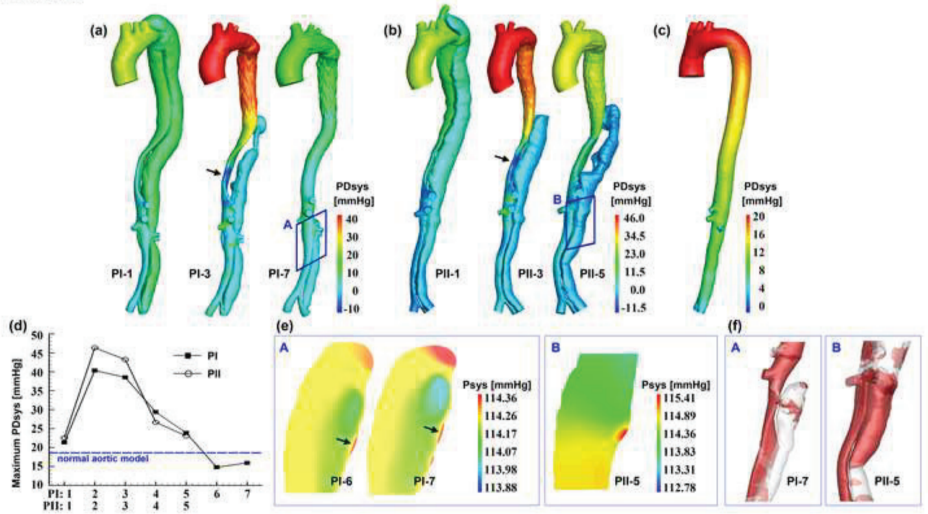


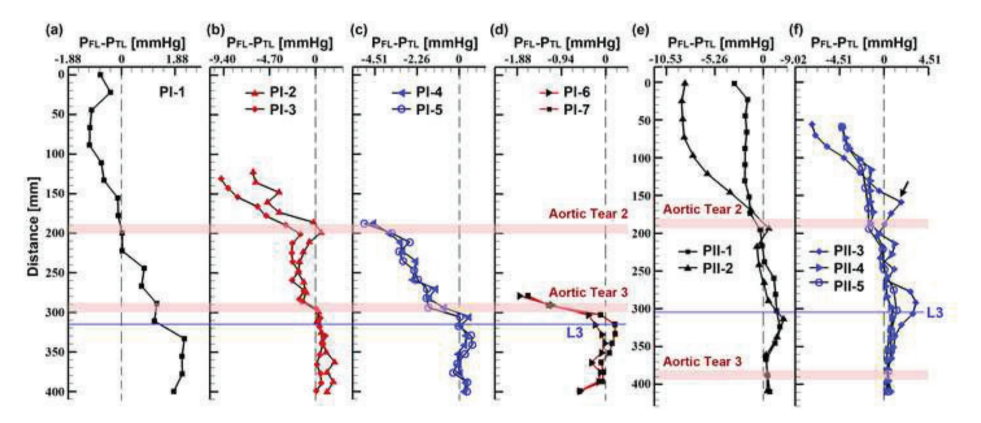


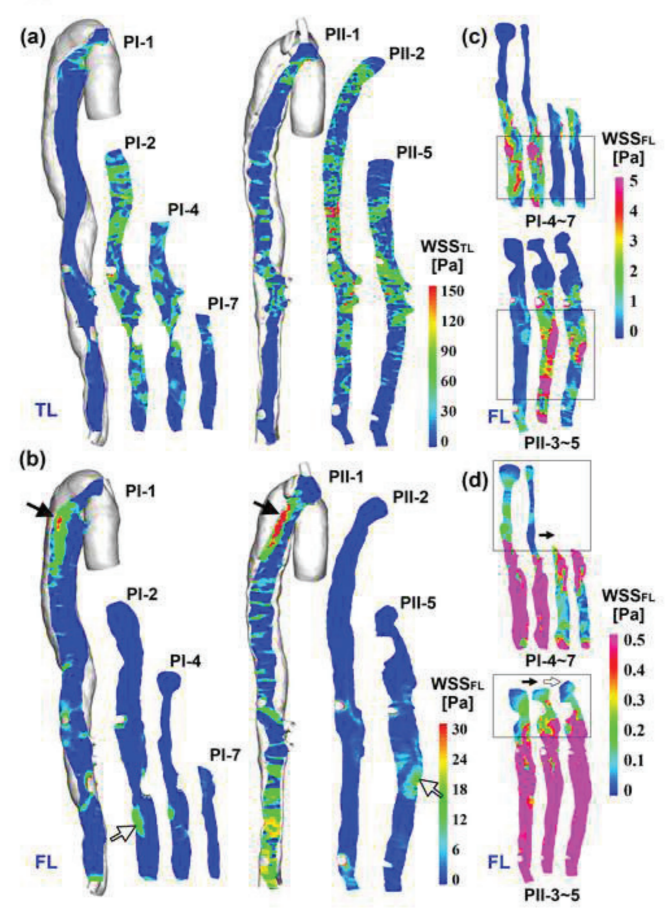


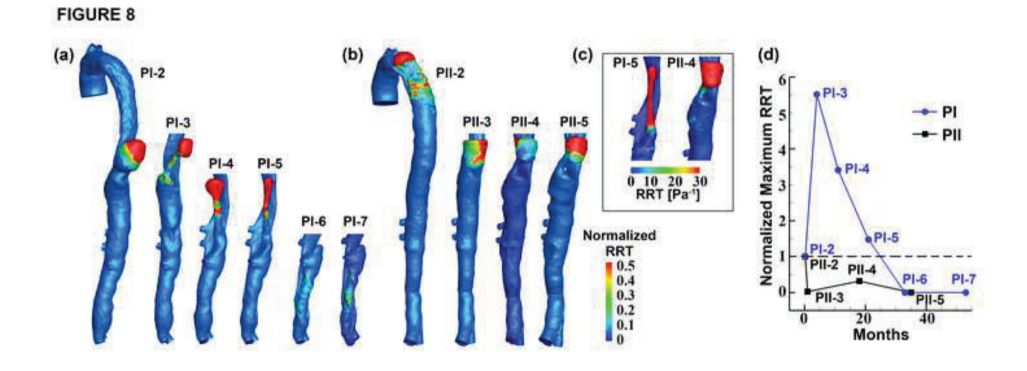












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