| 1 | A Comparison of Accuracy of Image- versus Hardware-based Tracking Technologies |
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| 2 | in 3D Fusion in Aortic Endografting |
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| 25 | Short Statement of Influence in Clinical Practice. |
| 26 | Fusion imaging is recognized as an important tool in complex aneurysm repair to |
| 27 | improve the success of implantation and decrease radiation dose and contrast use. |
| 28 | It has been previously impossible to compare accuracy of fusion systems because |
| 29 | they require fixed hardware, but a new cloud-based system is now available. We |
| 30 | compare the accuracy of two different types of fusion imaging. If confirmed, these |
| 31 | preliminary results could change clinical practice by encouraging further |
| 32 | development of automated image base tracking fusion process. |

35 ABSTRACT

36 Objectives

Fusion of three-dimensional (3D) computed tomography (CT) and intra-operative 2D imaging in endovascular surgery relies on manual rigid co-registration of bony landmarksand tracking of hardware to provide a 3D overlay (Hardware based tracking, HWT). An alternative technique (Imaged based tracking, IMT) uses image recognition to register and place the fusion mask. We present preliminary experience with an agnostic fusion technology that uses IMT, with the aim of comparing the accuracy of overlay for this technology with HWT.

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45 Method

Data was collected prospectively for 12 patients. All devices were deployed using 46 47 both IMT and HWT fusion-assistance concurrently. Post operative analysis of both 48 systems was performed by 3 blinded expert observers, from selected time-points 49 during the procedures, using the displacement of fusion rings, the overlay of vascular 50 markings and the true ostia of renal arteries. Mean overlay error as well as deviation 51 from mean error was derived using image analysis software. Comparison of mean 52 overlay error was made between IMT and HWT. Validity of the point-picking 53 technique was assessed.

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55 Results

IMT was successful in all of the first 12 cases, whereas technical learning curve challenges thwarted HWT in four cases. When independent operators assessed the degree of accuracy of the overlay, the median error for IMT was 3.9 mm (IQR; 2.89-6.24, max 9.5), versus 8.64 mm (IQR; 6.1-16.8, max 24.5) for HWT (p=0.001). Variance per observer was 0.69 mm² and 95% limit of agreement +/- 1.63.

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62 Conclusion

63 In this preliminary study, the error of magnitude of displacement from 'true 64 anatomy' during image overlay in IMT was less than for HWT. This confirms that

- ongoing manual re-registration, as recommended by the manufacturer, should be
- 66 performed for HWT systems to maintain accuracy. The error in position of the fusion
- 67 markers for IMT was consistent, thus may be considered predictable.

70 INTRODUCTION

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Endovascular interventions have expanded the treatment opportunities available to patients with aortic disease and have become progressively complex.^{1–3} When repair includes coverage of the visceral aortic segment, accurate device deployment and efficient catheterization of target vessels is critical. Fluoroscopic techniques require frequent contrast administration and high quality image recording (DSA) to visualize key structures, resulting in exposure of the patient and surgeon to considerable radiation,⁴ and may be associated with deterioration in renal function.^{5,6}

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80 Endovascular image fusion refers to the process of merging pre-operative imaging with intra-operative imaging to provide a 3D vascular mask.^{7,8} Several studies using 81 82 commercially available devices have documented variable reduction in radiation 83 exposure, and significant reduction in contrast usage.^{9–12} All commercially available 84 systems to date use hardware based tracking to position the mask on the 85 fluoroscopic image. Not all fixed imaging systems require cone beam CT (CBCT) to perform fusion imaging, but at our institution, CBCT is used to create an 86 87 intraoperative 3D volume that is co-registered with pre-operative imaging. The CBCT 88 data provides the basis for a 3D co-ordinate reference frame that is automatically 89 registered with fluoroscopic imaging, but also incorporates positional data for the 90 vascular landmarks acquired on pre-operative imaging. By combining both soft and 91 bony landmarks for registration, this technique should be superior to those using 92 registration of bony landmarks alone. The position of the image intensifier and 93 operating table are tracked with respect to the co-ordinate reference frame, allowing for appropriate vascular landmark representation when the fluoroscopic 94 image is changed.^{13,14} The reliability of this technique depends on the accuracy of 95 "hardware tracking" and the stability of the patient's position on the table once rigid 96 co-registration has been performed.¹⁵ Furthermore, considerable user interaction is 97 98 required to define the vascular landmarks on a workstation pre-operatively,

99 manually register the images, and correct registration errors intra-operatively that100 may arise from patient movement.

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102 A fully automated, image-based 2D-3D registration system that is independent of 103 imaging system manufacturer has been proposed by Carrell et al, and its initial use was described in 2010.¹⁶ This system provides several advantages including being 104 suitable for any theatre even those equipped with mobile C-arm; it is radiation and 105 106 contrast free for the initial registration; and being fully automated makes it "user 107 friendly" for the operator. The drawback of IMT is that it can only perform fusion on 108 +/-30 degree angles from a standard AP view, and it does require additional 109 equipment to be installed in theatre.

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111 The aim of this study is to compare the accuracy of the fusion overlay between two 112 systems that use different mechanisms to maintain accurate overlay of vascular 113 markings: hardware tracking, and image-based tracking. Because most fusion 114 systems are brand-specific, there has been no previous simultaneous comparison of accuracy between systems on the same patient in the same conditions. Thus, we 115 116 sought to compare the accuracy of an initial manual registration, followed by 117 hardware tracking using a commercially available device, against continuously 118 updated image-based matching in an investigative device.

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120 MATERIALS AND METHODS

All patients undergoing aortic repair between July 2015 and September 2015 by the Aortic Team at Royal Free London were included in this study. These patients were enrolled in a pre-market trial of CYDAR software and signed consent for involvement. The study was approved by NHS England (IRAS ID 158839) and was closed on September 30, 2015 in accordance with the approved protocol.

126

127 All patients with aneurysms underwent pre-operative high-resolution computed 128 angiography (CTA) as standard of care. All patients received stent-graft deployments 129 with fusion assistance using two different systems: a pragmatic application of a

130 commercially available device that uses hardware tracking (Siemens Artis Zeego, 131 Siemens Healthcare, Erlangen, Germany) and a novel image-based device the Cydar 132 EV system (Cydar Medical, Cambridge, UK) in order to allow for a comparison 133 between the two systems. Fusion would be considered successful if the initial images were available and assisted the surgical procedure. Fusion would be 134 135 considered a failure if no mask appeared on the screen, or if the position of the mask 136 was so far removed from reality that it was not helpful in the opinion of the 137 operating team.

- 138
- 139 Hardware-Tracking Fusion Protocol

140 A hardware-tracking fusion protocol for complex aortic repair has been used at Royal 141 Free London since October 2014. Prior to CBCT, the surgical team imports the pre-142 operative CTA onto the theatre-based workstation and marked the target vessels by drawing rings at the level of the vessel ostia using Syngo[™] (Siemens Healthcare, 143 144 Erlangen, Germany) software. After induction of general anaesthetic and after all adjustments are made to the patient's position, the patients are fully prepared and 145 146 draped to minimize any extraneous patient movement after registration. All staff retreat to a shielded and sterile control room prior to CBCT. A 5sDR (5 second 147 148 acquisition, taking 133 frames at 30 frames/sec) is used for all procedures. Rigid co-149 registration of the pre-operative CTA with the bony CBCT volume is then performed 150 by the surgeon or an expert radiographer through a manual process. Target vessel 151 rings are assessed intra-operatively on the fluoroscopy screen. Manual re-152 adjustments of the fusion overlay was not performed since we sought to compare 153 the accuracy of automatic image overlay in both systems after initial co-registration.

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155 Image-based Tracking Fusion Protocol

For each patient the Cydar EV system was also used to generate vascular landmarks which were viewable on an additional screen. Segmentation of the aorta and relevant visceral vessels was performed from the DICOM data of CTA using a semiautomatic method (thresholding followed by region growing), and then rings were manually drawn on the rendered surface by the software provider prior to the day of surgery. The software provider requires 24 hours to prepare the overlay mask. The

software then applies a computational algorithm on pre-operative CT volume to 162 163 generate a series of images (digitally reconstructed radiographs, [DRRs]) that mimic 164 fluoroscopic images across a range of C-arm rotations and magnifications to match vertebral bodies in both images. An intensity-based registration algorithm then 165 scans the DRR series for images with similar pixel distributions, and automatically 166 matches the most appropriate DRRs to the live fluoroscopic images throughout the 167 168 procedure. During each fluoroscopic position, the tracking software analyzes the 169 visualized field and attempts to identify vertebrae. If two or more vertebrae are 170 identified, the vascular overlay image created from CT angiography is projected. The 171 algorithm assumes there is a rigid relationship between CT and fluoroscopy, since 172 registration is based on vertebral bodies, and does not adjust for changes in spinal 173 position.¹⁷ The system works when the C-arm is angulated within 30⁰ craniocaudally and 40⁰ in an anterior-oblique direction, which is a range chosen by the 174 175 manufacturer that represents a balance between working range of the system and 176 speed of registration.

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178 Error Analysis

179 Evaluation of error in terms of displacement of fusion rings, or overlaid vascular 180 markings, and the true ostia of the renal arteries on the fluoroscopy screen was the 181 principal measure in this study, which required expert assessment of the true 182 anatomy. We enlisted the observations of three blinded expert observers to identify 183 the location of true renal arteries in each projection. For each case, fluoroscopic 184 screen shots containing representations of renal artery fusion markers for each 185 fusion system were saved and loaded for post-hoc analysis into RView imageanalysis software (https://www.doc.ic.ac.uk/~dr/software/; Imperial College 186 187 London) which was provided to us by the engineers at CYDAR. In order to provide data in millimetres, calibration was performed in each case against longitudinal rigid 188 189 landmarks on either a calibrated catheter or between two gold markers on a 190 fenestration. For patients receiving fenestrated grafts, conversion of pixels into 191 millimetres was performed using known diameters of fenestrations (either 6 or 8 192 mm), by measuring the number of pixels against this known dimension (figure 1a).

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194 For patients receiving standard infra-renal grafts conversion was performed in the 195 same manner using the longitudinal markings of a standard measuring pig-tail 196 catheter (distance between each marker: 10 mm). In both scenarios, fluoroscopic 197 images were chosen with measurement markings close to the centre of the screen. 198 Since soft tissues are not visible on fluoroscopy, observers were asked to pick their 199 best estimate of the centre of the renal ostia using images of the fully deployed graft 200 with bridging stents in situ (figure 1b). They were then asked to pick the centre of 201 the fusion markers derived from both the hardware-tracking (figure 1c) and image-202 based matching (figure 1d) systems. This process was performed for both renal 203 arteries in each case. For the standard infra-renal endovascular aneurysm repair 204 (EVAR) cases, three endovascular observers independently selected the centre of the 205 renal vessel ostium on the basis of the pre-deployment digital-subtraction 206 angiogram. They then selected the centre of the fusion marker. This point picking 207 procedure was repeated with three different observers to provide three error 208 recordings. The RView analysis software provides positional data in the form of pixel 209 co-ordinates for each selected point (x and y). Euclidean principles were used to 210 calculate the distance, in pixels, between the centre of the renal ostia and the fusion 211 markers (as selected by the observers), and was referred to as "error". After 212 conversion to millimetres, each case therefore contained data for two renal arteries, 213 each of which was comprised of three error recordings per fusion system used, that 214 was averaged to give a single mean error value per renal artery, for each fusion 215 system used.

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217 Statistical Analysis

218 Analysis was performed on SPSS 22.0 (IBM corporation, Chicago, III). Data was 219 treated in the following manner: For each renal fusion marker (hardware- and 220 image-tracked), mean values for error in 'x' and 'y' dimensions and error magnitude 221 were calculated across the three observers. Additionally, the difference to mean 222 error magnitude (for all three observers) for each renal was calculated for each 223 observer. Analysis using Pearson's second skewness coefficient found the data to be 224 not-normally distributed. Therefore, in order to compare the distributions in mean 225 error magnitude between the two groups, a non-parametric test was used for un-

226 paired continuous variables (Mann-Whitney U). To determine if there were any 227 significant differences between the expert "point-pickers", or observers, a Kruskal 228 Wallis test was performed comparing all recorded errors grouped according to 229 expert observer. To determine inter-observer variance and limits of agreement, a Bland Altman-type analysis was used,¹⁸ plotting the mean magnitude of error across 230 the three observers against difference to mean magnitude for each observer. P 231 232 values of less than 0.05 were considered significant. Calculation of measurement 233 variations were performed with the assistance of engineers at CYDAR imaging.

234

235 **RESULTS**

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237 Between July 2015 and September 2015, twelve patients underwent endovascular 238 repair under general anaesthesia for aortic aneurysms of varying morphology (Table 239 1) and consented to inclusion in this trial. Seven patients underwent fenestrated 240 endovascular aneurysm repair (FEVAR). Of these patients, four were group IV 241 thoracoabdominal and three were juxtarenal aneurysms. Five patients received 242 infrarenal EVAR, of which three also received iliac branched devices for iliac artery 243 aneurysms. Two patients received coil embolization of the internal iliac on the 244 contralateral side to the branched device. One patient had an isolated iliac aneurysm 245 treated in the presence of a previous endograft with type Ib failure. Mean age for 246 the cohort was 71.9 years (Standard Deviation (SD) 9.7 years). Median aneurysm sac 247 size (aortic or iliac as appropriate) was 6.1 cm (SD 1.1cm). Details of preoperative 248 demographics and intraoperative variables are described in Table 1

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All patients underwent successful aneurysm exclusion. Mean procedure time from first entering theatre to leaving theatre was 373 minutes (SD 92m) for fenestrated cases and 220 minutes (SD 45m) for iliac branch cases. The best estimate of procedure time collected at our institution is time from first dose of heparin and first dose of protamine, and for fenestrated cases was 192 minutes (SD 63m); this data was not available for iliac branch cases.

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257 Image Fusion Reliability

258 The simultaneous overlay of image- and hardware-based tracking in the same 259 patient was feasible in most patients included in the study, and a representative 260 example is shown in figure 2. In the twelve patients included in this study, image-261 based tracking was successful in all cases and hardware-based tracking was successful in 8 patients. In two patients for which hardware-based tracking was 262 263 unsuccessful, no fusion overlay appeared intra-operatively, whilst in a further patient the fusion markers were grossly misaligned, and rotated by 90⁰ in relation to the true 264 265 orientation of the aorta. This required manual re-adjustment of the hardware-266 tracked overlay, and the data was therefore excluded from final statistical analysis. 267 In these cases, failure of hardware-based tracking was due to operator error, and not 268 manufacturing defect, during the workflow of manual registration. In a fourth 269 patient, hardware-based tracking was not possible due to a concurrent update in the 270 hospital's picture archiving and communication system (PACS), preventing image 271 transfer of the pre-operative CTA necessary for 3D rendering and drawing of fusion 272 markers. For IMT cases, all had successful overlay masks projected, and the delay for 273 each different projection was less than 10 seconds for 55% of registrations, and was 274 less than 14 seconds for 92% of registrations.

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276 The mean magnitude of error (figure 3) for the hardware-based tracking system was 8.64 mm (IQR; 6.1-16.8, max 24.5), compared with 3.9 mm (IQR; 2.89-6.24, max 9.5) 277 278 for the image-based system (p=0.001). Figure 4 gives a positional representation of 279 the distribution of overlay errors registered on the coordinate system described in 280 the methods section. The symbols indicate the direction in which the overlay needs 281 to move in order to match the intra-operative renal position. The image-based 282 overlay markers were consistently located below and mostly on the right side of the true vessel ostium. In contrast, the hardware-tracking based overlay errors were of a 283 greater magnitude, particularly in lateral directions, and located above and below 284 285 the true vessel ostium. The inter-observer reliability of the blinded "point-picking" technique used by expert observers was good, with the variance per observer in this 286 study being 0.69 mm and the 95% limit of agreement being +/- 1.63 mm, as 287 288 indicated in the Bland Altman-type plot in figure 5.

289 **DISCUSSION**

This is the first preliminary study to compare accuracy of two different types of fusion imaging techniques applied to the same patient undergoing aortic repair. We observe a significant reduction in fusion overlay error (3.9 mm (IQR; 2.89-6.24, max 9.5) compared with 8.64 mm (IQR; 6.1-16.8, max 24.5) (*p*=0.001)) using a technique that relies on image, rather than hardware for tracking in complex and simple endovascular aortic procedures. The agreement between observers for this error was good.

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298 The use of a similarity-based measure to match digitally reconstructed radiographs 299 to real radiographs, using lumbar vertebrae as a means for rigid 2D-3D image 300 registration was initially proposed by Penney et al in 1998,¹⁹ but was not used 301 clinically due to limitations in fluoroscopic imaging techniques. A more recent study 302 describes the use of a prototype version of the Cydar EV system in a series of 303 retrospective registrations of pre-operative CT-angiograms with archived 304 fluoroscopic images, again using lumbar vertebrae as anchor-points for rigid 2D-3D image registration.¹⁶ The authors observed a mean error of 4.5 +/- 2.8 mm across a 305 306 total of 98 registrations. Using a newer iteration of the software in this study, we 307 observed a median error of 3.99 mm across 21 renal targets, which was superior to 308 hardware-based image tracking in a pragmatic trial. In the current market place, 309 both GE and Phillips now have proprietary methods for performing 2D-3D fusion 310 without use of a CBCT. In contrast, the routine use of CBCT-based fusion in complex 311 aneurysm repair began as early as 2009 at the Cleveland Clinic, and has enjoyed clinical use in many centres since that time.⁹ Removing the CBCT from the process of 312 fusion imaging may have the benefit of decreasing radiation dose while continuing to 313 314 provide accurate image guidance, however all systems still base tracking on 315 hardware which is subject to inaccuracy if the patient moves.

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There was a consistency to errors in image-based tracking that did not appear in hardware-based tracking. All errors appear to portray a slightly lower level of renal arteries. This observation is consistent with findings by Maurel et al, who evaluated the displacement of key visceral arteries by comparing pre-operative CTA with intra-

321 operative contrast enhanced CBCT, and found that both renal arteries were 322 predominantly displaced in a superior and left direction following the introduction of stiff endovascular instruments.²⁰ This seems to be true independent of the side of 323 324 large sheath access, which was different between Maurel et al's experience and our 325 own.. The impact of endovascular tools on soft tissue deformation was suggested by 326 Carrell et al as a possible reason for increased error during image-based registration when the aortic neck was angulated beyond 30⁰, since these relatively inflexible 327 devices tend to "straighten out" the aorta.¹⁶ Parallax or differences in body position 328 329 compared with CT scan protocol could also account for this error. By comparison, 330 inaccuracies due to respiratory movement are thought to be of lesser significance, particularly at the vessel origin.^{21,22} 331

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We observed errors of greater magnitudes during hardware-tracked fusion, 333 334 particularly in lateral directions, as well as errors occurring above and below the 335 intra-operative renal artery origin. These may be accounted for by the fundamental 336 differences in registration technique utilized by the two systems. In theory, the 337 continued and automated image-based matching ought to prevent errors relating to patient movement from occurring, since the system corrects for this by 338 339 automatically overlaying the most appropriate DRR matched directly to the patient. 340 In contrast, the hardware-tracked system adjusts its representation of fusion markers according to tracked movement of the C-arm and table in a 3D coordinate 341 342 system, on the assumption that the patient has remained static within that 343 coordinate system after initial rigid co-registration of bony landmarks on has taken 344 place. Our practice was to perform CBCT and the initial registration prior to 345 performing open surgical groin cut-downs, thereby minimizing the risk of 346 contamination. It is plausible that patient movement during this phase and during other manoeuvres that move the patient, such as brachial punctures, may have 347 348 contributed to the broader distribution of registration errors. Where possible, the 349 team was cautious to maintain the position of the patient throughout the procedure, 350 but despite this attention to detail, the movement was still observed. The authors of 351 this study acknowledge that instructions for use for the Artis Zeego clearly 352 recommend manual adjustment of the overlay following any manipulation or

353 movement of the patient. In practice, this would require a surgeon to leave the 354 operating theatre and sterile field to use the workstation, or the continued presence 355 of a trained radiographer with experience of using the system, which is not 356 pragmatic in our practice. The work flow for such a protocol is less intuitive and may 357 introduce greater error. The intention of the study was to evaluate the overlay accuracy of both systems when a "hands-off" protocol was applied during the 358 359 procedure, and to describe the impact of insensible movement on the accuracy of 360 fusion. The finding that patient movement did likely effect hardware based tracking 361 to a greater extent than imaged based tracking suggests that image based tracking 362 may be more resistant to the patient movement in a non-anesthetized patient, 363 which will be a point for future research.

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365 Hardware-tracked fusion failed in 4 patients, which is due to user error, but reflects 366 the multistep process entailed in this technique. In two instances, fusion markers did 367 not appear on the fluoroscopic screen, whilst in a third case the markers were 368 grossly rotated by 90⁰ in relation to the orientation of the aorta. In the fourth 369 patient, loss of communication of the PACS system rendered the overlay 370 inaccessible. These failures reflect the cumbersome process that hardware-based 371 tracking currently involves, with many different variables that might impact the 372 workflow. Registration in these cases was performed by senior radiographers who 373 had received intensive training on two separate sessions, each of two days in length. 374 Despite adequate training, the complexity of the registration work-flow seemingly 375 requires operators with a large amount of experience and regular exposure for it to 376 run seamlessly.

377

It is possible to compare these systems on factors other than accuracy. Certainly in its current form, the HWT system has a larger working range and uses proprietary software which precludes the installation of additional hardware into the operating theatre. Drawbacks of the image-tracked system include the time the registration process takes intra-operatively: each change in C-arm rotation requires a new match to be made between the fluoroscopic image and a DRR which takes several seconds. In some instances this matching cycle needs to be repeated, resulting in a delay

385 between the change in view and an appropriate overlay of up to 14 seconds. At 386 present, the system does not automatically register when the C-arm is angulated beyond 30⁰ and 40⁰ in cranio-caudal and anterior-oblique directions, respectively. 387 388 Whilst sufficient for visualization and cannulation of renal targets, cannulation of 389 mesenteric and coeliac vessels using lateral views and fusion guidance is not possible 390 with the present iteration of the software. Work is presently in progress to expand 391 the scope of available DRRs to enable registration during more angulated 392 fluoroscopic acquisitions. Until that time, use of both systems to augment data 393 available intraoperative seems most prudent.

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395 This is one centre's first attempt at use of image-based tracking, and comprises very 396 early experience. As such, there are a number of limitations to this study. Small 397 numbers of patients in this study could have adversely affected the level of 398 significance observed, and a larger study may provide a more accurate evaluation of 399 the true benefit of this technology. Defining the true centre of the vessel ostia in the 400 fenestrated cases presented a challenge, since it is our practice not to perform pre-401 deployment DSA and to minimize the use of contrast injections when cannulating 402 the target vessels. We relied instead on using three expert observers to make best 403 estimates with the fenestrated piece fully deployed and the bridging stents in situ, 404 since this provided the most accurate representation of the position of the renal 405 arteries. However, placement of the device could have contributed to the movement of the vessel ostia. Analysis of the distribution of recorded errors in 406 407 relation to mean error, however, demonstrated a small amount of variance between 408 observers and narrow limits of agreement, suggesting reliability of this method. The 409 lack of data describing patient movement during the procedure is an unfortunate 410 weakness, since the affect of patient movement on the magnitude and direction of 411 overlay inaccuracies in both systems cannot be fully determined. We used as a 412 reference distance the known diameters of the fenestrations for the fenestrated 413 cases, and the calibrated pigtail catheter for the infrarenal cases. We believe in most 414 cases these were perpendicular to the angle of the beam. However, this technique 415 could have a lack of precision and be slightly shorter than expected. For instance for 416 the infrarenal cases if the pigtail catheter is not strictly vertical; or for the

fenestrated cases if the lateral anterior and posterior markers are not on an horizontal line, then the distance between the top and the bottom markers of the window may not correspond to the highest and the lowest points. Finally, we did not take into account neck angulation and renal ostia position on a clockwise that could modify displacement after the insertion of the delivery system and consequently the measurements.

423 CONCLUSION

Synchronous fusion using two different techniques was feasible, and allowed for a 424 425 direct comparison of overlay accuracy for image-based and hardware tracking 426 systems. In this very preliminary study, errors in fusion overlay associated with 427 image-based tracking seem predictable and are of a smaller magnitude compared 428 with those observed in a pragmatic application of a hardware-tracked device. 429 Additionally, a major benefit from the image-based fusion is that it does not require 430 a pre-operative CBCT and could help in decreasing the radiation exposure. Further 431 investigation with a larger series is warranted

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437

438 *Conflict of Interest*

TMM is a proctor and consults for Cook Medical, as well as speaking arrangements
with Maquet Getinge Group. JC has spoken on behalf of Maquet Getinge Group.
The CYDAR team helped facilitate the measurements performed.

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510 Figures and Tables

512 Table 1: Demographics for included patients. (*SD: standard deviation; HTN: Hypertension; DM: diabetes mellitus; CCF: congestive cardiac failure; BMI: body mass index ; GFR: glomerular filtration rate; DAP: dose area product; CAK: cumulative air kerma*)

| | N (12) |
|--------------------------|------------------------------------|
| Age | 71.9 years (SD 9.7 yrs) |
| Male | 11/12 |
| Medical Comorbidities | |
| HTN | 12/12 |
| Dyslipidemia | 11/12 |
| Current Smoker | 2/12 |
| DM | 4/12 |
| CCF | 2/12 |
| BMI | 26.6 (SD 4.9) |
| Pre op GFR | 67.7 (SD 24.7) |
| Post op GFR | 64.8 (SD 23.9) |
| Aneurysm Characteristics | |
| Aneurysm sac size | 6.1cm (SD 1.08 cm) |
| Infrarenal Aneurysm | 2/12 |
| Iliac Artery Aneurysm | 3/12 |
| Juxtarenal aneurysm | 3/12 |
| Type IV TAAA | 4/12 |
| Intraoperative Variables | |
| DAP | 91.7 Gy.cm ² (SD 67.92) |
| КАР | 0.78 mGy (SD 0.69) |
| Volume of Contrast | 46cc (SD 14.9) |





523 Figure 1



529 Figure 2









546 Table and Figure Legend

Table 1: Patient demographics and aneurysm morphology. IA- Iliac artery. IRAAAinfrarenal abdominal aortic aneurysm. EVAR- endovascular aneurysm repair. FEVARfenestrated endovascular aneurysm repair. IBD- iliac branched device. Results are expressed in mean and standard deviation (SD). (*SD: standard deviation; HTN: Hypertension; DM: diabetes mellitus; CCF: congestive cardiac failure; BMI: body mass index ; GFR: glomerular filtration rate; DAP: dose area product; CAK: cumulative air kerma*)

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556 Figure 1: Calibration and Point-selection a) Conversion of pixels into mm, in this case 557 the known dimensions of a fenestration. b) Centre of the renal ostium is selected. c) 558 Centre of the hardware-tracked fusion marker is selected d) Centre of the image-

559 tracked fusion marker is selected

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561 Figure 2: Hardware-based tracking and Image-based tracking fusion systems applied

to two cases, with each row representing the same case. a) and c)- hardware-based
tracking. b) and d)- image-based tracking

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565 Figure 3: Comparison of mean error magnitude for all three observers per renal 566 artery, by fusion system used.

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Figure 4: Scatter-plot showing mean error per renal artery in x and y coordinates.
The symbols pointing away from the origin represent the direction in which the
overlay would have to be moved to match the actual vessel ostium.

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573 Figure 5: Bland Altman-type plot showing deviation from the mean error for each 574 renal artery. Each data point represents the difference between an observer's 575 recorded overlay error during point selection to the mean overlay error recorded for 576 all three observers for a given renal artery. The dotted lines represent the 95% limits 577 of agreement.

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